Light Adaptation in *Drosophila* Photoreceptors: I. Response Dynamics and Signaling Efficiency at 25°C

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**ABSTRACT** Besides the physical limits imposed on photon absorption, the coprocessing of visual information by the phototransduction cascade and photoreceptor membrane determines the fidelity of photoreceptor signaling. We investigated the response dynamics and signaling efficiency of *Drosophila* photoreceptors to natural-like fluctuating light contrast stimulation and intracellular current injection when the cells were adapted over a 4-log unit light intensity range at 25°C. This dual stimulation allowed us to characterize how an increase in the mean light intensity causes the phototransduction cascade and photoreceptor membrane to produce larger, faster and increasingly accurate voltage responses to a given contrast. Using signal and noise analysis, this appears to be associated with an increased summation of smaller and faster elementary responses (i.e., bumps), whose latency distribution stays relatively unchanged at different mean light intensity levels. As the phototransduction cascade increases, the size and speed of the signals (light current) at higher adapting backgrounds and, in conjunction with the photoreceptor membrane, reduces the light-induced voltage noise, and the photoreceptor signal-to-noise ratio improves and extends to a higher bandwidth. Because the voltage responses to light contrasts are much slower than those evoked by current injection, the photoreceptor membrane does not limit the speed of the phototransduction cascade, but it does filter the associated high frequency noise. The photoreceptor information capacity increases with light adaptation and starts to saturate at ~200 bits/s as the speed of the chemical reactions inside a fixed number of transduction units, possibly microvilli, is approaching its maximum.

**KEY WORDS:** vision • retina • information • neural coding • graded potential

**INTRODUCTION**

The ability to adapt to mean illumination allows a photoreceptor to gather and process information about relative light changes (contrasts) over a vast range of intensities without saturating its steady-state membrane potential. The process of adaptation itself involves both the workings of the phototransduction cascade and the photoreceptor membrane. The phototransduction cascade is a signal pathway where a photoisomerized photopigment activates a cascade of intracellular biochemical reactions, which modulates the opening of light-sensitive ion channels on the photoreceptor membrane. Its output is the light (or transduction) current. In turn, the photoreceptor membrane has additional voltage-sensitive ion channels, which together with its capacitance shape the changes in the light current into a voltage response. There are many physical factors that can constrain the fidelity of the photoreceptor voltage responses: the physics of photon absorption, the delays and reproducibility of chemical reactions in the transduction cascade, and the stochasticity in the ion channel kinetics. So what kind of coding strategies do photoreceptors use against the noise, and how reliable are their graded voltage responses as neural representations of the dynamic contrast stimulation?

*Drosophila* photoreceptors have been successfully used as a model system for analyzing insect phototransduction. Recently, the transduction dynamics in dark-adapted photoreceptors have been extensively studied by patch-clamping dissociated cells (for reviews see Hardie and Minke, 1995; Scott and Zuker, 1998), but the physiology and response properties of light-adapted photoreceptors have been largely ignored. The reason for this is simple: the in vitro preparation does not readily survive prolonged light stimulation; on the other hand, although the in vivo intact fly preparation can survive hours of light adaptation, its small size has made intracellular recordings very difficult. Consequently, only limited data, such as some basic noise analysis of the elementary responses (i.e., quantum bumps) during light adaptation are available (Wu and Pak, 1978; Johnson and Pak, 1986). This report is an extensive in vivo study of the response and membrane properties and light adaptation dynamics in *Drosophila* photoreceptors at 25°C using linear signal and noise analysis with natural-like contrast stimulation. We found that the stronger light adaptation greatly improves the photoreceptors’ information capacity. At
low light intensity levels, the fidelity of photoreceptor responses is limited by the photon shot noise. Amplification of single photon responses into individual detectable events leads to noisy voltage responses, whose slow speed is set by the slow rate of the transduction reactions and matches the filter properties of the photoreceptor membrane. Such low frequency signaling keeps the photoreceptor information capacity low. On the other hand, in bright illumination, the Poisson properties of the light provide a high fidelity contrast stimulus. The voltage responses consist of a multitude of small and fast bumps, the photoreceptor membrane provides faster signaling, but the bump latency distribution remains relatively unaffected and this now sets the ultimate speed limit of the voltage responses. Further light adaptation does not improve the signaling fidelity when the rate of the chemical reactions is already at its maximum in the majority of the transduction units. Consequently, the photoreceptor information capacity starts to saturate \(~200–300\) bits/s at a mean photon absorption rate of \(3 \times 10^5\) photons/s.

Materials and Methods

Animals and Preparation

Flies, normal wild-type red-eyed *Drosophila melanogaster* were obtained from a laboratory culture and reared at a constant temperature of \(25^\circ\text{C}\). 4–7-d-old flies were mounted with their head protruding from the open tip of a conical holder, whose hollow copper core was shielded outside with a ceramic insulator. Flies were fixed by their backs to the copper tip with a mixture of beeswax and heat sink paste, and the proboscis was stretched to eliminate vergence eye movements. This left the abdomen intact for experiments, were used in a number of different experiments. Each of the animals stated otherwise. All the statistics are given as mean ± SD.

The fly's body temperature was measured with a thermocouple mounted in the copper core next to the fly and could be changed in seconds from 10 to 35°C by a custom-designed feedback-controlled power source driving the Peltier temperature. The room temperature was monitored with a separate thermocouple. Control measurements from the fly's head revealed that the head temperature followed the temperature of the copper holder with \(±1^\circ\text{C}\) accuracy over the temperature range. This paper presents photoreceptor data recorded at 25°C, whereas the companion paper (Juusola and Hardie, 2001 in this issue) investigates the effect of temperature on photoreceptor signaling over a 15°C temperature range.

Microelectrodes and Recording Criteria

The microelectrodes were pulled with horizontal pullers (model P-97 or P-2000; Sutter Instrument Company) from fiber-filled borosilicate or quartz glass capillaries (Clark) with an inner and outer diameter of 0.5 and 1.0 mm, respectively. Electrodes were back-filled with 3 M KCl and coated with Vaseline. The resistance of the microelectrodes inside a cell varied between 120 and 250 MΩ. Because recording from small photoreceptors requires very sharp microelectrodes with good electrical properties, the resistance of the electrodes with a suitable shape (short shank with rapid taper to minimize the effects of intramural capacitance) was measured before the actual experiments in a grounded salt drop. This selection greatly increased the odds of good recordings.

The room temperature was monitored with a separate thermocouple mounted in the copper core next to the fly and could be changed in seconds from 10 to 35°C by a custom-designed feedback-controlled power source driving the Peltier temperature. The fly's body temperature was measured with a thermocouple mounted on the back of the fly's head close to the eye.

Membrane potentials were recorded with the amplifier operating in the compensated current-clamp (CC) or balanced bridge mode. The recordings were carried out from green-sensitive RI-6 photoreceptor cells that are the dominant input to the *Drosophila* visual system (Strausfeld, 1989). Because we used red-eyed flies instead of the commonly used white-eyed mutations, which lack all the screening pigments, and provided the light stimuli through a small point source (see Light Stimulation), the effects of extracellular field potentials on the recordings were minimal. The maximum extracellular field potentials evoked by saturating light flashes measured in the retina were typically \(<5\) mV. A successful photoreceptor penetration was seen as a 60–75-mV drop in the electrode potential and vigorous responses to low intensity light pulses. The input resistance of the photoreceptors in the dark was \(700 ± 540\) MΩ \((n = 9)\) and in fully light-adapted conditions, which depolarized the membrane 25–40 mV above the resting potential, \(320 ± 100\) MΩ \((n = 4)\). These values are much higher than those previously reported from intracellular recordings (Wu and Pak, 1978; Johnson and Pak, 1986), but similar to those measured using patch-clamp electrodes (Hevers and Hardie, 1995). By injecting a pseudorandomly modulated current into the cell and calculating the resulting membrane impulse response, we could estimate the membrane time constant. In the dark, membrane charging could be approximated with a single exponential time constant, \(\tau_m \approx 20\) ms; when depolarized by a bright light background, \(\tau_m\) was reduced to between 1 and 3 ms. However, often in light-adapted conditions, membrane charging was better fitted with two exponentials, probably indicating activation of separate voltage- and/or light-sensitive conductances. Before stimulus modulation experiments, the cells were allowed to dark adapt and seal properly for 2–10 min. Only data from photoreceptors with saturating impulse responses \(>60\) mV, minimal dark input resistance of \(300\) MΩ, and dark resting potential below \(-60\) mV are presented in detail in this study. These photoreceptors allowed stable recordings sometimes for several hours and, therefore, were used in a number of different experiments. Each of the experiments was repeated with a minimum of five cells unless stated otherwise. All the statistics are given as mean ± SD.

Recording Procedures

The stimulus generation, data acquisition, and signal analysis was performed by a custom written program (BIOSSYSL, M. Juusola, 1997-9) based on the MATLAB programming language (Mathworks) and an interface package for National Instruments boards (MATDAQ, H.P.C. Robinson, 1997-9).

