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Nonlinear optical spectroscopy and two-photon excited fluorescence spectroscopy reveal the excited states of fluorophores embedded in a beetle’s elytra

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Upon illumination by ultraviolet light, many animal species emit light through fluorescence processes arising from fluorophores embedded within their biological tissues. Fluorescence studies in living organisms are however relatively scarce and so far limited to the linear regime. Multiphoton excitation fluorescence analyses as well as nonlinear optical techniques offer unique possibilities to investigate the effects of the local environment on the excited states of fluorophores. Herein, these techniques are applied for the first time to study of the naturally controlled fluorescence in insects. The case of the male Hoplia coerulea beetle is investigated because the scales covering the beetle’s elytra are known to possess an internal photonic structure with embedded fluorophores, which controls both the beetle’s coloration and the fluorescence emission. An intense two-photon excitation fluorescence signal is observed, the intensity of which changes upon contact with water. A third-harmonic generation signal is also detected, the intensity of which depends on the light polarization state. The analysis of these nonlinear optical and fluorescent responses unveils the multi-excited states character of the fluorophore molecules embedded in the beetle’s elytra. The role of form anisotropy in the photonic structure, which causes additional tailoring of the beetle’s optical responses, is demonstrated by circularly polarized light and nonlinear optical measurements.

1. Introduction

Fluorescence occurs in many living organisms ranging from insects and birds to amphibians, mammals and plants [1–7]. These organisms’ tissues emit longer-wavelength light (typically in the visible band) when they are illuminated by shorter-wavelength light (usually, blue or ultraviolet). Such a light emission process is due to embedded fluorescent pigments, called fluorophores, such as green fluorescent protein (GFP), psittacofulvin, biopterin and papiliochrome II, within
the biological integuments of the living organisms. It results from the transition of fluorophores’ electrons between real states (i.e. long-lived observable quantum states) with the same spin multiplicity. The colours of emitted light span the whole visible electromagnetic spectrum from blue and green to yellow, orange and red. This phenomenon is thought to play a role in some living organisms’ visual communication such as species recognition, mate choice, agonistic behaviour, prey detection and camouflage [5,6]. Despite these important roles, fluorescence emission still remains under-explored from the biological, chemical and physical points of view, specifically in insects. Of course, in many cases, discerning whether fluorescence emission plays a functional role in the species’ coloration and visual pattern or whether it is a non-functional consequence of the presence of specific molecules within the living organism’s tissues may be complicated. To date, only a handful of organisms have been demonstrated to display fluorescence emission associated with biological functions [5,6].

The male *Hoplia coerulea* beetle (figure 1a), investigated here, is known to display an iridescent blue-violet coloration due to a porous periodic multilayer made of chitin found in the scales covering its body (figure 1b) [8]. Upon contact with liquids and vapours, its colour was found to turn green [9–12] due to the penetration of liquids into the scales. Interestingly, this photonic structure was demonstrated to contain naturally embedded fluorophores [13–15] and to give rise to controlled fluorescence emission from these fluorophores, through modifications of the optical system’s local density of optical states [14–17]. Upon contact with water, the fluorescence emission caused by one-photon absorption from these confined fluorophores was observed to change from turquoise to navy blue [14]. These water-induced changes were found to be reversible, implying that the fluorophores were not significantly altered from a chemical point of view by exposure to water.

This male beetle reflects incident light specularly. Fluorescence emission from its elytra takes place in a much larger solid angle than the angle of the very directional light reflection and could play a role in the signalling behaviour of the species. However, this species could have simply evolved a diffusively scattering surface that would be more efficient and scatter more light than the light emitted by fluorescence due to the known poor conversion efficiency of such processes in natural fluorophores, and the fact that sunlight is less intense in the UV range with respect to the visible range. The role of these fluorophores may simply be a by-product of another biological function such as photoprotection due to the increased UV absorption induced by the presence of these molecules.

The chemical and physical aspects of this fluorescent process are far from understood. Fluorescence emission from natural organisms’ photonic structures has never been investigated, neither by multiphoton excitation fluorescence techniques, nor by nonlinear optical techniques, although both are known as invaluable tools for the imaging and spectroscopy analyses of biological tissues [18]. Using one-photon excited fluorescence (OPEF) and two-photon excited fluorescence (TPEF) microscopy and spectroscopy, in addition to co- and cross-circularly polarized (CP) reflectance spectrophotometry and third-harmonic generation (THG) spectroscopy, we investigated the optical response of the male *H. coerulea* beetle, unveiling new features such as the optical role of the anisotropy in its photonic structure and its fluorophores’ multi-excited states (i.e. states excited by several photons) within much of the human visible spectrum.

