Light Adaptation in Drosophila Photoreceptors: II. Rising Temperature Increases the Bandwidth of Reliable Signaling

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Abstract

It is known that an increase in both the mean light intensity and temperature can speed up photoreceptor signals, but it is not known whether a simultaneous increase of these physical factors enhances information capacity or leads to coding errors. We studied the voltage responses of light-adapted Drosophila photoreceptors in vivo from 15 to 30°C, and found that an increase in temperature accelerated both the phototransduction cascade and photoreceptor membrane dynamics, broadening the bandwidth of reliable signaling with an effective Q10 for information capacity of 6.5. The increased fidelity and reliability of the voltage responses was a result of four factors: (1) an increased rate of elementary response, i.e., quantum bump production; (2) a temperature-dependent acceleration of the early phototransduction reactions causing a quicker and narrower dispersion of bump latencies; (3) a relatively temperature-insensitive light-adapted bump waveform; and (4) a decrease in the time constant of the light-adapted photoreceptor membrane, whose filtering matched the dynamic properties of the phototransduction noise. Because faster neural processing allows faster behavioral responses, this improved performance of Drosophila photoreceptors suggests that a suitably high body temperature offers significant advantages in visual performance.

Key words: vision • retina • information • neural coding • graded potential

Introduction

Animals live in a noisy and changeable environment with a variety of internal noise sources also present in the nervous system. Yet, with adequate stimulus conditions, their responses are usually both reliable and accurate. How robust is the underlying neural code, and can it be influenced by general physical factors like temperature? In the visual system, the flow of information starts from an intricate network of photoreceptors and interneurons, many of which use graded potentials (analogue signals) for their communication. This communication has its limitations even in its initial stages. If photoreceptors cannot code incoming light information into voltage responses larger than their own intrinsic voltage noise, higher order neurons in the brain cannot usefully process the signals they are receiving, and the action potentials they generate will carry little information. Because of their intrinsic noise and limited dynamic range, photoreceptors also face the problem of gain control. They must have a high gain to respond to small light contrasts and, yet, be able to accommodate large stimulus changes associated with changes in mean illumination. The reliability of this complex gain control, which involves both the phototransduction cascade and the cell membrane (including voltage-sensitive conductances), may also be prone to temperature-sensitive changes in the reaction kinetics (Lamb, 1984).

Most animals, including insects and other invertebrates, cannot thermoregulate, but still are active over a wide range of temperatures. Studies in toads have emphasized how cooling can improve performance at absolute threshold by reducing the rate of thermal isomerization of rhodopsin (Aho et al., 1988). However, warming can also enhance performance since this accelerates the response kinetics and, hence, improves temporal resolution (Lamb, 1984). In invertebrate photoreceptors, the time course of both the impulse and elementary responses (i.e., quantum bumps) accelerates with warming (French and Järvilehto, 1978; Wong et al., 1980; Roebroeck et al., 1990; Tatler et al., 2000). Although the response dynamics of invertebrate photoreceptors are sensitive to normal fluctuations in body temperature, the effects of temperature on their signaling and coding efficiency have not been studied thoroughly.

Our previous study of the signaling efficiency in Drosophila photoreceptors (see Juusola and Hardie, 2001, in this issue) provided compelling evidence that the photoreceptor membrane does not restrict the speed of voltage responses, and that there are two separate processes operating in the phototransduction cascade, whose adaptive partnership defines the signaling fidelity of the photoreceptor (namely, the average bump waveform and its timing). The photoreceptor responses are simply a convolution of these processes...
(Wong and Knight, 1980; Henderson et al., 2000). By using a combined noise and signal analysis, we could isolate one from the other and investigate their individual dynamics at different mean light intensity levels. We discovered that most of the adaptive changes in the size of the photoreceptor voltage responses results from a logarithmic reduction in the bump waveform, whereas the speed of the voltage responses is limited by the timing delays in the reactions responsible for the latency dispersion of the bumps. The fact that bump latencies are only marginally affected by light adaptation leads to a direct restriction in the bandwidth of the photoreceptor signals and, hence, the information capacity. The results are consistent with the suggestion that each of the $3 \times 10^8$ rhabdomeral microvilli represents an individual transduction unit (Howard et al., 1987; Hochstrat and Hamdorf, 1990; Postma et al., 1999).

Physical chemistry tells us that the speed of chemical reactions can be increased by increasing the number of effective hits between the molecules taking part in the reaction. This can be achieved by increasing the concentration of the reactants or catalysts, or by increasing the thermal energy. The speed of chemical reactions approximately doubles with a 10°C rise in temperature; expressed as a $Q_{10}$ of 2. Our results from Drosophila photoreceptors at 25°C (see Juusola and Hardie, 2001, in this issue) suggested that the speed of the reactions limiting the bump latency is saturated, thus, implying that the concentrations of the reactants and catalysts cannot be elevated by light adaptation. If this is true, then the only plausible way of improving the signaling fidelity is to increase the photoreceptor temperature. More specifically, if the processes behind the light-induced voltage noise (i.e., the average bump waveform), have a $Q_{10}$ smaller than the processes behind the signal timing dynamics (i.e., bump latency distribution), or the effects of noise are limited by temperature-sensitive filtering, then the information capacity of the photoreceptors will increase with warming.

