Dynamic Relationship between the Slow Potential and Spikes in Cockroach Ocellar Neurons

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ABSTRACT The relationship between the slow potential and spikes of second-order ocellar neurons of the cockroach, Periplaneta americana, was studied. The stimulus was a sinusoidally modulated light with various mean illuminances. A solitary spike was generated at the depolarizing phase of the modulation response. Analysis of the relationship between the amplitude/frequency of voltage modulation and the rate of spike generation showed that (a) the spike initiation process was bandpass at ~0.5-5 Hz, (b) the process contained a dynamic linearity and a static nonlinearity, and (c) the spike threshold at optimal frequencies (0.5-5 Hz) remained unchanged over a mean illuminance range of 3.6 log units, whereas (d) the spike threshold at frequencies of <0.5 Hz was lower at a dimmer mean illuminance. The voltage noise in the response was larger and the mean membrane potential level was more positive at a dimmer mean illuminance. Steady or noise current injection during sinusoidal light stimulation showed that (a) the decrease in the spike threshold at a dimmer mean illuminance was due to the increase in the noise variance: the noise had facilitatory effects on the spike initiation; and (b) the change in the mean potential level had little effect on the spike threshold. We conclude that fundamental signal modifications occur during the spike initiation in the cockroach ocellar neuron, a finding that differs from the spike initiation process in other visual systems, including Limulus eye and vertebrate retina, in which it is presumed that little signal modification occurs at the analog-to-digital conversion process.

INTRODUCTION

In the visual system of animals, photic inputs are initially converted into slow (graded) potential signals and the graded signals are then converted into spike signals. Because the photic input that animals experience in the natural environment is a fluctuation of light intensity around a mean illuminance, the slow potential response and the spike response of visual neurons to light fluctuation have been studied (Fuortes and Hodgkin, 1964; Victor and Shapley, 1979; Tranchina et al., 1983; Chappell et al., 1985; Victor, 1987). However, little attention has been given to the dynamics of the cellular processes that convert the slow potential into spikes. A notable exception is the sine wave analysis of Limulus eccentric cells. Knight et al. (1970) found that the...
magnitude of slow response is linearly converted into a spike rate in eccentric cells. Knight (1972a, b) developed an “integrate-and-fire” model for the spike initiation process, which explained well the actual spike activity of *Limulus* eccentric cells. Recently, Sakuranaga et al. (1987) found that the model can be applied to the spike discharge of catfish retinal ganglion cells. These studies suggested that the spike initiation process produces a replica of the slow potential without any major modification of the signal.

The insect ocellus is a simple photoreceptor system advantageous for studying basic mechanisms of visual signal processing. The insect ocellus contains over 100 photoreceptors and these converge onto fewer than 12 large second-order neurons, called L-neurons (Ruck, 1961; Dowling and Chappell, 1972; Goodman, 1981). Intracellular recordings from insect ocellus were first made by Chappell and Dowling (1972), who showed that a light stimulus depolarized the ocellar receptors and hyperpolarized the L-neuron of dragonfly ocelli. Incremental responses of the L-neuron have been studied in dragonflies (Chappell and Dowling, 1972) and in cockroaches (Mizunami et al., 1986). It was concluded that the incremental sensitivity of the insect L-neuron is an exact Weber-Fechner function. In addition to the graded hyperpolarizing response, the L-neuron of the cockroach exhibits spikes at the decremental phase of step stimuli or sinusoidal stimuli (Mizunami et al., 1982, 1986), as in the case of the locust L-neuron (Wilson, 1978a).

In the present study, we analyzed the dynamic relationship between the slow potential and spikes in the cockroach ocellar L-neuron, using sinusoidally modulated light stimuli around various mean illuminances. The cockroach L-neuron is suited for such an analysis because (a) stable intracellular recordings of >60 min are feasible and (b) a sinusoidal light modulation produces almost sinusoidal voltage modulation, thereby allowing for a high quality of sine wave analysis of the spike initiation process. Our major findings are that (a) the spike initiation process has bandpass filtering characteristics; (b) the spike initiation process contains a dynamic linearity and a static nonlinearity; (c) the noise in the response lowers the spike threshold: it has facilitatory effects on the spike initiation; (d) the noise effects are prominent when there is a low-frequency potential modulation under a dim mean illuminance; and (e) the mean potential level, which changes depending on the mean illuminance, has little effect on the spike threshold. We conclude that (a) the spike initiation process in cockroach L-neurons can be modeled by an integrate-and-fire generator, and (b) the spike initiation process in the L-neuron is an important step in signal processing in the cockroach ocellar system.

