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Coexpression of Two Visual Pigments in a Photoreceptor Causes an Abnormally Broad Spectral Sensitivity in the Eye of the Butterfly *Papilio xuthus*

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The compound eye of the butterfly *Papilio xuthus* consists of three different types of ommatidia, each containing nine photoreceptor cells (R1–R9). We have found previously that the R5–R8 photoreceptors of type II ommatidia coexpress two different mRNAs, encoding opsins of green- and orange-red-absorbing visual pigments (Kitamoto et al., 1998). Do these cells contain two functionally distinct visual pigments? First, we identified the sensitivity spectrum of these photoreceptors by using combined intracellular recording and dye injection. We thus found that the R5–R8 of type II ommatidia have a characteristic sensitivity spectrum extending over an excessively broad spectral range, from the violet to the red region; the photoreceptors are therefore termed broadband photoreceptors. The spectral shape was interpreted with a computational model for type II ommatidia, containing a UV visual pigment in cells R1 and R2, two green visual pigments in cells R3 and R4, a far-UV-absorbing screening pigment (3-hydroxyretinol) in the distal part of the ommatidium, and a red-screening pigment that surrounds the rhabdom. The modeling suggests that both visual pigments in the R5–R8 photoreceptors participate in phototransduction. This work provides the first compelling evidence that multiple visual pigments participate in phototransduction in single invertebrate photoreceptors.

**Key words:** vision; color; optical filters; insects; rhodopsin; signal transduction

**Introduction**

It is a generally accepted concept in vision research that single photoreceptors contain a single type of visual pigment. However, recent studies have demonstrated that some photoreceptors violate this dogma: they coexpress multiple visual pigments (Röhllich et al., 1994; Sakamoto et al., 1996; Kitamoto et al., 1998; Glossmann and Ahnelt, 2002; Parry and Bowmaker, 2002). For example, a subset of murine cone photoreceptors contains both shortwave- and midwave-absorbing visual pigments (Applebury et al., 2000). An obvious question that arises is whether these visual pigments participate in phototransduction. Some electrophysiological measurements have been performed (Makino and Dodd, 1996; Lyubarsky et al., 1999), but such attempts are rather rare, and therefore additional proof would be desirable.

In the course of studies on the color vision of the butterfly *Papilio xuthus*, we discovered that some photoreceptors coexpress two mRNAs encoding different visual pigment opsins (Kitamoto et al., 1998). This report addresses whether these photoreceptors contain two physiologically active visual pigments.

An ommatidium of *Papilio* contains nine photoreceptors (R1–R9) (see Fig. 1d). The rhabdomeres of the photoreceptors, harboring the visual pigments, together form the fused rhabdom, a cylindrical structure that acts as an optical waveguide. Clusters of either red or yellow pigment, lining the rhabdom (see Fig. 1b) (perirhabdomeral pigmentation), act as filters of light propagating along the rhabdom. According to the pigmentation, *Papilio* ommatidia can be divided into three types: type I are strongly red pigmented and nonfluorescent, type II are pale-red pigmented and emit a whitish fluorescence under UV light (see Fig. 1c), and type III are yellow pigmented and nonfluorescent (Arikawa and Stavenga, 1997; see cover of this issue).

We found five classes of spectral receptors in the *Papilio* retina, peaking in the UV, violet, blue, green, and red wavelength regions (Arikawa et al., 1987). We localized the receptors in the three types of ommatidia by *in situ* hybridization (see Fig. 1, table): we found that the *Papilio* retina expresses five mRNAs of visual pigment opsins, termed *P. xuthus* UV (PxUV), blue (PxB), and long-wavelength 1–3 (PxL1–PxL3) (Kitamoto et al., 1998, 2000), creating UV-, blue-, and long-wavelength-absorbing visual pigments, respectively. By combining this finding with the electrophysiological results, we concluded that PxL1 and PxL3 construct green-absorbing visual pigments, whereas PxL3 constructs orange-red-absorbing visual pigment. Interestingly, we found a subset of ommatidia whose R5–R8 photoreceptors were positive for both PxL2 and PxL3 mRNAs (see Fig. 1d,e; highlighted in the table). If both visual pigments trigger the phototransduction process in a photoreceptor cell, the sensitivity spectrum should be broader than those of receptors containing only PxL2 or PxL3.