We have studied the response and membrane properties of light-adapted Drosophila photoreceptors in vivo over a 15°C temperature range using linear signal analysis with natural-like contrast and current stimulation similar to those of our companion paper (see Juusola and Hardie, 2001, in this issue). We found that warming light-adapted Drosophila photoreceptors increases their quantum efficiency and accelerates both the phototransduction cascade and cell membrane dynamics. The bump waveform, which is primarily responsible for noise processes on the cell membrane, is relatively temperature insensitive, whereas the bump latency distribution speeds dramatically by warming the photoreceptors. The faster and more precise timing of a larger number of bumps increases the size of the voltage responses to contrast stimuli, hence, expanding the bandwidth of reliable signaling and dramatically increasing the photoreceptor information capacity with a $Q_{10}$ of 6.5.

**Materials and Methods**

Flies, wild-type red-eyed Drosophila melanogaster (Oregon), were taken from a laboratory culture and reared at 25°C. Intracellular voltage responses of green-sensitive R1-6 photoreceptors to light and current stimuli were recorded over a range of temperatures from 15 to 30°C. The recording procedures, light and current stimulation, and data analysis are explained in the companion paper (see Juusola and Hardie, 2001, in this issue).

**Recording Criteria**

Only photoreceptors with saturating impulse responses ($V_{max}$) over 40 mV, minimum input resistance of 100 MΩ, and resting potential in the dark below $-50$ mV were selected for this study. These photoreceptors allowed stable recordings sometimes for several hours and, therefore, were used in a number of different experiments, each of which was repeated with a minimum of three cells, unless stated otherwise. However, we found the recording criteria to be temperature-dependent. In cooler temperatures, the photoreceptors required a significantly longer time to recover from the previous light exposure. Consequently, when a
dark recovery period was kept constant, the light responses of a cool photoreceptor were often smaller than those measured in the same cell in warmer temperatures. The impulse responses were largest in a temperature range from 22 to 27°C (50.2 ± 9.3 mV, n = 30). Altogether, the data were collected from 51 photoreceptors. All the statistics are given as mean ± SD.

R E S U L T S

Impulse Responses of Dark-adapted Photoreceptors at Different Temperatures

The photoreceptor voltage responses to light impulses were first studied after 10–30 min of dark adaptation. Fig. 1 A shows typical voltage responses to a saturating light flash at different temperatures. Warming accelerated the voltage responses, producing an earlier peak. The responses also terminated more rapidly at higher temperatures.

Light Adaptation Is Highly Temperature-sensitive

Fig. 1 B shows typical voltage responses of a photoreceptor to a prolonged light pulse at 28 and 18°C, both recorded after a 2-min dark adaptation period. Soon after the onset of the light pulse, the photoreceptor rapidly depolarizes from the dark-adapted resting potential (around −65 mV) to values close to zero, before declining in a multiphasic fashion to a much lower plateau potential. The decline of the responses from the initial transient to the lower steady-state potential, which is reached after ~10–20 s of continuous illumination, reflects the processes of light adaptation. Not only the timing and the shape of the initial voltage transients, but also the steady-state potential depends on temperature. At the brightest adapting background of 3 × 10^6 photons/s (BG0) the magnitude of the steady-state potential varies from 5–15 mV at 16°C to 20–40 mV at warmer temperatures.

The temperature also greatly affects the voltage responses to contrast steps of light-adapted photoreceptors. Fig. 2 A shows averaged contrast responses of two different photoreceptors recorded at BG0 at 29 and 24°C, respectively. Although light increments (positive contrasts) produce smaller responses than light decrements of the same amplitude, warming increases and accelerates voltage responses to any contrast stimulus. The faster time course of the contrast responses with warming is accompanied by accelerated photoreceptor membrane dynamics. This is shown in Fig. 2 B, which compares current-evoked voltage responses of the same cells at the same temperatures. Small depolarizing current pulses (up to 0.3 nA) produce voltage responses showing strong outward rectification because of the activation of voltage-sensitive potassium channels (Hardie, 1991; Hevers and Hardie, 1995). Hyperpolarizing pulses evoke slower responses, reaching voltages below the dark adaptation resting potential, which resemble RC charging. The time courses of these events are distinctively temperature-sensitive, however.

These findings were confirmed and further characterized in recordings from single light-adapted photoreceptors using more natural, time-dependent light and current stimuli, i.e., white noise-modulated light con-
Light Adaptation in Drosophila Photoreceptors II

In the next paragraph, we show: (I) how the light-adapted signal and noise dynamics of photoreceptor voltage responses change with temperature; (II) how the filtering properties of the photoreceptor membrane are tuned by the temperature; and (III) how the adaptive co-processing by the phototransduction cascade and membrane refines the photoreceptor signaling.

I: Light Adaptation and Photoreceptor Signaling at Different Temperatures

Light-adapted Drosophila photoreceptors produce slightly variable voltage responses to repeated presentations of an identical light contrast stimulus (Juusola and Hardie, 2001). To find out whether this variability depends on the ambient temperature, we recorded photoreceptor responses to a dynamically modulated wideband light contrast stimulus at different temperatures. By applying both signal and shot noise analysis to the data, we could characterize the temperature-induced changes in the photoreceptor signal and noise dynamics and compare the filtering dynamics of the photoreceptor membrane to those of the phototransduction cascade.

In the next paragraph, we show: (I) how the light-adapted signal and noise dynamics of photoreceptor voltage responses change with temperature; (II) how the filtering properties of the photoreceptor membrane are tuned by the temperature; and (III) how the adaptive co-processing by the phototransduction cascade and membrane refines the photoreceptor signaling.

\[ n_V(t) = r_V(t) - s_V(t). \]  

Voltage Responses to Dynamic Contrast Sequences at Bright Background Light

The signal and noise dynamics of light-adapted photoreceptors were studied using a 10-s-long Gaussian contrast stimulus at BG0. The stimulus was spectrally white up to \( \sim 200 \) Hz and had a mean value of 0.32 (Juusola and
Hardie, 2001). Therefore, it both covered the operational frequency range of *Drosophila* photoreceptors and mimicked the light fluctuations in a fly’s natural environment (Laughlin, 1981). Fig. 3 illustrates the key findings from a single photoreceptor at four temperatures (18, 22, 25, and 29°C) with a summary of the corresponding signal and noise statistics from 34 photoreceptors.

