Action Spectra and Chromatic Mechanisms of Cells in the Median Ocelli of Dragonflies

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ABSTRACT Spectral sensitivities were recorded intracellularly in median ocelli of Anax junius, Aeschna tuberculifera, and Libellula pulchella. All cells had peak sensitivities at 360 and 500 nm while UV-blue-green cells found only in Anax had a third peak sensitivity at 440 nm. Ratios of UV-to-green sensitivities varied from cell to cell in each ocellus, but no UV-only or green-only cells were recorded. Half of the cells tested had a reverse Purkinje shift: They were more sensitive in the green at low illuminations but more sensitive in the UV at high illuminations; their intensity-response curves at 370 and 520 nm crossed but became parallel for large responses. Wave-lengths 420 nm and shorter elicited a family of low intensity-response curves with one slope; wavelengths 440 nm and longer elicited a family of curves with another slope. Orange-adapting lights selectively adapted sensitivity in the green, but UV-adapting lights had little selective effect. Amounts of log-selective adaptation were proportional to log orange-adapting intensity. It is concluded that two spectral mechanisms can be recorded from each cell, possibly by coupling of UV and green cells or possibly because each cell contains two visual pigments. Selective chromatic adaptations may provide the ocellus with a kind of "automatic color control," while the reverse Purkinje shift could extend the ocellus' sensitivity to prevailing skylight.

INTRODUCTION
Many roles are ascribed to insect dorsal ocelli (cf. Goodman, 1970), but color vision has yet to be proven to be one of them. Electrophysiological studies, using the electroretinogram (ERG), have indicated potentialities for color vision: Goldsmith and Ruck (1958) observed that dorsal ocelli of honeybees had peak sensitivities in both the UV and the blue-green. However, Goldsmith (1960) found that the bees' compound eyes alone mediated a phototactic preference for UV vs. green light. Cockroach dorsal ocelli appeared to possess sensitivity only in the blue-green (Goldsmith and Ruck, 1958). More promising perhaps were recordings from the dorsal ocelli of the dragonfly Libellula luctuosa, where Ruck (1965) observed sensitivities in the visible and (to
an undetermined extent) in the UV, as well as differences in waveforms of ERG's at wavelengths above and below 410 nm. From these results he concluded that two (or more) cell types, sensitive to different parts of the spectrum, existed in dragonfly ocelli and could provide a basis for color vision.

In the present paper, we reassess this conclusion using intracellular recordings from single cells of the dragonfly median ocellus (Chappell and Dowling, 1972). Our results indicate not that there are populations of cells with sensitivities in different parts of the spectrum, but that all cells respond maximally in both the UV and the visible. In addition, from considerations of waveforms of responses, ratios of UV to visible sensitivities, intensity-response functions as functions of wavelength, and selective chromatic adaptations, we conclude that two (or three) chromatic mechanisms contribute to the spectral responses recorded from within single cells.

**MATERIALS AND METHODS**

**Preparation of the Animal**

Dragonflies of the species *Anax junius*, *Aeshna tuberculifera*, and *Libellula pulchella* were raised from nymphs in the laboratory (Chappell, 1970) and were used no sooner than 1 day after eclosion of the adults. Experiments on *Anax* and *Aeshna* were performed in early fall, while further experiments on *Aeshna* and one experiment on *Libellula* were made in late spring. An animal's head was cut off and a portion of the exoskeleton was removed from the front of the head, exposing the median ocellus. The head was then waxed to a cork such that the optical axis of the median ocellus was approximately horizontal. The whole head was next covered with a dragonfly Ringer's (Fielden and Hughes, 1962). Finally, a small portion of the pigmented capsule of the ocellus was torn away, using fine tweezers, and revealing the white, pigment-free interior. With this preparation, the microelectrode could approach the retinular cells of the ocellus perpendicular to their long axes and without having to penetrate the tough, ocellar capsule. All experiments were done at 20–22°C. Further details of the preparation are given in Chappell and Dowling, 1972.

**Intracellular Recording**

Successful micropipette electrodes were pulled from 1-mm OD Pyrex glass tubing containing soft glass fibers, were filled with 2 M KCl by injection, and had tip resistances of 25–200 MΩ in Ringer's. Cells were most easily penetrated by briefly applying excess negative-capacity compensation to the recording amplifier and allowing it to oscillate. The indifferent electrode was placed in the Ringer's bath and resting potentials were measured with respect to it using a calibrated bucking potential. Further details of the recording apparatus are given by DeVoe, 1972.

**Optical Stimulation**

The details of the automated monochromator system have been given previously (DeVoe et al., 1969; DeVoe, 1972). In essence, the system consists of a monochromator with a xenon arc light source and colored glass filters to reduce stray light; a mer-
cury arc lamp with a UV filter to isolate the 365-nm spectral line for UV adaptations; and a quartz-halogen lamp with a long-pass, orange-appearing filter to provide adaptation at visible wavelengths. All light sources were run from regulated power supplies. The UV and orange-adapting beams were combined by means of a dichroic beam splitter (Laser Energy Inc., Rochester, N.Y., type SWP-450). The adapting beams and the monochromator beam were combined with a quartz beam splitter of 45% reflection, 45% transmission (Esco Products, Oak Ridge, N.J.). All combined beams were then focused by a quartz objective lens (Esco) upon a 0.5 mm, near-UV conducting, glass fiber-optic (type ULGM, American Optical Corp., Southbridge, Mass.). The other end of this fiber optic bundle lay in the Ringer’s solution directly in front of the lens of the ocellus.