In our early recordings, we had encountered units with broad sensitivity spectra. We initially considered that they might originate from multiple photoreceptors, artificially coupled because of low-quality penetrations. However, detailed analysis of the...
units, combining electrophysiology, histology, and computational modeling, provided evidence that the recordings were in fact from single R5–R8 photoreceptors of type II ommatidia. In this report, we present the results that led to this conclusion, and we will argue that both PxL2 and PxL3 are physiologically active in these photoreceptors.

Materials and Methods

**Animals.** Spring-form males of the Japanese yellow swallowtail butterfly *P. xuthus* were used within 3 d of emergence. The butterflies were reared on fresh citrus leaves at 25°C under an 8/16 hr light/dark cycle. The pupae were stored at 4°C for at least 3 months and then allowed to emerge at 25°C.

We restricted the following analysis to the ventral two-thirds of the retina, because in the dorsal one-third, the cornea fluoresces under UV illumination, which obscures the ommatidial fluorescence in type II ommatidia. Type II ommatidia are presumably absent in the dorsal region (Arikawa and Stavenga, 1997).

**Electrophysiology.** A butterfly whose wings and legs were removed was fixed on a plastic stage with its dorsal side up and then mounted in the electrophysiological setup in a Faraday cage. A silver wire inserted in the stump of an antenna served as the indifferent electrode. To insert a glass micropipette into the eye, a hole covering ~10–20 facets was made in the dorsal region of the eye with a razor blade.

Monochromatic light stimuli were provided by a 500 W xenon arc lamp through 22 narrow-band interference filters ranging from 300 to 700 nm. The intensity of the light stimulus was attenuated with neutral-density filters and an optical wedge with a range of 4 log units. The light was focused on the tip of an optical fiber that led the light into the cage. The other end of the optical fiber was attached to a Cardan arm perimeter device, where it provided a point source of light (1.6° in diameter). The quantum flux of each monochromatic stimulus was measured by a radiometer (model 470D; Sanso, Tokyo, Japan), and the maximum quantum flux of each monochromatic stimulus at the corneal surface was adjusted with another optical wedge to $5.0 \times 10^{11}$ photons $\cdot cm^{-2} \cdot sec^{-1}$ ($-2.5 \times 10^8$ photons $\cdot facet^{-1} \cdot sec^{-1}$, with a facet diameter of 25 $\mu$m). A glass micropipette filled with 5% Lucifer yellow aqueous solution (resistance, $-150$ MΩ) was inserted into the retina through the hole made in the cornea. A successful penetration of a photoreceptor cell was deduced from a resting potential of approximately $-50$ mV and responsiveness to dim light. The optical axis of the responding unit was located by moving the tip of the optical fiber until dim white light flashes produced a maximal response.

We recorded from nearly dark-adapted photoreceptors. To minimize the effect of light adaptation during the experiment, we used light flashes that were limited to 30 msec in duration. This is the shortest light stimulus that still produces a receptor potential with an amplitude of the initial transient component that is the same as that elicited by longer (e.g., 1 sec) stimuli.