The magnitude of the photoreceptor voltage signal, $s_V(t)$, increases $\sim$4–15 times (Fig. 3 A) when warmed from 18 to 29°C. This is accompanied by an increasing signal mean, $\mu$ (or the BG0-induced steady-state depolarization), from $\sim$5 mV to 20–25 mV above the dark-adapted resting potential (Fig. 3 E) and by changes in the signal probability distribution (Fig. 3 C, continuous line) from Gaussian at the coolest temperatures ($\leq$18°C, including the cells in Fig. 3 D) to slightly negatively skewed at warmer temperatures (22, 25, and 29°C). In general, photoreceptors with the largest dark-adapted impulse responses also had the largest light-adapted steady-state potentials. Thus, the relation $\mu/V_{\text{max}}$ increases with warming, from 13% at 16°C ($n = 3$) to 43% at 27°C ($n = 4$), confirming that cooling reduces the steady-state potential of light-adapted photoreceptors. By contrast, the size of the contrast-evoked voltage signals, $s_V(t)$, of light-adapted photoreceptors is not proportional to their dark-adapted impulse responses, and photoreceptors with $V_{\text{max}}$ of $\sim$70 mV produce an $s_V(t)$ similar to that of photoreceptors with $V_{\text{max}}$ of $\sim$40 mV.

The voltage signal (Fig. 3 A) increases with temperature, being in two cells less than the noise at temperatures below 16°C (Fig. 3, D and F), but much greater under warmer conditions. Striking however, the average noise variance is virtually insensitive to the temperature (Fig. 3, B and F) and its probability distribution (Fig. 3 C, dotted line) is Gaussian. Since the warming-induced changes in the signal overwhelm those of the noise, an increasing temperature improves the photoreceptor SNR, often $\geq$10-fold over the studied temperature range (Fig. 3 G). At least three (not necessarily mutually exclusive) interpretations can be suggested to explain why both $s_V(t)$ and $\mu$ (steady-state potential) increase with warming, while $n_V(t)$, does not. First, the quantum efficiency of photoreceptors (i.e., the probability of producing an elementary voltage response, quantum bump, from an absorbed photon) increases with warming. Thus, the observed larger contrast responses and the steady-state potential at warm temperatures would simply result from the summation of a larger number of bumps. Second, the intracellular pupil mechanism (Lo and Pak, 1981) has temperature-dependent dynamics, so that in warmer temperatures a larger light flux would be allowed to enter the rhabdomeres. Again, more photons would sum up to produce larger voltage responses. Third, the synchronization of the bumps improves with warming. This would increase the time resolution of the voltage responses and expand the signaling bandwidth, so that $s_V(t)$ would enlarge with increasing temperature. To test these hypotheses, the corresponding signal and noise data were analyzed in the frequency domain.

### The Signal and Noise Dynamics in Frequency Domain

Fig. 4 shows the signal and noise power spectra of light-adapted *Drosophila* photoreceptors, and related functions at different temperatures. Because the photoreceptor signal power spectrum (Fig. 4 A) is calculated from the contrast-evoked voltage signals, it contains information about the mean bump waveform and its latency, whereas the noise power spectrum (Fig. 4 B), portraying the mean of the coding errors, is largely light-induced voltage noise (i.e., bump voltage noise; Juusola and Hardie, 2001). Warming increases the photoreceptor signal power, $|S_V(f)|^2$ (Fig. 4 A), and extends its bandwidth, whereas the peak and shape of the corresponding voltage noise, $|N_V(f)|^2$ (Fig. 4 B), changes relatively little. Assuming that the instrumental noise and light-induced voltage noise are independent but additive, we can estimate the bump voltage power, $|B_V(f)|^2$ (Fig. 4 F), at different temperatures by subtracting the photoreceptor voltage noise power spectrum measured in the dark, from the photoreceptor voltage noise power spectra, $|N_V(f)|^2$, measured at the adapting background of BG0 at different temperatures. Then, by fitting a single Lorentzian to the obtained bump voltage noise power spectra, $|B_V(f)|^2$:

$$
|B_V(f)|^2 = \frac{1}{(1 + (2\pi f)^2)^{n+1}},
$$

we obtain two parameters, $n$ and $\tau$ (Wong et al., 1980; Juusola and Hardie, 2001). These are used to calculate the average bump waveform (Fig. 4 G),

$$
b_V(t) \approx \Gamma_V(t; n, \tau) = \frac{1}{n! \tau^2} t^n e^{-t/\tau},
$$

and the effective duration, $T$ (Fig. 4 H), of the bump,

$$
T = \tau \left( \frac{n!}{2^n} \right) 2^{2n+1} (2n)!.
$$

The mean bump amplitude, $\alpha$ (used for scaling the bump waveforms in Fig. 4 G), is estimated from Campbell’s theorem (Wong and Knight, 1980):

$$
\alpha = \frac{\sigma^2}{\mu},
$$

where $\sigma^2$ is the noise variance and $\mu$ is the mean membrane potential. The bump rate, $\lambda$ (Fig. 4 I), is given by (Wong and Knight, 1980):

