Calcium imaging demonstrates colocalization of calcium influx and extrusion in fly photoreceptors

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During illumination, Ca\(^{2+}\) enters fly photoreceptor cells through light-activated channels that are located in the rhabdomere, the compartment specialized for phototransduction. From the rhabdomere, Ca\(^{2+}\) diffuses into the cell body. We visualize this process by rapidly imaging the fluorescence in a cross section of a photoreceptor cell injected with a fluorescent Ca\(^{2+}\) indicator in vivo. The free Ca\(^{2+}\) concentration in the rhabdomere shows a very fast and large transient shortly after light onset. The free Ca\(^{2+}\) concentration in the cell body rises more slowly and displays a much smaller transient. After ~400 ms of light stimulation, the Ca\(^{2+}\) concentration in both compartments reaches a steady state, indicating that thereafter an amount of Ca\(^{2+}\), equivalent to the amount of Ca\(^{2+}\) flowing into the cell, is extruded. Quantitative analysis demonstrates that during the steady state, the free Ca\(^{2+}\) concentration in the rhabdomere and throughout the cell body is the same. This shows that Ca\(^{2+}\) extrusion takes place very close to the location of Ca\(^{2+}\) influx, the rhabdomere, because otherwise gradients in the steady-state distribution of Ca\(^{2+}\) should be measured. The close colocalization of Ca\(^{2+}\) influx and Ca\(^{2+}\) extrusion ensures that, after turning off the light, Ca\(^{2+}\) removal from the rhabdomere is faster than from the cell body. This is functionally significant because it ensures rapid dark adaptation.

Many neurons, including sensory neurons, localize Ca\(^{2+}\) influx channels to small, often diffusely isolated compartments of the cell body (1). This localized Ca\(^{2+}\) influx into a small volume often results in sizeable but local changes in free Ca\(^{2+}\) concentration (Ca\(_m\); refs. 2 and 3). Photoreceptor cells of flies exemplify this strategy. On light stimulation, channels that are highly permeable for Ca\(^{2+}\) (4, 5) and exclusively located in the membranes of the microvilli (6, 7) are activated. The microvilli, tube-like protrusions of the plasma membrane with exceptionally small dimensions, are regularly packed together to form a dense stack, the rhabdomere (8). Ca\(_m\) in the rhabdomere rises up to 600 μM even when the cells are stimulated with bright light (9). Calculations indicate that similar or even higher concentrations also occur in one or a few microvilli already after the absorption of a single photon (10). From the rhabdomeric microvilli, Ca\(^{2+}\) diffuses into the cell body (11), where measured increases in Ca\(_m\) are more moderate but still exceed 10 μM (12, 13). These changes of Ca\(_m\) are functionally important, because they mediate light adaptation (14, 15).

Creating high local Ca\(_m\) values by confining the influx to specialized compartments potentially causes the problem that Ca\(_m\) in these compartments diminishes only slowly after the stimulus has ceased. This problem can be counteracted by colocalizing proteins that efficiently extrude Ca\(^{2+}\). The implementation of this design principle has been demonstrated, e.g., in stereocilia of haircells (16), in synaptic terminals of rods (17, 18), and in presynaptic boutons of hippocampal cells (19). In fly photoreceptor cells, highly active Na\(^{+}\)/Ca\(^{2+}\) exchangers extrude Ca\(^{2+}\) (20–23). Two genes encoding two different types of Na\(^{+}\)/Ca\(^{2+}\) exchangers have been shown to be expressed in the photoreceptor cells of Drosophila (24, 25), but the subcellular location of the proteins is unknown. In squid photoreceptor cells, however, the rhabdomeric membranes show Na\(^{+}\)/Ca\(^{2+}\) exchange activity (26). Using a combination of Ca\(^{2+}\) imaging and modeling, we show in this report that Ca\(^{2+}\) extrusion in fly photoreceptors takes place close to the location of Ca\(^{2+}\) influx, the rhabdomere, suggesting that Na\(^{+}\)/Ca\(^{2+}\) exchangers are located in or close to the rhabdomere. We show that this has profound physiological consequences, as it increases the speed of Ca\(^{2+}\) removal in the rhabdomere and avoids large gradients of Ca\(_m\) in the cells during continuous stimulation.

