Light Adaptation of the Cyclic GMP Phosphodiesterase of Frog Photoreceptor Membranes Mediated by ATP and Calcium Ions

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ABSTRACT The light-activated guanosine 3',5'-cyclic monophosphate (cyclic GMP) phosphodiesterase (PDE) of frog photoreceptor membranes has been assayed by measuring the evolution of protons that accompanies cyclic GMP hydrolysis. The validity of this assay has been confirmed by comparison with an isotope assay used in previous studies (Robinson et al. 1980. J. Gen. Physiol. 76: 631-645). The PDE activity elicited by either flash or continuous dim illumination is reduced if ATP is added to outer segment suspensions. This desensitization is most pronounced at low calcium levels. In 10⁻⁹ M Ca⁺⁺, with 0.5 mM ATP and 0.5 mM GTP present, PDE activity remains almost constant as dim illumination and rhodopsin bleaching continue. At intermediate Ca⁺⁺ levels (10⁻⁷-10⁻⁶ M) the activity slowly increases during illumination. Finally, in 10⁻⁴-10⁻³ M Ca⁺⁺ the sensitivity control exerted by ATP has almost disappeared, and PDE activity is more a reflection of the total number of rhodopsin molecules bleached than of the rate of the rhodopsin bleaching. At intermediate or low calcium levels a short-lived inhibitory process is revealed by observing a nonlinear summation of responses of the enzyme to closely spaced flashes of light. Each flash makes PDE activity less responsive to successive flashes, and a steady state is obtained in which activation and inactivation are balanced. It is suggested that calcium and ATP regulation of PDE play a role in the normal light adaptation processes of frog photoreceptor membranes.

INTRODUCTION
Increasing interest is focusing on the guanosine 3',5'-cyclic monophosphate (cyclic GMP) phosphodiesterase (PDE) complex of vertebrate photoreceptor membranes, because several lines of evidence suggest that it plays a central role in visual transduction. In isolated and fragmented rod outer segments activation of the enzyme occurs within milliseconds of light absorption (Yee and Liebman, 1978), and in intact isolated outer segments a correspondingly rapid drop in cyclic GMP levels is observed (Woodruff et al., 1977; Woodruff and Bownds, 1979). Cyclic GMP levels of isolated outer segments correlate
with their ionic permeability (Woodruff et al., 1977), and Miller and Nicol (1979) have shown that intracellular injection of cyclic GMP into toad rods causes their depolarization. These and other studies (for a review, see Hubbell and Bownds [1979]; Pober and Bitensky, 1979) raise the possibility that cyclic GMP might act as an internal transmitter that controls plasma membrane permeability. One suggestion has been that this control might occur through the phosphorylation-dephosphorylation of two minor proteins (Polans et al., 1979).

The plausibility of these suggestions rests on demonstrating appropriate correlations between the kinetics of the phosphodiesterase, cyclic GMP, and conductance changes measured in living receptor cells, and thus far definitive experiments have not been done. Kilbride and Ebrey (1979) and Kilbride (1980) have shown that illumination causes cyclic GMP levels of whole retinas to fall over a period of seconds, rather than milliseconds. Lowering calcium concentration in the Ringer's solution from 1.6 mM to a nanomolar level increases the cyclic GMP change. These experiments did not monitor the relevant conductance changes or determine whether the cyclic GMP changes occurred in the outer segments. It was suggested that the rapid light-sensitive decrease in cyclic GMP observed by Woodruff et al. (1979) in isolated rod outer segments might occur only at low calcium concentration and be much slower at millimolar calcium levels. Polans et al. (1981), however, have recently found that the cyclic GMP decrease in isolated outer segments is substantially complete within 1 s at calcium concentrations varying from $10^{-6}$ to $10^{-3}$ M.

A number of factors that contribute to the control of PDE have now been established. Wheeler and Bitensky (1977), Shinozawa et al. (1979), and Fung and Stryer (1980) have studied GTPase and guanosine nucleotide-binding proteins that appear to be obligatory for PDE activation. An elutable inhibitor has also been reported (Sitaramayya et al., 1977; Dumler and Etingof, 1976; Hurley and Ebrey, 1979; Baehr et al., 1979; Robinson et al., 1980). Liebman and Pugh (1979 and 1980) have found that ATP is used in reversal of activation. We and our co-workers have reported further controls of the PDE complex (Robinson et al., 1980). Crude frog rod outer segments contain labile components that mediate a light-induced increase in the Michaelis constant of the enzyme and also confer Ca++ sensitivity. The effect of increasing Ca++ is to make the PDE complex more sensitive to illumination. It was found during these experiments that a desensitization of PDE activity observed under low-Ca++ conditions required the presence of endogenous micromolar levels of ATP in the crude outer segment preparations. This was thought to be significant in light of the recent reports by Leibman and Pugh (1979 and 1980) that ATP causes an acceleration of the PDE activation-inactivation sequence.