$$
\lambda = \frac{\mu^2}{\sigma^2 \cdot T}.
$$
The derived light-induced voltage noise power, $|B_1(f)|^2$ (Fig. 4 F), appears to be virtually independent of the temperature. Accordingly, the effective bump duration (Fig. 4 G) shows, at most, a marginal decrease with warming. Although the bump amplitude (Fig. 4 H) appears to decrease slightly upon warming, this cannot account for the increased steady-state potential, or for the much higher $SNR_\delta(f)$ (Fig. 4 C) and information content (Fig. 4 D) of the photoreceptor voltage responses. The $Q_{10}$ for bump duration ($T$) and amplitude ($\alpha$) is $1.2 \pm 0.2$ and $1.8 \pm 0.1$, respectively ($n = 2$). On the other hand, the average bump rate, $\lambda$ (Fig. 4 I), increases with warming ($Q_{10} = 4.0 \pm 0.2$, $n = 2$). The quantum efficiency of light-adapted Drosophila photoreceptors, hence, increases with temperature, and this may be at least partially responsible for the observed increase in the light-induced steady-state potential at temperatures above 18°C.

We found that, on average, a 10°C warming leads to a 6.5 ± 2.8-fold increase in the photoreceptor information capacity ($n = 3$; Fig. 4, D and E). Since the average bump rate at different recording temperatures can vary >10-fold (Fig. 4 I), so that some photoreceptors with rates <10^5 bumps/s are still able to have information capacities >300 bits/s, it is very unlikely that the huge increase in the photoreceptor information capacity is solely due to the summation of a larger number of bumps. Presumably, the accelerated bump timing dynamics, as seen in the expanding signal bandwidth (Fig. 4 A), is also contributing.

### Frequency Response Analysis

The temperature dependence of the photoreceptor signal power spectrum suggests that photoreceptors can follow higher contrast stimulus frequencies at higher temperatures, but this provides no details about the underlying mechanisms. To characterize how warming influences the signaling dynamics, we calculated the photoreceptor frequency response at different temperatures.

The acceleration of signaling dynamics is apparent in the photoreceptor contrast gain, $G_c(f)$ (Fig. 5 A). The 3-dB cut-off frequency shifts towards higher frequencies ($Q_{10} = 2.3 \pm 0.3$, $n = 3$; Fig. 5 B), and the corresponding phase, $P_c(f)$ (Fig. 5 C), lags the stimulus less at higher temperatures. Superficially these heat-induced changes in the signal dynamics resemble light adaptation from very dim to very bright mean light, including a significant increase in the absolute contrast gain (Juusola and Hardie, 2001). Because the experi-
mental phase functions, $P_V(f)$, differ from the corresponding minimum phase functions, $P_{\min}(f)$, the light-adapted photoreceptors exhibit a pure time delay (Juusola and Hardie, 2001). The duration of this so-called photoreceptor dead-time (Fig. 5 D), computed from the difference between $P_V(f)$ and $P_{\min}(f)$, decreases more than threefold in a 10°C temperature increase ($Q_{10} = 3.1 \pm 0.1$, $n = 3$).

The photoreceptor coherence function, $\gamma^2_{\text{SNR}}(f)$ (Fig. 5 E), is close to unity at all the tested temperatures, reconfirming the earlier findings that the responses of insect photoreceptors to dynamic contrast stimuli are linear (Leutscher-Hazellhoff, 1975; French, 1980; Juusola et al., 1994; Juusola and Hardie, 2001). Accordingly, the coherence function, $\gamma^2_{\text{SNR}}(f)$, which was estimated from the photoreceptor signal-to-noise ratio (see Juusola and Hardie, 2001, in this issue), is also close to unity over the signal bandwidth (Fig. 5 F), thus, demonstrating the high reproducibility of the photoreceptor voltage responses to contrast modulation.

The effect of warming on the photoreceptor’s signal processing speed is also clearly seen in the first order Wiener kernels, $k_V(t)$ (Fig. 5 G). These impulse responses calculated via inverse FFT (see Juusola and Hardie, 2001, in
this issue) are larger, briefer, and show a reduced latency and shortened time to peak \(t_p\) at higher temperatures. \(t_p\) reduces from values >40 ms at <18°C to 15 ms at 30°C (Fig. 5 H), constituting a Q₁₀ = 2.5 ± 0.2 \((n = 3)\).

**Bump Latency Distribution Changes with Temperature**

The photoreceptor voltage responses to contrast stimulation at a particular mean light intensity level are generated by a summation of bumps. If the average waveform of the bumps is very short and small, the speed and time resolution of the voltage response depends critically on the synchronization of the bumps, i.e., their timing (see Juusola and Hardie, 2001). Since the bump waveform, \(b_v(t)\), changes only modestly with warming (Fig. 4 H), the greatly improved temporal resolution of the corresponding voltage responses is presumably caused by faster and more precise bump timing (i.e., the bump latency distribution, \(l(t)\)) narrows with increasing temperature. To verify this, we estimated the bump latency distribution, \(l(t)\), by deconvolving the bump shape, \(b_v(t)\), from the corresponding impulse response, \(k_v(t)\) (see Juusola and Hardie, 2001, in this issue). This was done at different temperatures; first, by using simple equations to fit both the impulse response and the bump shape so as to minimize the effects of noise. Fig. 6 (A–C) shows how the fitted log-normal expressions of the impulse responses, \(k_{\text{norm}}(t)\) (Fig. 5 G), the corresponding \(\Gamma\)-distribution–fitted bump waveforms, \(\Gamma_v(t)\) (Fig. 4 G), and the calculated latency distributions, respectively, vary with temperature. Fig. 6 D shows the normalized bump latency distribution. The bumps appear earlier and in greater synchrony at higher temperature levels. The same is confirmed by calculating the bump latency, \(l(t)\), from the photoreceptor frequency responses, \(T_v(f)\), and the corresponding voltage noise spectra, \(|N_v(f)|\) (Fig. 6 E; see also Juusola and Hardie, 2001, in this issue):

\[
l(t) = F^{-1}\left[\frac{T_v(f)}{|B_v(f)|}\right] = F^{-1}\left[\frac{T_v(f)}{|N_v(f)|}\right]
\]