**Materials and Methods**

High-Speed Fluorescent Imaging and Data Analysis. Single photoreceptor cells of female white-eyed mutant chalky blowflies (Calliphora vicina) were impaled with intracellular electrodes and iontophoretically injected with the fluorescent Ca\(^{2+}\) indicator dye Oregon green 5N (O5GN, Molecular Probes) as described previously (9, 13). The electrode was withdrawn, and the fly was placed on a goniometer that served as stage of a standard microscope (Nikon Optiphot-2, Nikon) connected to a confocal microscope (Odyssey XL, Noran Instruments, Middletown, WI), from which the confocal pinhole was removed to maximize light intensity. The fly was reoriented to ensure that the rhabdomere of the dye-filled cell was coaxial with the optical axis of the objective (×25 water immersion, NA 0.6; SW25, Leitz). Light came from the 488-nm line of a krypton–argon laser, and the emitted fluorescence light passed a triple-filter (Nikon) or a 510-nm fluorescence cube (Nikon). An LS2 shutter (Uniblitz, Vincent Associates, Rochester, NY) was added in the light path of the confocal microscope before the beam expander to rapidly turn the light on or off. Several sequences of images were recorded at a rate of 480 images per s and averaged off line; care was taken to ensure that the preparation did not move between the recordings. All findings were confirmed in at least three different preparations.

Modeling the Steady-State Distribution of Ca\(^{2+}\). For the calculations of the steady-state distribution of Ca\(_m\), we assume that a constant influx of Ca\(^{2+}\) (see Results) occurs in the rhabdomere homogeneously along the entire length of a photoreceptor cell (l = 225 μm; ref. 8), reducing diffusion to two dimensions. This assumption seems justified because the light intensities used for measuring the fluorescence saturate the light response: each microvillus was hit at least once in a few milliseconds. The cross section through the modeled cell was assumed square with side lengths a = 7.1 μm. The diffusion coefficient for free Ca\(^{2+}\) ions (D(Ca)) was assumed to be 220 μm\(^2\)s\(^{-1}\) (27). The diffusion coefficient of the mobile buffer, whether bound to Ca\(^{2+}\) or not, was equally assumed to be D(B) = 220 μm\(^2\)s\(^{-1}\) (28), as it is possible that most of the mobile buffer stems from the Ca\(^{2+}\) indicator. Accordingly, the dissociation constant of the buffer (Kd(B)) was taken to be 20 μM, the value reported for O5GN (29, 30). As only the steady-state distribution of Ca\(^{2+}\) was investigated here,
fixed (i.e., nondiffusing) Ca\(^{2+}\) buffers do not need to be considered (31). Equally, Ca\(^{2+}\) release (32) or Ca\(^{2+}\) uptake (33) into organelles that might occur in fly photoreceptor cells do not influence the steady-state distribution of free Ca\(^{2+}\).

The square cross section was divided by a grid into 40 \times 40 compartments of equal size. In the middle of one side, a 1.5 \mu m large region was designated to be the region of Ca\(^{2+}\) influx. The concentration of Ca\(^{2+}\) ions, either free or bound to a Ca\(^{2+}\) buffer, was calculated at the intersections of the grid lines. The volume assigned to each grid point was 1/1,600 of the total cell volume (\(V_{\text{cell}} = 11\) pl), but the points lying on one of the cell borders were assigned half that value, and the points in the corners a quarter. Diffusion was calculated to occur along the grid lines. The choice of either free or bound calcium at positions \(x\) and \(y\) during a time step (\(\Delta t = 28\) \mu s) caused by diffusion with coefficient \(D\) is then given by:

\[
\Delta C_{x,y} = \frac{D\Delta t}{\Delta x^2}(C_{x-1,y} + C_{x+1,y} + C_{x,y-1} + C_{x,y+1} - 4C_{x,y}),
\]