Light Stimulation

Light stimuli were provided with a green high intensity light-emitting diode (Marl Optosource) driven by a custom-built LED.
The light output of the LED was monitored continuously with a pin diode circuit. The light output range of 6 log units was calibrated by counting the number of single photon responses (bumps; Lillywhite and Laughlin, 1979) during prolonged dim illumination (Juusola et al., 1994). The LED light output was attenuated by neutral density filters (Kodak Wratten) to provide five different adapting backgrounds in z 1–log unit steps indicated by BG0, BG-1, BG-2, BG-3, and BG-4. The lowest adapting background applied, BG-4, was estimated to be z 300 effective photons/s and the highest intensity, BG0 (no filter), was z 3 x 10^6 photons/s. A Cardan arm system allowed free movement of the light source at a constant distance (85 mm) from the eye’s surface; the light source subtended z 2.8. Light contrast (c) was defined as a change in the light intensity (ΔY) divided by the mean light background (Y mean) (Fig. 1 A, a):

\[ c = \frac{ΔY}{Y_{\text{mean}}} \]  

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In case of pseudorandom contrast modulation (band-limited signal of a Gaussian amplitude distribution and spectrally white up to a 150 Hz; Fig. 1 B, a) \( \Delta Y \) is defined as the SD of the stimulus modulation \( \left( \nu \right) \), and here varied between \(-0.2\) and \(+0.2\) nA. (b) Voltage responses, \( v(t) \), were averaged to obtain (c) the signal, \( s(t) \), and (d) the noise, \( n(t) \), superimposed on it. \( s(t) \), contained any noise induced by the voltage-sensitive membrane and phototransduction noise. Sampling frequency was 1 kHz and the record duration was 10 s for 10 trials. (B) Because of the switched current clamp, we obtained true recordings of the current being injected into a photoreceptor and could calculate the variance of the current stimulus (i.e., stimulus noise). This variance was very small, again at the bit resolution limit of the A/D converter, and its power was \( \sim 10^{-6} \) of that of the average power of the injected current waveform. Current stimuli with different bandwidth produced similar results (data not shown). By taking the FFT from the stimulus, response, signal, and noise traces, we could calculate the corresponding power spectra (a, b, c, and d, respectively). (c) \( SNR(f) \) was calculated with Eq. 3. From \( SNR(f) \), we could estimate both (g) the linear coherence function, \( \gamma_{SNR}(f) \), and (f) the cell’s information capacity by using Eqs. 6 and 5, respectively. The information capacity of the membrane was much higher than that of transduction. See MATERIALS AND METHODS for more details. (C) From the signal and stimulus, we calculated (a) the coherence, \( \gamma^2_{exp}(f) \); the frequency response, i.e., (b) gain, \( Z(f) \), and (c) phase, \( P(f) \), and (d) the impulse response functions, \( z(t) \), as described in MATERIALS AND METHODS. From input impedance \( (Z(f), \text{ i.e., gain}) \) we took the DC value as the mean input resistance of the cell, here 450 MΩ. The membrane time constant \( \tau_m \) was approximated by fitting an exponential to \( z(t) \), here 1.98 ms.

**Current Stimulation**

To measure the light adaptational changes in the membrane impedance, we injected pulses or pseudorandomly modulated current into photoreceptors via the recording microelectrode (Weckström et al., 1992b) at all light intensity levels including darkness (Fig. 2 A, a). Electrodes that had suitable electrical properties (input resistance <180 MΩ) were used, and their capacitance was carefully compensated before the current injection experiments. Currents of up to 0.4 nA were injected though the electrodes to generate mean voltage changes \<0.2 mV. The use of a switched clamp amplifier allowed us to record and monitor the true intracellular photoreceptor voltage and current during current and/or light stimulation (Juusola, 1994).

**Data Acquisition**

Current and voltage responses were low-pass filtered at 0.1–2 kHz together with the corresponding LED output (model KEMO...
VBF/23 low pass elliptic filter). The signals were sampled at 0.5–10 kHz, digitized with a 12-bit A/D converter (model PCI-MIO-16E-4; National Instruments), and stored on a hard disk (Pentium II, 450 MHz). The sampling was synchronized to the computer-generated stimulus signal and records of the three signals were stored during each recording cycle. The length of records varied from 100 ms to 10 s, but during pseudorandom stimuli was ±4 s (see Figs. 1 and 2, which show 0.5-s-long samples out of 10-s-long stimuli). A 2-s steady light background stimulus was maintained between stimulus sequences to provide equal light adaptation conditions for each run. The recording system, including the microelectrode, had a frequency response with a 3 dB high frequency cut-off at 10 kHz or higher and, therefore, had negligible effect on the results.

At different mean light backgrounds, the photoreceptor performance was tested using repeated presentations of the same pseudorandom Gaussian stimulus (light contrast and/or current). Each experiment proceeded from the weakest to the strongest adapting background. After stimulation, cells were re-dark–adapted. Recordings were rejected if the same sensitivity was not recovered by dark adaptation.

**Processing of Voltage Responses in Time Domain**

Repeated presentations (10–30 times) of virtually identical pseudorandom light contrast, \( c(t) \), or current, \( i(t) \) (Figs. 1 A and 2 A, a) evoked slightly variable voltage responses, \( r_{\text{v}}(t) \). Subtraction of the signal, \( s(t) \), from the individual responses, \( r_{\text{v}}(t) \), gave the noise component of each individual response period (Figs. 1 A and 2 A, d; compare with Juusola et al., 1994):

\[
n(t) = r_{\text{v}}(t) - s(t).
\]

Additionally, to avoid a possible bias of the noise estimates by the relatively small number of samples, the noise was recalculated using a method that did not allow signal and noise to be correlated. For example, when an experiment consisted of 10 trials, 9 of the trials were used to compute the mean and the other to compute the noise. This was repeated for each possible set of 9 responses giving 10 noncorrelated noise traces. These two methods gave similar noise estimates with very low variance. Errors due to residual noise in \( s(t) \) and \( r_{\text{v}}(t) \) were small and proportional to (noise power)/\( \sqrt{n} \), where \( n \) is 10 (Kouvalainen et al., 1994). The signal-to-noise ratio in the time domain, SNRV, was estimated by dividing the signal variance by the corresponding noise variance.

**Signal and Noise Power Spectra and SNRV\(_f\) (f)**

\( s(t) \) and \( n(t) \) were segmented into 50% overlapping stretches and windowed with a Blackman-Harris four term window (Harris, 1978) before their corresponding spectra, \( S_f(s) \) and \( N_f(n) \), were calculated with an FFT algorithm. Signal and noise power spectra, \( |S_f(s)|^2 \) and \( |N_f(n)|^2 \), respectively, where \( || \) denotes the absolute value and \( \overline{c} \) denotes the average over the different stretches of the signal and noise data, were calculated as real-valued functions (see Figs. 1 B and 2 B, c and d). In the same way the stimulus presentations \( c(t) \) and \( i(t) \) and the individual voltage responses, \( r_{\text{v}}(t) \), yielded the power spectra \( |C_f(c)|^2 \), \( |D_f(i)|^2 \) and \( |R_f(r_{\text{v}})|^2 \) (see Figs. 1 B and 2 B, b and a, respectively). The variability in the stimulus was estimated by subtracting the average stimulus from the individual stimulus records (see above) and calculating the corresponding noise spectrum (Figs. 1 B and 2 B, a). It appears that the stimulus noise constituted \(<10^{-4}\) of the stimulus power. The variability in the photoreceptor responses was much larger and, hence, not caused by the variability in the stimulus.

The signal-to-noise ratio in the frequency domain, \( \text{SNRV}_f(f) \) (Figs. 1 Band 2 B, c), of the photoreceptor potential was determined by dividing its signal power spectrum, \( |S_f(s)|^2 \), by its noise power spectrum, \( |N_f(n)|^2 \) (Figs. 1 B and 2 B, c and d; Juusola et al., 1994):

\[
\text{SNRV}_f(f) = \frac{|S_f(s)|^2}{|N_f(n)|^2}.
\]

The shape of the derived signal power spectra showed some degree of ripple, following the slight unevenness in the stimulus power spectra. Since this effect can lead to reduction in the photoreceptor SNRV\(_f(f) \) at the stimulus frequencies that carry less power, the signal power spectrum was corrected by the stimulus power spectrum (Fig. 1 B, c, the dotted line):

\[
|S_f(s)|^2_{\text{corr}} = |S_f(s)|^2 \left[ \frac{|C_f(c)|^2}{|C_f(c)|^2_{\overline{c}}} \right].
\]

with \( |C_f(c)|^2_{\overline{c}} \) being the mean of the light contrast power spectrum over the frequency range investigated (i.e., 0–200 Hz). In most cases, the stimulus-corrected signal power spectrum overlapped smoothly that of the measured one. However, sometimes at low adapting backgrounds, we found that the stimulus-corrected signal power was noisier than the uncorrected signal power. In such cases, this smoothing procedure was not used.

Electrode recording noise power spectrum, \( |N_f(n)|^2 \), calculated from the voltage noise (measured in the extracellular space after pulling the electrode from the photoreceptor), was not routinely subtracted from the data as the levels were very low compared with signal power, \( |S_f(s)|^2 \), and noise power, \( |N_f(n)|^2 \), and therefore made little difference to estimates of the photoreceptor SNR or information capacity at the frequencies of interest.