The receptor potentials were recorded through a preamplifier (MEZ-7200; Nihon-kohden, Tokyo, Japan) connected to an oscilloscope (VC-11; Nihon-kohden). First, the spectral type of the unit was determined by a series of monochromatic flashes with an intensity of $5.0 \times 10^{10}$ $-$ $10^{11}$ photons $\cdot cm^{-2} \cdot sec^{-1}$. When the spectral response profile indicated a broadband unit, we measured the response–stimulus intensity ($V$–$log I$) functions at several wavelengths in the intensity range of $5.0 \times 10^{-1} - 10^{11}$ photons $\cdot cm^{-2} \cdot sec^{-1}$ ($-2.5 \times 10^8$ photons $\cdot facet^{-1} \cdot sec^{-1}$) in some initial cases. Data were fitted to the Naka–Rushton function: $V = V_{max} \cdot f(I) / (F + K)$, where $f(I)$ is the stimulus intensity, $V$ is the response amplitude, $V_{max}$ is the maximum response amplitude, $K$ is the stimulus intensity eliciting 50% $V_{max}$, and $n$ is the exponent. We thus confirmed that the $V_{max}$ and $n$ are not significantly different between $V$–$log I$ functions recorded at different wavelengths. We therefore recorded a single $V$–$log I$ function at 540 nm for each unit. The best fit of the Naka–Rushton function was taken as the $V$–$log I$ function of the given unit.

Recorded units were subjected to additional analysis only when the $V_{max}$ was $>40$ mV. Amplitudes of the spectral responses were extrapolated to the $V$–$log I$ function of a given unit to transform the amplitudes into photon numbers required for the responses. The normalized reciprocal of the relative photon numbers then yielded the sensitivity.

The recorded broadband units were marked by Lucifer yellow. We delivered Lucifer yellow from the tip of the electrode by applying a hyperpolarizing DC of 2–5 nA for 5–10 min. The animal was then unmouted from the Faraday cage and positioned under an epifluorescence microscope (BX-60; Olympus Optical, Tokyo, Japan). The ommatidia containing the Lucifer yellow-injected unit was identified and photographed under violet excitation (dichroic cube U-MNBV; excitation bandpass filter at 420 nm and emission cutoff filter at 470 nm). Subsequently, the excitation light was switched to UV (dichroic cube U-MWU; excitation bandpass filter at 350 nm and emission cutoff at 420 nm) to correlate the stained ommatidium with the array of fluorescing ommatidia.

**Histology.** After the electrophysiology and the fluorescence microcopy, the compound eye containing the Lucifer yellow-injected unit was processed for light-microscopic histology. The eye was isolated and fixed at room temperature for 30 min in 4% paraformaldehyde in 0.1 M sodium cacodylate buffer at pH 7.4 and then dehydrated with an acetone series and embedded in Spurr’s resin. Sections (10–14 $\mu$m thick) were cut with a rotary microtome. The sections were first observed under the epifluorescence microscope with violet excitation to localize the unit within the ommatidium, and subsequently normal transmission light was used to identify the color of the ommatidial pigmentation.

**Modelling.** The computational model used for calculating the sensitivity spectrum of the R5–R8 cells (see Figs. 2, 4) was essentially identical to that of Arikawa et al. (1999a,b), but the parameters were slightly adjusted. We calculated the absorption of light entering type II ommatidia on-axis in the individual photoreceptors (R1–R9). The change in light flux along the rhabdom was described by Snyder et al. (1973) as follows:

$$\eta(\lambda) = \left[1 - \eta(\lambda)f(z,\alpha,\sigma)I(z,\lambda)\right]^{1/10kat},$$

where $I(z,\lambda)$ is the light flux at a distance $z$ from the tip of the rhabdom, $\eta(\lambda)$ is the fraction of the light flux propagated within the rhabdom boundary, $\lambda$ is the wavelength of light, $f(z)$ is the fractional cross-section taken up by photoreceptor R1–R9, $\alpha$ is the peak absorbance coefficient of visual pigment $\beta$, $\sigma$ is the peak absorbance coefficient of the red-screening pigment in the photoreceptor, and $\alpha$ and $\sigma$ are the (normalized) spectral absorption coefficients of the visual and screening pigments, respectively. The light absorbed by the visual pigment in each photoreceptor, integrated over the length of the photoreceptor, yields its absorption spectrum. Normalization then yields the sensitivity spectrum.