for grid points that do not lie on the border of the cell; for grid points on the border, the coefficients need to be changed appropriately. The distance between two grid points \(\Delta x = a/40 = 177\) \text{nm}. After calculating \(\Delta C\) for every grid point, the new total Ca\(^{2+}\) concentration was obtained. From the total Ca\(^{2+}\) concentration, the concentrations of free and bound Ca\(^{2+}\) were determined with the assumption that the buffer reactions are in equilibrium. The calculations continued until a stable distribution of Ca\(_i\) was reached, i.e., until the amount of Ca\(^{2+}\) extruded differed less than 0.1\% from the amount of Ca\(^{2+}\) flowing in.

The activity of Na\(^{+}\)/Ca\(^{2+}\) exchangers, the proteins that extrude Ca\(^{2+}\) from fly photoreceptor cells (20–23), depends on many parameters, including the membrane potential and the intra- and extracellular Na\(^{+}\) and Ca\(^{2+}\) concentrations (for review, see ref. 34). As we analyze only the steady state, all parameters can be assumed constant except Ca\(_i\), Ca\(^{2+}\) binding to the transport site of the exchanger proteins is typically described with a Hill function (Hill coefficient \(h = 1\)), for which two parameters, the dissociation constant \(K_{\text{d,X}}\) and the maximal current \(I_{\text{max,X}}\), need to be determined (34). A rough estimate of the \(K_{\text{d,X}}\) value for the Ca\(^{2+}\) extrusion can be inferred from data by Hardie (22), who showed that the Na\(^{+}\)/Ca\(^{2+}\) exchanger is 10 times more active when Ca\(_i\) = 20 \mu M than when Ca\(_i\) = 1 \mu M. This yields 18 \mu M for \(K_{\text{d,X}}\), which, however, is rather high compared with the values reported for other preparations (34). The maximum calcium current transported by the Ca\(^{2+}\) extrusion process was set to \(I_{\text{max,X}} = 1.3\) nA to ensure that the calculated Ca\(_i\) values at the location of extrusion lie between 12 and 40 \mu M, as found experimentally for Ca\(_i\) in strongly light-stimulated cells (13). The choice of parameters for the Ca\(^{2+}\) extrusion is rather uncritical for the conclusions drawn in Results, because changing \(K_{\text{d,X}}\) or \(I_{\text{max,X}}\) affects the average Ca\(^{2+}\) concentration strongly but the calculated gradients only very weakly.

**Results**

**Imaging the Ca\(^{2+}\)-Induced Change in Fluorescence in a Cross Section of a Photoreceptor Cell.** We imaged the distal cross section of a blowfly photoreceptor cell iontophoretically injected with the fluorescent Ca\(^{2+}\) indicator OGSN in vivo by optically neutralizing the natural optics of the cornea [ref. 35; the principle of this approach is outlined in a schematic drawing provided as supplementary material (see www.pnas.org)]. Fig. 1 shows false-color images representing raw intensity values taken from a time series recorded with high temporal resolution (480 images per s). The image at 2 ms is the first image recorded after the light was turned on. Because the photoreceptor cells have a latency period of \(\approx 4\) ms before an increase in Ca\(_i\) can be detected (9, 12), this first image represents the autofluorescence of the tissue together with the fluorescence of the Ca\(^{2+}\) indicator at low values of Ca\(_i\).

The intense light used for measuring the fluorescence strongly activates the phototransduction cascade, resulting in opening channels that are permeable for Ca\(^{2+}\) (4, 5). This leads to a Ca\(^{2+}\) influx, visible in the image taken 10 ms after light onset. The fluorescence intensity has increased in the area that corresponds to the rhabdomere but not in the rest of the cell. This observation confirms that the influx of Ca\(^{2+}\) through the light-activated channels is localized exclusively to the rhabdomere (6, 7, 11). The subsequent images show that the Ca\(^{2+}\) increase spreads into the cell body and concomitantly reduces in the rhabdomere. A movie showing the initial 200 ms of the measurement depicted in Fig. 1 is provided as supplementary material.

