pH Changes in the Invaginating Synaptic Cleft Mediate Feedback from Horizontal Cells to Cone Photoreceptors by Modulating Ca\textsuperscript{2+} Channels

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Abstract Feedback from horizontal cells (HCs) to cone photoreceptors plays a key role in the center-surround–receptive field organization of retinal neurons. Recordings from cone photoreceptors in newt retinal slices were obtained by the whole-cell patch-clamp technique, using a superfusate containing a GABA antagonist (100 \mu M picrotoxin). Surround illumination of the receptive field increased the voltage-dependent calcium current (I\textsubscript{Ca}) in the cones, and shifted the activation voltage of I\textsubscript{Ca} to negative voltages. External alkalinization also increased cone I\textsubscript{Ca} and shifted its activation voltage toward negative voltages. Enrichment of the pH buffering capacity of the extracellular solution increased cone I\textsubscript{Ca}, and blocked any additional increase in cone I\textsubscript{Ca} by surround illumination. Hyperpolarization of the HCs by a glutamate receptor antagonist-augmented cone I\textsubscript{Ca}, whereas depolarization of the HCs by kainate suppressed cone I\textsubscript{Ca}. From these results, we propose the hypothesis that pH changes in the synaptic clefts, which are intimately related to the membrane voltage of the HCs, mediate the feedback from the HCs to cone photoreceptors. The feedback mediated by pH changes in the synaptic cleft may serve as an additional mechanism for the center-surround organization of the receptive field in the outer retina.

Key words: retina • synapse • lateral inhibition

Introduction

The center-surround organization of the receptive field is one of the most important characteristics of vertebrate retinal neurons. Since its discovery by Kuffler (1953), many investigators have attempted to unravel the neural mechanisms underlying the center-surround antagonism. In 1971, Baylor et al. showed that the cone photoreceptors of the turtle responded by hyperpolarization to direct illumination of the recorded cone, but by delayed depolarization to surround illumination of an area comparable to the receptive field size of the horizontal cells (HCs). In a more direct experiment, Baylor et al. (1971) also showed that hyperpolarization of the HCs by extrinsic current injection depolarizes the nearby cones. Thus, it has been widely accepted that negative feedback from the HCs plays a key role in the center-surround antagonism of retinal neurons.

There is considerable evidence to suggest that the feedback from the HCs to cones is mediated by GABA. In many animal species, the HCs have been shown to be GABAergic; they have glutamic acid decarboxylase (GAD, Lam et al., 1979), a GABA-synthesizing enzyme. They accumulate GABA via a selective uptake mechanism (Lam and Steinman, 1971; Marc et al., 1978), and release (Schwartz, 1987) it when they are depolarized. It has also been shown in isolated turtle cones that these photoreceptors have a high sensitivity to the presence of GABA at their terminal (Tachibana and Kaneko, 1984; Kaneko and Tachibana, 1986). The presence of GABA receptors has also been shown in mammalian cones (Picaud et al., 1998; Pattnaik et al., 2000).

On the other hand, there are several reports that argue against the GABA feedback hypothesis, because the surround response of the cone persists regardless of the presence of GABA antagonists (Thoreson and Burkhardt, 1990; Burkhardt, 1993; Verweij et al., 1996) or agonists (Thoreson and Burkhardt, 1990; Burkhard, 1993). H3-type cone HCs in the teleost retina show a triphasic chromatic response; it is believed that the formation of the triphasic chromatic response is modulated by feedback from H2 HCs to green cones (Stell et al., 1975). However, in the goldfish retina, while GABA uptake into HCs could not be confirmed in H2/3 cone HCs, H1 HCs showed GABA uptake (Marc et al., 1978). Thus, the mediator of the triphasic feedback response of the H3 HCs remains unknown.

Modulation of the calcium current (I\textsubscript{Ca}) in cones have also been proposed as a mechanism for feedback. Verweij et al. (1996) reported that surround illumination shifts the activation voltage range of the cone I\textsubscript{Ca} in the goldfish retina. Feedback by modulation of the
cone $I_{Ca}$ is also consistent with the observation in the turtle retina that surround illumination causes calcium-dependent spikes in the cones (Piccolino and Gerschenfeld, 1978, 1980; Gerschenfeld and Piccolino, 1980). Furthermore, Verweij et al. (1996) also proposed that GABA is not likely to be the main mediator of this feedback mechanism, since this modulation of the cone $I_{Ca}$ was resistant to GABA receptor and GABA transporter antagonists.

Byzov and Shura-Bura (1986) proposed an electrical (ephaptic) feedback mechanism by which a local current in the cone-HC synapse causes a voltage drop in the cone synaptic clefts and results in modulation of the transfer function between the cones and bipolar cells. More recently, Kamermans et al. (2001) suggested that an ephaptic effect of the current flowing through the hemigap channel of the dendritic tips of the HCs shifts the activation voltage of the cone $I_{Ca}$, based on evidence that (a) hemichannel proteins exist in the dendritic tips of HCs and (b) carbenoxolone, a blocker of hemichannels, suppresses the surround response of the cones. However, the validity of the hemichannel-mediated feedback hypothesis is still under debate, since the pharmacological specificity of carbenoxolone is uncertain.

This study was conducted with the aim of examining the mechanisms of feedback from HCs to cone photoreceptors. The surround responses of cone photoreceptors in newt retinal slices were recorded with whole-cell patch-clamp recording. We found that surround illumination shifts the activation voltage of the cone $I_{Ca}$ toward negative voltages, as reported by Verweij et al. (1996), and that the current-voltage relation of $I_{Ca}$ did not merely shift along the voltage axis as suggested by an ephaptic effect, but was very similar to the changes caused by alkalinization of the extracellular space. Enrichment of the pH buffering capacity of the extracellular solution increased cone $I_{Ca}$ and suppressed the light-induced surround effect. Hyperpolarization of the HCs induced by CNQX increased cone $I_{Ca}$, whereas depolarization of the HCs induced by kainate decreased cone $I_{Ca}$. The present data thus strongly suggest that feedback from HCs to cone photoreceptors is mediated by pH changes in the synaptic cleft.