We have modeled the rhabdom as a circular cylinder. On the basis of detailed anatomy, the assumptions were as follows. The total length of the rhabdom is 500 $\mu$m; a distal layer of 260 $\mu$m is fully occupied by the R1–R4 photoreceptors; in a transition zone of 70 $\mu$m, the R1–R4 rhabdome reduces in size, whereas the participation of the R5–R8 cells increases; the latter cells fully occupy the proximal 140 $\mu$m; the basal 30 $\mu$m is taken up by the R9 photoreceptor; and each of the R1–R4 and R5–R8 occupy one-quarter of the rhabdomere cross-sections in the distal and proximal layers, respectively. The fluorescing, UV-absorbing 3-hydroxyretinol is homogeneously distributed in the distal 70 $\mu$m, with an absorption coefficient of 0.2 $\mu$m$^{-1}$. The red-screening pigment is present in cells R3–R8, with an absorption coefficient of 0.005 $\mu$m$^{-1}$. The visual pigment peak wavelengths are 365 nm (PxUV), 530 nm (PxL1), 515 nm (PxL2), and 575 nm (PxL3); the photoreceptors R1 and R2 both contain exclusively PxUV; R3 and R4 contain equal amounts of PxL1 and PxL2; R5–R8 contain PxL2 and PxL3 in a concentration ratio of 3:1 (see below); and R9 contains both PxL1 and PxL2. The visual pigment spectra are described by template formulas (Stavenga et al., 1993). Contribution of metarhodopsin was neglected here, because the photoreceptors were nearly dark-adapted in the present experimental condition. The photoreceptors R3–R9 are polarization sensitive, with a polarization...
Anatomy of the compound eye of *P. xuthus*; the dorsal side corresponds to the top edge of the micrographs. 

(a) Diagram of three types of ommatidia (from left to right, types I–III). Each ommatidium contains nine photoreceptors, R1–R9 (1–9). R1–R4 are distal photoreceptors, which form the distal two-thirds of the rhabdom; R5–R8 are proximal photoreceptors, forming the proximal one-third of the rhabdom; and R9 is a basal cell. R1 and R2 contain distally purple pigment granules (distal pigment). Types I and II have red pigmentation around the rhabdom, whereas type III has yellow pigmentation (perirhabdomeral pigment).

(b) Transverse, unstained section through the proximal tier of the retina, showing red (black arrowhead; type I or II ommatidium) or yellow (white arrowhead; type III ommatidium) pigmentation around the rhabdom. c, UV-induced fluorescence caused by 3-hydroxyretinol marks the type II ommatidia (from eye in intact, living animal).

(d) In situ hybridization of PxL2 (green-absorbing visual pigment) (d) and PxL3 (orange-red-absorbing visual pigment) (e) mRNA in adjacent sections through the proximal tier. R5–R8 are exclusively labeled with either the PxL2 probes (solid circles; type III) or the PxL3 probe (dotted circles; type I). R5–R8 of type II ommatidia are labeled by both PxL2 and PxL3 probes (dashed circles). The table is a summary of the characteristics of the three ommatidial types. In addition to the perirhabdomeral pigmentation and the autofluorescence, it presents the spectral sensitivity of the photoreceptors.

<table>
<thead>
<tr>
<th>Type of ommatidia</th>
<th>Type I</th>
<th>Type II</th>
<th>Type III</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pigmentation</td>
<td>red</td>
<td>red</td>
<td>yellow</td>
</tr>
<tr>
<td>Fluorescence</td>
<td>open</td>
<td>open</td>
<td>No</td>
</tr>
<tr>
<td>Photoreceptors</td>
<td>[S(j)]</td>
<td>[S(j)]</td>
<td>[S(j)]</td>
</tr>
<tr>
<td>R1</td>
<td>UV*</td>
<td>V</td>
<td>B</td>
</tr>
<tr>
<td>R2</td>
<td>B*</td>
<td>Y</td>
<td>PxB</td>
</tr>
<tr>
<td>R5, R8</td>
<td>PxL2</td>
<td>PxB</td>
<td>PxB</td>
</tr>
<tr>
<td>R9</td>
<td>PxL3</td>
<td></td>
<td>PxB</td>
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</table>

The rhabdomere diameter is 2.6 μm. The first waveguide mode was assumed to be the dominant component of the light flux propagated in the rhabdom, because the wavelength dependence of the second-order mode is very similar to that of the first mode, and the relative contributions of the modes to the spectral sensitivity have not yet been established. The fraction of light within the rhabdom ([η](A)) was calculated using the waveguide optics formulas for insect photoreceptors (Snyder et al., 1973; Stavenga, 2003).