One advantage of the two-photon excitation imaging and spectroscopy methods (that are based on a third-order optical process) with respect to one-photon excitation techniques lies in the nonlinear intensity-dependent absorption [19]. Therefore, two-photon absorption is limited to the focal volume within the investigated sample. In theory no photo-induced bleaching occurs outside the focal volume, in contrast with one-photon absorption, which usually gives rise to related strong bleaching in the whole illuminated volume [19]. In addition, TPEF allows for an extended penetration depth compared to OPEF (from a few tens to hundreds of micrometres since the excitation wavelengths are in the near-infrared [19], a range for which biological samples are known to be more transparent.

THG is a nonlinear process in which three photons combine, through three virtual excited states (i.e. very short-lived and practically undetectable quantum states) of the material molecules, into one single photon with a tripled frequency with respect to the incident light (i.e. a third of the incident wavelength). The main advantage of this technique, which has not been fully explored, lies in the possibility to perform label-free analysis both in imaging and spectroscopy [20].

Owing to its recently revealed fluorescent properties and its well-known optical linear response, *H. coerulea*’s photonic structures are a perfect case study for nonlinear optical investigations of fluorescent natural photonic architectures in both dry and wet states. TPEF and THG techniques were employed here for the first time in the context of natural photonic structures. The present circularly polarized light and nonlinear optical study, together with the steady state and time-resolved fluorescence investigation, reveals the difference between the exited states of the beetle’s fluorophores embedded within the photonic structure following one-photon and two-photon absorption as well as the effect of the local environment.
on their fluorescence emission properties. In addition, they unveil the role of the form anisotropy (i.e. direction-dependent morphology displaying structures smaller than incident wavelengths) of the photonic structure inside the beetle’s scales, which influences both reflected and emitted light.

2. Methods

2.1. Beetle samples

Male H. coerulea specimens were collected in Saumane and Auzon (France) in June 2014. No further sample preparation was necessary. All the analyses were performed on the beetle’s elytra.

2.2. Co- and cross-circularly polarized reflectance spectrophotometry

The beetle elytron was characterized using co- and cross-circularly polarized reflectance spectrophotometry. The experimental set-up was described in a previous work [21] and consisted of an Ocean Optics (Delray Beach, FL, USA) HR-2000 broadband fibre-coupled light source and an Ocean Optics USB2000+ spectrometer. CP light beam was generated by a rotatable polarizer and an achromatic Fresnel rhomb quarter-wave retarder, oriented at 45° azimuth. The CP light handedness was chosen by orientating the polarizer azimuth at either 0° or 90°. The incident light was focused onto a single beetle elytron at normal incidence using an achromatic 10 × objective lens, producing a beam spot diameter of approximately 30 μm. This lens also collected light reflected by the sample, which was routed by a beam-splitter through a CP analyser comprising the Fresnel rhomb and a second rotatable polarizer. The calibration was performed using a plane aluminium mirror.

2.3. Optical modelling

The CP reflectance of the structure was modelled for normal incidence using a one-dimensional 4 × 4 matrix method for anisotropic multilayers [22]. The structural dimensions upon which this representative model is based were identified in [9,14] and are as follows. The multilayer contained 12 bilayer periods, each comprising a 35 nm thick chitin layer (isotropic, n = 1.56) and a 140 nm thick air–chitin layer. The chitin was assumed to be lossless. The properties of the air–chitin layers were based on a laterally periodic assembly of rectangular chitin rods of width 90 nm, spaced by air gaps of 85 nm width (electronic supplementary material, figure S1). The air–chitin layers were modelled with form-birefringent refractive indices (i.e. refractive indices depending on the polarization and propagation direction of the light, due to form anisotropy) n∥ = 1.32 and n⊥ = 1.20, which were calculated using the equations given in [8]. A 100 nm thick cap layer of chitin (isotropic, n = 1.56) was applied, which represented the enclosing permeable envelope [11]. Disorder in the azimuthal orientation of the chitin rods was accounted for in the model firstly by applying a random azimuth orientation of the in-plane birefringent axes for each period of the structure. The result was averaged over 50 iterations, with a different set of random axis orientations applied to the structure in each iteration.