In the studies reported in this paper we have investigated the effect of ATP and calcium in more detail. The results reveal a control system that may be relevant to the in vivo physiology of frog rod outer segments. This work has utilized an assay for PDE that measures the protons evolved by cyclic GMP hydrolysis (Yee and Liebman, 1978) and has demonstrated the validity of this
assay for the conditions used. It has been found that the proton assay, carried out with appropriate controls, is faster, more convenient, and more reproducible than the isotope assay used in most of the published studies on this enzyme.

**MATERIALS AND METHODS**

In these experiments we used rod outer segments freshly shaken from frog retinas and not subjected to further washing, to avoid loss of controls of the PDE enzyme complex (Robinson et al., 1980). The structures were, however, extensively fragmented by stirring during the proton assay for PDE (see below). In preparation for each experiment retinas were removed from one or two dark-adapted bullfrogs (*Rana catesbeiana* or *Rana grylio*) as described previously (Woodruff et al., 1977; Woodruff and Bownds, 1979). The retinas were then gently rinsed in 10 ml of Ringer's solution (115 mM NaCl, 2.5 mM KCl, 10 mM HEPES, 2 mM MgCl₂, 1 mM dithiothreitol, pH 7.8) and then transferred into 0.5 ml of the same Ringer's solution. Outer segments were detached by shaking the retinas with forceps for 2-3 min. The suspension was diluted fivefold to eightfold to bring the rhodopsin concentration to 7-15 μM and then kept at 0°-4°C until use. All manipulations were carried out under infrared illumination (Robinson et al., 1980). The calcium buffers used in these experiments were made by adding various amounts of EGTA to 0.1 mM Ca²⁺ solutions (Polans et al., 1980). This calcium level was in excess of the endogenous calcium levels introduced by glassware and reagents (~10⁻⁶ M, determined by atomic absorption spectroscopy and ion-specific electrode).

In each assay of PDE activity a 200-μl portion of the outer segment suspension was made 1 mM in GTP (by adding 5 μl of 40 mM GTP), placed in a small vial, and rapidly stirred at 20°C with a magnetic stirrer. In experiments using continuous illumination, the calibrated orange light source described previously was used (Brodie and Bownds, 1976) and intensity was expressed as rhodopsin molecules bleached per outer segment per second (rhod./o.s.-s). Cyclic GMP (10 μl of an 80 mM stock solution) was then added shortly after the onset of illumination to make a final concentration of 4 mM (chosen to be above the Kₘ of the light-activated PDE—cf. Robinson et al. [1980]). When flash illumination was used, the cyclic GMP was added just before the flash. The flash source (Sunpak Auto 411, Berkey Marketing Co., Woodside, N. Y.) used the orange filter described by Brodie and Bownds (1976), and the flash duration was 0.36 ms.

In each experiment the pH was allowed to drop from 7.8 to 7.7 as protons were released by cyclic GMP hydrolysis. The pH drop was monitored with a pH microelectrode (MI 410, Microelectrodes Inc., Londonderry, N. H.) and displayed on a pen chart recorder. Tracings of typical recordings are shown in Fig. 3. Proton release was determined by back-titrating the suspension with known amounts of 0.1 N NaOH, and PDE activity was then expressed as moles of cyclic GMP hydrolyzed per mole of rhodopsin present-minute (see below). In each assay the PDE activation caused by some level of dim continuous illumination was first determined, and then a saturating light (bleaching 3 × 10⁶ rhod./o.s.-s) was turned on so that the maximum rate of proton evolution for that sample could be determined. The ratio of the rate of proton evolution at a given intensity to that just after presentation of the saturating light is referred to as relative activity (expressed as percent of maximum). This relative activity was found to be more constant from preparation to preparation than the absolute activity and did not vary as much with the age of the outer segment preparation (see discussion of Fig. 4).
Ringer's solutions were made with ultrapure NaCl and KCl purchased from Alfa Inorganics (Beverly, Mass.) and MgCl₂ was obtained from Johnson Matthey Chemicals (Royston, England). Other chemicals were obtained from Sigma Chemical Co. (St. Louis, Mo.). Purity of the cyclic GMP, GTP, and ATP used was found to be >98% by analysis with high-pressure liquid chromatography (Biernbaum and Bownds, 1979).