**Information Capacity**

From the signal-to-noise ratio, the information capacity \( (H) \) can be calculated (Shannon, 1948; Figs. 1 B and 2 B, f):

\[
H = \int_{-\infty}^{\infty} (\log_2[\text{SNRV}_f(f)] + 1) \, df.
\]

The dimension of the information capacity is bits/s. Because of the unreliability of the signal at frequencies above \( j = 150 \) Hz, the upper frequency limit of the integral was not taken to infinity \( (\infty) \) but \( j \).

Because the voltage responses at high adapting backgrounds are not purely Gaussian, but slightly skewed towards hyperpolarizing values (see results), the information capacity estimates determined here can only be considered as upper bounds of the true information capacity (Juusola and French, 1997). On the other hand, at low adapting backgrounds, where the voltage responses are dominated by large and slow elementary responses, the signal is Gaussian, whereas the noise distribution is slightly skewed towards depolarizing values, resulting in an underestimation of the true information capacity. The information capacity estimates are further influenced by the fact that, as explained in the previous section, the photoreceptor noise power includes the electrode noise. This causes a slight underestimation of the true information capacity values.

The information capacity calculated from the input-corrected signal power spectra (Fig. 1 B, c, and see Eq. 4) was only slightly larger than the uncorrected value, on average less than 10% (Fig. 1 B, f: dotted line versus continuous line).
**Coherence**

The coherence function for a purely linear coding scheme is calculated from the signal-to-noise ratio (Bendat and Piersol, 1971; Theunissen et al., 1996; Haag and Borst, 1997):

\[
\gamma_{SNR}^2(f) = \frac{SNR(f)}{SNR(f) + 1^2}.
\]  

(6)

In a perfectly linear, noise-free system, the coherence is expected to equal 1 for all frequencies. Here, we have a case where noise is added to the signal as it travels through the photoreceptor filter to produce a response. The coherence function, \(\gamma_{SNR}^2(f)\) (see Figs. 1 and 2, B, g), follows the changes in its signal-to-noise ratio, \(SNR(f)\) (see Figs. 1 B and 2 B, c). On the other hand, the coherence function for the noise-free voltage signal, \(\gamma_{exp}^2(f)\) (see Figs. 1 C and 2 C, a), is calculated as (Bendat and Piersol, 1971):

\[
\gamma_{exp}^2(f) = \frac{\abs{\langle S_k(f) \cdot C^*(f) \rangle}^2}{\langle C(f) \cdot C^*(f) \rangle \cdot \langle S_k(f) \cdot S_k^*(f) \rangle}.
\]

(7)

Hence, we can compare the linear coherence, \(\gamma_{SNR}^2(f)\) (Eq. 6), to the noise-free coherence, \(\gamma_{exp}^2(f)\) (Eq. 7) and, thus, expose any nonlinearities of the dynamic voltage responses.

**Frequency and Impulse Responses**

After frequency domain averaging of the stimulus and signal spectra of different segments, the photoreceptor frequency response, \(T(f)\) (Eq. 8), and impulse response, \(k(t)\) (or first-order Wiener kernel; Eq. 9), as well as membrane impedance and impulse response, \(Z(f)\) and \(z(t)\), respectively, and coherence function, \(\gamma_{exp}^2(f)\) (Eq. 7; Figs. 1 C and 2 C, a-d), were calculated from the autospectrum of the corresponding input (contrast, \(C(f)\) - \(C^*(f)\)) or current (\(I(f)\) - \(I^*(f)\)) stimulus and output (photoreceptor signal, \(\langle S_k(f) \cdot S_k^*(f) \rangle\) and their cross-spectrum \(\langle S_k(f) \cdot C^*(f) \rangle\) or \(\langle S_k(f) \cdot I^*(f) \rangle\)), where the asterisk denotes the complex conjugate, and \(\langle \cdot \rangle\) is the average over the different stretches of the input and output data. For voltage signals to light contrasts to:

\[
T(f) = \frac{\langle S_k(f) \cdot C^*(f) \rangle}{\langle C(f) \cdot C^*(f) \rangle}.
\]

(8)

The frequency response, \(T(f)\), is a complex-valued quantity that can be expressed in terms of a gain, the ratio of the photoreceptor response amplitude to the stimulus amplitude (contrast gain: \(G_c(f) = \frac{\langle C(f) \rangle}{\langle S_k(f) \rangle}\) (Fig. 1 C, b) or injected current: impedance, \(Z(f) = \frac{\langle I(f) \rangle}{\langle S_k(f) \rangle}\) (Fig. 2 C, b)), and a phase, \(\phi(f)\), the phase shift between the stimulus and the response (Figs. 1 and 2, Cc):

\[
P(f) = \tan^{-1}\left[ \frac{\Im \{S_k(f) \cdot C^*(f)\}}{\Re \{S_k(f) \cdot C^*(f)\}} \right].
\]

(9)

where \(\Im\) is the imaginary and \(\Re\) is the real part of the cross-spectrum. Photoreceptors are not minimum phase systems, but include a pure time delay, or dead-time (French, 1980; Juusola et al., 1994; de Ruyter van Steveninck and Laughlin, 1996b; Anderson and Laughlin, 2000). The minimum phase of a photoreceptor is calculated from the Hilbert transform, \(F_{Hi}\), of the natural logarithm of the contrast gain function \(G_c(f)\) (de Ruyter van Steveninck and Laughlin, 1996b):

\[
P_{\text{min}}(f) = -1 \cdot \Im(F_{Hi}[\ln(G_c(f))]).
\]

(10)

(For more details see Bracewell, 2000). The frequency-dependent phase shift caused by the dead-time, \(\phi(f)\), is the difference between the measured phase and the estimated minimum phase (see Fig. 1 C, c):

\[
\phi(f) = P(f) - P_{\text{min}}(f).
\]

(11)

The dead-time was estimated over the flat frequency range (here 10–90 Hz) of \(\phi(f)/(2\pi f)\), where \(f\) is the frequency in Hz.

The impulse responses, \(k(t)\) or \(z(t)\), which characterize the linear filtering properties of a photoreceptor to contrast or current stimulation in the time domain, were calculated as an inverse FFT of the corresponding frequency responses. For voltage signals to light contrasts:

\[
k_v(t) = F^{-1}(T_v(f)).
\]

(12)

**Light Current**

Because in the light-adapted state both the membrane impedance and photoreceptor voltage responses behave linearly (as judged by the near unity coherence functions in Figs. 1 and 2, C; see also Results) we can calculate the phototransduction cascade’s (or light current’s) frequency response, \(T_v(f)\), and impulse response, \(k_v(t)\), using linear systems analysis techniques. A first order approximation of the light current signal, \(s_l(t)\), can be derived by deconvolving the impulse response of the membrane, \(z(t)\) (Fig. 2 C, d), from the corresponding contrast-evoked photoreceptor voltage signal, \(s_v(t)\) (Fig. 1 A, c), both recorded in the same photoreceptor at the same mean light intensity and temperature:

\[
s_v(t) = \int_0^\infty z(\tau) \cdot s_l(t-\tau) d\tau.
\]

(13)

Then \(T_v(f)\) and \(k_v(t)\) can be computed from the light contrast stimulus, \(C(f)\), and the light current signal, \(S_l(f)\), as described in Eqs. 8 and 12, respectively.

**Results**

We investigated the response properties of Drosophila photoreceptors to light contrast and current stimulation in the dark and at five different adapting backgrounds at different temperatures. We show here data measured at 25°C (Figs. 1 and 2). This was the rearing temperature of the pupae but, more importantly, in temperature gradient tests Drosophila have shown strong behavioral preference to dwell at ambient temperatures between 23 and 25°C (Sayeed and Benzer, 1996). We found that the general adaptational changes in photoreceptor response dynamics, as described below, were not restricted to a certain temperature (see also companion paper Juusola and Hardie, 2001, in this issue). Here our aim was to fold: (1) to define the light adaptation dynamics of Drosophila photoreceptors as a reference database for future studies of Drosophila eye mutations, and (2) to illustrate how the phototransduction cascade and photoreceptor membrane coprocess the photoreceptor voltage responses. To accomplish the latter task properly, the voltage responses of a photoreceptor to light contrast stimulation and current injection were measured in the same cell at the same mean light background. As will be
demonstrated below, the dynamic response characteristics of light-adapted photoreceptors vary relatively little from one cell to another and are very similar across animals under similar illumination and temperature conditions. We illustrate our data and analysis with results from typical experiments starting with impulse and step stimuli and progressing to more natural-like stimulation. The data are from five photoreceptors, whose symbols are maintained throughout the figures of this paper.

I: Voltage Responses of Dark-adapted Photoreceptors

The photoreceptor voltage responses to light stimuli were first studied after 5–30 min of dark-adaptation. Fig. 3 A shows typical voltage responses to 1-ms light impulses of increasing relative intensity: (0.093, 0.287, 0.584 and 1, where 1 equals 10,000 effectively absorbed photons; note that the light intensity of the brightest impulse is 3.3 times that of BG0). Photoreceptors respond with increasing depolarizations, sometimes reaching a maximum size of 75 mV, before returning towards the dark resting potential (60 to 75 mV). The latency of the responses decreases with increasing stimulus intensity, and often their early rising phases show a spikelike event or notch similar to those reported in the axonal photoreceptor recordings of blowflies (Weckström et al., 1992a).