The automated monochromator system utilized paper tapes which were preprogrammed to provide flashes eliciting approximately equal amplitudes of response during scans of 18–20 wavelengths from 350 to 650 nm and in the reverse direction. In addition, the tapes provided a range of stimulus intensities at least one wavelength in the UV (370 nm) or in the visible (520 nm). In this way, nearly equal-amplitude responses could be obtained at all wavelengths, while the exact intensities that should have been used to elicit equal, criterion amplitudes of response (usually 4 mV) could be interpolated from the slopes of the intensity-response curves at 370 and/or 520 nm. When necessary, flash wavelengths and/or intensities could also be set by hand. Unless otherwise specified, flash durations were 100 ms, which were found to be long enough to elicit maximum amplitudes of response from most cells. However, for a few cells, which responded exceptionally slowly, flash durations had to be lengthened to 200 ms. With a timer, intervals between flashes were set in the range of 5–10 s, sufficiently long that we observed no progressive light adaptation.

Calibrations

The quantum fluxes, in quanta per second emerging from the end of the fiber optic bundle (in air), were measured using a PIN-10UV photodiode (United Detector Technology Inc., Santa Monica, Calif.) calibrated by the factory and operated in the photovoltaic mode. These are the fluxes we cite as quanta per second upon the eye, as we do not know the illuminations of the receptor cells within the eye. Flux densities, as quanta per second per square centimeter at the end of the fiber optic bundle, would be about 500 times greater than the numbers we give. Between experiments, the calibrated fiber optic bundle was stored in a tube to reduce breakage to its individual fibers. This necessitated refocusing of the light beams onto the fiber optic bundle at the beginning of each experiment. Initially, it was thought to be sufficient to align the fiber optic bundle each time so as to maximize output in the UV (at 370 nm), as had been done during the original calibrations. As it was more convenient, these subsequent realignments were made using a cadmium-sulphide photometer (Science and Mechanics, New York, N.Y.), rather than the calibrated photodiode. However, it was found later that this realignment method did not inevitably result in the same fluxes at all wavelengths as in the original calibrations. Rather, near the point of optimum focus of UV wavelengths upon the fiber optic, output in the visible was very sensitive to small changes in focus, output in the UV, hardly at all. Presumably, this was because we used a nonachromatic, quartz condensor. In any event, the effect was to in-
crease the quantum fluxes at visible wavelengths by up to 0.5 log units more than in the original calibrations. This may mean that the quantum fluxes at visible wavelengths given in some of the figures are too low, which is to say that sensitivities in the visible are too high, relative to the UV, by an unknown amount of up to 0.5 log units. In later experiments, the fiber optic bundle was realigned before each experiment using the calibrated photodiode, so that the UV intensity (at 370 nm) and the visible intensity (at 520 nm) were in the same ratio as during the calibrations. Intensities at other wavelengths were then found to be in the correct ratios also. As expected, average relative green sensitivities were then somewhat lower in these later experiments, but otherwise we obtained the same results.

RESULTS

Properties of the Cells.

Some impaled cells survived long enough after penetration for us to determine sensitivities at 18–20 wavelengths, usually increasing from 350 to 650 nm, but sometimes decreasing from 650 to 350 nm as well. These we call fully analyzed cells. Other cells survived impalement only long enough for us to measure their spectral sensitivities at two wavelengths, 370 and 520 (or 500) nm. These we call partially analyzed cells. Table I lists, for both kinds of cells, average minutes held, initial resting potentials recorded, and mean resting potentials averaged over the number of minutes held. Cells from Libellula had significa-

| TABLE I |
| SUMMARY OF DATA ON RESTING POTENTIALS AND SURVIVAL TIMES OF ALL IMPALED CELLS IN DRAGONFLY OCCELLI |

|                      | No. of cells | Average min held | Average initial resting potentials | Mean resting potentials averaged over min held |
|----------------------|--------------|------------------|-----------------------------------|---------------------------------------------|
| Anax and Aeschna     |              |                  |                                   |                                             |
| Fully analyzed cells | 32           | 25 (range 3–106) | −60 ± 15 (range 41–97)            | −55 ± 16 (range 31–95)                      |
| Partially analyzed cells | 38          | 4 (range 1–12)   | −58 ± 17 (range 33–102)           | −57 ± 17 (range 31–95)                      |
| Libellula            |              |                  |                                   |                                             |
| Partially analyzed cells | 8           | 22 (range 1–68)  | −41 ± 10 (range 20–52)            | −43 ± 17 (range 26–73)                      |

For technical reasons, no full analyses were made of cells from Libellula. Resting potentials were measured relative to the Ringer’s solution bathing the preparation, using a calibrated bucking potential. For Anax and Aeschna, these resting potentials are somewhat higher than previously measured (Chappell and Dowling, 1972), when a calibrated bucking potential was not available and resting potentials were measured as differences of potential between the tissue outside a cell and the potential in the cell just after penetration. Resting potentials of cells from Libellula were significantly lower (P < 0.01) than those from Anax and Aeschna, although survival times after impalement were equally good. There were no significant differences between initial and mean resting potentials.