**MATERIALS AND METHODS**

*Preparation*

Adult males of the cockroach, *Periplaneta americana*, reared in our laboratory at Kyushu University, were studied. The whole animal was mounted dorsal side up on a Lucite stage and fixed with beeswax. The compound eyes and one of two ocelli were shielded from light by beeswax mixed with carbon black. The dorsal part of the head capsule was removed and the dorsal surface of the brain was exposed. Saline containing 1% Actinase (type E, Kaken Seiyaku, Tokyo, Japan) was applied to the brain to facilitate insertion of the electrode. The saline solution was that described by Yamasaki and Narahashi (1959).
Intracellular recordings from L-neurons were made using glass microelectrodes filled with 2 M potassium acetate and having a DC resistance of ~50–80 MΩ. These electrodes were inserted into L-neurons at the ocellar tract of the brain, an area at which the spikes of L-neurons initiate (Mizunami et al., 1987). Stable recordings of >60 min were feasible. These neurons were identified as L-neurons from their responses, in particular: (a) a hyperpolarizing response of >30 mV to a bright light stimulus, and (b) a large voltage fluctuation during dim light stimulation. In some preparations, the neurons were stained by injecting cobalt ions through the recording electrode and identified anatomically (see Mizunami et al., 1982). The electrodes were connected to a high-impedance, negative-capacity compensated preamplifier (MEZ-8201, Nihon Kohden, Tokyo), which was equipped so that a constant current could be passed through an active bridge circuit. The magnitude of the stimulus current depended linearly on the driving voltage applied to the current-passing circuit. A small piece of platinum in the bathing solution served as an indifferent electrode.

Current-voltage relationships of L-neurons were measured using double-barreled electrodes; one barrel was used to inject the current and the other was used to record the voltage. The electrodes had a small coupling resistance of <0.4 MΩ; therefore, the current-voltage plot was obtained by subtracting the voltage drop due to electrical coupling from the recorded voltage change.

A light-emitting diode (LED; Sharp Corp., Tokyo) was used as a light source. The LED had a spectral peak at 560 nm. The LED was driven by a sinusoidal current provided by a function oscillator (ET1101, NF Design Block, Tokyo). The illuminance of the stimulus depended linearly on the magnitude of the driving current. The stimulus light was monitored by a photodiode (TFA1001W, Siemens-Allis, Inc., Cherry Hill, NJ) before being attenuated by filters. The light stimulus and cellular response were observed on an oscilloscope and stored on analog tape. Some analyses were made using a VAX 11/780 computer (Digital Equipment Corporation).
Corp., Maynard, MA) with an AP120B array processor (Floating Point Systems, Portland, OR). All experiments were done at a room temperature of 20–24°C.

**Analytical**

The sinusoidal light stimulus consisted of two components, a steady mean, \( I_0 \), and a dynamic component, \( I(f) \), as shown in Fig. 1. \( I(f) \) was defined by the modulation frequency (Hertz) and the depth of modulation. The depth (percentage) is defined in the conventional fashion: 

\[
\text{Depth} = \frac{I_{\text{max}} - I_{\text{min}}}{I_{\text{max}} + I_{\text{min}}} \times 100
\]

where \( I_{\text{max}} \) is the maximum illuminance and \( I_{\text{min}} \) is the minimum illuminance. The depth of modulation represents the “contrast” between the stimulus and the adapting light. The mean illuminance, \( I_0 \), was represented by log10 attenuation (0 log is 20 \( \mu \)W \( \cdot \) cm\(^{-2} \) throughout this study). In the actual experiments, neutral density (ND) filters were interposed between the light source and the preparation to attenuate \( I_0 \). The depth of modulation of the stimulus remained unchanged by interposing ND filters.

The resulting response recorded intracellularly from cockroach ocellar L-neurons consisted of a sinusoidal slow potential modulation and spikes. The slow potential response contained three components, a steady mean potential, \( V_0 \), a voltage noise, \( V_n \), and a modulation response, \( V(f) \). \( V_0 \) and \( V_n \) were related to the mean illuminance, \( I_0 \), and \( V(f) \) was related to light modulation, \( I(f) \). \( V(f) \) was defined as \( V_{\text{peak}} - V_{\text{bottom}} \), where \( V_{\text{peak}} \) is the potential at the peak and \( V_{\text{bottom}} \) is the potential at the bottom of the voltage modulation.

The magnitude of spike response could be defined as a probability of spike generation for one voltage cycle, \( S(f) \), as will be discussed in the Results. The L-neuron had no maintained discharge under steady illumination or in the dark; the spike response had no steady component. We analyzed the manner in which the spike response, \( S(f) \), is related to parameters of the slow potential response, \( V(f), V_n, \) and \( V_0 \). We also describe briefly how each component of the slow potential response and the spike response relates to each parameter of the light stimulus. Some aspects of the relationship between the light stimulus and the slow potential response have been reported (Mizunami et al., 1986).