Validation of the pH Assay for PDE

The proton evolution which results from cyclic GMP hydrolysis can be used as a rapid and convenient assay for PDE (Yee and Liebman, 1978), and is much less cumbersome than the assay that analyzes the products of hydrolysis of radioactive cyclic GMP (Robinson et al., 1980). Use of the proton evolution assay requires that one know the buffering properties of each solution employed, and the possible sources of proton evolution other than cyclic GMP hydrolysis must be excluded. Figs. 1 and 2 address these two points. Fig. 1 shows the titration of the outer segment suspension used in the PDE assay, which contained 1 mM GTP and was made ~10⁻⁹ M in Ca²⁺ by the addition of 0.1 mM CaCl₂ and 2.78 mM EGTA. (These experiments were performed in the light, with PDE fully activated, so that all of the 1 mM cyclic GMP added was hydrolyzed within 1 min.) At pH values of 7.7 or higher, hydrolysis of 1 mM cyclic GMP produces almost the same pH change as the addition of an equivalent amount of acid. This result is not obtained when 5'-GMP, GTP, or ATP are substituted for cyclic GMP. The deviation between cyclic GMP hydrolysis and the pH change that occurs at pH values lower than 7.3 can be explained by the appearance of 5'-GMP, which has an ionizable group with pK 6.3 (Yee and Liebman, 1978). In the experiments described in this paper the pH was allowed to drop from 7.8 to 7.7, and the amount of NaOH required to return the pH to 7.8 was taken as a
measure of the number of cyclic GMP molecules that had been hydrolyzed. The raw data are shown in Fig. 3 with the amount of NaOH added converted to moles cyclic GMP hydrolyzed. A pH range of 7.8-7.7 was chosen to ensure that proton evolution would accurately reflect cyclic GMP hydrolysis and also to stay within the pH optimum of the enzyme. The pH optimum in these studies was found to be broad, centering at pH 7.9. This confirms previous reports (Miki et al., 1973; Yee and Liebman, 1978). Titration curves such as the one shown in Fig. 1 were determined for each of the different solutions used in this work, since the buffering properties of the various solutions were slightly different.

Fig. 2 illustrates a comparison of the pH assay with the isotopic assay used in the experiments of Robinson et al. (1980). The two procedures are equivalent at early stages of reaction (which were studied in the experiments of this paper) but diverge after several minutes of illumination. This suggests that some portion of the hydrolyzed cyclic GMP cannot be measured by the pH assay. Similar results were obtained when either GTP or both ATP and GTP were present.

**RESULTS**

The basic observation with which this paper deals is shown in Fig. 3. A comparison of parts a and b demonstrates that the presence of ATP during continuous illumination causes a suppression of PDE activity. This is consistent with the original observation of Liebman and Pugh (1979) that ATP diminishes the PDE response elicited by flashes of light. In the experiment
shown in Fig. 3a two portions of an outer segment suspension, containing 1 mM GTP and $10^{-9}$ M Ca$^{++}$, were illuminated with dim light. (GTP was included in all experiments, for it is required for PDE activation [Wheeler and Bitensky, 1977]). After 1 min of continuous illumination, cyclic GMP was added to one portion, and its hydrolysis was monitored as proton evolution. The data shown are a direct tracing of the chart recording of the pH drop, with the ordinate showing the pH change converted to moles cyclic GMP hydrolyzed per mole rhodopsin present (see Methods). At the time indicated by the arrow, saturating illumination was given to determine the maximum enzyme activity. The second portion of outer segments, which had been kept in the dim illumination for 4 min, was then assayed by adding 4 mM cyclic GMP. The rate of cyclic GMP hydrolysis was larger, reflecting the fact that increased rhodopsin bleaching results in increased activity. Again, saturating light was presented, and the maximum rate was shown to be similar for both portions of outer segments.

A similar experiment was performed (Fig. 3b), but 0.5 mM ATP and 0.5 mM GTP instead of 1 mM GTP were used. (This was done to keep total nucleotide triphosphate concentration constant; see Fig. 5, legend). A first observation was that enzyme activity elicited by the same dim level of illumination is lower. Further, the activity assayed after 4 min of continuous...
illumination was similar to that assayed after only 1 min, in spite of the fact that more rhodopsin had been bleached during this period. Finally, saturating illumination elicited activity similar to that observed in the absence of ATP. Thus, ATP appears to diminish the effectiveness of dim light but does not alter the maximal enzyme activity which can be caused by saturating illumination (see also below). ADP, adenosine, and β,γ-methylene-ATP do not substitute for ATP. (During experiments of the sort shown in Fig. 3, ATP and GTP levels were followed by high-pressure liquid chromatography [Biernbaum and Bownds, 1979] and found to decline <20% in the 4-min period. Further, the addition of 100 µM ATP and GTP had no effect. Thus, adequate levels of triphosphates were present; the GTP and ATP effects on PDE require only micromolar amounts of these components [Wheeler and Bitensky, 1977; Liebman and Pugh, 1980]).