**MATERIALS AND METHODS**

**Preparation of Retinal Slices and Superfusion**

The experimental procedures conformed to the Guidelines for the Care and Use of Laboratory Animals, Keio University School of Medicine, and our experiments were approved by the University Animal Welfare Committee. Retina specimens obtained from newts (Cynops pyrrhogaster, 10–12 cm in length) were cut into slices by a method originally reported by Werblin (1978), as follows: The animals were kept in a room maintained under a natural 12 h/12 h light/dark cycle. Before the experiment, the newts were dark-adapted for 2 h. Subsequent manipulations were performed under dim red light. The animals were decapitated, the eyes enucleated and hemisected, and the frontal part, including the lens, was removed. The retina was detached from the eye-cup on to a piece of filter paper (pore size 0.45 or 0.80 μm; Toyo Roshi, Ltd.) and sectioned into 300-μm thick slices using a custom-made tissue slicer. Slices were superfused with Ringer’s solution containing (in mM): 110 NaCl, 2.6 KCl, 2 NaHCO$_3$, 3 CaCl$_2$, 10 glucose, adjusted to pH 7.4, saturated with a 95% O$_2$–5% CO$_2$ gas mixture. In this study, experiments were performed with perfusates containing 100 μM picrotoxin, to exclude any confounding effects by GABA-activated chloride currents in the cones (Kaneko and Tachibana, 1986), although picrotoxin did not change any current whose reversal potential was equivalent to the equilibrium potential of the chloride ions or suppress the surround responses of the cones in the retinal slices (three cones: unpublished data). 6-Cyano-7-nitroquinazoline di-sodium (CNQX, an antagonist of non-NMDA receptors) and kainate were added directly to the superfusate in some experiments. The pH of the HEPES- and Tris-containing solutions was adjusted to the same value (within 0.1 pH unit) as that of Ringer’s solution, using NaOH or HCl. The tonicity of the solutions was adjusted with sucrose. All the drugs, unless otherwise specified, were purchased from Sigma-Aldrich. The volume of the recording chamber was 0.75 ml, and the superfusate flowed continually at the rate of 0.9 ml/min. Plastic tubing (Tygon, Norton Co.) and Teflon tubing were used as conduits for the solutions, to minimize the loss of CO$_2$. The chamber containing the retinal slices was mounted on the stage of an upright microscope (BX50WI; Olympus), which was equipped with an infrared (>800 nm) illumination system and a viewing system with an CCD camera (ICD-47AC; Ikegami Tsushinki Co., Ltd.). The microscope was mounted on a Gibraltar platform and an X-Y stage (Burleigh Instruments, Inc.). The images of the slices were monitored on a CRT display. All the experiments were conducted in a dark room maintained at a room temperature of 22°C.

**Light Stimulation**

Light stimuli were supplied to the retinal slices from two 100-W halogen lamps. A small spot light with a diameter of 30 μm was projected through the 40× water immersion objective lens and a diffuse light with a diameter of 3,500 μm was projected through the condenser lens. For the recordings made from the bipolar cells (BCs), the diameter of the spot light was increased to 50 μm, to cover the entire receptive field center. The timing and duration of illumination with the spot light and diffuse light were controlled by an electrical shutter (Melles Griot) interposed in each of the light paths and controlled by computer. The shutter opened from 10% to 90% in ~20 ms. The light intensity was attenuated by inserting a neutral-density filter and by adjusting the supply voltage to the lamp. The intensity of the small spot light was $3 \times 10^{-5}$ μW/μm$^2$, which evoked a saturating response in the cones, but did not bleach them completely. The intensity of the large spot light was $5 \times 10^{-6}$ μW/μm$^2$, which evoked the maximum surround response, but did not affect the state of adaptation. The relative intensity of the 450-nm spot light was ~27% that of the 650-nm spot light, and 10% that of the 650-nm diffuse light. The spectrum of the light was measured with a monochromator (Shimadzu), and the intensity was measured with a calibrated silicon photodiode (Centronic). To stimulate the receptive field surround, a large (3- to 4-mm diameter) spot was superimposed concentrically over a small (30-μm diameter) steady light spot. These stimuli will hereafter be referred to as surround illumination, and the responses evoked by such stimuli, as surround responses.

**pH-mediated Feedback to Cone Photoreceptor**
Newt cones express three kinds of opsins, Cp-LWS, SWS-1, and SWS-2 (Sakakibara et al., 2002). Thus, three kinds of cones (red-, blue-, and ultraviolet-sensitive cones) are expected. In this study, we identified the subtype of 39% of the cones (70/179), according to their responses to red, green, and blue light–emitting diodes (LED’s; DHR6610, SBY 5710, and GNB4710; Iwasaki Electric Ltd.). The remaining 109 cones could not be classified. However, since we did not find any differences in the responses among the cone subtypes, all the cones were treated equally for the data analyses.

**Whole-cell Recording**

Pipettes for whole-cell recordings were fabricated from standard-wall borosilicate glass tubing, and their tips were heat-polished. The Cs+-based pipette solution contained (in mM): 90 Cs methanesulfonate (CsMeSO_4_), 10 TEA-Cl, 20 BAPTA, 10 HEPES, 1 MgCl_2, 2.5ATP-Mg, 1 GTP-Na, 1 cGMP, 10 phosphocreatine, and 50 μM creatinephosphokinase, adjusted to pH 7.3 with CsOH. Since cone photoreceptors have a high voltage–activated (HVA) Ca^2+ current and Ca-activated K^+ and Cl^- currents (Barnes, 1994), we added 20 mM BAPTA to minimize the Ca-activated currents in the cones.