**RESULTS**

**Responses to Sinusoidal Light Stimulus**

The cockroach ocellus contains ~10,000 photoreceptors, which converge on four large second-order neurons, the L-neurons (Weber and Renner, 1976; Toh and Sagara, 1984). The axon of the L-neuron exits the ocellus and projects into the ocellar tract of the brain through the ocellar nerve (Mizunami et al., 1982). In the ocellar tract, L-neurons make output synapses onto several types of third-order neurons (Toh and Hara, 1984; Mizunami and Tateda, 1986).

Fig. 2 shows typical records of responses from an ocellar L-neuron to sinusoidal light modulation. Responses to stimuli with various modulation frequencies (A) and various modulation depths (B) are shown. The stimuli had a mean illuminance of 0.2 \( \mu \)W \( \cdot \) cm\(^{-2} \) (−2 log units). The response of the L-neuron consisted of two components, a graded voltage fluctuation and the spikes. We refer to the former as the slow potential response and to the latter as the spike response. The waveform of the slow potential response to sinusoidal light was roughly sinusoidal, which suggests a quasilinear response. A small nonlinearity was detected in the slow potential response: the peaks of the sine wave were sharper than the trough. The nonlinearity reflects a compression of the response amplitude to an incremental (hyperpolarizing) light stimulation (Mizunami et al., 1986). The nonlinearity may also reflect a subthreshold active (regenerative) response. A solitary spike was evoked at the peak
or depolarizing phase of the modulation response. One modulation cycle of stimulus usually evoked only one spike, even when a stimulus with a large modulation depth (75%) was applied. When the modulation frequency was low (less than ~0.5 Hz) and the mean illuminance was low (less than about ~3 log units), two to four spikes were frequently seen within a cycle of voltage modulation, as will be discussed later (see Fig. 6). We defined the magnitude of spike response as the rate of spike generation for one cycle of modulation stimulus. The rate of spike generation was measured as follows. The responses to 30–80 cycles of stimulus were recorded, and the rate of spikes was calculated by dividing the number of voltage modulation cycles in which spikes were generated by the total number of the cycles. Both the peak-to-peak amplitude of the slow response and the rate of spikes depended on the modulation frequency and the modulation depth of the stimulus.

Fig. 2 shows the peak-to-peak amplitude of the slow response of an L-neuron plotted against the modulation depth of the stimulus. To simplify the figure, only five series of responses to different modulation frequencies are shown. The stimulus had a mean illuminance of 2 μW·cm⁻² (~1 log unit). The amplitude of the slow response depended almost linearly on the modulation depth of the stimulus, at least within the range of modulation depths of 80%. A similar linear relationship was observed over a wide range of frequencies (0.1–30 Hz), and also over a 3.6-log range of mean illuminance (0.005–20 μW·cm⁻²). The linear nature of the slow potential response facilitated analysis of the dynamic relationship between the slow potential and the spikes, using a sine wave–modulated light. If the slow potential response were
nonlinear and the response to sinusoidal stimulation had a serious nonlinear
distortion, the light stimulation would be inadequate for a quantitative analysis of the
relationship between the slow potential and the spikes.

**General Characteristics of the Slow Spike Conversion Process**

Fig. 4 A shows the relationship between the peak-to-peak amplitude of the slow
potential response and the rate of spike generation at a modulation frequency of 1
Hz. The mean illuminance of the stimulus was -2 log units. There was a dead zone in
which spikes were not generated. Beyond that zone, the spike rate increased with the
increase in the amplitude of the slow response. The generation of the spike response
is probabilistic, showing that the spike initiation has an internal noise. The plot is
almost sigmoidal, with a linear part covering the spike rate ranging from 10 to 90%.
Similar sinusoidal or quasilinear relationships between the amplitude of voltage
modulation and the spike rate were obtained over a frequency range of 0.1–20 Hz
and over a mean illuminance range of 3.6 log units.

Fig. 4 B shows the relationship between the peak-to-peak amplitude of the slow
potential response and the spike rate obtained at different frequencies. The results
at a spike rate of between 10 and 90% are shown. The extrapolated straight lines are
regression lines for each frequency. The lines cross the vertical axis at almost the same
point. This suggests that the nonlinear threshold is frequency independent: the
nonlinearity of the spike initiation process is static. On the other hand, the slope of
the lines changes with the frequency, which indicates that the spike initiation process
contains a dynamic linearity. In short, the spike initiation process contains a dynamic
linearity and a static nonlinearity. A simple model for the spike initiation process will
be proposed, based on these observations, in a later section (see Fig. 12).