Fig. 3 B shows voltage responses of a dark-adapted photoreceptor to 100-ms-long current pulses (maximum magnitude 0.4 nA). The photoreceptors demonstrate strong, time-dependent, outward rectification, because of the increased activation of voltage-sensitive potassium channels starting approximately at the resting potential (Hardie, 1991b). The depolarizing pulses elicit voltage responses with an increasingly square wave profile, with the larger responses to stronger currents peaking and rapidly returning to a steady depolarization level. By contrast, hyperpolarizing pulses evoke slower responses, which resemble passive RC charging. The input resistance appears to vary from 300 to 1,200 MΩ between cells, yielding a mean cell capacitance of 52 ± 18 pF (n = 4).

II: Voltage Responses to Mean Light Intensities

Fig. 3 C shows 10-s-long traces of the membrane potential recorded in darkness and at different light intensity levels 20 s after stimulus onset. Because of the high membrane impedance (300 MΩ), dark-adapted photoreceptors appear relatively noisy. Some of this voltage fluctuation represents instrumental noise due to using high resistance electrodes, but most is photoreceptor noise, possible sources being stochastic channel openings, noise from feedback synapses in the lamina, or spontaneous photoisomerizations. This was concluded because the electrode noise measured in extracellular compart-
ments was much smaller than that of the photoreceptor dark noise. No further attempts were made to identify the dark noise source. Dim light induces a noisy depolarization of a few millivolts because of the summation of irregularly occurring single photon responses (bumps). At higher light intensity levels, the voltage noise variance is much reduced and the mean membrane potential saturates at 25–40 mV above the dark resting potential. The steady-state depolarization at the brightest adapting background, BG0 (~3 × 10^6 photons/s), is on average 39 ± 9% (n = 14) of that of the photoreceptor’s maximum impulse response in darkness.

**III: Voltage Responses to Dynamic Contrast Sequences**

Since a fly’s photoreceptors in its natural habitat are exposed to light intensity fluctuations, the signaling efficiency of *Drosophila* photoreceptors was studied at different adapting backgrounds with repeated presentations of an identical Gaussian light contrast stimulus, here with a mean contrast of 0.32. Although the contrast in natural sceneries is non-Gaussian and skewed, its mean is close to this value (Laughlin, 1981; Ruderman and Bialek, 1994). Averaging 10–30 voltage responses gives a reliable estimate of the photoreceptor signal for a particular background intensity. The noise in each response is determined by subtracting the average response (the signal) from the individual voltage response.

Fig. 4 shows 1-s-long samples of the 10-s-long contrast stimulus (sampling at 500 Hz, filtering at 250 Hz), photoreceptor voltage signal (Fig. 4 A) and noise (Fig. 4 B) with their corresponding probability distributions (Fig. 4 C) at different adapting backgrounds. The size of the voltage signal measured from its variance (Fig. 4 D; the

![Figure 4](image-url)
symbols depict individual photoreceptors) increases $(5 \pm 1)^2$ times when the mean light intensity increases $10^4$ fold, before it saturates as does the mean membrane potential (i.e., $\mu$ (in millivolts); Fig. 4 E). Concurrently the signal resolution for finer temporal details in the stimulus also improves greatly, seen as the increasing transients in the signal waveform (Fig. 4 A). As the signal content changes, so does its spread. The signal probability distribution (Fig. 4 C, continuous line) is Gaussian under dim light conditions, but slightly skewed to hyperpolarizing values at brighter adapting backgrounds (BG-1 and BG0), suggesting that compressive nonlinearities either in the phototransduction cascade or membrane dynamics affect depolarizing voltage responses (see later in the phototransduction cascade or membrane dynamics affect depolarizing voltage responses (see later IV: Photoreceptor Membrane during Natural-like Stimulation).

The photoreceptor voltage noise (Fig. 4 B) increases with the mean light intensity until around BG-3 or BG-2, showing some cell to cell variability (Fig. 4 F), initially exceeding the corresponding signal, before rapidly diminishing at bright adapting backgrounds, BG-1 and BG0. The variance and power spectrum of the voltage noise in a single photoreceptor behaves alike whether the cell is stimulated only with a constant light background or with a Gaussian contrast stimulus superimposed on it (Fig. 4 B and Fig. 3 C are from the same cell; the thorough examination of the noise power spectra is shown later in Fig. 8). The probability distribution of the voltage noise is positively skewed (Fig. 4 C, dotted line) under dim light conditions, most likely because of infrequent photon absorption, seen as bursts of responses rising from near dark-adapted potentials, but is Gaussian at brighter backgrounds, where the noise is dominated by small, but numerous bumps (see later Bump Noise Analysis).

Because the photoreceptor voltage response to the contrast stimulus increases with the adapting light intensity while the noise decreases, the signal-to-noise ratio (Fig. 4 G), $\text{SNR}_V$, calculated by dividing the signal variance by the corresponding noise variance, improves in the different investigated photoreceptors between 30 to 90 times with intensifying light adaptation. As previously reported in larger flies (Howard et al., 1987; Anderson and Laughlin, 2000) the increase in $\text{SNR}_V$ is roughly proportional to the square root of intensity, which is consistent with a photon noise-limited Poisson process. However, at the highest intensities the $\text{SNR}_V$ flattens, presumably because of biological constraints such as the restricted number of transduction units, attenuation by the intracellular pupil (Howard et al., 1987), and the saturating speed of the phototransduction reactions (see also Juusola and Hardie, 2001, in this issue).

The Signal and Noise Dynamics in the Frequency Domain
To see how the frequency content of the photoreceptor voltage signal and noise changes during light adaptation, the signal and noise power spectrum, and their derivatives (signal-to-noise ratio and information capacity) were compared at different adapting backgrounds.

Fig. 5 A illustrates the light adaptational changes in the photoreceptor signal power spectrum, $|S_V(f)|^2$. Under dim light conditions, most of the signal power occurs at low frequencies, but brightening the adapting background shifts the power towards high frequencies and attenuates its low frequency end. The shape of the corresponding photoreceptor noise power spectrum, $|N_V(f)|^2$ (Fig. 5 B), is dominated by the frequency domain characteristics of the average bump waveform (the elementary response dynamics are explained later in Bump Noise Analysis), but also includes a small contribution of instrumentation noise and channel noise. At dim light conditions (BG-4), $|N_V(f)|^2$ resembles $|S_V(f)|^2$ but has more power. In brighter conditions, the noise power sinks over the whole signal bandwidth and at bright light intensities (from BG-2 to BG0) is less than the signal power over all frequencies from 1 Hz to the steep roll off. The general signal and noise dynamics during light adaptation closely resemble those reported by Juusola et al. (1994) in Calliphora photoreceptors, but are shifted to a much lower frequency range.

The photoreceptor signal-to-noise ratio spectrum, $\text{SNR}_V(f)$, is calculated by dividing the signal power spectrum by the noise power spectrum. The photoreceptor performance improves with increasing mean light intensity, with the bandwidth of high $\text{SNR}_V(f)$ (Fig. 5 C) and information, $H$ (Fig. 5 D), progressively shifted towards high frequencies. As light adaptation expands the bandwidth of reliable signaling, the average information capacity increases from ~30 bits/s at the background of BG-4 to ~200 bits/s at BG0 (Fig. 5 E). At the brightest adapting background, the average information capacity hence is ~0.2 times that measured by de Ruyter van Steveninck and Laughlin (1996a) at 20–22°C in Calliphora photoreceptors under similar illumination conditions, which is consistent with the suggestion that Drosophila processes visual information more slowly than the fast-flying flies (Skingsley et al., 1995; Weckström and Laughlin, 1995).

Bump Noise Analysis
$|N_V(f)|^2$ contains information about the average waveform of discrete voltage events caused by the single photon absorptions, i.e., quantum bumps (compare with Wong and Knight, 1980). To reveal how the average bump shape changes with light adaptation, the photoreceptor noise power spectrum at different adapting backgrounds was analyzed as follows.

We assume that the measured voltage noise of light-adapted photoreceptors contains light-induced noise and instrumental as well as intrinsic noise, which are independent and additive. Hence, by subtracting the...
photoreceptor noise power spectrum estimated in darkness, $\langle |N_v(f)|^2 \rangle$, from the photoreceptor noise power spectra at different adapting backgrounds, $|\langle N_v(f) \rangle|^2$, we can estimate the light-induced voltage noise power, $|\langle B_v(f) \rangle|^2$, at the different mean light intensity levels (Fig. 5 F):

$$ |B_v(f)|^2 \equiv |N_v(f)|^2 - |N^D_v(f)|^2. \quad (14) $$

From this voltage noise power the effective bump duration ($T$) can be calculated (Dodge et al., 1968; Wong and Knight, 1980; Juusola et al., 1994), assuming that the shape of the bump function, $b_v(t)$ (Fig. 5 G), is proportional to the $\Gamma$-distribution:

$$ b_v(t) \propto \Gamma_v(t;n,\tau) = \frac{1}{n!\tau^n} \left( \frac{t}{\tau} \right)^n e^{-t/\tau}. \quad (15) $$