The behavior shown in Fig. 3 has been noted in over 100 separate experiments utilizing outer segments which have been detached from the retina for >20 min. Outer segments assayed in the absence of ATP within 5 or 10 min of their detachment show desensitized behavior similar to that shown in Fig. 3 b, and then over a period of 10–20 min shift to the more sensitized state shown in Fig. 3 a. During this period endogenous ATP is being hydrolyzed (Biernbaum and Bownds, 1979), and it seems most likely that the disappearance of this endogenous ATP causes the sensitization that subsequently can be reversed if excess ATP is added back to the system.

Another parameter must be mentioned as background for the figures that follow, detailing the effects of ATP and calcium. In many experiments of the sort shown in Fig. 3 we noted that the maximum PDE activity obtained with saturating illumination increased with the age of the outer segment suspension. The data of Fig. 4 (solid line) indicate that this increase in maximal activity is complete in ~1.5 h. The cause of the increase is not known. It could be due either to proteolysis that activates PDE (Miki et al., 1973) or to elution of inhibitor(s) (see above and references in Robinson et al. [1980]). In spite of an increase in maximum activity, the relative effect of a given level of illumination remains constant for 2 h after outer segments are shaken from the retina. This is shown by the dashed line in Fig. 4, which is a plot of the enzyme activity elicited by 1 min of dim light bleaching 1.4 × 10⁷ rhod./o.s.-s expressed as a percent of the maximum activity elicited by saturating illumination. This remains constant, and it is for this reason that relative activities are expressed in the remaining figures in this paper. In all experiments the actual maximum PDE activities were in the range shown in Fig. 4, and all experiments were completed within 2 h after rod outer segments were shaken from the retina.

**Regulation of PDE in Dim Light by an ATP-dependent Process**

The effect of ATP on light activation of PDE mentioned above is shown in Fig. 5, which illustrates the behavior of the system in low calcium (10⁻⁶ M) Ringer's solution under several different conditions. The data are plotted in a form which permits one to summarize many experiments of the sort shown in Fig. 3. Rod outer segments were exposed to continuous illumination, and
each point in the figure represents a determination of PDE activity (as percent of maximum; see above) made after the number of rhodopsin molecules indicated on the abscissa had been bleached by light. PDE activity elicited by the various levels of illumination in the absence of ATP is indicated by the open symbols, and activity in the presence of ATP is noted by the closed symbols.

Note that for a given level of rhodopsin bleaching the PDE activity is always lower when ATP is present. (Compare, for example, the open and closed circles in Fig. 5). This is compatible with the observations of Liebman and Pugh (1979 and 1980), using flash illumination. This process is referred to here as desensitization, and we have determined in separate experiments that it is complete within seconds of ATP addition. Second, if PDE is assayed without added ATP, the data points for all light intensities (open symbols) lie on the same straight line. Very different behavior is shown by outer segments exposed to both GTP and ATP (closed symbols). At different continuous light intensities, PDE activity is initially different, and there is less change in activity as each level of illumination continues. The curves for different levels of illumination in the presence of ATP are not superimposable, whereas those in the absence of ATP are. Thus, ATP has the effect of making PDE activity a better reflection of the level of continuous illumination under all conditions shown. This behavior was also noted by Robinson et al. (1980), using the isotope assay for PDE. The maximal activity obtained with saturating illu-
mination is not influenced by ATP. In 10 separate experiments the ratio of maximum PDE activity with and without added ATP was 1.06 ± 0.06 (SEM).