Values are given as means ± SD.
significantly lower resting potentials than did cells from *Anax* and *Aeschna* (*P* < 0.01); they also had significantly different spectral properties (see Table II, and below).

Waveforms of intracellularly recorded responses from the ocellus to white light have been illustrated previously (Chappell and Dowling, 1972). Available information about waveforms of responses to colored lights comes only from studies on the ERG. On the one hand, Ruck (1961) observed identical responses from ocelli of "various species" to red light (from a Wratten no. 25 filter) and blue light (from a Wratten no. 47 filter). On the other hand, in *Libellula luctuosa*, he found differences between waveforms of responses of the ERG at wavelengths shorter and longer than 410 nm (Ruck, 1965). We have been able to duplicate both results with intracellular recordings, as illustrated in Fig. 1. Waveforms of intracellular responses from *Anax* (and also from *Aeschna*) to red and blue, broad-band lights are the same, as they are also for monochromatic lights in any part of the spectrum. Those waveforms of response illustrated at the left in Fig. 1 are typical, but we have at times in one and the same ocellus, but from different cells, recorded responses with slower rise and fall times and which lacked the initial spike. They too were wavelength independent. Conversely, as seen at the right in Fig. 1, intracellular responses from some (but not all) cells in an ocellus of *Libellula pulchella* were faster at 370 nm than at 500 nm, thus possibly accounting for the faster "on" transients observed by Ruck (1965) in response to UV light. The presence of such wave-

![Figure 1](image-url)

**Figure 1.** Waveforms of intracellular responses at different wavelengths. (Left) Responses from the median ocellus of *Anax junius*, with indicated Wratten filters (Eastman Kodak) of bandpasses shown, placed in the beam from a 45-W tungsten halogen lamp. The intensities of the 200-ms flashes (duration indicated by horizontal bar beneath the responses) were adjusted to give equal responses. From Chappell, 1970. (Right) Responses from the median ocellus of *Libellula pulchella* to 100-ms flashes (duration shown by horizontal bar) of 370- and 500-nm wavelength. See text for other details.
length-dependent differences in waveforms of intracellular responses does speak against Ruck's conclusion that such differences in the ERG, by themselves, indicate the existence of two, spectrally different types of cells in the ocellus.

**Spectral Sensitivities**

Our recordings (of spectral sensitivities) from within single cells distinguished two broad but overlapping classes of cells: UV-green and UV-blue+green. Within each class, there was large variability in relative short-to-long wavelength sensitivities. Some typical examples of relative spectral sensitivities of fully analyzed, UV-green cells from *Anax* and *Aeschna* are shown in Fig. 2.

![Figure 2](image_url)

**Figure 2.** Relative spectral sensitivities of dark-adapted, UV-green cells of *Anax junius* (top curve) and *Aeschna tuberculifera* (bottom two curves). (Ordinate) Log units of relative spectral sensitivity for 4-mV criterion responses; curves have been displaced vertically an arbitrary amount for clarity. (Abscissa) Wavelengths of 100-ms test flashes, in nanometers. From top to bottom: dashed lines fitted to points in the visible represent nomogram curves (Dartnall, 1953) at 513, 504, and 502 nm, respectively. The full extents of the nomogram curves are not drawn, since their flanks were generally broader at long wavelengths than the data. Solid lines through the remaining points were drawn by eye. In the top two curves, data points came from duplicate spectral scans on the two cells, first from 350 to 640 nm, and then from 540 to 350 nm; they show the reproducibility of the data. The points on the bottom curve came from one spectral scan, 350-640 nm. The log ratios of UV sensitivity to green sensitivity in the three curves are shown; a negative ratio indicates that the cell was more sensitive in the green. The log quantum fluxes on the eye for a 4-mV response at 520 nm were 7.41, 11.6, and 10.04 for the upper, middle, and lower curves, respectively.
One example from a fully analyzed, UV-blue + green cell, (found only in *Anax*) is shown in Fig. 3. Dashed lines representing nomogram curves (Dartnall, 1953) have been drawn in the visible part of the spectrum in both figures where they fit most closely. At long wavelengths where the nomogram curves are a bit too broad, solid lines through the data points have been drawn instead for clarity. All points were considered, however, in making the best fit by shifting the Dartnall nomogram to obtain the location of the peak. Locating the blue peak in this manner was clearly an approximation but gave more re-
producible results than trying to choose the peak by eye. Table II shows wave-
lengths of peak sensitivities, determined in the visible by means of the nomo-
gram curves and in the UV by inspection. It can be seen that all cells have the
same wavelengths of peak sensitivity in the UV and in the green, while the
UV-blue+green cells have a third, consistent peak of sensitivity at about 440
nm.