Fig. 6 D shows the normalized bump latency distribution. The bumps appear earlier and in greater synchrony at higher temperatures. The Q₁₀ for the bump latency distribution, \(Q_{\text{bump}}\), which was calculated from the peak time and half-width values, are 3.4 ± 0.2 and 3.5 ± 0.5 \((n = 3)\), respectively, coinciding with the \(Q_{\text{bump}}\) of the dead-time and similar to previously reported \(Q_{\text{bump}}\) values for the bump latency distribution.

\[
k_v(t) = b_v(t) \otimes l(t).
\]

All the methods revealed that warming shortens and narrows the dispersion of bump latencies. Warming shortens and narrows the dispersion of bump latencies.

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**Figure 6.** Bump latency distribution at the adapting background of BG0 at different with temperatures. Removing the bump shape from the corresponding impulse response by deconvolution reveals the timing of the elementary responses, i.e., the bump latency distribution. (A) The log-normal approximations of the photoreceptor impulse responses, (B) the normalized \(\Gamma\)-distribution–fitted bump waveform, and (C) the corresponding bump latency distributions at different temperatures. The normalized bump latency distributions (D) and those calculated from recorded voltage and light data as explained in Eq. 5 (E) and Eq. 6 (F) at different recording temperatures. Warming shortens and narrows the dispersion of bump latencies.
Such a high value is a strong indication that the latency is not limited by diffusion, but suggests that the enzymatic steps of phototransduction determine its time course.

II: Cell Membrane during Natural-like Stimulation

In our companion paper (see Juusola and Hardie, 2001, in this issue), we showed that the bump latency and not the filtering properties of the photoreceptor membrane limit the speed of the photoreceptor voltage response. The present paper shows that increasing the temperature reduces the bump timing delays (Fig. 6 D). Thus, the question arises whether the photoreceptor membrane dynamics are accelerated to allow transmission of the faster phototransduction signals, or whether the time constant of the photoreceptor membrane might limit the speed of the voltage responses at warmer temperatures. To investigate how the temperature dependence of the voltage responses is related to the dynamic properties of the membrane, we recorded photoreceptor voltage responses to both Gaussian current injections and contrast stimulation at different temperatures in single cells (Fig. 7).

**Warming Accelerates Both Light Responses and Photoreceptor Membrane**

Fig. 7 A shows samples of 1 s of the photoreceptor voltage signal and noise evoked by 10-s-long current modulation at different temperatures. Increasing the temperature reduces the amplitude modulation of the voltage signal. In the particular photoreceptor of Fig. 7, warming from 22 to 29°C reduces the signal variance by 26%. Since the current stimulus is unchanged, the diminishing average responses at higher temperatures result from reduced membrane impedance. The variance of the voltage noise is small, compared with the signal, and unaffected by the temperature in the studied range. These findings are compared with those of contrast stimulation at the same test temperatures (Fig. 7 B). In contrast to the current stimulation and in line with the previous experiments (Fig. 3, D and F), the photoreceptor signals to light contrast stimulation increase with the temperature.
Again, the variance of the photoreceptor voltage noise has no obvious temperature dependence.

Fig. 7 C shows typical probability distributions of voltage signals to dynamic contrast stimulation and current injection, $\langle N_V^2(f) \rangle$ and $\langle N_I^2(f) \rangle$, respectively at the test temperatures. Warming broadens the signal distribution and increases its skewness to Gaussian light contrast, but symmetrically reduces the depolarizing and hyperpolarizing voltage signals to Gaussian current injection. This confirms our previous findings (see Juusola and Hardie, 2001) that the response rectification during light contrast stimulation is not caused by the voltage-sensitive membrane (Fig. 2), but is due to some asymmetry in the phototransduction cascade’s response to light increments and decrements (for a first suggestion of this mechanism in Calliphora photoreceptors see Juusola, 1993).

Between 3 and 200 Hz, the photoreceptor voltage noise power spectrum appears not to be affected by temperature and current-induced voltage modulation (Fig. 7 D), indicating that the processes responsible for the bump shape are fully determined by the adapting light background. Consequently, although different in size, the voltage responses to dynamic light contrast and current stimulation at BG0 are equally reproducible, as seen in their highly similar noise power spectra, $\langle N_V^2(f) \rangle$ and $\langle N_I^2(f) \rangle$, respectively. However, the noise power in the low frequency range below 3 Hz is the highest at 29°C (Fig. 4 B). During some experiments at higher temperatures, we observed that the center of the photorecep-

![Figure 8](image-url)
ceptor’s receptive field shifted sporadically, causing slow fluctuations in the steady-state potential and, thus, affecting the power of the low frequency voltage noise. These movements may be due to an increased rate of intracapsular muscle activity at higher temperatures.

The effects of increased temperature on the photoreceptor membrane are very similar to those of increasing light adaptation shown in our companion paper (see Juusola and Hardie, 2001, in this issue). Fig. 8 (A and B) shows that warming reduces the mean photoreceptor membrane impedance and that its bandwidth is extended to higher frequencies. The photoreceptor impedance closely approximates a minimal phase system (Fig. 8 C). Therefore, the estimated photoreceptor dead-time should not be affected by charging the light current into voltage responses. According to the near-unity coherence values (\( \gamma \approx 0.98 \); Fig. 8, D and E), the light-adapted photoreceptor membrane produces linear voltage responses to current injection. The normalized impedance (Fig. 8 F) and the normalized gain of the light responses (Fig. 8 G) show that warming accelerates both the membrane impedance and the light responses. At all tested temperatures, the bandwidth of the membrane impedance is much broader than that of the light responses, as was also found previously for all different states of light adaptation (Juusola and Hardie, 2001). The \( Q_{10} \) values for the 3-dB cut-off frequency of the membrane impedance and the light responses are \( \approx 1.7 \) and \( \approx 2.3 \), respectively, in the photoreceptor of Fig. 8 (F and G).