**Effects of Mean Illuminance on the Spike Threshold**

Here we define 50% threshold as the peak-to-peak amplitude of the slow response at
a spike rate of 50%. Further analysis was made using the 50% threshold. Fig. 5 A
shows the relationship between the 50% threshold and the modulation frequency, obtained at a mean illuminance range of 3.6 log units. In this experiment, a neuron was impaled, the ocellus was dark-adapted for 5 min, and the test began with -3.6-log ND filters interposed. After each sinusoidal test run (started after 60 s of adaptation), the density of the ND filter was decreased. After a test at the maximum illuminance (0 log), the sequence was reversed. This series was repeated four times. The 50% threshold was smallest at frequencies of ~0.5–5 Hz: the slow spike conversion process was bandpass. The 50% threshold at optimal frequencies, where the 50% threshold was smallest (~0.5–5 Hz), was unchanged over a mean illuminance range of 3.6 log units. However, the 50% threshold at frequencies of <0.5 Hz did change depending on the mean illuminance levels. The threshold was lower at a
dimmer mean illuminance. Similar observations of eight L-neurons were made repetitively, although the absolute value of the 50% threshold differed slightly (<30%) from preparation to preparation.

Fig. 5 B shows the phase characteristics of spike generation, measured from the peak of the potential modulation. Measurements were done from the records of responses in which the spike rate was ~50%. The phase of spike generation progressively led that of the slow potential with decreases in the frequency, which suggests that the process has a differential or change-sensitive nature. The variance of the phase was larger at a dimmer mean illuminance. The variance of the phase at -3.6 log units was about twice of that at 0 log units.

Fig. 6 shows actual recordings from an L-neuron exposed to low-frequency stimulation, at mean illuminances of -3.6 (upper trace) and of 0 (lower trace) log
Figure 5. (A) 50% threshold of spike response, defined as the peak-to-peak amplitude of the potential modulation at a spike rate of 50%, plotted against the modulation frequency. The plots are from the responses of an L-neuron to sinusoidal lights with a mean illuminance of 0.005 (−3.6 log), 0.2 (−2 log), and 20 μW·cm⁻² (0 log). (B) Phase characteristics of the spike response measured from the peak of the potential modulation. Measurements were made from the records of responses in which the spike rate was ~50% (40–60%), and the results at a mean illuminance of 0 log and −3.6 log are shown. The phase variance is larger at a dimmer illuminance (−3.6 log). A and B are from the same L-neuron.

units. The modulation depth of the stimulus was kept at 60% and the modulation frequency was kept at 0.2 Hz. Spikes could be seen in the −3.6-log record but not in the 0-log record, although the waveforms and the averaged peak-to-peak amplitudes of the modulation response were similar. This observation indicated that the 50% spike threshold is lower; i.e., a smaller amplitude of voltage modulation can produce spikes at a dimmer mean illuminance. In addition, more than one spike was frequently observed within one cycle of the potential modulation, when the mean illuminance was less than −3 log units and the frequency of modulation was below ~0.5 Hz. The increase in the spike number was accompanied by an increase in the variance of the phase of spike initiation.

Note that our analysis was designed for a system in which a single stimulus (single

Figure 6. Responses of an L-neuron to sinusoidal lights with a mean illuminance of 0.005 (−3.6 log) or 20 μW·cm⁻² (0 log). The stimuli had a modulation frequency of 0.2 Hz and a modulation depth of 60%. The dashed lines indicate the mean membrane potential in the dark (about −45 mV). Note that spikes are seen from the −3.6-log record but not from the 0-log record.
voltage modulation) produces a single effect (single spike), and had a limitation when multiple spikes occurred during a single modulation cycle. We defined the rate of spike generation as the number of modulation cycles in which spikes were generated, divided by the total number of cycles: the spike rate in this study does not reflect the number of spikes during a single modulation cycle. Thus, the 50% threshold, defined on the basis of the spike rate, is less informative about the actual spike response when multiple spikes occur during a cycle. However, our extended analysis (noise current stimulus combined with the sine wave light stimulus; see below) helped us understand which features of slow potential are relevant in the production of multiple spikes during a single modulation cycle.

Fig. 7 shows the gain and phase portion of the transfer characteristics from light to the slow potential response and of those from light to the spike response. The gain of the slow potential response was defined as the slope of the depth-response curve, i.e., the magnitude of the slow response (millivolts) divided by the depth of stimulus modulation (percentage). This definition is based on the observation that the slow potential response is almost linear (see Fig. 3). Because the depth represents the contrast between the stimulus and the adapting light, the gain is also the contrast sensitivity. The gain of the spike response was defined as the inverse of the depth of stimulus modulation (percentage) at a spike rate of 50%. We found for the gain...
portion of the transfer characteristics (Fig. 7, A and C) that: (a) both the slow potential and spike responses had bandpass characteristics with optimal frequencies of ~0.5–5 Hz; (b) the spike response is more sharply bandpass than is the slow potential response; (c) the gain of the slow potential response is practically unchanged over a mean illuminance range of 3.6 log units: the gain is independent of the mean illuminance levels; (d) the gain of the spike response at optimal frequencies remains unchanged over a mean illuminance range of 3.6 log units; however, (e) at lower frequencies, the gain of the spike response changes depending on the mean level of illuminance. These differences reflect dynamic properties of the spike initiation process. The phase of the slow potential response is about -180 degrees at low frequency, which reflects that the sine of the response is negative-going (hyperpolarizing) and lags with increases in the stimulus frequency. A comparison of the phase of the slow potential and spike response shows that (a) the phase of the spike response leads that of slow potential response, and (b) the phases of both responses are affected little by the mean illuminance change, except that variance of the phase of the spike response changes depending on the mean illuminance, as shown in Fig. 5 B. We conclude that (a) the dynamics (frequency selectivity) of the slow potential response are independent of the mean illuminance levels, but (b) there is an enhancement of low-frequency sensitivity in the spike response at a dim mean illuminance: the dynamics of the spike response depend on the mean illuminance.