The resistance of the filled pipettes was usually in the range of 9–12 MΩ. The cells were usually voltage clamped at −40 mV. The mean input resistance of the cones in the saturating spot light, calculated from the linear leak conductance, was 686 ± 7 MΩ (mean of 47 cells analyzed). The patch pipettes were connected to an Axopatch 1-D amplifier (Axon Instruments, Inc.). No series resistance compensation was employed. Signals were low-pass filtered (Bessel filter) at 2 kHz, sampled (12 bit) at 5 kHz, and then stored on the hard disk of a personal computer.

**Isolation of the Calcium Current in Cones**

The I_Ca of the cones was identified as the current selectively blocked by 3 mM cadmium (Cd). The I-V relations of I_Ca were obtained by subtracting the I-V curves recorded in a 3-mM Cd-containing solution from those recorded in control solution (four cones). Another method used to obtain the I-V relations of I_Ca was by subtraction of the leakage current. The leakage current was estimated by extrapolating the linear part of the I-V curve, between −50 and −36 mV. Since the I-V curve obtained by the leakage subtraction method showed good agreement with that obtained by the Cd-subtraction method (Fig. 2, boxed inset), we determined the I-V relations of I_Ca by the leakage subtraction method in the subsequent experiments.

**Analysis**

The digitized data were analyzed and plotted using Origin 6.1 software (Microcal Ltd.). The digitized waveform data were finally low-pass filtered (<1 kHz) with an FFT-smoothing algorithm. The statistical data are presented as means ± SEM. The statistical significance of the differences among the data was tested by Student’s t test.

**RESULTS**

**Response of Cone Photoreceptors in Newt Retinal Slices to Surround Illumination**

A voltage-dependent surround response of the cones in newt retinal slices was obtained in the current-clamp mode (Fig. 1). Spot illumination hyperpolarized the cones, while surround illumination depolarized them (Fig. 1, middle trace, no extrinsic current injection). The size of the surround response was dependent on the membrane voltage. Hyperpolarization of the cones by extrinsic current injection (−0.03 nA current injection) suppressed the surround response without reducing the amplitude of the response to spot illumination. Depolarization (+0.03 nA) of the cones also reduced the size of the surround response. The amplitude of the surround response was maximal at around −30 mV. Cones that were hyperpolarized up to −50 mV by spot illumination did not show any surround response, but the surround response appeared when the membrane voltage was brought to near −30 mV by extrinsic current injection (unpublished data).

A voltage-dependent calcium current (I_Ca) in the newt cone was activated by depolarization to voltages more positive than −30 mV, similar to activation of I_Ca in tiger salamander rods (Barnes et al., 1993). The I-V curve of the cone I_Ca was obtained by the linear leak current subtraction method. Under voltage-clamp recording, surround illumination evoked an inward current at voltages more positive than −30 mV, while no inward current was evoked at voltages more negative than −40 mV (Fig. 2 A). Surround illumination augmented the cone I_Ca measured in the presence of spot illumination by subtraction of the leakage current. The leakage current subtraction method showed good agreement with that obtained by the Cd-subtraction method (Fig. 2, boxed inset), we determined the I-V relations of I_Ca by the leakage subtraction method in the subsequent experiments.

**Figure 1.** The response of a newt cone photoreceptor recorded in the current-clamp mode. The outer segment of the cone was illuminated by a spot (diameter, 30 μm; duration, 3,380 ms; timing indicated by the top horizontal line). A diffuse light (diameter, 4,000 μm; duration, 1,250 ms) was superimposed on the spot as indicated by the shorter horizontal line. The retinal slice was superfused with control Ringer’s solution buffered with bicarbonate and containing 100 μM picrotoxin. Under control condition (when no current was injected from the recording pipette: 0 nA), illumination with the spot evoked hyperpolarization, and the surround illumination evoked depolarization in the cone. Both hyperpolarization and depolarization of the cone induced by current injection (−0.03 and +0.03 nA) from the recording pipette abolished the surround response. The vertical scale on the left indicates the absolute membrane voltage. Recovery at the spot offset was slow (1 s), probably due to blockade of the calcium feedback to the phototransduction cascade in the cones (Lamb et al., 1986; Nakatani and Yau, 1988), because the intracellular Ca^{2+} level was maintained at a low level due to the addition of 20 mM BAPTA in the pipette solution. Lowering the BAPTA concentration in the pipette solution (5 mM) accelerated the recovery (0.5 s; unpublished data).
Figure 2. Surround illumination augments the cone Ca\(^{2+}\) current A. The Ca\(^{2+}\) current (I\(_{Ca}\)) in the cone photoreceptors of the newt retinal slice was recorded under the whole-cell voltage clamp condition. The retinal slice was superfused with control Ringer’s solution buffered with bicarbonate and containing 100 μM picrotoxin. The cone was held at −40 mV and polarized to voltages ranging from −50 mV to +8 mV in 2-mV steps. Five representative traces, voltage-clamped at −40, −26, −24, −16, and −4 mV, are shown. During the command voltage, surround illumination (diameter, 4,000 μm; duration, 400 ms; shorter bar) was applied every 4 s, while the spot illumination (diameter, 30 μm; top bar) remained. An additional 2-mV depolarization was applied to mimic an ephaptic effect (external voltage drop) after withdrawing the surround illumination. Note that at −4 mV (pink trace); surround illumination evoked an inward current, while a 2-mV pulse evoked an outward current. The current amplitude was sampled at the time indicated by the symbols, to construct the IV curves shown in B a and B b. (B a) Leak-subtracted IV curve of the cone I\(_{Ca}\) in the presence of the spot (filled squares) and during surround illumination (open squares). The data are from the same cone as in A. The leakage current amplitude (conductance, 0.59 nS), determined by extrapolation of the linear portion of the IV curve between −50 and −32 mV, was subtracted from the measured current amplitude at each voltage. Inset shows activation curves fitted to the Boltzmann function derived from the IV curves. The midpoint of the curve as obtained under the control condition (−15.5 mV; black line) was shifted by 1.8 mV in the negative direction during surround illumination (red line). The maximum conductance was calculated from the slope of the IV curve between +2 and +8 mV and normalized to 1.0. (b) Leak-subtracted IV curve of the cone I\(_{Ca}\) in the presence of the spot light (filled circles) and during a 2-mV depolarizing pulse (open circles). Inset shows activation curves fitted to the Boltzmann function derived from the IV curves. The midpoint of the curve as obtained under the control condition (−16.2 mV; black line) was shifted by 2.3 mV in the negative direction during the 2-mV induced depolarization (red line). The 0.3 mV discrepancy in the curve shifting is probably due to a curve fitting error or a voltage clamp error. (Boxed inset) Isolation method of IV relations of cone ICa. These data were obtained from a different cone in A. Top) The IV relations of the cone were obtained in the control solution (filled squares (1)) and in a 3 mM Cd-containing solution (open squares (2)). The leakage conductance (2.7 nS) was estimated by extrapolating the linear part of the IV curve between −50 and −36 mV (solid line (3)). Bottom, open circles) IV relations of the cone ICa obtained by subtracting the IV curve recorded in a 3 mM Cd-containing solution from that recorded in the control solution ((1) − (2)). (filled circles) IV relations obtained by subtracting the IV curve from the extrapolated leakage current from that recorded in the control solution ((1) − (3)).