A relative amount of PxL2 and PxL3 of 3:1 was chosen, because this ratio gave satisfactory outcomes in the computer simulations. We have noticed that the *in situ* hybridization results did not necessarily match this choice: the labeling density of PxL2 is comparable with or maybe even weaker than that of PxL3 (Fig. 1d,e). However, taking equal concentrations of PxL2 and PxL3 yielded a much too high sensitivity in the long-wavelength range in the simulations. This may imply that the labeling density of *in situ* hybridization does not provide a direct measure of the expressed visual pigment concentration in photoreceptors. Another possible cause of the discrepancy is the limited accuracy of the predicted absorption peak wavelength of the PxL2 and PxL3 visual pigments. We predicted absorption peaks of 515 nm for PxL2 and 575 nm for PxL3 only by model calculation (Arikawa et al., 1999a), not by direct measurements. Direct proof of the absorption spectra requires expression experiments (Feiler et al., 1992; Shaaban et al., 1998; Townson et al., 1998; Jacobs et al., 1999).

**Results**

The proximal photoreceptors of the fluorescent type II ommatidia express a broadband spectral sensitivity

In the course of previous studies on the photoreceptors of *P. xuthus*, we sometimes encountered units with abnormally broad sensitivity spectra, but we could not unequivocally prove that the recordings were from single photoreceptors. We therefore designed the experimental procedure described below.

With the electrode advancing in the retina, we searched for units with broad sensitivity spectra. Broadband units were detected only when the electrode track ran through the proximal tier of the ventral retina. Figure 2a shows a typical recording...
trace, which contains two series of spectral responses (300–700 nm; with equiquantal monochromatic flashes) and \( V = \log I \) functions (recorded at 560 and 440 nm in this case). Figure 2b is a set of \( V = \log I \) functions recorded from a single broadband unit at 420, 440, 540, and 620 nm, with the best fits of the Naka–Rushton function. The maximum response amplitude, \( V_{\text{max}} \), and the exponent, \( n \), are virtually identical for the different recording wavelengths (for values, see Fig. 2 legend). The sensitivity spectrum presented in Figure 2c shows the average of 10 cells from nine butterflies.

We identified and localized all of the recorded units histologically, as follows, to determine whether the units correspond to single photoreceptors. After the electrophysiological recordings, we delivered Lucifer yellow from the electrode tip by applying a hyperpolarizing DC. The minor amount of Lucifer yellow that may have leaked into the recorded photoreceptor during the spectral measurements before applying the current did not perturb the sensitivity spectra, because the spectra of known spectral receptors encountered in the course of the present study were identical to those of our previous recordings with 3 m KCl-filled electrodes (Arikawa et al., 1987; Bandai et al., 1992).

The ommatidium containing the Lucifer yellow-injected unit was subsequently localized by fluorescence microscopy, applying violet epi-illumination (Fig. 3a). By switching the excitation light to UV, we found that the ommatidium emitted whitish fluorescence, proving that the unit was a member of a type II ommatidium (Fig. 3b). The eyes were then further processed for histology to identify the receptor type. In the particular case of Figure 3c, showing a transverse section through the proximal tier of the ommatidia, the unit appeared to be a single R8 photoreceptor. Regular transmission microscopy of the same frame revealed that the Lucifer yellow-injected photoreceptor contained red pigment (Fig. 3d), in agreement with the previous observation that type II ommatidia are red pigmented (Arikawa and Stavenga, 1997).