**A Sizeable Ca\(^{2+}\) Current Flows into the Photoreceptor Cells During the Steady State.** The electrical response caused by bright illumination consists of a fast transient depolarization (on average 65 mV) that quickly decays to a steady-state depolarization of about 30 mV (e.g., ref. 13). During this steady state, a continuous current flows through the light-activated channels, which is counterbalanced by the strong nonactivating currents through voltage-dependent potassium channels (36). The voltage-dependent potassium current at 30 mV depolarization is \(\approx 3\) nA (36). This value can be used to estimate the Ca\(^{2+}\) current through the light-activated channels, as the current through the light-activated channels (\(I_l\)) needs to balance the currents through the voltage-dependent K\(^+\) channels. Assuming the validity of the Goldman–Hodgkin–Katz theory (ref. 37; for application to fly photoreceptors, see refs. 10, 23, and 38), the current carried by Ca\(^{2+}\) ions (\(I_{\text{Ca}}\)) through the light-activated channels can be calculated as (10):

\[
I_{\text{Ca}} = I_l\nu_{\text{Ca}}\frac{f_{\text{Ca}}}{f_{\text{Na}} + f_{\text{K}} + f_{\text{Ca}} + f_{\text{Me}}},\text{ whereby}
\]

\[
f_q = z_q^2\nu_{\text{Ca}}\frac{C_{\text{Ca}} - C_{\text{Ca}}e^{-z_q\nu_{\text{Na}}}}{1 - e^{-z_q
u_{\text{Na}}}}.
\]

The index \(q\) denotes the four cations considered, \(z_q\) the valence, \(C_{\text{Ca}}\) the intracellular, and \(C_{\text{Na}}\) the extracellular concentration of ion sort \(q\). \(F = F/(RT)\), with \(F\) the Faraday constant, \(R\) the molar gas constant, and \(T = 293\) K. The membrane potential (\(V_m\)) is taken to be \(-30\) mV, the steady-state value of strongly stimu-
lateralized cells (13). The relative permeabilities ($w_i$) of the light-activated channels have been measured in *Drosophila* photoreceptor cells (4, 5); here we use $w_{Na} = w_K = 0.02, w_{Ca} = 0.85$ and $w_{Mg} = 0.11$, the values described for wild-type flies (ref. 5; see also ref. 10). The following ion concentrations (in mM) are typically found in insect photoreceptor cells and retinas: $C_{Na,i} = 10, C_{K,i} = 140, C_{Na,o} = 120, C_{Mg,i} = 4, C_{Mg,o} = 2$ (39, 40). In strongly stimulated photoreceptor cells, rather high values for the intracellular Ca$^{2+}$ concentration ($C_{Ca,i}$) have been reported (12, 13), and the extracellular Ca$^{2+}$ concentration ($C_{Ca,o}$) is probably reduced from its resting value of 1.4 mM (40). Here we use $C_{Ca,i} = 20$ μM and $C_{Ca,o} = 1.2$ mM.

By using these values, $I_{Ca}$ works out to be 1.4 nA when $I_I$ is assumed to be 3 nA. This, however, neglects the contribution of Na$^+$/Ca$^{2+}$ exchange that can depolarize the cells because of its electrogenicity (21, 23). Furthermore, currents through the voltage-dependent K$^+$ channels might have been overestimated, as we neglected possible light-induced accumulation of K$^+$ in the extracellular space (39). In the following, we take $I_{Ca} = 0.7$ nA, in order not to overestimate the Ca$^{2+}$ current through the light-activated channels.

Our previous measurements (13) and the data presented in Fig. 1 indicate that Ca$^{2+}$ does not increase substantially after ~200–500 ms of light stimulation. This shows that Ca$^{2+}$-extruding mechanisms must generate a Ca$^{2+}$ current of (at least) 0.7 nA during the steady state to balance the influx.