The two parameters $n$ and $\tau$ can be obtained by fitting a single Lorentzian to the experimental power spectrum of the bump voltage noise (Fig. 4 F):

$$ |B_v(f)|^2 \propto |\tilde{\Gamma}_v(f;n,\tau)|^2 = [1 + (2\pi f)^2\tau^2]^{-(n+1)}, \quad (16) $$

where $\tilde{\Gamma}$ indicates the Fourier transform. The effective bump duration, $T$ (i.e., the duration of a square pulse with the same power), is then:

$$ T = \tau \frac{(n!)^2 2^{2n+1}}{(2n)!}. \quad (17) $$
Fig. 5 H shows how light adaptation reduces the bump duration from an average of 50 ms at the adapting background of BG-4 to ~10 ms at BG0. The mean bump amplitude (α) and the bump rate (λ) are estimated with a classic technique for extracting rate and amplitude information from a Poisson shot noise process called Campbell’s theorem. The bump amplitude is as follows (Wong and Knight, 1980):

$$\alpha = \frac{\sigma^2}{\mu}. \quad (18)$$

Consequently, this means that the amplitude-scaled bump waveform (Fig. 5 G) shrinks dramatically with increasing adapting background. This data is used later to calculate how light adaptation influences the bump latency distribution. The bump rate, λ (Fig. 5 I), is as follows (Wong and Knight, 1980):

$$\lambda = \frac{\mu^2}{\sigma^2} \cdot T. \quad (19)$$

In dim light conditions, the estimated effective bump rate is in good agreement with the expected bump rate (extrapolated from the average bump counting at BG-5 and BG-4.5; data not shown), namely 265 bumps/s vs. 200 bumps/s, respectively, at BG-4 (Fig. 5 I). However, the estimated rate falls short of the expected rate at the brightest adapting background (BG0), possibly because of the increased activation of the intracellular pupil mechanism (Franceschini and Kirschfeld, 1976), which in larger flies (compare with Lucilia; Howard et al., 1987; Roebroek and Stavenga, 1990) limits the maximum intensity of the light flux that enters the photoreceptor.

**Frequency Response Analysis**

Because the shape of photoreceptor signal power spectra, $[S(f)]^2$ (i.e., a frequency domain presentation of the average summation of many simultaneous bumps), differs from that of the corresponding bump noise power spectra, $[KB(f)]^2$ (i.e., a frequency domain presentation of the average single bump), the photoreceptor voltage signal contains additional information that is not present in the minimum phase presentation of the bump waveform, $|F_V(f)|$ (in this model, the bump starts to arise at the moment of the photon capture; Wong et al., 1980). This information, which includes the bump latency distribution and possible dynamic nonlinearities in light adaptation, can be extracted by calculating the photoreceptor frequency response, $T_V(f)$, and coherence, $\gamma^2(f)$, functions at different mean light intensity levels.

The gain part of the frequency response function, $G_V(f)$ (Fig. 6 A), resembles the corresponding signal power spectrum (Fig. 5 A) at the same adapting background, indicating that the photoreceptor is operating linearly. As the photoreceptor signal shows increased contrast gain and broadened bandwidth with increasing mean light intensity, its 3-dB cut-off frequency (the point at which the gain falls to half of the maximum) shifts towards higher frequencies (Fig. 6 B) saturating on average ~25 Hz at the brightest adapting background.

The corresponding phase, $\phi_V(f)$ (Fig. 6 C), shows that the voltage signal lags the stimulus less as the mean light intensity increases. Furthermore, by comparing $P_V(f)$ to the minimum phase, $P_{min}(f)$ (Fig. 6 C), derived from the gain part of the frequency response function, it becomes obvious that the photoreceptor voltage signals contain a pure time delay. This pure time delay, i.e., dead-time (Fig. 6 D), depends on the mean light intensity. It is largest (~25 ms) at the dimmest adapting background of BG-4 and exponentially reduces to ~10 ms at BG0. Similar adaptive dead-times have been observed in Calliphora photoreceptors (Juusola et al., 1994; de Ruyter van Steveninck and Laughlin, 1996b), but with twice as fast dynamics as in the Drosophila eye.

The coherence function, $\gamma^2_{\text{exp}}(f)$ (Fig. 6 E), an index of the system’s linearity, is close to unity over the frequency range at BG0, indicating that the photoreceptor signals are approximately linear under these conditions. The low coherence values at low mean intensity levels are largely a result of the noisiness of the signal estimates when the rate of photon absorptions is low, since the coherence improves with increased averaging or selecting more sensitive photoreceptors. However, because the photoreceptor signal bandwidth is narrow at low adapting backgrounds, the coherence values are already near zero at relatively low stimulus frequencies. The high degree of linearity at bright illumination, as seen in the coherence, indicates that the skewed distribution of the signals causes a small nonlinear effect on the signal amplification during dynamic stimulation. A similar behavior has been encountered in the blowfly (Calliphora) photoreceptors (Juusola et al., 1994). There, it was later shown that adding a nonlinearity (second-order kernel or static polynomial component) into a dynamic linear photoreceptor model (linear impulse response) causes no real improvement as judged by the mean square error (Juusola et al., 1995).

When a photoreceptor operates as a linear system, one can calculate the coherence function from the SNR$_V(f)$. As shown above (Fig. 4), at low adapting backgrounds, the photoreceptor voltage responses are small and noisy. Accordingly their linear coherence estimates, $\gamma^2_{\text{SNR}}(f)$ (Fig. 6 F), are significantly lower than the coherence, $\gamma^2_{\text{exp}}(f)$ (Fig. 6 E), calculated from the signal (i.e., the averaged voltage response). At the brightest adapting backgrounds, the photoreceptor voltage responses are highly reproducible, having significantly reduced noise content. The discrepancy between the two independent coherence estimates is then much less. From such a comparison, one can deduce,
for example, that the drop in the low frequency coherence is a consequence of both the significant low frequency noise content and the speed of adaptation (a dynamic nonlinearity), which progressively reduces the gain of the low frequency voltage responses, as the photoreceptor adapts to higher mean light intensity levels.

The linear impulse response, \( k_v(t) \), defined as the photoreceptor voltage responses to a pulse of unit contrast given at various backgrounds, was calculated from the same data (Fig. 6 G). Its amplitude increases with the mean light intensity, appearing to saturate at the adapting backgrounds above BG-2, whereas its latency and total duration are reduced. The time to peak of the impulse response \( t_p \) is halved from \(~40\) ms measured at the lowest mean light intensity to \(~20\) ms at the brightest adapting background (Fig. 6 H). Also, the rise time of the impulse response decreases with the increase in the adapting background.

**Bump Latency Distribution**

Because of the dead-time and the variance in timing of individual bumps, the shape and the time course of the impulse response and the average bump are different. These timing irregularities form the bump latency distribution, which can be estimated accurately from the existing data at different adapting backgrounds (see also Henderson et al., 2000, who describe the bump dynamics in dark-adapted photoreceptors). The adapting
bump model (Wong et al., 1980), assuming that the processes are linear, states that convolving the bump waveform, \( b(t) \), measured at a certain light intensity level, by its corresponding latency distribution, \( l(t) \), produces the photoreceptor impulse response, \( k_V(t) \):

\[
  k_V(t) = b_V(t) \otimes l(t),
\]

(20)

where \( \otimes \) denotes convolution. Above, we have calculated the linear impulse responses (Fig. 6 G) and estimated the corresponding bump waveforms (Fig. 5 G) of individual photoreceptors at different adapting backgrounds. Thus, the bump latency distributions can be reconstructed by removing, or deconvolving, the bump waveforms from the impulse responses. To minimize the effects of voltage noise on the recordings, the bump latency distributions were first calculated by using fitted expressions for both the impulse response and bump waveform data. The normalized photoreceptor impulse response, \( k_{V,norm}(t) \) is well fitted by a log-normal function, (Payne and Howard, 1981):

\[
  k_{V,norm}(t) \equiv \exp \left\{ \frac{-(\ln(t/t_p))^2}{2a^2} \right\},
\]

(21)

where \( t_p \) is the time to peak of the impulse response, and \( a \) is the width factor. Fig. 7 A shows typical log-normal approximations of the photoreceptor impulse responses (Fig. 6 G) and the corresponding bump latency distributions at different mean light intensity levels. (D) The normalized bump latency distributions (as seen in C). Additionally, these were calculated from the voltage and light recordings as explained in Eq. 22 (E) and Eqs. 23 and 24 (F).

**Figure 7.** The bump latency distribution stays relatively unchanged at different adapting backgrounds. Removing the bump shape from the corresponding impulse response by deconvolution reveals the bump latency distribution. (A) The log-normal approximations of the photoreceptor impulse responses. (B) The normalized \( \Gamma(t) \) distribution fits of the bump shape; and (C) the corresponding bump latency distributions at different mean light intensity levels. (D) The normalized bump latency distributions (as seen in C). Additionally, these were calculated from the voltage and light recordings as explained in Eq. 22 (E) and Eqs. 23 and 24 (F).

The bump latencies appear to have a rather similar distribution at different adapting backgrounds. This becomes more obvious when the latency distributions are normalized (Fig. 7 D). According to these estimates, apart from the lowest adapting background, where the original photoreceptor data is too noisy to provide accurate results, the first bump starts to appear \( \sim 10 \) ms after the flash with a peak in the distribution \( \sim 8 \) ms later. The peak and the width of these latency distribution estimates vary relatively little, suggesting that the general shape of the bump latency distribution was maintained at different adapting backgrounds. Because the fitted expressions could only estimate the true bump and impulse waveforms, these findings were further checked against the latency distributions calculated from the raw data using two different techniques described below.