Where cells do differ from one another is in the ratios of UV-to-green (or
blue) sensitivities. The variability of the log ratios of UV-to-green sensitivity is
illustrated in Fig. 4 for all cells from Anax and Aeschna. Numerical values are
given in Table II. What is important here is that these log ratios do not ap-
pear to subdivide into three main groups, as might have occurred had each of
the three cells usually making up a retinula (Chappell and Dowling, 1972) had
its own characteristic spectral sensitivity. In addition, no cell had such a low
relative UV sensitivity or low relative green sensitivity that it could be con-
sidered to be essentially a green cell or a UV cell, respectively. Uncertainties
in light calibrations (see Methods) may have affected the absolute values of

| Wavelengths of peak sensitivities | Log sensitivity ratios UV/green | Log sensitivity ratios UV/blue | Log sensitivity ratios UV/green |
|-----------------------------------|--------------------------------|-----------------------------|--------------------------------|
| Fully analyzed | 0.36±0.33 Dark-adapted 360±7 | 498±2 | 358±6 |
| Partially analyzed | 0.44±0.41 UV-adapted 361±6 | 499±4 |
| UV-blue+green cells, Anax | 0.39±0.50 Dark-adapted 354±6 | 442±12 | 503±5 |
| Partially analyzed | UV/green | 0.57±0.46 | UV/blue | 0.36±0.33 |
| Libellula cells | UV/green | 1.49±0.88 |

Wave lengths of peak UV sensitivity read by eye to nearest 5 or 10 nm; wavelengths of peak blue and green sensitivities read from nomogram curves fitted to the data.

All values are given as means ± SD.

Log UV/green sensitivity ratios of UV-blue+green cells and Libellula cells are both significantly different from those of UV-visible cells (P < 0.05 and P < 0.01, respectively). Log UV/green sensitivity ratios of fully and partially analyzed UV-green cells are not significantly different from each other (P > 0.4). UV-green cells and UV-blue+green cells do not have significantly different wavelengths of peak sensitivity in the UV and green (P > 0.3).
some of the ratios in Fig. 4, but as variabilities of ratios were found within each experiment, this rules out calibration errors being the cause of such variabilities. Thus, the appearance of dual spectral sensitivities recorded from within single cells suggests that variable sums of two spectral mechanisms, each with a constant wavelength of maximum sensitivity are involved, and that in the green, at least, most likely being due to the absorption properties of a single visual photopigment (as determined by the fit of the nomogram curves). The possibility of a third spectral mechanism in the blue is indicated and requires further study.

**Intensity-Response Curves**

In most studies of spectral properties of single arthropod visual cells (cf. Wasserman, 1973), the finding has been that intensity-response curves recorded from within single cells have had shapes independent of wavelength, that is, they have been parallel to one another. For this reason, it is generally necessary to record an intensity-response curve at only one wavelength in order to convert amplitudes of responses at other wavelengths into equivalent sensitivities (see Methods). We initially assumed this would be true for cells of dragonfly ocelli as well and so accompanied each of our spectral scans of these cells with measurements of one intensity-response curve, either at 370 or 520 nm. However, in our early experiments, we did hold 12 of the 30, fully analyzed cells long enough to measure intensity-response curves in both the UV and the green. For 6 of these 12 cells, the two intensity-response curves were not parallel: The curve measured at 370 nm was steeper and crossed the curve measured at 520 nm. The insert in Fig. 5 illustrates a case where the curves so crossed. The cell from which these data came was more sensitive to UV light at higher intensities but was more sensitive to green light at lower intensities. Clearly, the shape of the spectral sensitivity curve of this cell is dependent on the criterion amplitude of response chosen, and so too is the ratio of its UV-to-green sensitivity. Possibly this might have been true for half of all our cells, so that the histogram in Fig. 4 might have been dependent on criterion amplitudes of response used for measuring spectral sensitivities.

From two cells, we were able to obtain intensity-response curves for intensities up to saturation. Fig. 5 (main part) shows the results from the one of these two cells whose curves would have crossed at low intensities. However, above 8 mV of response, the curves measured at wavelengths of 370 and 520 nm became parallel to each other. This is shown by the same solid line drawn through both sets of data. The other cell, stimulated over an equally large range of re-

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¹ During review it was pointed out to us that these data can be fitted by power functions, the lower part of the data at 500 nm having an exponent half that of the upper part and of the data at 370 nm. This relationship might be helpful in developing a model of the system although its meaning is unclear at present.
FIGURE 5. UV and green intensity-response curves over large ranges of intensities. At responses 8 mV and over, the intensity-response curve at 370 nm (filled circles) and at 520 nm (open circles) were parallel up to and including saturation at 33-mV response. The ordinate gives the millivolts of peak response to 100-ms flashes whose intensities are given on the abscissa in quanta per second upon the eye. The solid lines represent the same curve, which was fitted by eye to both sets of data where they were parallel and to the data at 370 nm alone where the responses were less than 7 mV. Below 7 mV, the intensity-response curves converge; however, we did not measure responses at amplitudes less than the crossing amplitude, estimated to be 2.6 mV for this UV-blue + green cell from *Anax*. (Inset) Crossing of UV and green intensity-response curves. In 6 of 12 cells for which intensity-response curves were measured in the UV (at 370 nm) and in the visible (at 520 nm), the curves crossed at an average amplitude of 3.3 mV (range: 1.8–4.4 mV). The solid and filled circles here represent one such cell's amplitudes of response (ordinate) at 370 and 520 nm, respectively, to 100-ms flashes whose intensities in quanta per second are given on the abscissa. The solid curves were drawn by eye. UV-green cell from *Aeshna.*