**The Steady-State Distribution of Ca$^{2+}$ Contains Information About the Localization of Ca$^{2+}$-Extruding Proteins.** The Ca$^{2+}$ ions, flowing in through the light-activated channels located in the rhabdome, have to diffuse to the location of the Ca$^{2+}$-extruding proteins to be extruded. The subcellular location of the Ca$^{2+}$-extruding proteins therefore has a profound influence on the shape and size of the gradients that build up in the cytosol. This is illustrated in Fig. 2, where we calculated distributions of Ca$^{2+}$ in the cross section of a square model cell for four different locations of Ca$^{2+}$-extruding proteins, as indicated in Fig. 2 (Top). It was assumed that a continuous Ca$^{2+}$ current ($I_{Ca}$) of 0.7 nA flows into the model cell at the center of one side, corresponding to the place where the rhabdomeric microvilli are attached. The influence of mobile Ca$^{2+}$ buffers was initially neglected. When assuming that the Ca$^{2+}$ extrusion takes place on the basolateral sides (Fig. 2a), where the Na$^+$/K$^+$ pumps (41) are located, Ca$^{2+}$ in the rhabdome should be as high as 97 μM, leveling off to reach 12–40 μM at the basolateral sides of the cell. Placing the Ca$^{2+}$ extrusion to the apical side (except the rhabdome) results in a much flatter distribution in the cell body (Fig. 2b). Still, Ca$^{2+}$ in the rhabdome would be at 73 μM and on the apical side as low as 14 μM. Modeling extrusion on all sides except the rhabdome (Fig. 2c) results in an intermediate between the situations shown in Fig. 2a and b. Only when the Ca$^{2+}$ extrusion is modeled to take place exclusively at the location of Ca$^{2+}$ influx, the rhabdome, a homogeneous distribution of Ca$^{2+}$ is found throughout the cell (Fig. 2d).

The gradients that can possibly be expected to be measured experimentally, however, are not as large as those depicted in Fig. 2. The addition of fluorescent Ca$^{2+}$ indicators, highly mobile Ca$^{2+}$ buffers that are necessary for the measurements, will reduce those gradients (31, 42). To estimate more realistically the measurable gradients, we repeated the calculations with varying concentrations of a highly mobile Ca$^{2+}$ buffer (Fig. 3). Increasing the buffer concentration flattens the expected distribution of Ca$^{2+}$. For the extreme case of 5.0 mM Ca$^{2+}$ buffer (the concentration of the indicator in the recording electrode), barely any gradients are discernible, no matter where the extrusion is assumed to take place (Fig. 3 a and b). These calculated values of Ca$^{2+}$ can be used to derive the expected fluorescence intensity of a Ca$^{2+}$ indicator with $K_{d,B} = 20$ μM. The expected difference in indicator fluorescence intensity between the rhabdomeric region and the side of the cell body opposite the rhabdome is plotted as a function of the buffer concentration in Fig. 3c. It shows that even with buffer concentrations of 750 μM, the intensity difference should be 18% if the Ca$^{2+}$ extrusion is located at the basolateral sides. When the Ca$^{2+}$ extrusion takes place on the apical side, the intensity difference should amount to 9%. For lower buffer concentrations, the fluorescence difference is predicted to be even larger. Importantly, no matter how large the buffer concentration, no gradient will arise if Ca$^{2+}$ extrusion is confined to the place of Ca$^{2+}$ influx, the rhabdome (not shown). These considerations show that measurements of Ca$^{2+}$ gradients in the cell body can yield information about the location of Ca$^{2+}$ extrusion, provided the concentration of the Ca$^{2+}$ indicator is not too high.