2.4. Spectrofluorimetry

Fluorescence measurements were performed using an Edinburgh Instruments (Livingston, UK) FLS920 UV–vis-NIR spectrofluorimeter equipped with a Hamamatsu (Hamamatsu City, Japan) R928P photomultiplier tube. Excitation and emission spectra were recorded using a 450 W xenon lamp as the steady state excitation source. The incident light formed a 45° angle with the direction normal to the surface of sample and the emitted light was detected at a 45° angle on the other side of the normal direction. The sample was excited at different excitation wavelengths within the range of the spectrofluorimeter used for these measurements: 350 nm, 375 nm, 400 nm, 450 nm, 500 nm and 550 nm. Time-resolved dynamics of fluorescence emission from the sample were recorded using a μF920H xenon Flashlamp light source, operating at a frequency of 100 Hz, and with pulse width of 1 μs. The same excitation wavelength as for the steady state measurements were used. The emission dynamics were observed at the maximum of the emission peak. The recorded time-resolved dynamics were fitted to double exponential functions, with coefficients of determination R² ranging from 0.985 to 0.990 for all fits. The polarization steady state and time-resolved measurements were recorded on the same spectrofluorimeter which is equipped with Glan–Thompson polarizers. The polarization state of excitation light for both measurements was selected by inserting a polarizer in the beam path before the sample. The polarizer was computer-controlled and the angles of light polarization could be varied between 0° and 90°. The excitation light polarization angle was tuned by steps of 15°. The unpolarized emission spectra were corrected for detector sensitivity.

2.5. Multiphoton microscopy and spectrometry

The multiphoton microscopy images were taken with an Olympus (Münster, Germany) BX61 W1-FV1200-M system. A Spectra-Physics (Santa Clara, CA, USA) InSight D5+ laser (82-MHz repetition rate, 120 fs pulse width, p-polarized) was used for excitation, at 800 nm, 900 nm and 1000 nm fundamental wavelengths. To regulate the intensity of the laser beam, an achromatic half-wave plate and an s-directed polarizer were placed directly behind the laser, which resulted thus in a s-polarized laser beam, which was used for all related experiments. A 15° LMV objective (NA 0.3) and 50° SLMPlan N (NA 0.35) were used for magnification, signal detection was non-descanned in backwards reflection through a Hamamatsu R3896 photomultiplier tube. Three detection cubes were used, each for a different fundamental wavelength. For 800 nm, the second-harmonic generation (SHG) and TPEF signals were divided through a 425 LPXR dichroic mirror. The SHG signal was further filtered with a 405/10 bandpass filter. Similar detection cubes were used with a 470 LPXR dichroic mirror and 450/7 × bandpass filter (900 nm) and a 525 LPXR dichroic mirror and 500/20 nm bandpass filter (1000 nm). The signal at each pixel is depicted in the images as different intensities in a false red colour. Owing to the elytra curvature and roughness, an extended depth of focus function was used in order to obtain in-focus microscopic images. No bleaching was observed, as the fluorescence signal stayed stable over time.

The spectral measurements were performed using the same laser as for the multiphoton microscopy observations at a 45° incidence angle, with different excitation wavelengths: 800 nm, 900 nm, 1000 nm, 1100 nm, 1200 nm and 1300 nm. An achromatic lens (60 nm EFL) was placed in front of the beetle’s elytron in order to focus the laser beam. The spectra were recorded in reflection and collected by using an achromatic aspheric condenser lens (focal length 30 mm). The beam was focused on the entrance slit of a Bruker (Billerica, MA, USA) 500 is/sm spectograph, and then recorded by the Andor Solis (Belfast, UK) iXon Ultra 897 EMCCD camera. In order to eliminate laser light entering from the detector, additional SCHOTT (Mainz, Germany) BG39 and KG5 filters were positioned in the beam path in front of the sample. Additionally, a long-pass filter was placed in front of the first lens to remove the higher-order harmonics of the laser itself. The linear polarization THG measurements were performed by placing and rotating a second half-wave plate in the beam path, behind the polarizer. The 0° and 90° angles of light polarization corresponded respectively to s- and p-polarizations. After subtracting the background, the THG peak was fitted to a Gaussian curve and
3. Linear optical and fluorescence properties