Identical observations were made for all of the broadband-sensitive units analyzed in the present study (i.e., the broadband units were invariably identified as belonging to the set of proximal photoreceptors, R5–R8, of the fluorescent type II ommatidia).

**Modeling of the broadband sensitivity spectrum**

In modeling the broadband sensitivity spectra, we have calculated the fraction of light entering the rhabdom of type II ommatidia that is absorbed by the visual pigments as well as by the screening pigments to be in the wavelength range of 300–700 nm. We have assumed that both Pxl2 and Pxl3 visual pigments in R5–R8 photoreceptors contribute to the spectral sensitivity in an identical manner (i.e., each absorbed photon by one or the other visual pigment contributes equally to phototransduction or opens the same number of channels). Figure 2c shows the results of the model calculations of the normalized absorption spectra of the R5–R8 receptors, together with the average of the broadband sensitivity spectra. We calculated the spectra for incident light with different polarization angles (i.e., 0, 90, 35, and 125°, respectively). The calculated and measured spectra closely match at the short and long wavelengths, but the spectra vary in the wavelength range of 480–580 nm. The variation can be attributed primarily to the polarization sensitivity of the distal photoreceptors, R1–R4, which acts as a polarization-dependent absorption filter for the proximal photoreceptors, R5–R8. For example, the R3 and R4 cells are single-peaked green receptors, peaking at 520 nm with maximum polarization sensitivity at 90°. When the ommatidium is stimulated with 90° polarized light, the light is strongly absorbed by R3 and R4, thus reducing the light reaching the proximal tier and lowering the response of proximal photoreceptors in the middle wavelength region.

Figure 4 presents the calculated and recorded spectra of the proximal as well as distal photoreceptors in type II ommatidia. The R1 and R2 photoreceptors, which contain PxUV, a visual
pigment absorbing maximally at 360 nm, become violet receptors, with sensitivity spectrum peaking at 400 nm, because of the abundant 3-hydroxyretinol concentrated in the distal part, which acts as a UV filter for the underlying photoreceptors (Arikawa et al., 1999b). The R3 and R4 photoreceptors, which contain PxL1 and PxL2, green visual pigments absorbing maximally at 530 and 515 nm, respectively, become single-peaked green receptors, because they lose the usual β-band as a result of the UV filter, confirming our previous study on the type II ommatidium (Arikawa et al., 1999b). Similarly, the R5–R8 have negligible sensitivity in the ultraviolet because of the strongly UV-absorbing 3-hydroxyretinol. The R5–R8 contain PxL2 and PxL3, visual pigments absorbing maximally at 515 and 575 nm, respectively, which are together responsible for the broadband spectral sensitivity, assuming that both visual pigments trigger phototransduction.

One might argue that the fluorescence of 3-hydroxyretinol could excite the photoreceptors in type II ommatidia. However, the recordings clearly indicate that this is not the case: the photoreceptors in type II ommatidia have virtually no sensitivity in the UV wavelength region (Fig. 4) in which the absorption of 3-hydroxyretinol exists. Although the fluorescence seems very prominent in micrographs (Fig. 1c), it requires extremely bright illumination provided by a focused mercury lamp of an epifluorescence microscope. Because of the low quantum yield, fluorescence will therefore have a negligible contribution to photoreceptor excitation.

Discussion

In this study, we describe a novel type of photoreceptor featured in the *Papilio* retina with an abnormally broad sensitivity spectrum, accordingly termed broadband receptor. Anatomically, the broadband-sensitive cells are the proximal photoreceptors (R5–R8) in the type II ommatidia [i.e., the red-pigmented ommatidia that fluoresce under UV excitation (Figs. 1, 3)]. The photoreceptors were shown to express two mRNAs, encoding the visual pigment opsins PxL2 and PxL3 (Fig. 1de). The absorption characteristics of the corresponding visual pigments are spectrally quite different: the absorption peak wavelengths are in the green (515 nm) and orange-red (575 nm) regions, respectively (Arikawa et al., 1999a).