**The Steady-State Distribution of Ca$^{2+}$ in a Cross Section of the Photoreceptor Cells Is Homogeneous.** Fig. 4 shows the quantitative analysis of the data presented in Fig. 1. To obtain a reasonable signal-to-noise ratio, we defined three regions of interest in the
The concentration of the Ca$^{2+}$ buffer strongly influences the modeled distribution of the free Ca$^{2+}$ concentration. (a and b) The free Ca$^{2+}$ concentration profile along the symmetry line through the cell body, from the Ca$^{2+}$ influx region to the opposite side of the cell body, is plotted. The extrusion was assumed to take place only at basolateral sides (a; as in Fig. 2a) or only at the apical side (b; as in Fig. 2b). Increasing the buffer concentration (indicated by the numbers, in millimolars) reduces the size of the predicted gradients. (c) The relative fluorescence intensity difference of a Ca$^{2+}$ indicator with $K_D = 20 \mu M$ between the Ca$^{2+}$ influx region ($x = 0 \mu m$ in a and b) and the opposite end of the cell body ($x = 7.1 \mu m$ in a and b) is plotted as a function of the buffer concentration, when assuming extrusion on the basolateral sides (C') or on the apical side (B'). Below a buffer concentration of 0.75 mM, the fluorescence intensity between the two points differs by more than 10% (dashed line).

**Fig. 3.** The concentration of the Ca$^{2+}$ buffer strongly influences the modeled distribution of the free Ca$^{2+}$ concentration. (a and b) The free Ca$^{2+}$ concentration profile along the symmetry line through the cell body, from the Ca$^{2+}$ influx region to the opposite side of the cell body, is plotted. The extrusion was assumed to take place only at basolateral sides (a; as in Fig. 2a) or only at the apical side (b; as in Fig. 2b). Increasing the buffer concentration (indicated by the numbers, in millimolars) reduces the size of the predicted gradients. (c) The relative fluorescence intensity difference of a Ca$^{2+}$ indicator with $K_D = 20 \mu M$ between the Ca$^{2+}$ influx region ($x = 0 \mu m$ in a and b) and the opposite end of the cell body ($x = 7.1 \mu m$ in a and b) is plotted as a function of the buffer concentration, when assuming extrusion on the basolateral sides (C') or on the apical side (B'). Below a buffer concentration of 0.75 mM, the fluorescence intensity between the two points differs by more than 10% (dashed line).

The intensity vs. time plots of the normalized fluorescence (Fig. 4 b and c) are fully consistent with our earlier measurements (9, 13). In the rhabdomere, very fast and large Ca$^{2+}$ transients occur; 10 ms after the onset of the light stimulation, the normalized fluorescence in the rhabdomere has reached its maximum and stays at a plateau for about 100 ms before declining again. The plateau at the highest level of the normalized fluorescence is caused by the saturation of the Ca$^{2+}$ indicator OG5N (9). The further away from the rhabdomere the region of interest is chosen, the slower the increase of the normalized fluorescence is found to be. Also, the transient at light onset is much reduced in the cell body as compared with the rhabdomere. About 400 ms after onset of light stimulation, however, the normalized fluorescence, and hence Ca$_{i}$, has reached the same value in all regions of interest, i.e., the distribution of Ca$_{i}$ in the steady state is homogeneous. This observation argues that the Ca$^{2+}$ extrusion is closely colocalized with the Ca$^{2+}$ influx, assuming that the concentration of highly mobile Ca$^{2+}$ buffers is below 750 $\mu M$ (Fig. 3). Evidence presented below shows that this condition is likely to be met in our experiments.

**Fig. 4.** The steady-state distribution of Ca$_{i}$ in a cross section of a photoreceptor cell homogeneous. (a) Raw intensity image (gray scale) of 100 successive images averaged during the steady state. The colored squares indicate the regions of interest quantitatively analyzed in b and c; red, rhabdomere; green, center of cell; blue, side of cell opposite the rhabdomere; yellow, background, outside the cell that was injected with the Ca$^{2+}$ indicator. (b and c) Normalized fluorescence traces (the first value obtained after turning on the light was used for normalization; arrows). The normalized fluorescence rises sharply in the rhabdomere (red line) and displays a large transient. After the decay of the transient, the normalized fluorescence shows the same values, independent of location of the region of interest. The distribution of Ca$_{i}$ during the steady state thus is homogeneous. The traces are averages of five experiments.