The male _H. coerulea_ beetle displays a vivid blue-violet iridescent colour (figure 1a) due to a porous multilayer structure (figure 1b) located inside the circular shaped scales covering its elytra and thorax [8]. The photonic structure is enclosed by a 100 nm thick envelope [11] and consists of a periodic stack of thin cuticle layers and mixed air–cuticle porous layers (figure 1b). The latter exhibit rod-like spacers giving rise to some anisotropy. In total, the structure comprises 12 periods. Using linear optical and OPEF techniques, the anisotropy of the beetle’s photonic structure (figure 1c) is shown to play a role in the reflected CP light, and the fluorophores embedded within this structure are found to possess several excited states, the decay time of which depends on the linear polarization state of the excitation light.

Previous linear optical studies of the photonic multilayer structure of _H. coerulea_ have included measurements of the unpolarized reflectance [8,15], with corresponding simulations that applied isotropic approximations of the structure [8,13,14]. In this work, polarized-light techniques are employed in order to identify the influence of anisotropy on the reflectance. The response of the structure to incident CP light was measured in terms of CP reflectance components (see §2). Both cross-CP and co-CP configurations were included, where cross-CP denotes either left-handed CP (LCP) light reflected from right-handed CP (RCP) incident light or vice versa, and co-CP refers to the incident and reflected CP components having the same handedness. The results (figure 2a) show two distinct pairs of spectra. The cross-CP spectra both have a peak centred on 500 nm. A cross-CP response is similar to that of a planar mirror. Additionally, this structure exhibits conspicuous reflectance spectra in both co-CP configurations, with a smaller peak centred on 450 nm. Such a co-CP reflection response is a feature that has not been extensively observed in scale-bearing insects, and is a strong indication of the presence of anisotropy within the photonic structure. Both intrinsic and form birefringence can be anticipated to confer optical anisotropy upon this structure. Firstly, the cuticular material could have intrinsic birefringence due to the anisotropic self-assembly of the cuticular macromolecules within the multilayer structure [23,24]. Secondly, the presence of form birefringence [25–27] is suggested by the morphological characterization (figure 1c), arising due to the rod-like inter-layer spacers forming the mixed air–cuticle layers. The azimuthal directions of these rods typically differ from one such layer to the next, similarly to a woodpile structure [28].

The CP reflectance of the structure was modelled for normal incidence using a one-dimensional matrix method for anisotropic multilayers [22]. The structural dimensions upon which the representative model is based were identified in [9,14] and are detailed in §2. In the absence of direct evidence of intrinsic birefringence for this system, these anisotropic refractive indices were obtained from a calculation of the form birefringence, using the equations given in reference [8]. Morphological studies [8,15] have indicated that the azimuth of the chitin rods is disordered, with variations both from layer to layer and with spatial position within each layer (figure 1c). This disorder was accounted for in the model firstly by applying a random azimuth orientation of the in-plane birefringent axes for each period of the structure. Secondly, the result was averaged over multiple simulations, with a different set of random axis orientations applied to the structure in each iteration. The results (figure 2b) show that the model reproduces the main features of the experimentally observed CP spectra (figure 2a), including the presence of the co-CP signals displaying the same shape. This result and its match with the experimental measurement supports a model of the multilayer
in which the air–chitin layers have form anisotropy that contributes to the CP reflectance.

The morphology identified here in *H. coerulea* offers comparison with alternative reflecting structures displaying anisotropy in scale-bearing beetles. The examples of *Lepidota stigma* and *Cyphochilus* spp. scarab beetles obtain a diffuse white appearance through incoherent scattering using a highly disordered network of chitin rods [29–31]. There, the anisotropy of the system enhances the brightness by favouring out-of-plane scattering. The case of *H. coerulea* is different since its optical response is angle- and wavelength-dependent due to coherent scattering.

