Molecular Basis for Ultraviolet Vision in Invertebrates

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Invertebrates are sensitive to a broad spectrum of light that ranges from UV to red. Color sensitivity in the UV plays an important role in foraging, navigation, and mate selection in both flying and terrestrial invertebrate animals. Here, we show that a single amino acid polymorphism is responsible for invertebrate UV vision. This residue (UV: lysine vs blue: asparagine or glutamate) corresponds to amino acid position glycine 90 (G90) in bovine rhodopsin, a site affected in autosomal dominant human congenital night blindness. Introduction of the positively charged lysine in invertebrates is likely to deprotonate the Schiff base chromophore and produce a UV visual pigment. This same position is responsible for regulating UV versus blue sensitivity in several bird species, suggesting that UV vision has arisen independently in invertebrate and vertebrate lineages by a similar molecular mechanism.

Key words: Drosophila; ERG (electroretinogram); photoreceptor; retina; vision; rhodopsin

Introduction

Color vision is one of the most familiar forms of stimulus discrimination and is dependent on an organism having at least two different classes of photoreceptor cells of different spectral sensitivities (SS) (Jacobs, 1981). Except in eyes using screening pigments or oil droplets, this is based on the expression of spectrally distinct forms of the visual pigment rhodopsin, in different photoreceptor cells. Invertebrates are sensitive to a broad spectrum of light that ranges from UV to red in some species (Gärtnér, 2000; Briscoe and Chittka, 2001). UV vision is used in foraging, navigation, and mate selection in both flying and terrestrial invertebrate animals (Tovee, 1995). UV vision is mediated by a form of rhodopsin, the visual pigment of both vertebrate and invertebrate organisms. Rhodopsin is composed of an 11-cis-retinal chromophore that is attached covalently through a Schiff base linkage to a lysine residue in the opsin apoprotein (Fig. 1a). The absorption spectrum of the visual pigment determines the SS of the photoreceptor in which it is expressed, and this in turn is dependent on specific interactions between the opsin apoprotein and the retinal chromophore. Drosophila melanogaster uses six visual pigments that range in absorption from UV to green (Fig. 1b) (Salcedo et al., 1999). These diverse pigments provide an opportunity to examine the basis for spectral tuning in insects and other invertebrates. Furthermore, transgenic flies can be used as an expression system to examine the function of novel or modified visual pigments in vivo (Feiler et al., 1988, 1992; Britt et al., 1993; Townson et al., 1998; Salcedo et al., 1999).

Previous studies of bovine rhodopsin have shown that a chromophore/apoprotein interaction occurs between the protonated Schiff base nitrogen and a negatively charged counter-ion at position glutamate 113 (E113) (Fig. 1a) (Sakmar et al., 1989; Zhukovsky and Oprian, 1989; Nathans, 1990). This interaction stabilizes the protonated Schiff base and is responsible for the opsin red shift that converts both UV absorbing chromophore and apoprotein into a visible absorbing complex. Removal of the negative charge of the counter-ion results in deprotonation of the chromophore and yields a UV-absorbing pigment. In invertebrate visual pigments, the position homologous to bovine rhodopsin E113 is occupied by a tyrosine (Y) or phenylalanine (F) residue in the visible and UV-absorbing invertebrate pigments, respectively (Fig. 2). This site has been proposed as the invertebrate counter-ion, and the F/Y polymorphism may be responsible for the difference in absorption between the visible and UV pigments of invertebrates.

In the present study, we performed a series of experiments to test the hypothesis that the F/Y polymorphism is responsible for the difference in absorption between UV and visible absorbing pigments. We found that this polymorphism is not responsible for UV sensitivity and then conducted a comparative phylogenetic analysis of a group of closely related visible and UV-absorbing invertebrate rhodopsins in an effort to identify additional candidate residues. We show that one of these polymorphisms is responsible for UV vision in invertebrates and present both a model of the underlying mechanism of wavelength regulation and a test of the model that indicates the effect of the polymorphism is to deprotonate the Schiff base chromophore and produce a UV visual pigment.