**Ca$^{2+}$ Removal in the Rhabdomere Is Faster than in the Cell Body.** After turning off the light stimulation, Ca$^{2+}$ influx through the light-activated channels ceases, and Ca$_{i}$ diminishes because of the ongoing action of the Ca$^{2+}$ extrusion mechanisms. When Ca$^{2+}$ extrusion takes place in the rhabdomere, it can be expected that Cai reduces faster in the rhabdomere than in the rest of the cell. We therefore exposed a cell to bright light for 1 s, which allowed Ca$_{i}$ to reach a homogeneous steady state (as in Fig. 4) and a subsequent dark period of 800 ms (Fig. 5a). At the end of the dark period, the normalized fluorescence, and hence Ca$_{i}$, in the rhabdomere (red trace) had reached lower values than in the other parts of the cell (green and blue traces, Fig. 5a). In a different experimental protocol (Fig. 5 b-d), we stimulated a
dark-adapted cell for only 200 ms and then investigated how Cai developed during the subsequent dark period. Fig. 5b shows that after 200 ms of darkness Cai has diminished in the rhabdomere, whereas it has continued to rise in the cell body, consistent with Ca\(^{2+}\) being redistributed by diffusion. After 400 ms of darkness (Fig. 5c), Cai in the cell body has started to decline; by the same time, Cai in the rhabdomere has fallen to a value lower than that found in the cell body. The same is seen more clearly after 600 ms of darkness (Fig. 5d). Hence, the Ca\(^{2+}\) gradient in the photoreceptor cell reverses its direction: after 200 ms of darkness, Cai in the rhabdomere is still higher than in the cell body, whereas the opposite is found after 400–600 ms of dark adaptation. To analyze the extent of the Ca\(^{2+}\) gradient after 600 ms of dark adaptation, the normalized fluorescence intensity at the onset of the second light stimulation in Fig. 5d was plotted as a function of the distance to the rhabdomere (Fig. 5e). Converting these values to free calcium concentrations (as described in ref. 9) shows that in this particular cell the gradient is substantial, with Ca\(^{2+}\) concentrations ranging from \(\approx 7\ \mu M\) in the rhabdomere to \(\approx 14\ \mu M\) at the opposite side of the cell body (Fig. 5f). Together, these results show that Ca\(^{2+}\) removal from the rhabdomere is faster than from the cell body. Consequently, Ca\(^{2+}\) diffuses from the cell body into the rhabdomere during the recovery from light stimulation.

We cannot directly control or determine the indicator concentration used in the experiments. The measurements shown in Fig. 5 demonstrate that Ca\(^{2+}\) gradients can exist under our experimental conditions after the Ca\(^{2+}\) transients in the rhabdomere at the onset of illumination has ceased. Therefore, the concentration of the Ca\(^{2+}\) indicator used in the experiments was sufficiently low to enable the measurements of Ca\(^{2+}\) gradients. This argues that the homogeneous distribution of Cai in the steady state (Fig. 4) cannot be explained solely with a high concentration of Ca\(^{2+}\) buffer but must indeed be caused by colocalization of Ca\(^{2+}\) influx and extrusion in or close to the rhabdomere (Figs. 2, 3).

**Discussion**

An increase in Cai in fly photoreceptor cells is brought about by an influx of Ca\(^{2+}\) through the light-activated channels that are exclusively located in the rhabdomere (6, 7, 11). In this report, we show that also Ca\(^{2+}\) extrusion takes place in or close to the rhabdomere. Ca\(^{2+}\) influx and Ca\(^{2+}\) extrusion are therefore colocalized.

Two independent arguments indicate that the homogeneous distribution of Cai we observe in the steady state is not caused by a large mobile Ca\(^{2+}\) buffer capacity: (i) Ca\(^{2+}\) gradients exist in the cell body during periods of dark adaptation (Fig. 5). High concentrations of a Ca\(^{2+}\) buffer would also strongly reduce these gradients, probably to a degree that makes it impossible to measure them. (ii) Using blunt electrodes very easily overloads cells with the Ca\(^{2+}\) indicator. Recordings from those cells show profoundly modified kinetics of the membrane potential (15) and the fluorescence (data not shown) and hence were discarded. Consequently, in not-overfilled cells the concentration of the indicator must have been much lower than 5 mM (the concentration of the indicator in the electrode). Therefore, our measurements exclude the possibility that Ca\(^{2+}\) extrusion is located only on the basolateral membranes (Fig. 2a) and make it unlikely that Ca\(^{2+}\) extrusion takes place only in the apical membrane excluding the rhabdomere (Fig. 2b).