Illumination at all holding voltages (Fig. 2 B a). This augmentation was voltage dependent; greater augmentation was seen at voltages more negative than −15 mV, at which the standing inward current was maximal, whereas little augmentation was seen at voltages between 0 and +10 mV. In 24 cones sampled, surround light illumination shifted the midpoint of the cone I\(_{Ca}\) activation curve by −2.55 ± 0.32 mV, within the range of −6.5 and −0.6 mV. These data suggest that surround illumination augmented cone I\(_{Ca}\) and shifted its activation voltage, similar to the observations in goldfish cones (Verweij et al., 1996). The cone surround response disappeared after rundown of the cone I\(_{Ca}\) (unpublished data), which also suggests that the cone surround response is cone-I\(_{Ca}\) dependent. Even in the absence of picrotoxin, surround illumination did not evoke any current whose reversal potential was equal to the equilibrium potential of chloride ions (three cones).

It was hypothesized recently that a current flowing into HCs creates an ephaptic effect (a field effect) that causes a drop of the voltage in the intersynaptic cleft at the cone terminal, resulting in an enhancement of the cone I\(_{Ca}\) (Kamermans et al., 2001). The ephaptic effect would be expected to shift the voltage dependence of the cone I\(_{Ca}\), parallel to the voltage axis, in the negative direction. To mimic the ephaptic effect, the cones were depolarized by 2 mV after switching off the surround illumination. During the 2-mV depolarization, the IV curve of the cone I\(_{Ca}\) clearly shifted by 2 mV in the neg-
ative direction (Fig. 2 B b). The change was clearly different from that induced by surround illumination. A difference was especially apparent at membrane voltages between −15 and +8 mV, where surround illumination increased the current amplitude (while 2-mV depolarization decreased it). The differences between these two conditions led us to wonder whether the surround response of the cones is mediated by a mechanism other than the ephaptic effect.

**pH Change Modulates the Cone Current in a Voltage-dependent Manner**

It has been demonstrated that extracellular acidification regulates the voltage-dependent Ca\(^{2+}\) current (I\(_{\text{Ca}}\)) (Tombaugh and Somjen, 1998) by several mechanisms, including (a) charge-screening effects of protons on the Ca channels (Krafte and Kass, 1988; Prod’hom et al., 1989; Klöckner and Isenberg, 1994) and (b) proton-binding effects on the Ca channels (Chen et al., 1996; Zhou and Jones, 1996). The effects of pH change on cone-I\(_{\text{Ca}}\)-influenced transmitter release have been examined (Barnes and Bui, 1991; Barnes et al., 1993; DeVries, 2001). We therefore confirmed the effects of extracellular pH change on the cone I\(_{\text{Ca}}\) in the newt retina. Pressure ejection of high pH Ringer’s (pH 9.0) solution onto the outer plexiform layer enhanced an inward I\(_{\text{Ca}}\). The high pH-induced inward current was voltage dependent; the inward current was more pronounced at voltages between −10 and −30 mV (Fig. 3 A). Thus, the external alkalinization augmented the peak of the cone I\(_{\text{Ca}}\) (Fig. 3 B, top; +39.5 ± 8.5%, n = 6), as reported in other vertebrate cones (Barnes and Bui, 1991; DeVries, 2001). Moreover, the midpoint of the activation curve of the cone I\(_{\text{Ca}}\) was shifted in the negative direction (Fig. 3 B, bottom; 8.2 ± 2.0 mV, n = 6), similar to the effects of surround illumination (Fig. 2 B a). Voltage-dependent calcium channels are localized in the terminal regions of the photoreceptors (Morgans et al., 1998; Nachman-Clewner et al., 1999), and are thus presumably localized near the center of the invaginating synapses. Therefore, it could be considered that surround illumination alkalinized the intersynaptic clefts of the invaginating synapses, to augment the cone I\(_{\text{Ca}}\) locally.