Effects of Noise and Mean Potential Level on the Spike Threshold

Fig. 6 also shows that (a) the magnitude of the spontaneous voltage fluctuation (voltage noise) is smaller under brighter illumination and (b) the mean potential level is more negative under brighter illumination (the dashed lines indicate the mean membrane potential in the dark). We considered that these may be the factors responsible for changes in the 50% spike threshold by changes in the mean illuminance.

The dependence of the mean potential level and the noise magnitude on the mean illuminance levels was examined. The L-neuron responded to steady illumination with a hyperpolarizing potential with a transient peak, and the potential reached a steady level within 30–40 s (Fig. 8 A). In the steady state, the membrane potential was more negative and the noise was less prominent under brighter illumination. Fig. 8 B shows the magnitude of the steady state hyperpolarization from the dark potential, plotted against the stimulus illuminance. Averages from five L-neurons are shown with the standard deviations. The membrane potential of L-neurons hyperpolarized ~0.8–1 mV for a 1-log increase in the stimulus illuminance, within a range from -4 to -1 log units. The membrane potential seemed to approach a constant level at the illuminance exceeding -1 log unit. Fig. 8 C shows the probability density functions (PDFs) of the noise obtained at a 4-log range of illuminance levels. The zero potential of the plot indicates the level of mean potential. The noise was reduced when the illuminance of the stimulus was increased. The noise reached a minimum at an illuminance of about -1 log unit, at which the half-width of the PDF was ~0.5 mV. Fig. 8 D shows power spectra of the noise obtained at a 4-log range of illuminance levels. The power spectra looked like a simple Lorenzian function, and most of the power was restricted to frequencies of less than ~50 Hz. This power was greatly reduced, with increases in the stimulus illuminance.
At least two possible sources for the noise were considered. First, the noise may reflect the electrical properties of the L-neuron: the noise may be due to a voltage-dependent noise source. In the dark or under dim illumination, the neuron may be more active than under brighter illumination because the membrane potential is more positive; thus, a larger voltage-related noise may be generated. Second, the noise may reflect that contained in the synaptic potential from photoreceptors. If the noise reflects electrical properties of the L-neuron, significant differences in the noise magnitude would be expected for an extrinsic current applied through the recording electrode. For example, when the dark-adapted L-neuron is hyperpolarized 3 mV by the extrinsic current, the noise would decrease to a value under a steady illumination of ~0 log units. Also, when the L-neuron is depolarized 3 mV under conditions of steady illumination of 0 log units, the noise would increase to a value seen in the dark. Before the current-injection experiments were done to test these possibilities, we measured the current-voltage relationship of L-neurons using double-barreled electrodes. Examples of recordings of the voltage responses to current stimuli are shown in Fig. 9 B, and a typical current-voltage relationship is shown in Fig. 9 C. The input resistance, the slope of the current-voltage plot, was ~1.6 MΩ in the dark and ~0.8 MΩ in the presence of a steady illumination of 0 log units. On the basis of these observations, we injected a negative current of 2 nA into a dark-adapted L-neuron to hyperpolarize it ~3 mV, or a posi-
ative current of 4 nA under illumination of 0 log units to depolarize the neuron ~3 mV (Fig. 9A). In both cases, no prominent changes in the noise magnitude were observed. We thus conclude that the contributions of the voltage-dependent noise source to the noise seen in the L-neuron are small. The noise reflects that contained in synaptic potentials from photoreceptors.