Fig. 7 E shows normalized bump latency distributions at different adapting backgrounds calculated by first dividing the photoreceptor frequency response, \( T_V(f) \), by the corresponding photoreceptor noise spectrum, \( |N_V(f)| \), and taking the inverse Fourier transformation of this product:

\[
  l(t) = \mathcal{F}^{-1} \left[ \frac{T_V(f)}{|N_V(f)|} \right] \equiv \mathcal{F}^{-1} \left[ \frac{T_V(f)}{N_V(f)} \right].
\]

(22)
This approximation is justified because the bump noise clearly dominates the photoreceptor noise, as was shown by the noise power spectra in the Fig. 5 B. Furthermore $F^{-1}[[B_V(f)]]$ provides a minimum phase representation of $b_V(t)$ (Wong and Knight, 1980). Here, the shape of the bump latency distribution was free of any systematic error relating to the data fitting, but was influenced by the low level of instrumental noise remaining in the noise spectra. The noisy data at the lowest adapting background did not allow a reasonable estimate of the latency distribution, and this trace was not normalized. Since these estimates closely resemble those of the other methods, any transducer noise and instrumental noise in $[S_N(f)]$ could only have had a marginal effect on the calculations.

Another way to calculate the bump latency distribution is shown in Fig. 7 F. First, the estimated $\Gamma_V(t)$-bump waveform (Fig. 7 B) was deconvolved from the actual 10–30 nonaveraged traces of the recorded voltage response data, $r_V(t)$, to produce corresponding timing trails, $d_V(t)$, of the bump events:

$$r_V(t) = \Gamma_V(t) \otimes d_V(t).$$  \hspace{1cm} (23)

Then the impulse, $l(t)$, calculated between the corresponding contrast stimulus and the bump timing cross-spectrum, is the bump latency distribution (see Eqs. 8 and 12):

$$l(t) = F^{-1}\left[\frac{\langle D_V(f) \cdot C^*(f) \rangle}{\langle C(f) \cdot C^*(f) \rangle}\right].$$  \hspace{1cm} (24)

Once again the bump latency distribution estimates (Fig. 7 F) showed relatively small differences from one light intensity level to another, being in line with the other estimates. Again, the data at the lowest mean light were too noisy for a reasonable estimate.

**IV: Photoreceptor Membrane during Natural-like Stimulation**

In *Drosophila* and many other insect photoreceptors, the interplay between the opening and closing of light channels (Trp and Trpl) and voltage-sensitive ion channels (for $K^+$ and $Ca^{2+}$) shapes the voltage responses to light. The more open channels there are at one moment on a cell membrane, the lower its impedance, the lower the voltage responses. On the other hand, when the mean light intensity is increased, the contrast evoked $s_V(t)$ increases (Fig. 8 B). This is due to the logarithmic increase in the bump number, although the average size of bumps is reduced. During both the current and light contrast stimulation, $n^{l}_{V}(t)$ and $n^{c}_{V}(t)$ were about the same size and behaved very much in the same way, getting smaller with increasing mean light intensity (Fig. 8 A and B).

Fig. 8 C shows typical probability distributions of voltage signals to dynamic contrast stimulation and current injection at the three selected adapting backgrounds. Since the current injection produced signals that had always purely Gaussian distributions (scattered squares fitted with Gaussian; $n = 15$), the skewness seen in the corresponding light contrast—evoked signals (filled histograms) is unlikely to have originated from voltage-dependent ion channels on the membrane (delayed rectifier and A-type potassium channels; Hardie, 1991b), but presumably mirrors some earlier asymmetry in the phototransduction cascade’s response to light increments and decrements.

Since the process of driving the photoreceptor voltage with dynamic stimulation may itself add or reduce noise and nonlinearities to the signaling (as reported in spider mechanoreceptors by Juuola and French, 1997), we checked the measured photoreceptor voltage noise during dynamic stimulation against that during constant light stimulation. No such discrepancy is found here. The photoreceptor voltage noise power spectra, evoked either by a specified mean light background solely, i.e., $[N_V(f)]^2$; a dynamic light contrast superimposed on the same light background, $[\langle N_{V}^c(f) \rangle]^2$;
or a pseudorandom current injection at the same light background, $|\langle N_{v}(f) \rangle|^2$, are remarkably similar (Fig. 8 D). Since the shape of the noise power spectra changes with the increasing mean light intensity (Fig. 5 B), this indicates that the photoreceptor voltage noise is dominated by the bump noise.

**Light Adaptation Accelerates the Dynamics of Both Light Responses and Photoreceptor Membrane**

To establish how the signal conduction properties of the photoreceptor membrane compare to the speed of the light contrast-evoked voltage responses, the membrane impedance, $Z(f)$, and the corresponding light frequency response, $T_v(f)$, were calculated at different adapting backgrounds from the previous data. The photoreceptor membrane impedance function (Fig. 9 A) is reduced at brighter backgrounds, covers a broader frequency band, and lags the stimulus slightly less (Fig. 9 C) than at dim illumination. This is of course due to the increasing depolarization, or rather the increased number of open light- and voltage-sensitive ion channels, lowering both the membrane resistance and its time constant, thus, allowing faster signaling. This is further illustrated in Fig. 9 B, which shows how the 3-dB cut-off frequency of the photoreceptor membrane impedance increases with the mean light intensity. In darkness, the mean cut-off frequency of the membrane is $45 \pm 36$ Hz ($n = 5$) and shifts to $145 \pm 32$ Hz ($n = 6$) at $3 \times 10^6$ photons/s.

The photoreceptor membrane operates linearly when converting the Gaussian current modulation into voltage responses. The membrane coherence function,
calculated between the current stimulus and the corresponding voltage signal (Fig. 9 D), approaches unity (from 0.84 to 0.9999), being lowest at frequencies below 5 Hz. On the other hand, the coherence estimates, which are calculated from the photoreceptor membrane signal-to-noise ratios (Fig. 9 E; see also Fig. 2 B, g), are affected by the corresponding photon noise of each particular adapting background, but still maintain values close to unity indicating highly linear operation.

Fig. 9 (F and G) compares the gain parts of the light contrast- and current injection-induced photoreceptor frequency responses measured at different mean light intensities. The traces are normalized to show how the increased light adaptation speeds up the response dynamics. Light adaptation accelerates the dynamics of both the light responses and membrane impedance, as seen by their expanding frequency ranges. However, the bandwidth of the photoreceptor membrane always extends further than that of the corresponding light response. This suggests that during normal diurnal light conditions, the speed of the chemical reactions in the phototransduction cascade is not limited by the electrical properties of the light-adapted photoreceptor membrane. To demonstrate this more directly, we attempted to derive the light-induced current by deconvolving the effect of the membrane impedance.

Adaptive Filtering by Phototransduction Cascade and Membrane Improves Signaling

Because light-adapted photoreceptors respond to both dynamic contrast and current stimulation with linear voltage responses, we can derive the output of the phototransduction cascade, i.e., the light current by deconvolving the membrane impedance impulse response, \( z(t) \) (Fig. 2 C, d), from photoreceptor voltage responses.

**Figure 9.** The photoreceptor membrane characteristics at different light adaptation levels. The photoreceptor impedance, \( Z(f) \), calculated from the current injection and the resulting voltage responses, is reduced (A, gain), it is accelerated (B, 3-dB cut-off frequency) and it lags the stimulus less (C, phase) when it is shifted towards higher frequencies with increasing light adaptation. Regardless of the adapting background the membrane operates linearly over the studied frequency range (in D, \( \gamma^2_{\text{resp}}(f) \) and in E, \( \gamma^2_{\text{SNR}}(f) \); coherence close to unity). Both the normalized impedance (F) and the gain of the contrast-induced voltage responses (i.e., light response), (G) demonstrate a gradual shift of their bandwidth towards high frequencies. The cut-off frequency of the impedance is always much higher than that of the light responses in the same photoreceptor at the same adapting background; in this particular photoreceptor, 3.1, 1.9, 3.2, 5.1, and 4.2 times higher, when going from BG-4 to BG0.
to light contrast stimulation, measured in the
same cell at the same mean light:

\[ r_v(t) = r_d(t) \otimes z(t). \] (25)

The light current frequency response, \( T_I(f) \), is then calculated between the contrast stimulus, \( c(t) \), and the current signal, \( s_I(t) \) (i.e., the mean \( \langle r_I(t) \rangle \)). Fig. 10 (A–C) shows the normalized gain parts of the photoreceptor impedance \( (Z(f)) \), light-current \( (G_I(f)) \), and voltage response \( (G_v(f)) \) frequency responses at three different mean light intensities. The high impedance photoreceptor membrane acts as a low-pass filter for the phototransduction signal, effectively filtering the high frequency content of the light current, which might also include high frequency ion channel noise. This inevitably makes the voltage response slightly slower than the corresponding light current. The membrane dynamics speeds progressively when the mean light increases, so that its cut-off frequency is always much higher than that of the light current, and only under the dimmest (Fig. 10 A) conditions does the membrane significantly limit the frequency response of the voltage signal. Furthermore, the high mean impedance in dim light conditions causes small changes in the light current to charge relatively larger voltage responses than those under brighter conditions as seen in the corresponding voltage, \( k_v(t) \), and light current, \( k_I(t) \), impulse responses (Fig. 10 D).