**Strong pH Buffering Suppresses the Surround Response of the Cones**

If pH changes in the synaptic clefts were considered to mediate the surround response of the cones, fixation of the extracellular pH by enrichment of the pH-buffering capacity of the external solution would be expected to prevent the generation of the cone surround response. In fact, changing the superfusate from control Ringer’s solution to one supplemented with 10 mM HEPES (having an identical pH value) increased cone I\(_{\text{Ca}}\) and prevented any additional inward current potentially caused by surround illumination (Fig. 4 A). The addition of 10 mM HEPES (pKa = 7.5) was estimated to increase the buffering capacity of the solution by 5.75 mM at pH 7.4 (Chesler, 1998). The effect of HEPES enrichment of the superfusate on the membrane current of the cones was examined at several holding voltages, from −50 mV to +6 mV. When cone I\(_{\text{Ca}}\) was not activated, however, HEPES had no significant effect on the membrane conductance (traces at −34 mV in Fig. 4 B a). The use of HEPES-enriched solution reversibly increased cone I\(_{\text{Ca}}\) and prevented any additional inward current potentially evoked by surround illumination (traces at −18 and −10 mV in Fig. 4 B a). Such changes in I\(_{\text{Ca}}\) were seen throughout the entire voltage range at which I\(_{\text{Ca}}\) was activated (Fig. 4 B a).
b). As a result, use of HEPES (10 mM) shifted the Boltzmann-fit activation curve of cone $I_{Ca}$ by 1.8 mV in the negative direction (Fig. 4 B c), similar to the shift induced by surround illumination of the retinal specimens in control Ringer’s solution (1.6 mV). When HEPES-enriched solution was used, surround illumination induced little additional shift of the activation curve. On average, use of HEPES (10 mM)-enriched buffer shifted the activation curve of cone $I_{Ca}$ by 1.2 ± 0.3 mV in the negative direction ($n = 5$), which was al-
most identical to the shift caused by surround illumination of the retinal specimens in control solution (1.1 ± 0.3 mV), with peak I\(_{Ca}\) increased by 15.5 ± 1.0%. HEPES did not block the light response of the HCs to diffuse light illumination (Fig. 4 A, inset, \(n = 4\)). In 10-mM HEPES-enriched buffer, the surround responses were suppressed completely in three out of eight cones, and partially in the remaining five cones at −26 mV (Fig. 4 B b, inset, \(V_h = −26\) mV). In 20-mM HEPES-enriched buffer, the surround response was suppressed in the four cones tested.

HEPES, MES, and TAPS, commonly used “Good buffers,” have an aminosulfonate moiety, and it has been reported that these protonated aminosulfonate compounds decrease the permeability of connexin-26 hemichannels (Bevans and Harris, 1999). To exclude the possibility that HEPES closes the connexin-26 hemichannels on HCs to cause blockade of the ephaptic effect (Kamermans et al., 2001), the effects of Tris buffer that does not possess an aminosulfonate moiety were also examined. Tris (15 mM)-supplemented external solution as the superfusate also enhanced cone I\(_{Ca}\).
and suppressed any potential additional increment of $I_{Ca}$ caused by surround illumination (Fig. 5 A a), indicating that the effects of pH buffering on cone $I_{Ca}$ and surround response were not unique to HEPES. When large surround responses of 20 pA were recorded in the control solution, diffuse light elicited a current dip at the time of switch-off of the illumination (but not at the beginning) when the specimens were superfused with enriched buffer solution (arrow in Fig. 5 A). The addition of 15 mM Tris (pKa = 7.8) was estimated to increase the buffering capacity by 8.59 mM at pH 7.4. On average, Tris-enriched buffer as the perfusate shifted the activation curve of the cone $I_{Ca}$ by 3.7 ± 0.7 mV, which was almost identical to the shift caused by surround illumination of the retinal specimens in control solution (3.3 ± 0.6 mV) (Fig. 5 A, c), with peak $I_{Ca}$ increased by 26.6 ± 1.8% ($n = 4$). In Tris-enriched buffer, surround light illumination did not evoke any additional shift of the I-V curve of cone $I_{Ca}$. Tris-enriched buffer solution increased the amplitude of the light response of the HCs (by 32 ± 11%; $n = 3$) (Fig. 5 B).

When the leak conductance of the cones exceeded 2 nS, the inward $I_{Ca}$ could be masked by an outward leak current. Thus, in some cones, the net current before subtraction appeared to be moderately outward (Fig. 4 B a, control [at −18 mV]), or slightly inward (Fig. 5 A a, control [at −18 mV]).

HEPES did not change the spatial summation of the HCs. The ratios of the amplitudes of the HC light responses to spot (50-μm diameter) versus diffuse lights (4,000-μm diameter) were 0.78 ± 0.08 in control Ringer’s solution, and 0.77 ± 0.10 in HEPES-enriched buffer ($n = 4$) (unpublished data). Tris had no significant effect on the spatial summation of the HCs, either. Thus, suppression of the surround response of the cones by solutions with a high buffering capacity cannot be due to a reduction of the spatial summation of the HCs.

Hare and Owen (1998) studied the effects of substituting bicarbonate for HEPES in the perfusate solution on the light responses of the retinal neurons. The substitution with an external pH buffer could have changed the intracellular pH-buffering capacity in several retinal neurons, because HEPES buffer is membrane impermeable. In contrast to the experiments conducted by Hare and Owen (1998), we examined the effects of supplementation of either HEPES or Tris (with identical pH), using an identical protocol, only allowing for a change of the pH-buffering capacity in the extracellular space.

**Horizontal Cells Mediate the Surround Response of Cones**

It is widely accepted that the surround response of cones has a close relationship with the membrane voltage of the HCs. The preceding experiments strongly suggest that the pH in the invaginating synaptic clefts is modified by the membrane voltage of the HCs. To verify this hypothesis, we performed two experiments. First, we examined the effect of a small (0.25 mm) diameter surround light, since it is known that the response amplitude of HCs depends on the illuminated
area of the retina. In the presence of a small spot (30-
\(\mu m\) diameter), the surround response of the cones to a
small diffuse light became smaller than that to a large
(4 mm) diffuse light (Fig. 6 A). As a result, cone IC\(_a\) be-
came smaller (Fig. 6 B) and the activation curve of IC\(_a\)
shifted in the positive direction (Fig. 6 C). The re-
sponse of the HCs to a small surround light also be-
came smaller (top right). This observation is a
good indication that the surround response of the
cones depends strongly on the membrane voltage of
the HCs (Baylor et al., 1971).

In the second experiment, we examined the effects
of pharmacological depolarization and hyperpolariza-
tion of the HCs. 20 \(\mu M\) kainate, an agonist of the
postsynaptic glutamate receptors of the HCs, depo-
larized the HCs from their dark membrane voltage
(\(-49.7 \pm 6.4 \text{ mV}\)) to \(-1.3 \pm 0.8 \text{ mV}\), and blocked their
light responses (bottom Fig. 7 A, \(n = 3\)), similar to the
responses of HCs of the tiger salamander (Yang and
Wu, 1989).