**Dependence of Desensitization on ATP Concentration**

Fig. 6 demonstrates that micromolar levels of ATP are sufficient to influence PDE activity. In the experiment shown, outer segments were exposed to dim illumination bleaching $2.6 \times 10^2$ rhod./o.s.-s, in the presence of the indicated concentrations of added ATP, and the enzyme activity was determined 1 and 4 min after the onset of illumination. ATP acts to suppress both the initial PDE activity (Fig. 6a, 1 min) and the increase in activity caused by continuing illumination (Fig. 6a, 4 min). Fig. 6b shows that the effect of ATP on the activity of PDE after 4 min of illumination is not further enhanced as its concentration is increased from 10 µM to 1 mM. In these experiments endogenous ATP levels were analyzed by high-pressure liquid chromatography (Biernbaum and Bownds, 1979) and found to be ~0.2 µM in the sample.
to which no ATP was added. Note also that endogenous ATPase activity present in these crude outer segment preparations (Biernbaum and Bownds, 1979) would be expected to hydrolyze some of the added ATP after 4 min of incubation, but the extent of this hydrolysis was not determined.

Calcium Reduces the Effect of ATP on PDE

Robinson et al. (1980) observed that PDE activity is less sensitive to illumination at low Ca$^{++}$ concentrations. In that work it was also found that increasing Ca$^{++}$ concentration to millimolar levels causes a sensitization of the enzyme. The data of Fig. 7, when compared with Fig. 5, confirm that observation, examined now in the context of the ATP effect. A first observation is that in 1 mM Ca$^{++}$, in the presence of GTP alone or GTP plus ATP, PDE activity is more nearly a linear function of the number of rhodopsin molecules bleached than is the case in Fig. 5. This agrees with the previous published work on PDE (cf. Yee and Liebman [1978]), which indicates that PDE activation is a linear function of the amount of rhodopsin bleached.
Another obvious difference when high and low calcium conditions are compared is the way in which PDE activity changes during continuous illumination (cf. closed symbols of Figs. 5 and 7). In low calcium, PDE activity increases slowly as illumination continues, whereas in high calcium the activity increases more rapidly. This increase in the slopes of the curves in Fig. 7 compared with Fig. 5 is found for both the absence (open circles) and presence (closed circles) of ATP. Finally, in the presence of ATP and GTP the activity of the PDE is significantly lower in low calcium than in high calcium, for all corresponding intensities and durations of illumination.

A more direct comparison of the effect of low and high calcium on PDE activity is shown in Fig. 8, in which enzyme activity measured 1 min after the onset of illumination is plotted as a function of the continuous light intensity.

![Figure 7. PDE activity in Ringer's solution containing $10^{-3}$ M Ca$^{2+}$. PDE activity was measured as in Fig. 5. Data were taken from three separate experiments, and points with bars (mean ± SEM) represent more than three samples. Curves were drawn by hand.](image)

(as in Robinson et al. [1980]). The desensitization that occurs in the presence of ATP and GTP (cf. solid lines) as calcium concentration is lowered from $10^{-3}$ to $10^{-9}$ M is shown as a shift to the right of the intensity-response curves of the enzyme. The dashed lines indicate that a small decrease in sensitivity also is observed when calcium concentration is lowered in the absence of added ATP (data replotted from Figs. 5 and 7).

There is some interest in examining what effects phosphodiesterase inhibitors might have on PDE intensity-response curves of the sort shown in Fig. 8, because several recent electrophysiological studies have examined the effect of these inhibitors on the rod photoresponse (cf. Lipton et al. [1977]). In the current experiments it has been found that 500 µM isobutylmethylxanthine, a concentration that exerts almost the maximum inhibition effect observed,
causes a twofold inhibition of both dark and maximally light-activated PDE. It does not shift the intensity-response curve to lower or higher light intensities (data not shown).

**PDE Activity as a Function of Ca**<sup>++</sup> **Concentration**

Fig. 9 demonstrates the variation in PDE activity over a range of Ca**<sup>++</sup> concentrations in response to illumination bleaching $2.6 \times 10^8$ rhod./o.s.-s, an intensity at which the effect of Ca**<sup>++</sup> in the presence of ATP could most clearly be seen (cf. Fig. 8). Two increases in relative enzyme activity were observed, one occurring between $10^{-9}$ and $10^{-7}$ M Ca**<sup>++</sup> and the second between $10^{-5}$ and $10^{-3}$ M Ca**<sup>++</sup>. In these experiments the maximum PDE activity in saturating illumination was $275 \pm 10$ (±SEM) moles of cyclic GMP hydro-

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**Figure 8.** PDE activity as a function of light intensity in low and high Ca**<sup>++</sup> solutions. PDE activity was measured after 1 min of continuous illumination at the indicated intensities in low ($10^{-9}$ M) or high ($10^{-3}$ M) Ca**<sup>++</sup> with 0.5 mM GTP and 0.5 mM ATP present (solid lines). Each point represents the mean (±SEM) of three or four separate experiments. Intensity-response curves under the same conditions, but without added ATP, (1 mM GTP, dashed lines), were calculated from Figs. 5 and 7, assuming that PDE activity was determined by the total number of rhodopsin molecules bleached per outer segment.