In the modeling, we incorporated the light-filtering effects of a number of optical absorbers, namely 3-hydroxyretinol in the distal part of the type II ommatidia, the red-screening pigment surrounding the rhabdom, and the visual pigments in the distal photoreceptors R1–R4, of which the visual pigments in the R3 and R4 photoreceptors filter specifically in the green wavelength region. The calculated absorption spectra reasonably fit to the broadband sensitivity spectra in the short- and long-wavelength regions. However, the modeling does not yet satisfactorily reproduce the bumpy profile of the sensitivity spectrum in the wavelength region between 480 and 580 nm (Fig. 2c). As described in Materials and Methods, we calculated the spectrum by taking only the first-order waveguide mode into account. The second mode presumably contributes to the bumpy profile of the spectrum (Stavenga, 2003). To incorporate higher-order modes and additional details of spectral filtering, more detailed anatomy of the rhabdom geometry and fine structure will be necessary. Of course, some of the discrepancy of the model calculations and the recorded spectra may be caused by experimental errors. For instance, the alignment of the polarization angle of the stimulus light and the ommatidial dorsoventral axis is uncertain because of the slight distortion in the ommatidial lattice arrangement [Arikawa and Uchiyama (1996), their Fig. 3c]. Nevertheless, the general correspondence of the measured broadband sensitivity spectrum with the computational results strongly suggests that both visual pigments contribute to phototransduction.

To the best of our knowledge, the broadband receptor of *P. xuthus* is the first clear example of a class of invertebrate photoreceptors with multiple visual pigments functioning simultaneously. It extends the as-yet-limited list of similar vertebrate photoreceptors (Makino and Dodd, 1996; Lyubarsky et al., 1999). The fact that the *V* max value and *n* value remain virtually constant between different stimulus wavelengths suggests that
the response kinetics is wavelength independent (Fig. 2a,b). Therefore, it is likely that the two distinct visual pigments trigger the same transduction pathway. Of course, it is still possible that there are two independent transduction pathways, triggered by each visual pigment, whose kinetics is very similar.

The biological function of the broadband receptor for butterfly vision must remain a matter of speculation. It seems unlikely that it plays a role in color vision, a function that is logically inferred to the R5–R8 photoreceptors of the type I ommatidia: they have a narrow-band sensitivity spectrum, distinctly peaking in the red (Arikawa and Uchiyama, 1996; Arikawa et al., 1999a). Also, the R5–R8 photoreceptors of the type III ommatidia may well participate in color vision, because they have a clear spectrum peaking in the green. The broadband R5–R8 photoreceptors of type II ommatidia rather could play a role as general light intensity (or luminosity) detectors. Another possible function of broadband photoreceptors is related to the function of type II ommatidia themselves. The photoreceptors in type II ommatidia are virtually insensitive to UV because of the filtering effect of 3-hydroxyretinol [(Arikawa et al., 1999b), their Fig. 5]. Together with type I ommatidia, containing UV receptors (Fig. 1, table), the UV-insensitive type II ommatidia probably participate to enhance UV contrasts in the visual scene. Detecting UV contrasts is particularly important for flower visitors, because many flowers provide information on nectar localization by so-called nectar guides, which involve UV reflection/absorption (Heiling et al., 2003). The broadband photoreceptors are sensitive over the entire spectral range of butterfly vision, except in the UV range. Slight changes in UV intensity can therefore be efficiently analyzed by a combination of the broadband and UV receptors.

Butterfly eyes have a strikingly heterogeneous organization, with specific sets of spectral detectors combined in different types of ommatidia. Although the overall lattice of ommatidia is of crystalline regularity, the different types are randomly arranged within that regular lattice (Arikawa and Stavenga, 1997). Our working hypothesis is that the regular ommatidial lattice is connected to spatial vision and that the heterogeneous organization is related to color processing. Additional study is required to resolve in which way and to what extent the signals of the different photoreceptors feed into either of both visual modes.

References