responses, had curves parallel to each other at both low and high intensities. From the results on these two cells, it appears that spectral sensitivities would have been criterion independent had we chosen criterion amplitudes of response of 8 mV or greater. In our experiments, we generally used 4-mV cri-
terion responses, both to avoid progressive light adaptations that brighter flashes might have caused and because during chromatic adaptations discussed below, 4 mV was often the maximum response we could elicit at all. This 4 mV criterion response is near the average of 3.3 mV of response at which intensity-response curves crossed, when they did. Thus the effect of our choice of criterion response was to minimize the ratios of UV-to-green sensitivities, compared with having used a much larger or much smaller criterion amplitude of response.

Subsequent to the above experiments, we measured the intensity-response curves in one cell at various wavelengths randomly presented, in order to see how many families of such curves there might be. There were systematic deviations from originally fitted, parallel, common curves, such that the data could be fitted better by two families of curves. This can be seen in Fig. 6. The change between the two families of curves occurs between 420 and 440 nm, which is to say at approximately the wavelengths of minimum sensitivity in UV-green cells (cf. Fig. 2). The solid lines in Fig. 6 have been fitted by eye and seem to have different shapes than in the less extensive results in the inset in Fig. 5, although we do not know what this means. As it happened, the

![Figure 6](https://example.com/figure6.png)

**Figure 6.** Comparisons of intensity-response curves measured across the spectrum. Data were taken from the most stable of cells in *Aeschna* (impalement time: 106 min; resting potential: 95-97 mV). The relative log quantum flux of 100-ms flashes is given on the abscissa vs. peak amplitudes of response, in millivolts, on the ordinate. Data points for the wavelength ranges 350-420 nm and 440-620 nm were arbitrarily slid along the log intensity axis for best fit by eye; solid lines were drawn by eye. Wavelengths were presented in random order, while several control curves were measured at 520 nm, interspersed among the other measurements as a check on the reproducibility of the preparation. The spectral sensitivity of this cell, measured at a 4-mV criterion amplitude of response, had the same shape as the bottom curve in Fig. 2 (log UV-to-green sensitivity ratio of 0.8). For the data of this and Fig. 5, care was taken to avoid spuriously large readings of amplitudes caused by abortive initial spikes.
curves at 370 and 500–520 nm did not cross for this cell, the cell being so much less sensitive (by five times) in the green at all intensities.

In summary, then, we find that intensity-response curves at low amplitudes of response of half the cells tested are parallel to each other only within given regions of the spectrum and that these regions correspond to those spectral mechanisms with constant wavelengths of peak sensitivity found during determinations of spectral sensitivity. At large amplitudes of response, however, intensity-response curves became parallel to each other and independent of wavelength; or, if they were parallel at low amplitudes of response, remained parallel at high amplitudes as well.

In our early determinations of spectral sensitivities, we did not have available the results shown in Fig. 6, although in effect we anticipated them: For most of those cells whose intensity-response curves at 370 and 520 nm did cross, we used the curve at 370 nm for converting amplitudes of responses at 350–420 nm into sensitivities, and the curve at 520 nm for converting amplitudes of responses at 440–650 nm. At most, there would have been errors of up to 0.2 log units in relative UV-to-visible sensitivities had we not done this.

**Selective Chromatic Light Adaptations**

Finally, we tested to see if we could selectively reduce the sensitivities recorded in single cells to one or the other end of the spectrum by light adapting the ocelli with constant background illuminations of UV or long wavelengths. For six cells with log UV-to-green sensitivity ratios of from 0.0 to 0.8, we measured complete spectral sensitivities when adapted to the dark and to backgrounds of 365 nm or to backgrounds with wavelengths 550 nm and longer (orange adaptations) (Fig. 7 illustrates results obtained when adapting lights were adjusted in intensity to reduce the sensitivities equally in the UV (left) or in the green (right). We have also used the maximum available intensities of both adapting lights, with equivalent results (not shown). The points in Fig. 7 are given in terms of absolute sensitivities, as measured, and the distances between curves give the amounts of log adaptation. Nomogram curves (dashed lines) have been drawn through the points in the visible, where they fit.

The results of these and similar experiments are clear-cut: Spectral sensitivities do not remain the same for both UV and orange adaptations. In general, UV adaptations reduced sensitivities equally across the spectrum, so that the top curves in Fig. 7, representing spectral sensitivities in the dark, were simply shifted downwards along the ordinate (sensitivity on a log scale). In only one cell did UV adaptation cause a small decrease in relative UV sensitivity. With orange adaptations, it was otherwise: Strong orange-adapting lights reduced sensitivity more in the visible than in the UV. This is especially clear at the left in Fig. 7, where the adapting lights used adapted the cell equally in the UV but unequally in the visible. It is also true, but not as obvious, at the right
in Fig. 7; when an orange-adapting light was as effective as a UV-adapting light in reducing sensitivity in the visible, it was less effective in reducing sensitivity in the UV. However, despite these asymmetries in adaptation, there were no significant shifts in wavelengths of peak sensitivity in the UV or in the visible, as can be seen in Table II. These results come only from UV-green cells; we have not tried chromatic adaptations of UV-blue+green cells of Anax.