Materials and Methods

Ectopic expression of native and modified Drosophila opsin genes. Flies expressing the Rh3 and Rh5 opsin genes in the R1–6 photoreceptor cells of ninaE mutants have been described previously (Feiler et al., 1992; Chou et al., 1996; Salcedo et al., 1999). Flies expressing modified forms of Rh1, Rh3, and Rh5 were generated in a similar manner. Briefly, the gene encoding the opsin of interest was modified by site-directed mutagenesis using inverse PCR with Pfu DNA polymerase and DpnI digestion of the methylated template (Weiner et al., 1994; Fisher and Pci, 1997). The sequence of the mutagenized fragment was confirmed, and the fragment
was then subcloned into an expression cassette containing the Rh1 promoter (2.4 kb including 33 bp of the 5′ untranslated region) and the remainder of the opsin gene of interest. The transcriptional fusion encoding modified Rh3, Rh5, or the modified form of Rh1 was subcloned into the y′-marked P-element vector “C4” obtained from Pam Geyer (University of Iowa, Iowa City, IA) (Patton et al., 1992). The construct was injected into yw; sr ninaE17 mutant embryos, and multiple independent P-element-mediated germ line transformants were obtained (Kress and Rubin, 1984).

**Electrophysiology.** Electrotetrogram (ERG) recordings were obtained from immobilized white-eyed (w) flies using glass microelectrodes filled with normal saline (0.9% NaCl, w/v) (Salcedo et al., 1999). Flies were stimulated with light from a xenon arc lamp (350 W; Osram, Oriel, Stratford, CT), using interference and neutral density filters. Light intensity was measured using a calibrated silicon photodiode (model 71883; Oriel) and an optical power meter (model 70310; Oriel).

SS was measured using a modification of the voltage-clamp method of Franceschini (1979, 1984), which we have described in detail previously (Townson et al., 1998; Salcedo et al., 1999). Briefly, the amplitude of the ERG response to a flickering (10 Hz) monochromatic stimulus was maintained at a criterion level by adjusting the light intensity, whereas the wavelength of stimulating light was varied during a scan. In the measurements in this study, for each ~0.3 sec window (ERG response to approximately three flickers), we averaged all of the ERG voltages during this period and used the average deviation as an estimate of the amplitude (width) of the response (Press et al., 1992). The average deviation is a function of the variance between the individual data points and the mean voltage during the sinusoidal ERG response. Thus, the average deviation is related to the response amplitude, although much less sensitive to baseline drift and noise.

During an experiment, as the monochrometer was stepped through a scan in 0.1 nm increments, the average deviation of the ERG response was calculated and compared with a criterion set point. The ERG response was maintained near the set point during the scan by constantly adjusting the light intensity using a proportional–integral–derivative algorithm (Corripio, 1990). SS was defined as the inverse of the light flux required to produce the criterion response, taking into account the wavelength and intensity of the stimulating light [i.e., SS \(= 1/(\text{light intensity} \times \text{wavelength})\)]. Raw sensitivity data were normalized to an amplitude of 1.0 at the wavelength of maximum sensitivity, multiple individual measurements were averaged, and values that differed by >10% from the mean value within a 10 nm window were filtered out. The filtered spectra were then smoothed with a window of 10 nm.

Experiments were performed on a minimum of four individual flies, two each from two different transgenic lines for each construct. We typically record from many more animals and lines as well, although these may not all have been included in the calculations. Occasional discrepancies occur between recordings that we have attributed to individual differences (health, age, light exposure, etc.). We have never observed any instances in which one transgenic line with a particular construct was consistently different from another. The sampling interval of the instrument was ~0.2 nm, taking into account the scan rate of the monochrometer (0.5 nm/sec) and the sampling rate for determining the response amplitude (three per second). The monochrometer was calibrated using an Oriel 6025 mercury (argon) calibration lamp with spectral lines at 126.2, 365.0, 404.7, 546.1, 577.0, and 579.1 nm. Monochromator error was within the level of resolution of the instrument; over the wavelength range from 300 to 600 nm (±1.0 nm at a slit width of 280 μm). Curve fitting of spectral sensitivity recordings to rhodopsin nomograms (see below) was within ±1.0 nm. Variation between individual measurements was typically within 1–2 nm.