To establish how effectively the photoreceptor membrane filters the transduction noise, we calculated the phototransduction bump noise by removing (deconvolving) the photoreceptor impedance, \( Z(f) \) from the \( \Gamma \)-distribution estimate of the normalized bump voltage noise spectrum, \( |\Gamma_V(f)| \), measured at the same mean light intensity level:

\[ |B_v(f)| = \frac{|B_r(f)|}{Z(f)} = \frac{|\Gamma_V(f)|}{Z(f)} = |\Gamma(f)|. \] (26)

Fig. 10 (E–G) compares the normalized photoreceptor impedance to the corresponding normalized spectra of the phototransduction bump noise, \( |\Gamma(f)| \), which now presents the minimum phase shape of the elementary transduction event, i.e., light-current bump, at three different adapting backgrounds. Although the membrane impedance’s cut-off frequency is much higher than the corresponding light current signal, \( G_I(f) \), at all light intensity levels, the corresponding phototransduction bump noise spectrum, \( |\Gamma_I(f)| \), and membrane impedance, \( Z(f) \), show considerable overlap. These findings indicated that the transfer characteristics of the photoreceptor membrane serve a dual function. By tuning to the mean light intensity levels, the photoreceptor membrane provides a fast conduction path to the phototransduction signal and concurrently; and improves the reproducibility of the photoreceptor voltage responses by removing the high frequency noise in the light current, associated with the shortening of the bump duration (compare with Fig. 5 H).

**DISCUSSION**

The results presented here characterize the light adaptation dynamics of *Drosophila* photoreceptors in unprecedented detail. The experiments, in which photoreceptor voltage was modulated with dynamic contrast and current stimuli at various mean light intensity levels, allowed us to quantify the increase in signaling efficiency with light adaptation and demonstrate that it is the product of the following three factors: (1) bump compression of several orders of magnitude; (2) fast but relatively constant bump timing; and (3) accelerated membrane dynamics. In the following discussion, we attempt to relate these findings to the current ideas of the *Drosophila* phototransduction and then go on to consider their functional significance in terms of coding strategies.

Phototransduction in *Drosophila* occurs through a phospholipase C–mediated and Ca\(^{2+}\)-regulated reaction cascade, where light-activated rhodopsin leads to sequential activation of a heterotrimeric G-protein and eye-specific PLC. This hydrolyzes the minor membrane phospholipid, PtdIns(4,5)P\(_2\) to generate InsP\(_3\) and diacylglycerol (DAG) leading finally to the opening of the light-sensitive channels. Although a large number of fly phototransduction mutants have been identified and analyzed, much of the molecular interactions inside the pathway are incompletely understood. In particular, events linking activation of PLC to the opening of the light-sensitive channels still remain enigmatic: recent evidence suggests that InsP\(_3\) is not involved (Acharya et al., 1997; Hardie and Raghu, 1998; Raghu et al., 2000), suggesting that PtdIns(4,5)P\(_2\), DAG, or its metabolites may be the key messenger(s) of excitation (Chyb et al., 1999; Raghu et al., 2000). However, there is general consensus that events determining the response latency occur largely at or upstream of PLC, whereas events determining bump shape and amplitude are generated downstream of PLC. This is because hypomorphic mutations of PLC and G-protein affect quantum bump latency but not bump shape (Pak et al., 1976; Scott et al., 1995; Scott and Zuker, 1998). Recently, it has been demonstrated that many proteins central to the phototransduction cascade are coupled into a supramolecular signaling complex, through association with the PDZ-domain scaffold protein, INAD, including PLC and at least one of the two types of light-sensitive channel, (Trp) (Shieh and Zhu, 1996; Huber et al., 1996; Tsunoda et al., 1997; Scott and Zuker, 1998; Montell, 1999). It has been speculated that this may minimize diffusional delays involved in excitation (Scott...
and Zuker, 1998; Montell, 1999), though experimental verification of this proposal is lacking.

I: Light Adaptation Affects the Bump Waveform but Has a Relatively Small Influence on Bump Timing

It is widely accepted that the steady-state photoreceptor potential represents a summation of single photon-evoked elementary voltage responses, i.e., quantum bumps. In line with previous formulations (compare with Wong and Knight, 1980) we applied shot noise analysis to reconstruct the bump event rate, effective duration and mean amplitude from the photoreceptor voltage noise at different adapting backgrounds. In agreement with the earlier Drosophila studies (Wu and Pak, 1978) we found that the average size of the bumps decreases markedly and their time course becomes briefer as the mean light is increased. Since bump rate rose linearly with intensity up to intensities of BG-2 or -1, this means that, in keeping with other preparations, the reduction in bump amplitude and duration is the major process underlying the reduction in the absolute gain associated with adaptation, which is therefore presumably mediated downstream of PLC. The overall bump amplitude, estimated via noise analysis in combination with Campbell’s theorem, decreases $\sim 50$–100-fold over the measured range of intensities, which does not in-

![Figure 10. General comparison of the transduction signal and noise and membrane bandwidth at different adapting backgrounds. (A–C) The dynamics of the corresponding light current, voltage response, and membrane impedance displayed as their normalized gain. (A) Under dim conditions, the light current is noisy and the low-passing membrane removes the high frequency noise, producing slow voltage responses to light contrasts. When the mean light intensity is increased, both the transduction cascade and photoreceptor membrane allows faster signaling leading to accelerated voltage responses (B and C). The corresponding impulse responses (D), calculated from the same data, show how the light current and voltage responses quicken with increasing mean light intensity, but the light current is always peaking before its respective voltage response. Because of the large membrane impedance under dim conditions, the small light currents can charge relatively large voltage responses. The responses are normalized by the maximum value of each series. (E–G) The transduction bump noise, $\tilde{F}_I(f)$, was calculated by deconvolving the photoreceptor membrane impedance, $\tilde{Z}(f)$, from the respective voltage bump noise, $\tilde{F}_V(f)$. From dim light conditions (E and F) to the bright adapting backgrounds (G) $\tilde{F}_I(f)$ shows a considerable overlap with the corresponding membrane impedance.](image-url)
Role of Ca\textsuperscript{2+} as Messenger of Adaptation

Numerous studies have shown that calcium is the major mediator of adaptation in invertebrate and vertebrate photoreceptors (for reviews see Hardie and Minke 1995; Montell, 1999; Pugh et al., 1999). It is the obvious candidate for regulating bump shape and size as well as the modest changes in latency. Indeed, a recent study showed that Drosophila bump waveform and latency were both profoundly, but independently, modulated by changing extracellular Ca\textsuperscript{2+} (Henderson et al., 2000). In Drosophila, the vast majority, if not all, of the light-induced Ca\textsuperscript{2+} rise is due to influx through the highly Ca\textsuperscript{2+} permeable light-sensitive channels (Peretz et al., 1994; Ranganathan et al., 1994; Hardie, 1996; but see Cook and Minke, 1999). Recently, Oberwinkler and Stavenga (1999, 2000) estimated that the calcium transients inside microvilli of blowfly photoreceptors reached values in excess of 100 µM, which then rapidly (<100 ms) declined to a lower steady state, probably in the 10–50-µM range; similar steady-state values have been measured in Drosophila photoreceptor cell bodies after intense illumination (Hardie, 1996). Hardie (1991a; 1995a) demonstrated that Ca\textsuperscript{2+} mediated a positive, facilitatory Ca\textsuperscript{2+} feedback on the light current, followed by a negative feedback, which reduced the calcium influx through light-sensitive channels. Stieve and co-workers (1986) proposed that in Limulus photoreceptors, a similar type of Ca\textsuperscript{2+}-dependent cooperativity at light-sensitive channels is responsible for the high early gain. Caged Ca\textsuperscript{2+} experiments in Drosophila have demonstrated that the positive and negative feedback effects both take place on a millisecond time scale, suggesting that they may be mediated by direct interactions with the channels (Hardie, 1995b), possibly via Ca\textsuperscript{2+}-calmodulin, CaM, as both Trp and Trpl channel proteins contain consensus CaM binding motifs (Phillips et al., 1992; Chevesich et al., 1997). Another potential mechanism includes phosphorylation of the channel protein(s) by Ca\textsuperscript{2+}-dependent protein kinase C (Huber et al., 1996) since null PKC mutants show defects in bump termination and are unable to light adapt in the normal manner (Ranganathan et al., 1991; Smith et al., 1991; Hardie et al., 1993). However, until the identity of the final messenger of excitation is known, it would be premature to conclude that these are the only, or even major, mechanisms by which Ca\textsuperscript{2+} affects the light-sensitive conductance.

II: The Photoreceptor Membrane Does Not Limit the Speed of the Phototransduction Cascade

To characterize how the dynamic membrane properties were adjusted to cope with the light adaptational changes in signal and noise, we deconvolved the membrane from the contrast-induced voltage signal and noise data to reveal the corresponding phototransduction currents. This allowed us to compare directly the spectral properties of the light current signal and noise to the corresponding membrane impedance. At all adapting backgrounds, we found that the cut-off frequency of the photoreceptor membrane greatly exceeds that of the light current signal. Therefore, the speed of the phototransduction reactions, and not the membrane time constant, limits the speed of the resulting voltage responses. By contrast, we found a close spectral overlap between the high frequency roll-off of
the light current noise and the membrane impedance at all the tested light intensity levels. Overall, these results suggest that the increase in the signaling bandwidth of the photoreceptor membrane with the mean light intensity functions to accommodate any acceleration in the phototransduction kinetics while simultaneously filtering the increasing high frequency phototransduction noise. Suppression of photon shot noise by membrane filtering also has been reported in photoreceptors of the crane fly *Tipula* (Laughlin, 1996); however, the strategy there appeared rather different since the membrane in *Tipula* also significantly limits the frequency range of the light current signal.