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lyzed/mole rhodopsin-min in $10^{-3}$ M Ca**<sup>++</sup> (19 determinations) and $219 \pm 15$ (±SEM) in $10^{-9}$ M Ca**<sup>++</sup> (14 determinations).

To examine the consequence of these transitions, PDE activity was measured with 0.5 mM GTP and 0.5 mM ATP in $10^{-6}$ M Ca**<sup>++</sup>, using the procedures of Figs. 5 and 7. These data are shown in Fig. 10 (solid line), with the corresponding data from Figs. 5 and 7 shown by dashed lines. The slope of the curve showing PDE activity plotted against the number of rhodopsin molecules bleached per outer segment is intermediate between the slopes shown for $10^{-3}$ and $10^{-9}$ M Ca**<sup>++</sup>. In the presence of ATP and GTP, PDE activity in $10^{-9}$ M Ca**<sup>++</sup> remains almost constant as dim illumination continues. At a Ca**<sup>++</sup> concentration of $10^{-6}$ M the activity slowly increases during illumination. Finally, in $10^{-3}$ M Ca**<sup>++</sup>, PDE activity becomes more a function...
of the number of rhodopsin molecules bleached than of the rate of bleaching, and the sensitivity control exerted by the ATP has almost disappeared.

Response of PDE to Flashes of Light

The experiments described thus far have examined the responses of PDE to continuous illumination of the sort used in previous studies of cyclic GMP and permeability changes in isolated outer segments (Bownds and Brodie, 1975; Woodruff and Bownds, 1979). It should be possible to explain the behavior of PDE during continuous illumination as a summation of its discrete responses to closely spaced flashes of light. Fig. 11 presents the beginning of such an analysis by showing PDE responses to single flashes of light that bleach $3 \times 10^4$ rhod./o.s. (0.001% of the total rhodopsin present). This

![Figure 9](image)

**Figure 9.** Dependence of PDE activity on Ca$^{++}$ concentration. PDE activity was measured 1 min after the start of continuous illumination bleaching $2.6 \times 10^3$ rhod./o.s.-s. Outer segment suspensions containing 0.5 mM ATP, 0.5 mM GTP, and 0.1 mM Ca$^{++}$ were buffered to the indicated Ca$^{++}$ levels by the addition of different amounts of EGTA or CaCl$_2$ (see Materials and Methods). The data are taken from seven separate experiments (i.e., outer segment suspensions were obtained from seven different frogs). The curve was drawn by hand.

represented the lowest light level at which reproducible responses could be easily obtained.

One sees in Fig. 11 the increase and decrease (activation and inactivation) in PDE that follows a flash delivered under a variety of conditions. Several effects that may contribute to explaining how changing ATP or calcium concentration alters PDE activity can be seen. Note that Fig. 11 b is distinctively different from Fig. 11 a, c, and d: in $10^{-8}$ M Ca$^{++}$ with ATP present, less PDE activity is elicited by the dim flash and inactivation is relatively rapid. This is compatible with Fig. 5, which shows that under these conditions the response of PDE to continuous illumination is most highly suppressed. If the ATP is removed more PDE activity is generated by the flash, and its decay is slower (cf. Fig. 11 a and b). The rate constant for inactivation changes from
0.061 to 0.017 s\(^{-1}\). (The rate constants are best-fit values obtained by assuming
first-order reactions.) The reduced effect of ATP in \(10^{-3}\) M Ca\(^{++}\) noted in the
previous figure (cf. Fig. 7) is also clear from examining Fig. 11 c and d. ATP
has little effect on the peak activity generated by the flash, and its addition
causes a small but significant increase in the inactivation rate constant (from
0.02 to 0.03 s\(^{-1}\)). This is compatible with the shift shown in Fig. 7 (closed vs.
open symbols). A further point is that the time to reach peak activity after a
flash in the presence of ATP (Fig. 11 b and d) is shorter than in its absence
(Fig. 11 a and e). The response time of the pH electrode used (1–2 s) did not
permit accurate measurements of the time delay before peak activity.