**Results.**

**ERG and SS recordings were performed on transgenic animals expressing native or modified forms of rhodopsin in either a ninaE17 background or in a modified norpA[ninaE17] mutant background. The latter strain also contained an additional transgene driving the norpA CDNA in the R1–6 photoreceptor cells under the control of the Rh1 promoter. This background strain allows the activity of the modified pigment to be examined without interference from the R7 and R8 cells that are not affected by the ninaE mutation (Salcedo et al., 1999).**

Figure 1. Chromophore structure, rhodopsin absorption, and spectral sensitivity. a, 11-cis-Retinal chromophore attached to lysine 296 as the Schiff base with the counter-ion (glutamate 113) and glycine 90 are shown, from the bovine rhodopsin crystal structure (Palićzewski et al., 2000). Gray, Carbon; blue, nitrogen; red, oxygen. b, Calculated absorption profiles of Drosophila rhodopsins Rh1 (\(\lambda_{\text{max}} = 480 \) nm), Rh2 (\(\lambda_{\text{max}} = 430 \) nm) (Feiler et al., 1988), Rh3 (\(\lambda_{\text{max}} = 347 \) nm), Rh4 (\(\lambda_{\text{max}} = 375 \) nm) (Feiler et al., 1992), Rh5 (\(\lambda_{\text{max}} = 436 \) nm), and Rh6 (\(\lambda_{\text{max}} = 508 \) nm) (Salcedo et al., 1999). Rh1, Rh3, and Rh5 are shown in red. c, d, Measured SS of flies expressing Rh1 (c) and Rh3 (d) in the R1–6 photoreceptor cells. A curve fit of the rhodopsin nomogram is shown in red. Flies expressing Rh1 (c) in the R1–6 photoreceptor cells display a dual peak of sensitivity. The peak in the UV region occurs because of the action of a sensitizing pigment that absorbs in the UV and activates the Rh1 rhodopsin through energy transfer (Kirschfeld et al., 1977). The peak in the visible region (with red curve fit) is attributable to direct absorption by Rh1. e, f, SS of flies expressing rhodopsin mutants (c, Rh1Y126F; f, Rh3 F313Y) compared with flies expressing the unmodified pigments (red traces). Each sensitivity spectrum obtained in this study was fit to a rhodopsin absorption nomogram. The \(\lambda_{\text{max}}\) correlation coefficient, and number of flies examined are shown in Table 1.
Rhodopsin nomogram modeling. Rhodopsin absorption spectra were calculated from the SS recordings using the exponential function described by Stavenga et al. (1993). Briefly, the spectral shape of the rhodopsin α-band absorption can be described by the following lognormal function:

$$\alpha = A \exp\{-(a_0 x^2 + a_1 x + a_2 x^3)\},$$

where $x = 10\log(\lambda/\lambda_{max})$, $A = 1$, $a_0 = 380$, $a_1 = 6.09$ and $a_2 = 3a_1^2/8$.

A curve-fitting routine was implemented in KaleidaGraph (version 3.51; Synergy Software, Reading, PA) using the Levenberg–Marquardt (nonlinear least-squares) algorithm (Press et al., 1992). The computer solved for the $\lambda_{max}$ and amplitude of the rhodopsin absorption spectra and calculated the SD for each variable and the correlation coefficient (Pearson’s $r$). Each sensitivity spectrum obtained in this study was fit to a rhodopsin absorption nomogram. The $\lambda_{max}$ correlation coefficient, and number of flies examined are indicated in Table 1.

**Results**

In our first series of experiments, we tested the hypothesis that the F/Y polymorphism between UV and visible absorbing pigments, which occurs at the same position as the vertebrate counter-ion (E113), is responsible for UV vision. We constructed mutant forms of a blue-absorbing (Rh1) and UV-absorbing (Rh3) *Drosophila* rhodopsin in which the residues at this position were replaced with that found in the other class of pigments (i.e., Rh1 Y126F and Rh3 F133Y). To determine the effect of the amino acid change on the absorption of the visual pigment, the genes encoding these modified pigments were introduced into the germ line of transgenic *Drosophila* containing the ninaE mutation, which is a deletion in the endogenous Rh1 gene that is expressed in the R1–6 photoreceptor cells (O’Tousa et al., 1985; Zucker et al., 1985). We found that the SS of transgenic animals expressing either Rh1 Y126F or Rh3 F133Y showed very little change from that of the unmodified pigments (Fig. 1c–f). These results indicate that the F/Y polymorphism at the position of bovine E113 is not responsible for the difference in SS between the invertebrate UV and visible absorbing pigments and suggests that the invertebrate counter-ion is located elsewhere. These results are also consistent with the finding that the invertebrate tyrosine is not protonated Schiff base (Hashimoto et al., 1996; Nakagawa et al., 1999).