Importantly, the peak width and position for OPEF emission strongly depends on the excitation wavelength (figure 2c). The peak position shifts from 434 nm to 661 nm as the excitation wavelength is increased from 350 nm to 550 nm. This excitation wavelength dependence of the emission peak position suggests that the embedded fluorophores, the molecular composition of which remains unknown, exhibit complex dynamics with several excited states. In complex organic fluorescent molecules, the presence of several such excited states possibly indicates exciton resonances (namely, energy transfers between different donor and acceptor states) [32,33]. The presence of mixtures of different fluorophores is less likely as this would imply multi-peaked fluorescence emission spectra (at some excitation wavelengths), whereas here we observe single peaks.

In addition, the OPEF emission spectra depend on the linear polarization state of the excitation light (figure 2; electronic supplementary material, figure S2). As the rotatable polarizer angle $\phi$ is increased from $0^\circ$ to $90^\circ$, the emission intensity increases by a factor 1.55, independent from the excitation wavelength. Owing to the weak refractive index contrast between the incident medium (air) and the cuticle material, such an increase in intensity cannot only be explained by a difference of light transmission as a function of the linear polarization state [24]. Furthermore, the radiative decay time $\tau$ (i.e. decay time related to light emission) is also a function of the excitation light polarization (table 1), specifically at emission wavelengths far from the photonic band gap of the beetle’s scale structure (with a $45^\circ$ incidence, it ranges from about 410 nm to about 470 nm). For instance, at 456 nm, $\tau$ increases from 6.6 ns to 9.3 ns when $\phi$ is increased from $0^\circ$ to $90^\circ$. At emission wavelengths far from the photonic band gap, $\tau$ is almost constant. Similar but non-polarized measurements reported previously gave rise to much shorter radiative decay times [14] ranging from 3.9 ns in the dry state and inside the gap to 1.4 ns in the wet state and outside the gap (electronic supplementary material, table S1). This means that the real excited states and their related radiative decay times are affected by the polarization state of the excitation light.

The photonic structure anisotropy gives rise to an additional moulding of the reflected light, namely through the selection of reflected light wavelengths depending on the CP light handedness. The embedded fluorophores are found to possess various excited states, the emission spectra and the decays of which are functions of the polarization states of the excitation light.

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4. Two-photon excited fluorescence emission

Upon excitation with an incident light beam with a wavelength ranging from 800 nm to 1300 nm (figure 3), a TPEF signal is measured from the beetle’s scales, even at relatively low incident laser power (20 mW), producing high-resolution images (figure 3a–c; electronic supplementary material, movie S1). These measurements unveil the presence of multi-excited states of the fluorophores.

Using 800 nm excitation a TPEF signal with a peak located around 523 nm is detected (figure 3c). No SHG response (expected at 400 nm) could be detected. TPEF emission recorded with different excitation wavelengths (figure 3c) confirms the existence of multi-excited states in the fluorophore molecules. Furthermore, the contrasting nature of one- and two-photon absorption processes is expressed in the differences between the TPEF and OPEF spectra in terms of their spectral shapes, peak positions and full widths at half maximum (FWHM) (figure 4a). States excited by one-photon absorption with a given wavelength give rise to emission spectra that differ significantly from the states excited by two-photon absorption with twice the corresponding wavelength. For instance, in OPEF and with a 400 nm excitation wavelength, the emission spectrum peaks at 474 nm, while in TPEF and with a 800 nm excitation wavelength, the emission peak is located at 537 nm. In the case of two-photon excitation, some vibrational transitions are additionally enhanced [34], which affect non-radiative relaxation pathways (i.e. relaxation in which the energy is not released as photons but as molecular vibrations), while in the case of one-photon excitation, simple electronic transitions between real states dominate.

Upon contact of the elytron surface with water, the TPEF response decreases in intensity and red-shifts by about 6 nm (figure 3d). Similarly to previously reported OPEF measurements [14], the intensity reduction can be explained by a change in the scattering efficiency at the sample surface due to the presence of water. The peak shift may seem, however, to contrast with the previously reported 17 nm blue-shift of