**III: Temperature-sensitive Signal and Noise Properties of Light Current**

Because light-adapted photoreceptors respond to both the dynamic contrast and current stimulation with linear voltage responses, we can derive the output of the phototransduction cascade (light current), \( \gamma I(t) \), by deconvolving the contrast-induced voltages, \( \gamma V(t) \), from Figure 9. General comparison of the phototransduction signal and noise and membrane bandwidth at BG0 at different temperatures. (A–C) The dynamics of light current, voltage responses to light, and membrane impedance are displayed as their normalized frequency responses: \( G_I(f) \), \( G_V(f) \), and \( Z(f) \), respectively. At BG0, regardless of the tested temperature the light current was smooth and had a much narrower bandwidth than the corresponding photoreceptor membrane. As the temperature is increased, both the phototransduction cascade and photoreceptor membrane allows faster signaling leading to accelerated voltage responses (i.e., \( G_I(f) \) hardly differs from \( G_V(f) \)). The corresponding impulse responses (D), calculated from the same data, show how the light current and voltage responses quickens with warming, but the light current is always peaking before its corresponding voltage response. Because of the large membrane impedance at 22°C, the ratio between the corresponding light current and voltage response is larger than at warmer temperatures, where the photoreceptor impedance is less. The responses are normalized by the maximum value of each series. (E–G) The phototransduction bump noise, \( \gamma I(f) \), was calculated by deconvolving the photoreceptor membrane \( Z(f) \) from the respective voltage bump noise, \( \gamma V(f) \), measured at different temperatures. From 22°C to 29°C \( \gamma I(f) \) shows a considerable overlap with the corresponding membrane impedance indicating that the temperature-dependent shift in the membrane bandwidth effectively matches the temperature-dependent speed of the noise generating phototransduction processes.
Given that the Fourier transform of the membrane impedance impulse response, $z(t)$, measured in the same cell at the same mean light and temperature (see Juusola and Hardie, 2001, in this issue):

$$r_c(t) = z(t) \otimes r_v(t).$$

The light current frequency response, $T_f(f)$, is calculated from the contrast stimulus, $c(t)$, and the deconvolved current response, $r_c(t)$, (as shown in Juusola and Hardie, 2001, in this issue). Fig. 9 (A–C) compares the gain parts of the frequency responses of the photoreceptor membrane impedance, $Z(f)$ (dash/dotted line), light current, $G_l(f)$ (continuous line), and voltage responses, $G_v(f)$ (short dots), at three different temperatures. The high impedance membrane clearly acts as a low-pass filter for the light current, filtering specifically the high frequency content of the phototransduction signal, which is dominated by the bump voltage noise. Warming progressively adjusts the filtering properties of the membrane so that the cut-off frequency of its impedance never significantly limits the frequency response of the voltage signal. The higher mean impedance under cool conditions, results in small light currents charging relatively larger voltage responses compared with the situation at warmer temperatures (Fig. 9 D).

The phototransduction noise spectrum, $|\tilde{\Gamma}_{f}(f)|$, is estimated by deconvolving the normalized bump noise spectrum, $|\tilde{\Gamma}_{v}(f)|$, with the corresponding membrane impedance $Z(f)$:

$$|B_{v}(f)| = \frac{|B_{v}(f)|}{Z(f)} \propto \frac{|\tilde{\Gamma}_{v}(f)|}{Z(f)} = |\tilde{\Gamma}_{f}(f)|.$$

Given that the Fourier transform of $\Gamma_{v}(t)$ is its minimum phase representation in the frequency domain (Wong and Knight, 1980), we can compare the normalized $|\tilde{\Gamma}_{v}(f)|$ at different temperatures to the corresponding normalized membrane impedance function, $Z(f)$ (Fig. 9, E–G). In general, warming shifts the bandwidth of $|\tilde{\Gamma}_{v}(f)|$ in tune with $Z(f)$ so that they overlap at all the test temperatures (Fig. 8, E–G; Juusola and Hardie, 2001).

As the timing of the bumps becomes faster and more precise (as seen in the bump latency distribution, Fig. 6) and the changes in the bump waveform are matched by the temperature-sensitive filtering by the photoreceptor membrane (Fig. 9), the bandwidth of reliable signaling and the photoreceptor information capacity improves with warming (Fig. 10 A). The information capacity increases from values of below 50 bits/s at dim illumination (Fig. 10 A) to >100 bits/s during 0.32 dynamic contrast stimulation at BG0 at 29°C. Since the quantum efficiency of the light-adapted photoreceptors drops at cool temperatures, possibly because of some saturation related processes (see DISCUSSION), we also calculated the $Q_{10}$ for the average information capacity estimates at different mean light intensity levels at 25 and 29°C (where the estimated photoreceptor bump rates ($\nu$) for the same adapting backgrounds are very similar). We found that the $Q_{10}$ for information capacity at different mean light intensity levels. The consistently high $Q_{10}$ indicates that the large improvement in photoreceptor signaling is not a consequence of some light-dependent processes at bright light conditions, but reflects the increased speed and precision of the phototransduction reactions.