To identify additional candidate residues responsible for UV versus visible sensitivity, we undertook a phylogenetic analysis of a group of well-characterized invertebrate visual pigments. Figure 2 shows that one of UV-absorbing visual pigments (e.g., Rh5) is closely related to, and seem to share a common ancestor with the UV-absorbing pigments of invertebrates (e.g., Rh3) (Salcedo et al., 1999). This suggests that during evolution one or both of these groups acquired amino acid changes that are responsible for the differences in their spectral properties. Figure 2 shows a group of polymorphic amino acids that are conserved within each group (UV or visible absorbing) but differ between them and might be expected to affect chromophore absorption based on side chain position and differences in polarity or charge. Nine modified forms of the gene encoding Rh5 (blue absorbing) containing these 12 amino acid changes, as single or double amino acid substitutions, were introduced into ninaE mutant flies. Electrophysiological analysis of the animals expressing the transgenes showed little or no change in SS for virtually all of the Rh5 mutants. One modified pigment, Rh5 N104K, showed a dramatic shift in SS (from $\lambda_{max} = 346–377$ nm) (Fig. 3a). To confirm that the lysine (K) versus asparagine (N) polymorphism is responsible for a large component of the difference between UV and visible absorbing pigments in invertebrates, we replaced the existing lysine residue in Rh3 with the corresponding amino acid found in Rh5. Transgenic flies expressing the modified Rh3 K110N opsin showed a complementary shift in SS (from $\lambda_{max} = 347–420$ nm) (Fig. 3b). This confirms that the K versus N polymorphism is responsible for UV sensitivity.

The blue-absorbing pigments have either an asparagine (in the case of Rh5) or a glutamate (E) (in the case of the Bee blue and other members of this class) at this position (Fig. 2). To test whether the asparagine (neutral) versus glutamate (negatively charged) polymorphism may also affect spectral tuning, we generated the Rh3 K110E mutant and found that its SS is identical to Rh3 K110N (Fig. 3b). This indicates that replacement of the lysine with a neutral (asparagine) or negatively charged (glutamate) residue has the same effect, suggesting it is the removal of the positively charged lysine that converts Rh3 to a visible absorbing pigment. Because the mutant pigments are not fully shifted to the $\lambda_{max}$ of the related pigment, these experiments also demonstrate that additional amino acid polymorphisms must contribute to differences in SS. This is expected given that the *Drosophila* UV-absorbing and visible absorbing pigments differ considerably among themselves (e.g., the $\lambda_{max}$ of UV pigments Rh3 and Rh4 differ by 28 nm) (Fig. 1b).

The position within bovine rhodopsin corresponding to this site is glycine 90 (G90), which has been associated with the human congenital night blindness mutation G90D (Rao et al., 1994). In addition, a cysteine-serine (C/S) polymorphism at this site is responsible for ~35 nm difference in absorption between the UV and blue-absorbing visual pigments of birds, although the precise molecular mechanism underlying this effect is not known (Wilkie et al., 2000; Yokoyama et al., 2000). An analysis of struc-
tural models of these pigments based on the bovine rhodopsin crystal structure (Palczewski et al., 2000) has suggested that, in the UV pigments, the thiol group of C90 is positioned very close to E113 and may exist as a thiolate ion (Hunt et al., 2001). The authors propose that this may serve to further stabilize the proton on the Schiff base in the ground state, with a resulting shift of absorption into the UV. Alternatively, it has also been suggested that the cysteine residue may remove a water molecule from the region near the Schiff base and displace its positive charge. Deprotonation would be expected to produce a large shift into the UV region as well (Yokoyama et al., 2000).

Our finding that the introduction of a lysine residue at this position is sufficient to shift the absorption spectrum into the UV can be more easily interpreted. Comparative modeling of the helices of the Rh5 and Rh3 visual pigments reveals a similar geometry for the position of the N/K polymorphism with respect to the Schiff base nitrogen, as that of G90 in bovine rhodopsin (data not shown). Although the identity of the invertebrate counter-ion is not known, insertion of a positively charged lysine into the region of the positively charged Schiff base and negatively charged counter-ion would be expected to disrupt the interaction between them and lead to the deprotonation of the Schiff base. Although the phenylalanine/tyrosine at the position of the vertebrate counter-ion (E113) is apparently not the invertebrate counter-ion, this residue is likely to be in close proximity to the Schiff base in the pigments. If this is the case, one would predict that introduction of an additional negative charge at this site would block the effect of the lysine (at G90) in the UV pigment.