In Fig. 10A, steady or noise current was injected into an L-neuron during light stimulation, and the 50% threshold was measured. The mean illuminance of the stimulus was 0 log units, at which the mean membrane potential was ~3 mV negative to the dark level. There the noise was slight. A steady depolarizing current of 4 nA, estimated to depolarize the neuron ~3 mV based on the input resistance data (see Fig. 9C), had no significant effect on the 50% threshold. We conclude that the change in the mean potential level is not the major cause of changes in the 50% threshold by changes in the mean illuminance. The noise current stimulation was made as follows. The potentials of an L-neuron were recorded under conditions of dim illumination (~3.6 log units) and then stored on analog tape. These stored
potentials were passed through an active low-cut filter to subtract the DC components and were used to drive a current-passing circuit. The output current of the circuit, which was applied to an L-neuron, had a waveform similar to that of the driving noise potential. The 50% threshold at low modulation frequencies became smaller when a noise current that had a peak-to-peak intensity of ~4 nA was injected (Fig. 10 A, open circles). The curve of the plot in the presence of noise current was similar to that observed under dim illumination (see Fig. 5 A). Similar results were repeatedly obtained with four L-neurons. In addition to the change in the 50% threshold, we observed that the noise current produced (a) an increase in the variance of the phase of spike generation and (b) an increase in the spike number for one modulation cycle when the frequency was low. These findings were similar to those observed with a dim mean illuminance. We concluded that the decrease in the spike threshold to a low-frequency potential modulation by the decrease in the mean illuminance was due to the increase in the noise variance: the noise had a facilitative effect on the initiation of the spike.

Fig. 10 B shows the frequency selectivity of the facilitative effect of the noise on the spike initiation. In the figure, the magnitude of the decrease in the 50% threshold, induced by a noise current, is plotted against the modulation frequency. The procedure for the measurements was the same as that of Fig. 10 A, and averaged data...
from four L-neurons are shown. The facilitative effect of the noise on the spike initiation was more prominent at a lower modulation frequency and less prominent at a higher frequency. This can be explained as follows. When there is a subthreshold low-frequency potential modulation, the potential will remain just below the firing level for a longer period than at a higher frequency. Thus, the probability that the potential will reach the firing level by the superimposed noise will be higher. Similarly, the superimposed noise will produce repetitive firing if the potential remains near the firing level for a longer time than the refractory period of spike initiation.

Note that although the noise effect on the 50% spike threshold was highly frequency selective, the effect on the spike frequency might not be frequency selective, because of multiple times that a potential is near its peak at a high-frequency potential modulation. Our analysis was done not on the spike frequency but on the (50%) spike threshold, because L-neurons rarely produced trains of spikes and appeared to code signals in the form of solitary spikes (discrete events) rather than as the frequency of spike trains (analog signals).

**Steady State Inactivation of Spike Generation**

It was an unexpected observation that the change in the mean membrane potential levels was not accompanied by a change in the 50% spike threshold. We did current-injection experiments to confirm this point. A hyperpolarizing prepulse was applied for ~10 s in a dark-adapted L-neuron to maintain the membrane potential at different levels; then a depolarizing test pulse was applied to trigger a spike. In Fig. 11, the threshold depolarizing current, which is the amount of current of the test pulse necessary to trigger a spike (see inset B), was plotted against the maintained current (A). We could expect that the relationship between threshold depolarization, the magnitude of step depolarization necessary to produce a spike, and maintained potential would be similar to the data in Fig. 11, because the input resistance was fairly constant in the range of current magnitude used (see Fig. 9 C). At a dark
potential, i.e., at a maintained current of 0 nA, depolarizing current pulses failed to evoke spikes: the neuron was completely inactivated at a dark potential (the same can be seen in Fig. 9 B). In the presence of a hyperpolarizing prepulse, the neuron generated a spike in response to a depolarizing current pulse. The membrane potential, under steady illumination of 0.005–20 μW·cm⁻², was ~0.8–3 mV negative to the dark potential (see Fig. 8 B). This potential range corresponds to the maintained current range between about −0.5 and −2 nA (marked as “light” in Fig. 11). In this range of maintained current, the threshold depolarizing current was fairly constant. This observation confirms that the change in the mean potential level had little effect on the spike threshold.

The observation in Fig. 11 suggests that there is a high degree of steady state inactivation in light-adapted L-neurons. The L-neuron was completely inactivated in the dark, and the membrane potential of the light-adapted L-neuron was <3.5 mV negative to that in the dark. Apparently the steady state inactivation produced a seemingly paradoxical relationship between the spike threshold and the mean potential levels; i.e., the spike threshold is independent of the mean potential levels.

DISCUSSION

Most studies on visual systems have been concerned with the response to steps of light given in the dark, and measurements have been made on the static aspects of the step-evoked responses. Therefore, little is known of the response dynamics to a light fluctuation around a mean illuminance. In particular, the dynamics of the cellular process that converts the slow potential into spikes are not well understood. In the vertebrate retina, dynamic properties of the spike response of ganglion cells have been studied (Schellart and Spekreijse, 1972; Victor and Shapley, 1979; Victor, 1987), but most of these studies concentrated on the dynamics of synaptic inputs into the cells, and less attention was directed to the dynamics of the spike initiation process.