Fig. 12 illustrates another interesting feature of the system. The response
of PDE to a series of flashes is not a linear summation of responses of the sort
shown in Fig. 11. Rather, as a series of flashes is presented at 15-s intervals, a
nonlinear summation is observed. Each succeeding flash is less effective in
stimulating PDE activity. Fig. 12 a illustrates data obtained at \(10^{-9}\) M Ca\(^{++}\).
The open symbols show the peaks of PDE activity elicited by each flash at 15-
s intervals (0, 15, 30 s, etc.). After the third flash, PDE activity no longer
increases above the level reached after the previous flash, and a steady state
is reached. Finally, a saturating flash is delivered (arrow) to demonstrate
maximum activation of the enzyme. The closed circles show the response of
PDE to flashes of light spaced at 1-min intervals. Now each response is similar;
there appears to be no residual effect of a flash presented 1 min previously. Delivery of a saturating flash (arrow) again demonstrates maximum activation. Fig. 12 b shows analogous data obtained using a calcium concentration of $10^{-6}$ M. Similar behavior is noted, except that individual responses are a bit larger because the calcium concentration has increased (cf. Figs. 9 and 10).

The data demonstrate that a short-lived process is triggered by illumination that renders a second flash less effective. The process decays within 1 min, so that the responses of PDE to flashes spaced at 1-min intervals are very similar.

**Figure 11.** Response of PDE to single flashes of light. A subsaturating flash bleaching 0.001% of the rhodopsin present was given to rod outer segments in the presence of 4 mM cyclic GMP with ATP (0.5 mM ATP + 0.5 mM GTP) and without ATP (1 mM GTP) in $10^{-9}$ M Ca$^{++}$ (a and b) and $10^{-8}$ M Ca$^{++}$ (c and d). Proton evolution was measured at the indicated times, activity of a dark control was subtracted, and net activity resulting from the flash was graphed as a function of time after the flash. Assuming first-order kinetics, best-fit lines were obtained (solid lines), and the rate constants are shown. The points are the mean ($\pm$SEM) of three (a), six (b), and two (c and d) separate experiments. Variation between the points of the two experiments of (c) and (d) was within $\pm5\%$.

**DISCUSSION**

An emphasis in this work has been the study of the activation of the PDE complex under conditions that minimally perturb the rod outer segments. The structures must be disrupted to permit access of substrate and controlling nucleotides, but purification and washing procedures have not been used because work in several laboratories (see references in Robinson et al., 1980)
has shown that controlling elements can be easily eluted. Thus, we hope that the experiments are as relevant as is possible to the in vivo expression of the enzyme. The Mg\(^{++}\) and GTP cofactors known to be required for PDE activation (Wheeler and Bitensky, 1977) have been kept constant at millimolar levels so that the Ca\(^{++}\) and ATP controls could be examined. Since both ATP and GTP are known to be present in intact outer segments at millimolar levels, and because light has been shown to decrease GTP but not ATP

![Figure 12](image-url)

**Figure 12.** Response of PDE to successive flashes of light. (a) Successive flashes, each of which bleached 0.001% of the rhodopsin present, were given to rod outer segments in 10\(^{-9}\) M Ca\(^{++}\) in the presence of 0.5 mM GTP and 0.5 mM ATP at intervals of 15 (open circles) and 60 s (closed circles). Finally, a saturating flash was given to each of the samples at the time indicated by the arrow. Proton evolution was measured, dark activity was subtracted, and net activity resulting from the flashes was graphed as a function of time. Curves were drawn by hand. (b) Similar experiment but in 10\(^{-6}\) M Ca\(^{++}\).

(Biernbaum and Bownds, 1979; Robinson and Hagins, 1979), it might be reasonable to assume that amounts of ATP sufficient to saturate the PDE controls shown in this paper are always present in living receptor cells. The present data are similar to those obtained by Woodruff and Bownds (1979) for the behavior of cyclic GMP in intact outer segments, supporting the possibility that ATP normally plays a role in controlling PDE sensitivity. In
Ringer's solution containing $10^{-9}$ M Ca++, 0.5 mM ATP, and 0.5 mM GTP, PDE activity reaches its maximum at a light intensity bleaching $2 \times 10^4$ rhod./o.s.-s, and half-maximum activation occurs at levels bleaching $3 \times 10^3$ rhod./o.s.-s (Fig. 8). This is in close agreement with the behavior of cyclic GMP levels in freshly isolated frog rod outer segments, which maintain the ability to perform a light-induced permeability decrease (Woodruff and Bownds, 1979).