**A Method for Locating Ca\(^{2+}\) Extrusion.** Imaging of cells injected with fluorescent Ca\(^{2+}\) indicator dyes is a well-established method for demonstrating the location of Ca\(^{2+}\) channels (e.g., refs. 11, 43, and 44). Here we show that measuring spatial Ca\(^{2+}\) gradients, in combination with modeling, can be used for demonstrating the location of Ca\(^{2+}\) extrusion in vivo. As this method characterizes the location of Ca\(^{2+}\) extrusion by its very function, it is not necessary to know which types of molecules extrude Ca\(^{2+}\). It contrasts in this respect with the traditional approach using antibodies. An intrinsic problem of the latter is that not every protein detected by an antibody might be functional, and proteins might be regulated differentially depending on where they are located. The approach used in this report therefore might provide a viable and direct alternative in systems where data on Ca\(^{2+}\)-extruding proteins are absent and might complement traditional approaches in other systems.
The Identity of the Ca$^{2+}$-Extruding Mechanism. Ample evidence suggests that Na$^{+}$/Ca$^{2+}$ exchanger proteins are present in fly photoreceptor cells (20–25) and in photoreceptor cells of other invertebrates (bee, ref. 45; *Limulus*, refs. 46 and 47; squid, ref. 26). Manipulating the function of the Na$^{+}$/Ca$^{2+}$ exchanger can augment Ca$^i$, in *Limulus* (46), which mimics the phenomena of light adaptation (14), generally believed to depend on an increase in Ca$^i$ (14, 15). The Na$^{+}$/Ca$^{2+}$ exchanger thus appears to be the main mechanism of Ca$^{2+}$ extrusion in invertebrate photoreceptor cells. Probably, therefore, the Na$^{+}$/Ca$^{2+}$ exchanger proteins contribute the largest part (or all) of the Ca$^{2+}$ influx observed in our experiments. From the experiments presented here, it follows that Na$^{+}$/Ca$^{2+}$ exchanger proteins are located in or close to the rhabdomere of fly photoreceptor cells. This is in agreement with Bauer et al. (26), who demonstrated Na$^{+}$/Ca$^{2+}$ exchange in the rhabdomeres of squid photoreceptor cells. Because squids and flies do not belong to phylogenetically closely related groups, this raises the possibility that colocalization of Na$^{+}$/Ca$^{2+}$ exchangers and light-activated channels in rhabdomeres is a general feature of invertebrate photoreceptor cells.

The Close Colocalization of Ca$^{2+}$ Influx and Ca$^{2+}$ Extrusion Has Important Functional Implications. Ca$^{2+}$ directly regulates light-activated channels (22, 48) and either directly or via calmodulin regulates many other proteins involved in phototransduction (49). Because most of those proteins are located in or close to the rhabdomere (49), Ca$^i$ in the rhabdomere rather than in the cell body may control the state of light adaptation. Rapid dark adaptation depends therefore on rapid removal of Ca$^{2+}$ from the rhabdomere. Localizing Na$^{+}$/Ca$^{2+}$ exchanger proteins to the rhabdomere helps achieve this (Fig. 5). Similarly, the colocalization of Ca$^{2+}$ influx channels and Ca$^{2+}$ extrusion mechanisms in small compartments observed in other cell types (16–19) might have the same function, i.e., to increase the speed of Ca$^{2+}$ removal.

Colocalization of Ca$^{2+}$ influx and extrusion results in a homogeneous distribution of Ca$^i$ in the steady state throughout the cell body (Fig. 4c). This ensures that structures located remotely from the Ca$^{2+}$ influx are also exposed to the same levels of Ca$^i$. This is potentially important for the regulation of mitochondria that are located near the basolateral sides of fly photoreceptor cells (33) and are known to be regulated by Ca$^i$ (50).

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