Kainate reversibly suppressed cone IC\(_a\) (Fig. 7 A) with-
out any significant effect on the leak conductance (from

![Diagram](https://example.com/diagram.png)
Kainate also suppressed the surround response of the cones (Fig. 7 B a, inset). The midpoint of the Boltzmann-fit activation curve of cone ICa shifted by 3.9 mV in the positive direction (Fig. 7 B b). The effect of kainate on cone ICa cannot be a direct effect on the Ca channels of the cones, since subsequent application of CNQX in the presence of kainate not only restored ICa, but eventually increased it to more than its original value (n = 2). Furthermore, subsequent application of 10 mM HEPES in the presence of kainate also restored the cone ICa (n = 2), lending support to the assumption that the effect of kainate is due to extracellular acidification.

In contrast to the effects of kainate, 20 μM CNQX hyperpolarized the HCs from -42.5 ± 7.8 mV to -59.7 ± 9.0 mV (n = 3; bottom of Fig. 8 A), and blocked their light responses. In cones, CNQX augmented ICa to such an extent that surround illumination did not induce any additional increase in ICa (Fig. 8 B a). Similar results with CNQX have also been reported in goldfish cones (Verweij et al., 1996). CNQX had no significant effect on the linear leak conductance. CNQX shifted the midpoint of
HEPES (10 mM)-enriched solution enhanced the maintained inward current (enhancement of the maintained inward current), while the surround illumination evoked an inward current (suppression of the maintained inward current). The current was recorded under the voltage-clamp conditions (holding voltage, −40 mV). In control Ringer’s solution, spot illumination evoked an outward current (suppression of the maintained inward current), while the surround illumination evoked an inward current (enhancement of the maintained inward current). The HEPES (10 mM)-enriched solution enhanced the maintained inward current (from −12 to −14 pA) and abolished the surround response. The transient inward current at the time of switch-off of spot illumination was prominently increased in HEPES buffer. The two traces were obtained from the same cell. (B) The same recordings as those shown in A in an expanded time and amplitude scale. Horizontal bars indicate the timing of the surround illumination.

**Figure 9.** Center and surround responses of an off-type bipolar cell recorded in a solution enriched with HEPES. (A) The current responses to a spot light (50 μm in diameter, indicated by the long horizontal bar above the response trace, duration 1,560 ms) and to surround illumination (diffuse light, 4,000 μm in diameter, superimposed on the spot light, duration 400 ms) of an off-type bipolar cell recorded under the voltage-clamp conditions (holding voltage, −40 mV). In control Ringer’s solution, spot illumination evoked an outward current (suppression of the maintained inward current), while the surround illumination evoked an inward current (enhancement of the maintained inward current). The HEPES (10 mM)-enriched solution enhanced the maintained inward current (from −12 to −14 pA) and abolished the surround response. The transient inward current at the time of switch-off of spot illumination was prominently increased in HEPES buffer. The two traces were obtained from the same cell. (B) The same recordings as those shown in A in an expanded time and amplitude scale. Horizontal bars indicate the timing of the surround illumination.

the Boltzmann-fit activation curve of the cone ICa by 6.0 mV in the negative direction (Fig. 8 B).

These observations support the contention that the surround response of cones is closely related to the membrane voltage of HCs. Depolarization of the HCs had the effect of suppressing cone ICa, and hyperpolarization of the HCs had the effect of enhancing cone ICa. Taken together with the results of experiments in which the membrane voltage of the HCs was controlled, it is tempting to speculate that the membrane voltage of the HCs controls the pH in the synaptic clefts of the cone terminal.

**Strong pH Buffering Suppresses the Surround Responses of the Bipolar Cells**

It is widely considered that feedback from the HCs to the cones is involved in the center-surround organization of the bipolar cell (BC) receptive field. Do pH changes in the synaptic clefts also generate receptive field organization in the BCs?

To answer this question, we examined the effects of HEPES-enriched buffer on the surround response of off-type BCs under voltage-clamp. When retinal slice preparations were superfused with control Ringer’s solution, spot illumination (50-μm diameter) evoked a sustained outward current in the BCs held at −40 mV. Surround illumination elicited an inward current, suggesting a surround-induced depolarization of the off-type BCs (Kaneko, 1970). Changing the superfusate to HEPES-enriched buffer solution slightly increased the holding inward current in the dark (mean increase by −2.6 ± 0.7 pA; n = 3), and suppressed the surround response (Fig. 9 A). The transient inward current at the time of switch-off of the spot illumination was prominently increased (n = 3). This effect of the HEPES-enriched buffer on the surround response was reversible (Fig. 9 B). Similar results were obtained from three off-type BCs.

**DISCUSSION**

We examined the mechanisms underlying the generation of the surround response in cone photoreceptors in retinal slice preparations of the newt. Under current-clamp recording, a surround response appearing at around −20 mV was suppressed by both depolarization and hyperpolarization of the membrane voltage, which is consistent with a number of previously published reports that have suggested that the surround response of the cone photoreceptors is voltage dependent (turtle, O’Bryan, 1973; tiger salamander cones, Skrzypek and Werblin, 1983; Wu, 1991). Current flow through gap junctions between neighboring cones (DeVries et al., 2002) did not generate a surround response in the present study, because the amplitude of the surround response was independent of the difference between the membrane voltage and the presumed resting potential of neighboring cones. In the case of voltage-clamp recording, the surround response appeared in the voltage range in which the cone ICa was activated, which indicates that the surround response of the cones is closely related to ICa. The contribution of cone ICa to the surround response has been demonstrated previously in turtle (Piccolino and Gerschenfeld, 1978, 1980; Gerschenfeld and Piccolino, 1980; Thoreson and Burkhart, 1991) and goldfish (Verweij et al., 1996; Kraaij et al., 2000) retinas. In this study, we focused on the mechanism by which surround illumination triggers activation of cone ICa.