Cockroach ocellar L-neurons do not discharge spontaneously, and a single cycle of voltage modulation produces either a single spike or very few spikes. The cockroach L-neurons probably encode signals in the form of a single spike or very few spikes; this is different from neurons in other visual systems, in which signals are, in most cases, encoded in the frequency of spike trains.

We examined the relationship between the slow potential and spikes of cockroach ocellar L-neurons, using a sinusoidally modulated light stimulus. We found that (a) the spike initiation process of the cockroach ocellar L-neuron is probabilistic (stochastic): the process has an internal noise; (b) the process has bandpass filtering properties; and (c) the process contains a dynamic linearity and a static nonlinearity. The observations suggest that a simple model, such as the integrate-and-fire model developed by Knight (1972a, b), may represent the actual spike initiation process. The final form of his model consists of a “forgetful integration” process, which is frequency dependent (dynamic) and linear, followed by a “stochastic firing” process, which is frequency independent (static) and nonlinear. Bryant and Segundo (1976) analyzed the spike initiation process of the *Aplysia* neuron and concluded that a similar model was quite accurate in predicting experimentally observed spike
discharges. Most recently, Sakuranaga et al. (1987) examined the spike discharge of catfish retinal ganglion cells and concluded that it can be modeled by an integrate-and-fire generator. These similarities among different preparations suggest that the integrate-and-fire model does represent the actual spike initiation process.

There is, however, some disagreement between our observations of the cockroach L-neuron and the integrator model. The forgetful-integrator model is nonadapting and always responds with tonic repetitive firing when stimulated with an adequate constant current. However, the cockroach L-neuron never generates tonic firing in response to a constant depolarizing current. This indicates that the spike initiation process of the L-neuron has a differential (or high-pass) property as well as an integrative (or low-pass) property. The model needs to include a bandpass filtering property in order to apply accurately to the actual spike initiation process of the cockroach L-neuron.

Knight (1972a, b) concluded that spike encoding in Limulus eccentric cells allows for a firing rate that is a replica of the shape of the stimulus. Sakuranaga et al. (1987) also concluded that signals contained in slow potentials remain substantially unchanged in the spike trains in catfish ganglion cells. In these neurons, little signal modification occurs during the spike initiation process. In the cockroach L-neuron, however, there are fundamental differences between the slow potential response and the spike response: (a) the slow potential response is linear, whereas the spike response is nonlinear; (b) the dynamics of the slow response are independent of the mean illuminance levels, whereas the dynamics of the spike response depend on the mean illuminance; and (c) the bandpass property of the spike response is sharper than that of the slow potential response. Therefore, we conclude that spike initiation in the L-neuron is an important step in visual signal processing. The function of the spike initiation process in the L-neuron is to filter out specific features from the slow potential signals (i.e., dimmings from the mean level), rather than to produce a replica of the slow potential.

There was a prominent enhancement of the low-frequency sensitivity in the spike response when the mean illuminance was low. A similar dependence of response dynamics (frequency dependence) on the levels of mean illuminance has been found in all the visual systems so far studied, including Limulus eyes (Fuortes and Hodgkin, 1964), insect compound eyes (Pinter, 1972; Dubs, 1981), and vertebrate retina (Naka et al., 1979, 1987; Tranchina et al., 1983; Chappell et al., 1985). In these visual systems, such a dependence is prominent at any level of the system, including photoreceptor cells, whereas in the cockroach ocellar system, the slow potential response of L-neuron exhibits no such dependence. It is a notable finding that in the cockroach ocellus, a coupling between the response dynamics and the mean illuminance levels is produced at the spike initiation process of L-neurons, not in the receptor cells.

The spike response of the L-neuron had a bandpass filtering property with an optimal frequency range of ~0.5–5 Hz, which reflects that both the slow potential response and the spike initiation (slow spike conversion) process had optimal frequency ranges of ~0.5–5 Hz. This optimal frequency range is much lower than that noted in vertebrate retinal ganglion cells (Victor and Shapley, 1979; Sakuranaga et al., 1987). The difference may reflect the fact that cockroach L-neurons are
designed to detect a change in the illumination averaged over a wide receptive field
(note that the 10,000 ocellar photoreceptors converge onto only four L-neurons),
whereas vertebrate retinal ganglion cells are designed to detect a small target within a
narrow receptive field.

The bandpass property of the spike initiation process apparently reflects the
activation and inactivation kinetics of the voltage-dependent current of the L-
neuron. If the rate of voltage change is slow with respect to the inactivation time
constant of the inward current, the conductance change for the inward current
would be low and no action potential would be generated. As the frequencies of
voltage modulation are increased, action potentials will be generated. However, at
high frequencies, when the rate of change of voltage is rapid with respect to the time
constant of activation of the inward current, there would not be sufficient time to
activate the inward current and thus no action potentials would be generated. In
short, an inactivation time constant limits the low-frequency response and an
activation time constant limits the high-frequency response. Another factor that
limits the low-frequency response is the activation time constant of the outward
current, because the effects of activation of the outward current are the same as for
the inactivation of the inward current. The optimal frequencies of the spike initiation
process of the cockroach ocellar L-neuron (0.5–5 Hz) are much lower than those of
the squid giant axon (50–80 Hz; Guttman et al., 1980). This difference indicates that
the activation time constant of the inward current of the cockroach L-neuron
(probably a calcium current; Mizunami et al., 1987) is slow in comparison with that of
the inward sodium current in squid giant axon.