**DISCUSSION**

The rate of photon absorption and the dynamics of the phototransduction machinery and the photoreceptor membrane together set the limits for the information capacity of photoreceptor voltage responses, and ultimately for the temporal resolution of neural images affecting visual behavior. Comparative studies of insect photoreceptors suggest that their phototransduction and membrane dynamics are tuned to accommodate the statistical properties of rapidly moving images encountered in the natural habitat (McCann and MacGinitie, 1965; Laughlin and Weckström, 1993; Weckström and
Laughlin, 1995). We find in *Drosophila* photoreceptors the fidelity of this neural code highly temperature-sensitive, with the different processes behind the signal and noise dynamics having distinctively different Q10 values.

In the next paragraphs, we will first summarize the general effects of temperature on the photoreceptor responses, compare these to the relevant literature, suggest possible mechanisms behind them, and correlate them to more general neural coding schemes.

Response Dynamics at Cold and Warm Temperatures

In cold environments (below 20°C), photoreceptor signaling is badly impaired. The light-induced steady-state potential is lower (Fig. 3 E), although the bump amplitude (Fig. 4 G) and duration (Fig. 4 H) increase slightly with cooling. The bandwidth of the signal-to-noise ratio and its absolute values (Fig. 4 C), even at low frequencies, are greatly reduced (sometimes >10-fold). Since the photon density is high (BG0) and the linear photoreceptor membrane filters the signal and noise of the light current equally (Fig. 9), cooling must increase the variability in the bump dynamics in some fundamental way. Indeed, we find that the two most crucial elements of the neural code, the rate of events, and their timing precision, are jeopardized. Thus, cooling not only reduces the quantum efficiency (Fig. 4 I), but it also smears out the latency distribution of the bumps in time (Fig. 6 D), so that the light contrasts produce smaller and less synchronized responses (Fig. 5 G).

In warm environments (above 25°C), photoreceptors produce large responses with little noise. Warming slightly reduces the average shape and size of the bumps (Fig. 4 G), yet, more importantly, it increases the effective rate of bump production (Fig. 4 I), shortens the time delays in phototransduction (i.e., the bump latency distribution; Fig. 6 D), and accelerates the photoreceptor membrane dynamics (Fig. 8 F). As photoreceptor signaling is faster and has a higher quantum efficiency, the phototransduction cascade can generate faster and larger receptor currents, which are accommodated by the accelerated membrane characteristics so that the resulting voltage responses are more reproducible.

Sensitivity to Temperature

The temperature sensitivity of biochemical reactions is expressed by *Q*10, which is the fractional increase in their speed per 10°C. In vertebrate photoreceptors, *Q*10 values of light response kinetics are ~2 (Lamb, 1984). In invertebrate photoreceptors, the reported *Q*10 values vary from 2.5 for frequency responses at low temperatures (French and Järviilehto, 1978) in blowfly *Calliphora* to *Q*10 values of 4 for bump latency dispersion in *Limulus* (Wong et al., 1980). Recently, Tatler et al. (2000) measured *Q*10 values of 1.9 for the corner frequency of blowfly *Calliphora* voltage responses in the light-adapted state and over 3 in the dark-adapted state. We find similar values in light-adapted *Drosophila* photoreceptors, where the *Q*10 for the time to peak (*t*50) of the impulse responses and the corner frequency of the frequency responses is 2.5 and 2.3, respectively.

Our study is unique in the sense that we could isolate the effects of temperature on the phototransduction signal and noise and the filtering properties of the photoreceptor membrane by applying combined shot noise and signal analysis. This investigation reveals that the phototransduction processes related to the signal timing and the quantum efficiency (QE) are highly temperature-sensitive. The *Q*10 values for the dead-time, bump latency distribution, and the bump rate are 3.1, 3.5, and 4.0, respectively. On the other hand, the stochastic processes responsible for the voltage noise of light-adapted photoreceptors and the filtering properties of the membrane are less temperature-sensitive. Here, in contrast to findings of Roebroek et al. (1990) who reported a high *Q*10 of 2.5 for bump duration in *Calliphora* photoreceptors, we found that the light-adapted *Drosophila* photoreceptors have a relatively temperature-insensitive voltage noise power spectrum giving a *Q*10 for the effective bump duration and bump amplitude 1.2 and 1.8, respectively. Similarly, the *Q*10 for the corresponding corner frequency of the membrane impedance is only modest, 1.7, an intermediate value between the signaling and noise-generating processes. The dynamic interaction between the corresponding bump timing, waveform, and filtering processes, each having a different *Q*10 value, results in the coding fidelity of the photoreceptor voltage responses improving with increasing temperature, as seen in an effective *Q*10 estimate for information capacity of 6.5. Such a conspicuously high value reflects, in part, its derivation from the integral (area) of the SNR(f), and demonstrates how critical the operating temperature is for this most fundamental measure of coding efficiency in the nervous system.

Early Enzymatic and Late Diffusion Reactions

In our companion article on signaling dynamics in *Drosophila* photoreceptors at 25°C (see Juusola and Hardie, 2001, in this issue), we demonstrated that increasing the mean light intensity shortens and reduces the size of the average bump waveform, but leaves the bump latency distribution relatively unaffected. Following the reasoning of Wong et al. (1980) in *Limulus* photoreceptors, we attribute the bump timing and waveform dynamics to two independent stages in the phototransduction cascade, whose convolution defines the photoreceptor response. Because hypomorphic *Drosophila* mutations of PLC and G-protein affect the bump latency but not the bump shape (Pak et al., 1976; Scott et al., 1995; Scott and Zuker, 1998), events determining
the timing of the response are believed to occur largely at, or upstream of, PLC, whereas events determining the bump shape and amplitude are generated downstream of PLC and mediated by intracellular calcium. Our present findings reinforce the separability of bump waveform and latency in the light-adapted state by showing that the former is relatively temperature insensitive, while the latter is highly dependent on temperature. A plausible explanation is that the speed of the calcium-mediated bump shape dynamics under fully light-adapted conditions is saturated with the high intracellular calcium level, whereas the time course of the earlier transduction reactions relating to bump timing is still very susceptible to even small changes in temperature. The very different Q_{10} of the bump latency distribution and bump shape suggests that the early phototransduction processes responsible for the timing of the elementary responses are enzymatic, whereas the latter membrane-bound reactions responsible for the amplification of the responses are diffusion limited.