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the OPEF peak wavelength (from 463 nm to 446 nm) [14]. In the dry state, with a 355 nm excitation wavelength, the emission peak was positioned within the photonic band gap, a range where the local density of optical state is very sensitive to changes in the refractive index contrast induced upon contact with water [14]. This opposite and smaller shift observed with TPEF can be explained by the fact that the selection rules are entirely different for the one- and two-photon absorption processes [34]. The corresponding excited states are different, as observed in figure 4a, and, therefore, their radiative and non-radiative relaxations are different in the presence of liquid. Additionally and in contrast with the OPEF emission peak, the TPEF emission peak is far from the photonic band gap, i.e. in a spectral range where the local density of optical states is not much affected by the presence of liquid in the pores of the structure [14]. All together this produces a 6 nm red-shift of the TPEF peak and a 17 nm blue-shift of the OPEF peak. Finally, the liquid-induced red- (blue-) shift of TPEF (OPEF) suggests that the polarities (i.e. electric charge separation) of the fluorophores in the excited states involved in one- and two-photon absorption processes are different. In the TPEF case, the presence of water modifies slightly the environment around the ground and real excited states making them slightly more polar in the wet sample than in the dry sample, resulting in a decrease of the energy difference between excited and ground states [32], as indicated by the red-shift of TPEF upon contact with water.

These observations of the TPEF signal have highlighted the multi-excited states nature of the fluorescent molecules. Upon contact with water, the difference between the measured red-shift of the TPEF peak and the blue-shift of OPEF may be explained by the emission wavelength band that is far from the photonic band gap, by the difference in selection rules for one- and two-photon absorption processes and by the possible difference in polarities of the excited states.

5. Third-harmonic generation

Using THG spectroscopy, the beetle’s elytra are found to give rise to a polarization-dependent signal which represents an additional consequence of the photonic structure’s form anisotropy.

In spite of the absence of a detectable SHG signal at 20-mW incident power (figure 3c), a THG signal with a peak around 433 nm was measured using a higher incident power (250 mW) and an excitation (fundamental) wavelength equal to 1300 nm (figure 4b). The peak positioned around 650 nm is

![Figure 3](image-url). Two-photon excitation fluorescence response of *H. coerulea*’s elytra. The elytra of the male *H. coerulea* beetle give rise to a strong TPEF response with (a) a 900 nm and (b) a 800 nm excitation light allowing one to observe the scales covering the beetle’s elytra (with a 75 mW incident power). The imaged area corresponds to the junction between both elytra. Similarly to the previously reported OPEF measurements [14], in the areas where scales overlap, the TPEF intensity is higher. (c) With a low incident power (20 mW), TPEF excitation spectra can be measured with different excitation wavelengths. (d) Upon contact with water, a small red-shift of the peak wavelength and a decrease in peak intensity are observed with an excitation wavelength equal to 800 nm and an incident power of 20 mW. Scale bars: (a) 200 μm and (b) 100 μm.
mainly due to TPEF signal, which may possibly obscure a much weaker SHG signal. The intensity of the THG peak depends on the polarization state of the incident light (figure 4c). THG can arise from any material regardless of the symmetry exhibited by the constituting molecules. In addition to the chemical composition of H. coerulea’s scales and the structure of molecules that form this biological material (which are far from perfectly known [11], like for any other beetle), the nonlinear optical response from the beetle’s elytra is likely to be affected by light interference within the host material, i.e. by the scale’s photonic structure. The polarization-dependent response could be attributed to the form anisotropy of the material, highlighted by the co- and cross-CP reflectance measurements (figure 2a,b), since the THG intensity of an isotropic material does not depend on the incident light polarization [20,35–37].

6. Conclusion

Beyond its structural blue-violet coloration, the male H. coerulea beetle is an instructive example of fluorescence emission controlled by natural photonic structures. Using fluorescence, linear and nonlinear optical techniques, we have gained a better understanding of the beetle’s elytra optical properties and of the excited states of the embedded fluorophores. Observations of a non-vanishing reflection in co-CP light configuration, as well as of a polarization-dependent THG signal, unveil the optical role of form anisotropy, potentially responsible for an additional control of the fluorescence emission. OPEF and TPEF observations show that one single type of fluorescent molecules having several excited and multi-excited states is present in the scales’ biological tissues. The corresponding one-photon and two-photon absorption excited states were observed to exhibit very different emission spectra. A spectral red-shift of the TPEF peak induced upon contact of the elytra with water suggests that the multi-excited states may have different polarizations. This investigation we hope will promote the study of photonic structures in natural organisms using multi-photon excited fluorescence and nonlinear optical techniques in order to derive a comprehensive picture of their fluorescent and optical properties. This novel approach is expected to result in a deeper questioning of the biological roles of fluorescence and other optical properties of living organisms.

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