These adaptive membrane dynamics result from the dynamic interaction between a variety of light- and voltage-sensitive ion channels. As previously described, *Drosophila* photoreceptors express at least three different voltage-sensitive potassium channels (slow delayed rectifier conductance \[I_{Ks}\], fast transient A-current \[I_A\], and a delayed rectifier with intermediate kinetics \[I_{Kr}\]), each with different activation and inactivation kinetics (Hevers and Hardie, 1995). In addition, two classes of light-sensitive channels (Trp and Trpl), each with a characteristic voltage dependence, contribute to the overall light-induced lowering of the membrane impedance. Specific channel mutants will have the potential to analyze the functional roles of such individual channel species in detail.

The idea of matching the dynamic membrane properties by voltage-sensitive ion channels to natural signal conditions is not new and has been explored in both fast and slow flying insects (Laughlin and Weckström, 1993; Weckström and Laughlin, 1995; Laughlin, 1996). However, our study was unique in the sense that we could derive an accurate representation of the transduction current dynamics and correlate these with the membrane dynamics in the same photoreceptor. The data are also the first to show that the skewness of the photoreceptor voltage responses to Gaussian contrast stimulation at bright adapting backgrounds is not caused by the voltage-sensitive membrane, but reflects either the opening dynamics of the light-sensitive channels or some compressive nonlinearity early in the phototransduction cascade. As the skewness of the responses mirrors the skewness of the contrast distribution in natural scenery (Laughlin, 1981), it should be beneficial to implement this feature in the early transduction rather than in later signal shaping to keep the coding machinery as energetically efficient as possible.

**III: The Photoreceptor Signaling Works Competently within the Imposed Physical Limits**

The photoreceptor responses are a product of individual bump waveforms and their timing, i.e., the bump latency distribution. In near darkness and in dim light conditions, photoreceptors are adapted to processing signals of low signal-to-noise ratio, where the sparse and random arrival of photons restricts the signal fidelity. The enzymatic reactions transduce and amplify the single photon absorptions into voltage fluctuations, which vary in their size and timing, but can be separated reliably as discrete events. As pointed out by many (see van Hateren, 1992), the general coding task here appears to be to detect and count the photons rather than to characterize the light stimulus. Applying the classical Shannon expression (Eq. 5), we can make approximate calculations of the information content of the light stimulus at certain intensity levels knowing that the light itself is a Poisson process having a defined SNR\(_T\) = \(\bar{N}\) at all stimulus frequencies, and limiting the bandwidth to cover the photoreceptor’s operational range (see Eq. 27). This allows us to compare the photoreceptor’s information capacity estimates at a certain mean light intensity (\(Y\)) to the theoretical maximum over the bandwidth of the photoreceptor’s operation:

\[
H = W \cdot \log_2 \frac{s_v + n_v}{n_v},
\]  
(27)

where \(s_v\) and \(n_v\) are photoreceptor voltage signal and noise variance over the bandwidth, \(W\) (Shannon, 1948). Or similarly for the light stimulus:

\[
H = W \cdot \log_2 (\text{SNR}_T + 1) = W \cdot \log_2 (\bar{N} + 1) \tag{28}
\]

Because the adapting background of BG-4 contained \(~300\) photons/s, we have \(\log_2 (\sqrt{300} + 1) = 4.2\) bits distributed over the photoreceptor signal bandwidth, say 70 Hz (Fig. 5 A). The information content is 294 bits/s, indicating that every counted photon carries a bit. However, with light adaptation, the photoreceptor is shifting from counting photons to integrating them into a neural image. The irregular arrival of photons makes the neural integration noisy, and the estimated photoreceptor information capacity from the average photoreceptor SNR\(_T\) of 0.152 (Fig. 4 G) gives 14 bits/s. This is close to the photoreceptor information capacity calculated between the signal and noise power spectra at the same adapting background (Fig. 5 E, which varied from 15 to 34 bits/s). Whereas at the bright adapting background of BG0, the estimated LED output was \(3 \times 10^5\) photons/s. Yet, the photoreceptors could only detect a tenth of them (possibly because of the activated pupil mechanism; Fig. 5 I). This gives the information content for BG0:

\[
\log_2 (\sqrt{3 \times 10^7}) \cdot 70 = 1274\ \text{bits/s}.
\]

Again, from the corresponding mean photoreceptor SNR\(_T\) of 7.7, we have \(\log_2 [8.7] \cdot 70 = 218\ \text{bits/s},\) close to
the measured average of 216 bits/s (Fig. 5 E). This simple comparison between the information content of the light stimulus and the corresponding information capacity of the *Drosophila* photoreceptors suggests that the efficiency to code light information into a neural signal increases with the adapting background: from 5% under dim conditions to 17% during bright illumination.

Because imprecision either in the bump timing or summation can smear the voltage responses, any variability in one of these processes reduces the photoreceptor information capacity. It appears that, at low mean light intensity levels, the variability of the signal mostly reflects changes in the bump shape. On the other hand, when the physical limitations imposed by low numbers of photons vanish at brighter adapting backgrounds, the visual coding strategy changes accordingly. When the number of bumps is very large and the bumps themselves very small, the speed of synchronizing a large population of bumps becomes precision limiting. Although the bump shape can in principle be reduced to some extent by intensifying the mean light intensity level, the speed limit imposed by the dead-time in phototransduction prevents the signal bandwidth to grow accordingly. This restricts the time course of the voltage responses and starts to cause saturation of the photoreceptor information capacity at high light intensities.

What is the maximum number of photons that can be processed during intense light adaptation at 25°C? Following Hamdorf (1979), Howard et al. (1987), and Hochstrat and Hamdorf (1990), we assume that each microvillus is a transduction unit, and that a second photon being absorbed by a single microvillus within the dead-time cannot be detected. Either it will have no effect or, at best, it may shorten the latency of the bump as suggested to occur in *Calliphora* by Hamdorf and Kirschfeld (1980). The measured 10-ms dead-time would, therefore, restrict the bump rate to ~100 events/s/microvillus. From anatomical measurements of microvilli, we estimate that each rhabdomere has ~30,000 microvilli. This would mean that the phototransduction machinery saturate at levels of $3 \times 10^9$ absorbed photons/s (as suggested by recordings in the *Drosophila* mutants that lack the screening pigment; Juusola, M., and R.C. Hardie, manuscript in preparation). Here, the maximum rate of photon absorption appeared to saturate at ~0.1–0.2 of this theoretical maximum, at 3–5 $\times 10^6$ photons/s. However, this is almost certainly because of the activation of the intracellular pupill mechanism, which limits the amount of light that is absorbed by the visual pigment. Furthermore, the photoreceptors were clearly not truly saturated in that a natural contrast modulation of 0.22 around this mean photon absorption rate was still translated into unattenuated contrast responses that had a mean information capacity of $216 \pm 61$ bits/s ($n = 14$). The information capacity varied somewhat from one photoreceptor to another and was ~0.2–1/3 of the maximum information transfer rate measured under similar illumination in blowflies (*Calliphora vicina*; de Ruyter van Steveninck and Laughlin, 1996a; Juusola et al., 1996), which have approximately three times more microvilli in their rhabdomere (Hardie, 1985).

Comparison of bump waveform and latency distribution clearly indicates that the latter is the main determinant of the shape of the light-adapted impulse response and, consequently, represents the major constraint on the overall frequency response of the photoreceptor signal. By contrast, the bump duration reaches values of <10 ms. This generates high frequencies, which are negligibly represented in the signal power spectra, thereby allowing the associated stochastic noise to be filtered by the membrane impedance without a significant loss of information. Our current understanding of phototransduction suggests that bump latency is determined by events up to and including activation of PLC and may, for example, represent the time required for the accumulation of a significant amount of second messenger to reach the threshold for channel activation. It is interesting to speculate whether the broad latency distribution is an unavoidable constraint of the stochastic behavior of the underlying biochemical machinery or whether in fact it is designed to provide the photoreceptor with a particular frequency response optimized to its visual ecology and metabolic demands (van Hateren, 1992; Laughlin et al., 1998). Certainly, faster flying flies such as *Calliphora* have evolved significantly faster response kinetics; however, this comes with a price: namely lower membrane impedance and, consequently, greater energetic costs in restoring the ionic equilibria (Laughlin et al., 1998). What factors determine the variability in bump latencies are unknown. However, the relatively long and finite dead-time and skewed Gaussian shape of the latency dispersion exclude a simple first-order stochastic model, which would be expected to generate an exponential distribution of latencies. It is also not known, for example, whether the latency dispersion derives from different microvilli having different characteristic latencies or whether a given microvillus would generate quantum bumps with variable latencies to successive photons. Interestingly, the bump amplitude, which also shows stochastic variations, appears to be much more reproducible than the bump intervals in trains of bumps recorded after activation of one rhodopsin in a single microvillus (Scott and Zuker, 1998).

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