We found that surround illumination increased cone ICa and shifted the activation voltage of ICa in the negative direction. External alkalization also evoked a similar effect on cone ICa. Thus, we considered that pH changes in the cone synaptic clefts might mediate feed-
back from the HCs to cones by modulating Ca\(^{2+}\) channels.

**Feedback Mediated by pH Changes in the Outer Retina**

Several factors have been proposed as candidate feedback messengers from the HCs to cones. These include substances that modulate ICa of the cones, such as nitric oxide, glutamate, chloride, protons, and the ephaptic effect caused by current flowing into the HCs (for review see Kamermans and Spekreijse, 1999). Enrichment of the pH buffering capacity of the extracellular solution increased cone ICa and prevented any additional increase in ICa produced by surround illumination. Thus, we consider that protons are the most likely candidates among these substances. Moreover, HC depolarization by kainate suppressed cone ICa, whereas HC hyperpolarization by CNQX increased it. Therefore, it is highly plausible that the pH in the cone synaptic cleft is tightly controlled by the membrane voltage of the HCs.

The most plausible interpretation for the effects of enrichment of the pH buffering capacity is that the surround response of the cones is modulated by pH changes in the invaginating synaptic clefts, and thus the following sequences are considered. In a normal bicarbonate-buffer solution, pH in the synaptic cleft is slightly more acidic (by about pH 0.2; estimated in Discussion) than the pH in the large extracellular pool (pH 7.4). Thus, cone ICa is relatively suppressed. The HC hyperpolarization caused by surround illumination neutralizes this acidic condition, thus restoring cone ICa and the cone surround response. On the other hand, in an external solution with a high buffering capacity, pH in the synaptic cleft is already fixed to the same pH as that of the extracellular pool (pH 7.4). Therefore, the enhancement of cone ICa by surround illumination does not occur.

The assumed pH change of 0.2 U may be a large enough change for pH electrode to detect. Therefore, we measured the pH in the outer plexiform layer by inserting a pH-selective microelectrode (tip diameter: 2 \(\mu\)m) into the outer synaptic region in the retinal slices. The fabrication method of the microelectrode used was that described by Smith et al. (1999) and Molina et al. (2000). The smallest pH change that our pH micropipette could detect was 0.06 pH U. However, no pH changes were detected after the addition of kainate (synaptic clefts should be acidified) and CNQX (the clefts should be alkalinized) to the superfusate in 23 slices examined. However, the absence of detectable pH change may not mean that the pH of the synaptic clefts is fixed. Insertion of the pH-sensitive micropipette may open up the narrow space at the invaginating synapse and make a wide artificial channel to the bath of the recording chamber. Through this artificial channel, any local pH change, that might be large enough within a narrow space, may become undetectable.

Recently, DeVries (2001) demonstrated that exocytosed protons from the cone terminal mediate a feedback to block the cone ICa. The exocytosed protons can form a negative feedback loop to control glutamate release in a sustained manner. In contrast, synaptic acidification caused by HC depolarization can depend on the illumination area of the surround light, and thus contribute to receptive field surround formation in the outer retina.

If the pH in the synaptic cleft is controlled by the membrane voltage of the HCs, the mechanism by which surround illumination augmented the cone ICa can be interpreted easily. Kamermans et al. (2001) proposed hemichannel-mediated ephaptic feedback in the goldfish retina, based on following evidence. (a) The presence of hemichannel proteins in the dendrite tips of the HCs, and (b) the blocking effect of carbenoxolone, a blocker of hemichannels, on the feedback responses. If the ephaptic effect modulates cone ICa, surround illumination should be expected to shift the I-V curve of the cone ICa at all voltages (like in Fig. 2 B b). However, importantly, surround illumination never induced a parallel shift of the I-V curve of the cone ICa in the negative direction; rather it increased ICa even in the region of the positive slope of the I-V curve of the cone ICa (between −10 and 10 mV; Fig. 2 B a), which is inconsistent with the ephaptic feedback hypothesis.

Protonated aminosulfonate compounds, including HEPES, have been reported to decrease the permeability of connexin-26 hemichannels, but compounds without an aminosulfonate moiety, such as Tris, maleate, and bicarbonate, do not decrease the permeability of these channels (Bevans and Harris, 1999). The effects of HEPES on the surround response are not likely to be due to blockade of the connexin-26 hemichannels in HCs (Kamermans et al., 2001), because Tris also suppressed the surround responses. Furthermore, there are other phenoma which seem to lend great support to the pH-feedback hypothesis. First, in five of eight cones, 10 mM HEPES partially blocked the surround response at −26 mV (inset in Fig. 4 B b). Since cone ICa was most sensitive to external pH change at −26 mV (Fig. 3), even a slight change in pH was detectable. Second, when large surround responses of 20 pA were recorded in the control solution, diffuse light occasionally elicited a current dip at the offset of illumination (but not at the beginning) in the enriched buffer solution (Fig. 5 A, arrow). This may be interpreted to mean that the rate of proton buffering by Tris did not match up well with the rate of the sudden increment of protons at the time of switch-off of the surround light.
In this study, surround illumination shifted the activation curve of the cone ICa by ~2.5 mV, which corresponded to the alkalinization-mediated shift of pH by ~0.2 (estimated from Barnes et al., 1993) in a pH 7.4 external solution. In retinal slices, the surround response may be weakened due to the reduction of the receptive field size. Thus, the pH change by surround illumination in the retinal slice may be an underestimate of the value prevailing in vivo. In an isolated retina of the goldfish, surround illumination shifted the I-V curve of the cone ICa by 7.5 mV (Verweij et al., 1996), which corresponds to an alkalinization of ~0.7 pH U. It is possible that the surround illumination induced alkalinization by 0.2 pH U in the bicarbonate buffer solution. In the retina, light stimulation evokes alkalinization of up to 0.2 pH U in the intraretinal extracellular space (Yamamoto et al., 1992), probably due to the change of H+ release caused by the energy metabolism in retinal cells. The activity-dependent changes in the external pH can play an active role in neuronal activity, as well as in the basal metabolism of retinal systems.