Reduction of the Quantum Efficiency

An unexpected finding of this study is the high temperature sensitivity of bump production rate in bright light conditions. What kind of processes could reduce the effective QE of light-adapted photoreceptors? Saturation of the available transduction units, possibly individual microvilli, has been shown to occur under very bright daylight conditions in Calliphora (Howard et al., 1987), yet, the numbers do not add up here for Drosophila photoreceptors. Knowing that a photoreceptor rhabdomere consists of 30,000 microvilli, each containing a complete transduction cascade with the signaling speed limit set by the photoreceptor dead-time (Juusola and Hardie, 2001), we can calculate the maximum signaling rate of a photoreceptor at BG0 (with the mean light flux of \(3 \times 10^6\) photons/s). At 18°C, the photoreceptor dead-time of 25 ms restricts the bump rate of a microvillus to 40 photons/s. Therefore 30,000 microvilli, each coding 40 photons/s, can maximally process \(1.2 \times 10^6\) photons/s. Since this is \(\sim 10-30\) times the estimated rate of the photons absorbed at the same temperature (30,000–100,000 photons/s), it is very unlikely that the phototransduction machinery is operating right at its saturation level. Another evidence for this conclusion comes from the information capacity recordings at different mean light intensity levels (Fig. 10 A). We find a similarly steep temperature dependence of information capacity at the adapting backgrounds of BG-3 and BG-2, where the number of photons per second absorbed is less than the number of transduction units.

Further experiments will be required to determine the cause of the unexpected loss of QE at lower temperatures. It does not appear to occur at the level of the visual pigment, since the dark-adapted QE appears to be unaffected by cooling to 15°C (Hardie, R.C., unpublished observations); but, it might represent saturation/exhaustion of some limiting step in the phototransduction pathway such as the PLC substrate (PIP2), which must be continually recycled via a multi-enzyme resynthesis process.

Temperature-dependent Limitations on the Speed of the Voltage Responses

Because of the high cut-off frequency of the photoreceptor membrane, the speed of the photoreceptor voltage responses is not significantly limited by the membrane over the tested temperature range, but is set by the delays in the phototransduction cascade. However, since the Q_{10} for the membrane corner frequency is 1.7 and the Q_{10} of latency processes is over 3, in high temperatures the filtering effect by the membrane on the light current becomes gradually more important, but not limiting. On the other hand, at 35°C, the extrapolated average bump duration would be 7.5 ms, which is \(\sim 60\%\) larger than the estimated half-width of the latency distribution of 4.3 ms. Hence, at such high temperatures, the speed of the light current would depend at least as much on the average bump duration as it does on the bump timing.

Benefits of Accelerated Responses

Previous studies in toads on the effect of temperature on visual performance have emphasized how cooling can improve performance by reducing the rate of thermal isomerizations of rhodopsin, thereby lowering the absolute detection threshold (Aho et al., 1988). Although this may be true under dark-adapted conditions when spontaneous bumps set the absolute limit to detection, the present study demonstrates how warming results in a massive improvement in SNR_{V} and information capacity under generally more relevant light-adapted conditions. In the cool of the night, the heavily low-passed signaling system provides good amplification for objects coded by low numbers of photons, so that they can be separated from the background, but the accelerated photoreceptor dynamics are beneficial for high temporal resolution in the hot daytime. Improved temporal resolution gives behavioral advantages to an animal and may improve its survival odds. Therefore, it is not surprising to find that some invertebrates may benefit from the body fluid circulation, which cause part of the heat produced by the thoracic muscles to enter and heat up the head (and hence retinae) above the ambient temperature. Hoverflies, moths, and dragonflies can maintain a head temperature above 30°C, even when the ambient temperature is below 20°C (Heinrich, 1993; Tatler et al., 2000).
Conclusion

In summary, we have provided a comprehensive description of the temporal resolving capability in *Drosophila* photoreceptors under natural conditions and, to our knowledge, the first analysis of the effects of temperature on information capacity in the nervous system. Although it has often been reported that temperature accelerates the kinetics of light responses (Wong et al., 1980; Lamb, 1984), a priori, the conclusion that this improves signaling efficiency is by no means certain: the final voltage response represents a finely tuned balance between multiple components within the phototransduction cascade together with the passive and active properties of the plasma membrane. For example, the membrane impedance function speeds up in parallel with the phototransduction cascade, so that at no point does it seriously compromise the bandwidth of the light current. If this was not the case, the membrane would have seriously reduced the information capacity; e.g., if the membrane impedance determined at 17°C (Fig. 8 B) is convolved with the light current responses at 29°C (Fig. 9 C), then the estimated information capacity would be reduced fivefold. A future challenge will be to analyze how the different components of the cascade and membrane interact to produce this finely tuned performance. *Drosophila* is an excellent model in which to attempt this analysis because of the opportunities for genetic manipulation of the various elements including cascade enzymes, and both light- and voltage-sensitive ion channels.

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