Selective Adaptation as a Function of Adapting Intensities

One cell, light adapted by 1 log unit with UV and orange backgrounds, seemed to show none of the above, selective adaptation. All other cells were adapted by 1.5 log units or more and showed selective adaptation. Possibly, therefore,
there was a threshold amount of light adaptation needed for the selective effects illustrated in Fig. 7. To test for this, we measured sensitivity decreases of cells relative to sensitivity in the dark, as functions of intensities of adapting backgrounds. To minimize the dangers of losing a cell before a series of adaptations was complete, we saved time by measuring only the incremental sensitivities at 370 and 500 nm, wavelengths near the peaks of sensitivity in the UV and visible, respectively. The experiment was performed on five cells; typical results, from one cell, are shown in Fig. 8. There, it can be seen that sensitivities

\[
\begin{align*}
\text{quantum flux at } 365 \text{ nm} & \rightarrow \text{equivalent flux at } 500 \text{ nm} \\
\text{in UV-adapting beam} & \rightarrow \text{in orange-adapting beam}
\end{align*}
\]

at 370 and 500 nm fall in parallel with increasing amounts of UV-adapting intensities, but that sensitivity falls more rapidly at 500 nm than at 370 nm with increasing intensities of the orange-adapting light. Straight lines have been drawn by eye through the data in Fig. 8; the slopes of these lines (log ΔI vs. log I) give the Weber fraction. For the cell in Fig. 8, the Weber fraction was nearly 1 (0.95) for both 370- and 500-nm test flashes against the UV background and for the 500-nm test flashes against the orange background. The Weber fraction was only about half of this, or 0.55, for 370-nm flashes against the orange background. In the four other cells tested, the Weber fractions were smaller although in roughly the same proportions as in Fig. 8. For

FIGURE 8. Log incremental sensitivity changes in the UV- and green- vs. chromatic-adapting intensities on a log scale. For determinations of sensitivity decreases as functions of increasing adapting intensities, sensitivities for 4-mV criterion responses were determined first in the dark and then for each amount of adaptation, first for orange backgrounds and then for UV backgrounds. Additionally, the dark-adapted cell's responses to flashes from the adapting beams were used to calibrate these beams by comparisons with responses to known fluxes from the monochromator; the UV-adapting beam was also calibrated radiometrically with equivalent results. Intensities of the orange-adapting beam (wavelengths 550 nm and longer from a tungsten halogen lamp) are given as fluxes of equally bright 500-nm flashes. Slope of change of log sensitivity (the Weber fraction) at 370 nm as a function of log intensity of the orange background is 0.55, or about half the slope of 0.95 for the other three curves, through each of which a line having this slope has been drawn by eye. Similar results were found in two other cells from Aeschna and from two cells from Libellula, although the absolute values of the slopes varied from cell to cell and were usually less than those shown here. Data from same cell as in Fig. 6.
all cells, however, there had to be about 0.5–1 log unit of total adaptation before adaptation became selective, although it appears in Fig. 8 that there is no abrupt threshold for selective adaptation.

Finally, for the experiment illustrated in Fig. 8, we calibrated the background intensities using the cell itself to compare flashes of orange background lights with calibrated 500-nm flashes from the monochromator and flashes of UV background lights with 365-nm light from the monochromator. We also calibrated the UV background intensity radiometrically, with equivalent results. Then, both curves at 500 nm being parallel in Fig. 8, it can be determined at a UV-adapting background need be only about \( \frac{1}{10} \) (1.1 log units) as bright as an orange-adapting background to provide the same amount of adaptation at 500 nm. As the cell in question was about 10 times (0.8 log units) more sensitive in the UV, it appears that adapting efficiencies at 500 nm of the two background lights were approximately proportional to relative spectral sensitivities recorded in the cell. Since there was no selective adaptation for small amounts of light adaptation, the same proportionality would apply for adapting efficiencies at 370 nm at these small amounts of adaptation, but not at larger amounts.

All told then, it appears that the selective adaptation we have measured is a phenomenon graded with the intensity of the adapting light, that it works only upon that mechanism in the visible which has a peak sensitivity at 500 nm, that it is found only with orange-adapting lights that would be absorbed preferentially or exclusively by the mechanism in the visible, but that efficiency of adaptation in the visible by UV and orange-adapting lights is roughly proportional to the sensitivities recorded in the cell in the UV and visible, respectively.

**DISCUSSION**

The results of the present experiments demonstrate that two (and possibly three) spectral mechanisms contribute to the electrical responses recorded from within single cells in the median ocelli of dragonflies. The spectral sensitivities of all cells exhibited maxima in the UV and green regions of the spectrum (Figs. 2, 3, and 7), whereas 50% of those in *Anax* had, in addition, a discernible peak at about 440 nm (Fig. 3).