External protons also inhibit the glutamate response of AMPA receptors by increasing steady-state desensitization (Ihle and Patneau, 2000). However, the IC50 values for proton inhibition exceed the physiological range (from pH 5.7 to 6.3; Traynelis and Cull-Candy, 1991). Thus, during the generation of the cone surround response, pH changes in the cone synaptic clefts may affect the presynaptic cone calcium channels selectively, but not the postsynaptic AMPA receptors on the HCs.

In off-type BCs, 10 mM HEPES augmented the inward current response at the offset of a light spot, in addition to suppressing the surround response (Fig. 9). This is probably due to augmentation of transmitter release from the photoreceptors, and is consistent with the fact finding that HEPES augmented the cone ICa (Fig. 4). The sustained inward current in darkness, however, was not significantly augmented and this was probably due to desensitization of postsynaptic kainate receptors on off-BCs (DeVries and Schwartz, 1999). Under spot illumination (glutamate release is stopped), the kainate receptors would be released from a desensitized state. Thus, at light offset, the postsynaptic current evoked by abrupt glutamate release can be marked. A picrotoxin-resistant surround response of off-type BCs has also been reported in the tiger salamander retina (Hare and Owen, 1996). Thus, pH-mediated feedback to cones, rather than a GABAergic feedforward input to BCs, may be mainly involved in the receptive field organization of the BCs, at least in the amphibian retina. Modulation of the cone ICa is very strategic for direct control of the amount of transmitter release. Thus, the modulatory effect of pH on ICa at the cone synaptic terminal has a dual role, of contributing to the formation of the receptive field surround in the cones, and of controlling transmitter release from the cones to the BCs.

A Possible Mechanism of Control of the Extracellular pH Associated with Voltage Change in the HCs

There are several reports that neuronal activities evoke pH changes in the extracellular spaces in neuronal tissue (Kaila and Chesler, 1998). However, the mechanism by which HC depolarization acidifies the extracellular space in the cone synaptic clefts remains to be elucidated. An applicable model for the present hypothesis has been proposed as the model of neuron–glial cell interaction (Ransom, 2000). In this model, depolarization of glial cells caused by neuronal excitation activates Na+–HCO3− cotransport (stoichiometry of 2HCO3−: Na+) in the glia, resulting in the cellular uptake of extracellular HCO3− and acidification of the extracellular space (Ransom, 2000). If extracellular protons are increased in darkness, the extracellular space in a tightly packed invaginating synapse might be easily acidified. Also in isolated HCs of the skate retina, Na+– and HCO3−-dependent ion transport regulates the intracellular pH (Hauga-Schmedt and Ripps, 1998). Besides Na+–HCO3− cotransport in HCs, other ionic transporters or exchangers, such as acid loaders or extruders must also be considered. Moreover, proton release via vesicular release of GABA, which has been suggested in mammalian HCs (Cueva et al., 2002), must also be considered.

Molina et al. (2000) reported that application of glutamate to isolated HCs of the all-rod skate retina reduces the number of hydrogen ions on its surface, indicating that glutamate causes proton influx into the cytoplasm of the HCs. This glutamate-mediated proton influx into the HCs is consistent with the evidence that L-glutamate raises the intracellular proton concentration in isolated HCs (Dixon et al., 1993). The results of Molina et al. (2000) and Dixon et al. (1993) appear to be inconsistent with our conclusion that the extracellular space should be acidified in darkness (when glutamate is tonically released). However, since the experiments by Molina et al. (2000) and Dixon et al. (1993) were performed with an external solution buffered with HEPES (without bicarbonate), the bicarbonate-dependent system may not be applicable in their studies. Moreover, Molina et al. (2000) performed their study using rod-driven HCs, whereas our discussion is on pH regulation in cone systems. Therefore, these two studies are not necessarily inconsistent with our present results obtained using bicarbonate buffer in the external solution.

Contribution of GABAergic Feedback to Receptive Field Organization

Our present analysis was focused on GABA-independent components; therefore, all recordings were made...
in the presence of 100 μM picrotoxin, to exclude any possible effects of GABA. It has been a matter of debate whether GABA is also involved in the formation of the receptive field surround of cone photoreceptors. Our observations indicate clearly that there is a large component that does not require GABA. In fact, in preliminary studies, we found that the amplitude of cone surround response was not reduced by GABA antagonists (picrotoxin, bicuculine, SR95531). Moreover, the GABA antagonists did not evoke any current change whose reversal potential was equal to the equilibrium potential of chloride ions (unpublished data). Thus, it is unlikely that GABA plays a major role as a mediator of the feedback in the outer retina.

GABA may, however, play a substantial role in modulating the feedback response in the outer retina, rather than being the main mediator of the feedback. GABA can change membrane voltages of the HCs by modulating GABA-gated chloride currents and GABA transporter currents in the HCs (Kamermans and Werblin, 1992; Takahashi et al., 1995a,b), besides modulating GABA-gated chloride currents in cones. Thus, it is assumed that GABA may play some role in the information processing in the outer retina, but its role has to be reexamined in light of new data on the surround response of cones.

The synaptic structure of the invaginating synapse may be specialized for evoking pH changes in the intersynaptic clefts, because its highly packed structure can promote accumulation of protons. In contrast, GABAergic synapses between amacrine cells and bipolar and ganglion cells are conventional synapses, much different from the invaginating type of synapse. Thus, it may be reasonable to assume that lateral inhibition in the inner retina is mediated by GABA (Cook and McReynolds, 1998).

In summary, we have proposed an alternative hypothesis for the mechanism underlying the generation of the receptive field surround in the outer retina. We propose that intersynaptic pH changes in the invaginating synaptic clefts at the cone terminal contribute to the generation of the receptive field surround in the outer retina. Although the mechanism by which HCs regulate the extracellular pH is still uncertain, the present pH hypothesis should serve as the simplest description, so far, for the mechanism of feedback in the outer retina.

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