The noise in the second-order visual neurons has been noted in barnacle ocelli
(Stuart and Oertel, 1978), Limulus eyes (Dodge et al., 1968), insect ocelli (Wilson,
1978b), insect compound eyes (Laughlin, 1973), and vertebrate retina (Ashmore and
Copenhagen, 1983). In the present study, we found that the noise facilitates spike
initiation; that is, it lowers the spike threshold. Noise has usually been discussed in
relation to the signal-to-noise ratio, in which the noise was thought to disturb signal
detection. Our finding suggests that, on the contrary, the noise has a positive role in
signal detection.

The noise in the L-neuron was derived from synaptic inputs from the photorecep-
tors. The underlying mechanism remains to be determined. Possible sources of the
noise are: (a) random photon capture of the photoreceptor cells, which results in a
discrete bump response (photon noise), (b) random isomerization of the rhodopsin
(transducer noise), and (c) random release of the transmitter substances from
photoreceptors (synaptic noise). The first is attributed to properties of light input
(extrinsic noise) and the latter two are intrinsic noise.

In the dark, the L-neuron was completely inactivated and no spikes were induced
by extrinsic depolarizing current. Apparently there was a high degree of steady state
inactivation in the light-adapted L-neuron, because the hyperpolarization resulting
from the steady illumination was relatively small (<3.5 mV).

The change in the mean potential levels was not accompanied by a change in the
spike threshold, which reflects the presence of a steady state inactivation. Function-
ally, this keeps the spike threshold unchanged over a wide range of mean illuminance.
A similar relationship between maintained potential and threshold depolarization has
been noted in squid giant axon in the presence of a large depolarizing DC bias, in which there might be a steady state inactivation of the inward current (Holden, 1976).

The final goal of our analysis was to answer the question of which features of the slow potential are relevant in spike initiation. We were successful in answering this question in a quantitative manner. However, the complete answer to this question needs a prediction of the actual spike response when the history of slow potential response is known. For a further prediction of the actual spike response, we need to develop a model for the spike initiation process.

In conclusion, we propose a simple model for signal processing in the cockroach ocellus (Fig. 12). Light signals that enter the ocellus are passed through a bandpass linear filter and produce a slow potential response in L-neurons. The linear filter consists of photoreceptors and synapses between photoreceptors and L-neurons. The details of its filtering properties have been discussed (Mizunami et al., 1986). The slow potential contains noise. The noise, whose exact origin remains to be determined, reflects that contained in the synaptic potential from photoreceptors.

The slow potential is further passed through a linear/nonlinear cascade and produces a spike discharge. The linear filter is bandpass, and the nonlinear filter is a static threshold. For details, see text.

The slow potential is passed through a linear/nonlinear cascade and produces a spike discharge. The linear filter is bandpass, having both a differential and an integrative nature. The nonlinear filter is a static threshold with a sigmoidal (probabilistic) input/output relationship. Although the effectiveness of the model in predicting the actual responses of cockroach ocellar neurons remains to be determined, the model will be a good base for further understanding signal processing in the cockroach ocellar system.

Ocellar L-neurons make output synapses onto several types of third-order neurons in the ocellar tract (Toh and Hara, 1984; Mizunami and Tateda, 1986). An important question that remains unanswered is whether the slow potential signals or the spike signals of L-neurons or both are encoded in the third-order neurons. Fig. 13 shows typical responses of two types of third-order neurons, called OL-I neurons (type I neurons projecting into the optic lobe) and D-I neurons (type I neurons descending to the thoracic ganglia; Mizunami and Tateda, 1986), to sinusoidal light stimulation. One type of third-order neuron, the OL-I neuron, had a spontaneous spike discharge and exhibited a modulation of the spike frequency around a mean (Fig. 13 A). The pattern of the response was similar to, although not the same as, that of the slow potential response of the L-neurons. The other type, the D-I neuron, had no
spontaneous spike activity and exhibited single spikes at the decremental phase of light modulation (Fig. 13 B). The pattern of the response was similar to that of the spike response of L-neurons. The observations suggest that both slow potential signals and spike signals in L-neurons are encoded into the third-order neurons. We conclude that the role of ocellar L-neurons is to produce two kinds of signals, linear (slow potential) and nonlinear (spike). The details of signal processing in third-order neurons will be dealt with in a future study.

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