To test this hypothesis, we substituted a glutamic acid at the position of E113 in bovine rhodopsin into both Rh1 (in place of the native tyrosine Rh1 Y126E) as well as into Rh3 (in place of the native phenylalanine Rh3 F133E). Whereas introduction of this additional negative charge in Rh1 had only a small affect (Fig. 4a), its introduction into the UV pigment Rh3 (Fig. 4b) caused signifi-
icant broadening in the sensitivity spectrum. This change was the best fit by absorption nomograms corresponding to a mixture of two (UV and blue absorbing) pigments (Fig. 4c,d). An analysis of ERG responses to flashes at varying wavelengths and intensities also confirmed that the Rh3 F133E mutant has a significant increase in sensitivity in the visible region (430–470 nm) (Fig. 4e,f). Thus, the introduction of an additional negative charge was sufficient to shift the equilibrium in the protonation of the Schiff base and produce a mixture of visual pigment states. Substitution with the glutamate residue may, thus, serve as an intramolecular titration that partially stabilizes the protonated Schiff base. Alternatively, the mutant pigment may adopt two different conformations in which the Schiff base is protonated in one and not in the other. In either case, these findings are consistent with the model that UV absorption in the invertebrate pigments is based on the deprotonation of the Schiff base by the introduction of a positively charged lysine residue. We believe a similar deprotonation event is responsible for the absorption of the avian UV pigments.

Discussion

The principal result from this study is that the K (versus N or E) polymorphism present at the position of G90 in bovine rhodopsin is the basis for UV vision in invertebrates. The introduction of the lysine residue at this position seems to deprotonate the Schiff base and shift the absorption of the pigment into the UV. This deprotonation seems to change during photoactivation, in which photon absorption by rhodopsin leads to the isomerization of the 11-cis-retinal chromophore to the all-trans configuration and induces a conformational change in the apoprotein, leading to the formation of the activated form of the visual pigment metarhodopsin. Photoactivation of the Drosophila Rh3 and Rh4 UV-absorbing rhodopsins leads to the formation of metarhodopsin species that absorb in the blue region (Feiler et al., 1992; Salcedo et al., 1999). This likely results from the protonation of the Schiff base in the metarhodopsin form of the pigment (Pande et al., 1987) as is thought to be the case for the mouse UV pigment (Dukkipati et al., 2002). This would require a conformational change that either removes the lysine (at G90), introduces an additional negative charge into the region near the chromophore, or uses an alternate counter-ion site in metarhodopsin, as is thought to occur in the vertebrate visual pigments (Yan et al., 2003).

The G90 amino acid position has been the subject of extensive study in the vertebrate visual pigments. A C/S polymorphism at this position is responsible for UV vision in birds (Wilkie et al., 2000; Yokoyama et al., 2000), and amino acid substitutions at this site have also been shown to alter the absorption maxima of other visual pigments (Lin et al., 1998; Dukkipati et al., 2001; Janz and Farrens, 2001). Furthermore, this position is associated with the human congenital night blindness mutation G90D (Rao et al., 1994). The introduction of an aspartic acid residue has been shown to alter the absorption maximum of the pigment and to generate a constitutively activated opsin apoprotein (Rao et al., 1994). Consistent with this observation, expression of the mutant visual pigment in vivo reduces light sensitivity in a manner similar to light adaptation that can be suppressed with the addition of visual pigment (Jin et al., 2003). Thus, the introduction of an additional negative charge was sufficient to shift the equilibrium in the protonation of the Schiff base and produce a mixture of visual pigment states. Substitution with the glutamate residue may, thus, serve as an intramolecular titration that partially stabilizes the protonated Schiff base. Alternatively, the mutant pigment may adopt two different conformations in which the Schiff base is protonated in one and not in the other. In either case, these findings are consistent with the model that UV absorption in the invertebrate pigments is based on the deprotonation of the Schiff base by the introduction of a positively charged lysine residue. We believe a similar deprotonation event is responsible for the absorption of the avian UV pigments.

References