Cells in ocelli of dragonflies differ in their spectral properties in four ways from most other arthropod visual cells with peak sensitivities in two regions of the spectrum. First, there are large cell-to-cell variabilities in any one ocellus in the relative UV-to-green sensitivities (and in UV-to-blue-to-green sensitivities in some cells in *Anax*). This was illustrated in Figs. 2 and 4. In cells from most other arthropods, there are neither large differences between relative UV and visible sensitivities nor large variabilities from cell to cell (Wasserman, 1973). Possible exceptions are the UV-green cells in anterior median
eyes of wolf spiders, which have even larger variabilities and differences (DeVoe, 1972), and the rare UV-VIS cells in median ocelli of Limulus, where there are 2–3 log units of difference between UV and visible sensitivities (Nolte and Brown, 1969, 1972a).

Second, in some (but not all) ocellar cells of Libellula, we found wavelength-dependent waveforms of responses (Fig. 1), the responses at 370 nm being faster than those at 500 nm. We did not use intermediate wavelengths, so we do not know at which wavelength(s) the transition occurs. Wavelength-dependent responses have been detected in only a few other arthropod cells. Nolte et al. (1968) originally reported finding biphasic, wavelength-dependent responses in the UV and the VIS cells of the Limulus ocellus but have since concluded that the initial hyperpolarizations were due to pickup of extracellular currents (Nolte and Brown, 1972a). Long-lasting “tails,” dependent on wavelengths of stimulation with extremely bright flashes, have been found in ocellar cells of both Limulus (Nolte and Brown, 1969, 1972b) and the barnacle (Hochstein, Minke and Hillman, 1973). However, we did not find biphasic responses nor did we need bright lights to elicit wavelength-dependent waveforms of responses from cells of Libellula. Most similar are the UV-green cells in anterior median eyes of jumping spiders, where purely depolarizing responses to weak UV lights are slower than purely depolarizing responses to weak visible lights (DeVoe).

Third, in about half the cells tested in Anax and Aeschna, the intensity-response curves were steeper at 370 nm than 520 nm and crossed each other. This occurred both in UV-blue-green cells of Anax and in UV-green cells (large plot and inset, respectively, of Fig. 5). At low intensities, these cells were more sensitive to visible wavelengths, at high intensities, to UV wavelengths. In essence, these cells underwent a reverse Purkinje shift. The intensity-response curves of most arthropod visual cells have slopes independent of wavelength, the known exception being in the lateral eye of Limulus. There, Adolph (1968) found the frequency of quantum bumps to increase more slowly as a function of intensity at 397 nm than at wavelengths of 460 nm and longer. However, judging from the responses of one dragonfly ocellar cell (that in Fig. 5), we found that intensity-response curves from cells of dragonfly ocelli became parallel at high intensities; that is, their slopes too became independent of wavelength. The shift at low intensities from one wavelength-dependent slope of an intensity-response curve to the other slope seemed to occur (Fig. 6) between 420 and 440 nm, namely, in the region of the spectral sensitivity curves (Fig. 2) where there was a minimum of sensitivity between the UV and the visible peaks. Finally, slopes of intensity-response curves seemed to be independent of adaptation (not illustrated); they simply shifted

\(^2\) DeVoe, R. D. Ultraviolet and green receptors in principal eyes of jumping spiders. Manuscript submitted for publication.
for the most part along the log intensity axes (the abscissae in Figs. 5 and 6). For fear that we would lose a cell before completing all adaptations upon it, we never attempted to record complete intensity-response curves (in addition to the portion around 4 mV of response) at both UV and visible wavelengths during chromatic adaptations. Thus, we cannot say with certainty that intensity-response curves at 370 and 520 nm which were not parallel for the dark-adapted cell were also not parallel for the chromatically adapted cell.

Fourth, we have been able to change the relative spectral sensitivities of cells in dragonfly ocelli by chromatic adaptations with visible, but not UV, wavelengths. Such selective adaptations of single cells have previously succeeded only in UV-VIS cells of Limulus ocelli (Nolte and Brown, 1969) and UV-green cells in anterior median eyes of wolf spiders, Lycosa (DeVoe, 1972). Hence all cells which have been successfully selectively adapted have had great variabilities in ratios of UV to visible sensitivities and/or large differences in UV to visible sensitivities. Cells which consistently have almost equal UV and visible sensitivities have not been selectively adapted with chromatic lights (cf. Wasserman, 1973). Other features which selectively adaptable cells have had in common are, first, that wavelengths of peak sensitivity do not change during selective adaptations, and second, in none of the cells have adaptations been completely selective. That is, all adapting backgrounds have reduced sensitivities all across the spectrum; selectivity results because sensitivities are reduced more in one region of the spectrum than another. However, only in dragonfly ocellar cells have UV backgrounds usually failed to adapt more in the UV than in the visible, as they do in ocelli of Limulus and Lycosa (Nolte and Brown, 1969; DeVoe, 1972). Likewise, only for dragonfly ocellar cells have the adapting lights themselves been calibrated, with the resultant finding that efficiencies of adaptation by UV and orange background lights, tested with 500 nm incremental flashes, are roughly proportional to the spectral sensitivity of the dark-adapted cell.

Our demonstration of two discriminable spectral mechanisms in recordings from single cells in the median ocelli of dragonflies raises the question of whether the observations are accounted for by coupling between cells or by the presence of two photopigments in each cell. Couplings between cells could be electrical, as proposed in models for the Limulus lateral eye (Smith and Baumann, 1969) or ocellus (Nolte and Brown, 1972a), and for the compound eye of the honeybee (Shaw, 1969; Snyder et al., 1973; Menzel and Snyder, 1974). Alternatively, coupling in the dragonfly ocellus might occur via the receptor-receptor synapses found in the plexus (Dowling and Chappell, 1972), although to our knowledge no such couplings between visual cells of unlike spectral sensitivities have been suggested. The other basic explanation, two photopigments in a single receptor cell, has been proposed to account for
spectral sensitivities in the compound eye of the locust (Bennett et al., 1967) and in the anterior median ocelli of the wolf spider (DeVoe, 1972).

We have found no simple way to fit our findings into either of these explanations. We have considered a coupling model based on that of Shaw (1969) to see if dragonfly ocellar UV-green cells might be couplings of hypothetical UV and green cells, much as the UV-VIS cells of Limulus ocelli are the result of couplings between known UV and VIS cells (Nolte and Brown, 1972a). It appears to us that we would need many ad hoc assumptions to fit a coupling hypothesis to our data. Some of these assumptions cannot be justified by, and may even be in contradiction to, data from other visual systems, and for these reasons a discussion of them best awaits further experimentation. Similarly, there are neither kinetic models nor experimental data for interactions of two photopigments in a single cell against which we could test our results, and we have no basis for constructing such a model. The most reasonable course at this time is to decide which hypothesis should be pursued further by testing (a) if there is any coupling at all between cells and (b) the possibility that there are two photopigments in a single cell. Intercellular diffusion of Procion Dyes (Kaneko, 1971) and dual electrode studies (Shaw, 1969) can be used to test for electrical coupling between receptor cells. In the dual electrode studies, intracellular injections of different Procion Dyes into the two cells will show how close together they are (Kaneko, 1971; Mote and Goldsmith, 1971). Coupling via receptor-receptor synapses in the plexus can be tested pharmacologically (Klingman and Chappell, 1974). The other hypothesis, the presence of two photopigments in each cell, can be tested by finding the spectral sensitivity of the early receptor potential, as has been done for single photopigments and their photoproducts in other arthropod visual cells (Smith and Brown, 1966; Brown et al., 1967; Minke et al., 1973; Pak and Liddle, 1974).

Color Vision in the Ocellus

In this study of color sensitivities of cells in dragonfly ocelli, we have found both UV-green cells, with peak sensitivities at 360 and 500 nm, and (in Anax) UV-blue-green cells with peak sensitivities at 355, 440, and 500 nm (rounding off the wavelengths of peak sensitivities in Table II to the nearest 5 nm). The roles, if any, of such cells in color vision by the ocellus would thus appear to be based on broad-band cells as opposed to receptors turned to a selective wavelength region of the spectrum (see Wasserman, 1973). Cells individually sensitive to one or another of each of these wavelengths have been found in compound eyes of the same genera (Autrum and Kolb, 1968; Horridge, 1969; Eguchi, 1971). Ocellar cells thus provide no information about regions of the spectrum not already available, in a way more useful for color vision, in the compound eye. Moreover, poor imaging by the ocellar lens and high con-
vergence of receptors onto second-order fibers might be thought to obviate contributions of individual receptors to color vision. However, the observation that movement information is conveyed from the median ocellus of dragonflies to the cervical nerve chain (Zenkin and Pigarev, 1971) indicates that sensory information from individual receptors or from small groups of receptors may become available to the animal. Assuming this to be so, then neural differencing or comparing of cells (in groups or alone) with both high and low relative visible sensitivities could provide information about relative UV-to-visible intensities. This might be particularly useful for analyzing the skylight towards which the median ocellus looks and might be important to the animal if the ocellus played a role in diurnal rhythms, that is, to estimate the time of day (cf. Goodman, 1970).

The selective chromatic adaptations we observed could be interpreted as a kind of “automatic color control,” possibly useful in spectral analysis of skylight: During the day, the median ocellus would be strongly light adapted by the predominantly UV light scattered from the sky. Such UV adaptation keeps the relative UV-to-visible sensitivities of ocellar cells constant. At twilight, when the proportions of short-to-long wavelength light in the sky are lower, UV sensitivities of ocellar cells would be less strongly adapted, thus compensating for decreased amounts of UV light. However, selective adaptation would disappear at light levels lower still, and the reverse Purkinje shift would occur in some, perhaps half, of the cells: Due to crossing of intensity-response curves, these cells would change from more-sensitive-to-UV light to more-sensitive-to-visible light, of which the intensities are higher at first or last light and at night (Menzel and Knaut, 1973). Since the spectral mechanisms that we found in ocellar cells have not been found in cells of the compound eyes, this could mean that the ocellus might furnish to the animal color information unavailable from the compound eyes.

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