While the constant and high levels of ATP present in vivo suggest that changes in ATP concentration may not play a role in the in vivo control of PDE, it does seem reasonable to suppose that calcium concentration changes might be important, for recent work has shown that calcium is extruded from photoreceptors upon illumination (Gold and Korenbrot, 1980; Yoshikami et al., 1980). Whether this extrusion reflects an increase or decrease in cytoplasmic calcium levels is not known. Perhaps the central observation of this work is that a desensitization of light-activated PDE that depends upon ATP is most striking at low calcium concentrations and is almost completely removed as calcium approaches millimolar levels. Lowering calcium concentration causes a shift of the intensity-response curve for the enzyme to higher light levels (Fig. 8). Robinson et al. (1980) (see also Bownds [1980]) have suggested that illumination of the living photoreceptor might cause at some point a lowering of internal calcium levels (due to the calcium extrusion just mentioned) that causes a desensitization of PDE. This process then might constitute part of the mechanism of light adaptation. The data thus far do not suggest that light adaptation might be completely explained by calcium effects on PDE, for these effects shift the intensity-response curve for the enzyme by no more than one log unit in vitro, and normal light adaptation shifts the intensity response curve of the photoreceptor by 3-4 log units (Fain, 1976).

It is useful to consider possible analogies between the behavior of PDE and the light-suppressible conductance mechanisms of rod cells, for one speculation has been that PDE directly controls plasma membrane permeability through its influence on cyclic GMP levels (see Hubbell and Bownds [1979]). Such analogies must be approached with caution, for it is likely that there are significant differences between the activity of PDE in the disrupted outer segment suspensions used in these studies and the activity that occurs in living cells. The rise of PDE activity triggered by flash illumination of broken outer segments is slower than the conductance response (Baylor et al., 1979 a), indicating either that the enzyme is performing more slowly in vitro than in vivo, or, alternatively, that it is not appropriate to correlate its kinetics with the kinetics of the conductance change. It will be important to determine whether the duration of the in vivo conductance response to the levels of illumination used in this work (typically bleaching at least $10^4$ rhod./o.s.) is similar to the duration of the PDE activation (~40–60 s).

The effect of background light (light adaptation) on living rod cells is to reduce the sensitivity to a dim test flash and to shorten the time-course of the
response, suggesting a modification of the kinetics and gain of the transduction mechanisms within the outer segment (Baylor et al., 1979 a). The normal receptor response to continuous light appears to arise from a superposition of events with the shape of the incremental flash responses (Baylor et al., 1979 b). A decrease in PDE sensitivity and shortening of its time-course can be observed (as calcium concentration is lowered in the presence of ATP and GTP) that is similar in form to the change in the wave form of the current response during light adaptation. Fig. 12 further shows a superposition of individual PDE flash responses that generates a steady response, reflecting a process which may be analogous to the superposition of individual conductance responses noted by Baylor et al. (1979 b). Whether or not these similarities prove ultimately to be relevant, it seems likely that there are many reactions other than PDE activation-inactivation that play a role in permeability control, and efforts must also be made to match their properties to the physiology (cf. Bownds [1980]). One such reaction is a light-induced decrease in GTP levels (Biernbaum and Bownds, 1979; Robinson and Hagins, 1979).

The data presented in this paper demonstrate that PDE displays a wide range of behavior in vitro. Activity monitored in 10^-3 M Ca++ media in the absence of ATP is a function of the total amount of rhodopsin bleached. This suggests that each PDE activation event is relatively long lived, with the individual events summing. PDE activity in the presence of ATP and low Ca++, on the other hand, is more nearly a function of the rate of rhodopsin bleaching. This suggests that each photon absorption results in a more transient activation-inactivation sequence, so that the summing of these short lived sequences results in net PDE activity that is set more by the rate of rhodopsin bleaching than by the total number of rhodopsin molecules bleached.

More insight into this behavior is derived by comparing the responses of PDE to discrete flashes of light, rather than to continuous illumination. One expects, by observing first the discrete responses of the enzyme complex to single flashes of light (Fig. 11) and then to a series of closely spaced flashes (Fig. 12), to reproduce the behavior observed with continuous illumination. Flashes elicit a rapid activation (<10 s to peak) of PDE, followed by a slower inactivation. The most obvious effect of ATP is to hasten the inactivation of PDE after a flash, in both low and high Ca++ media (Fig. 11; cf. Liebman and Pugh [1979 and 1980]). Thus, one would expect the enzyme activity at a low level of continuous illumination to be lower in the presence than in the absence of ATP. A second possible control is indicated by Fig. 11. ATP in low-Ca++ media suppresses the peak activity observed after a dim flash of light. Either the lifetime of the substance responsible for activating PDE has actually been shortened or a rapid inactivation process is established. Ca++ inhibits this control (Fig. 11 c and d), and this probably explains why there is only a slight difference in PDE sensitivity in 10^-3 M Ca++ in the presence and absence of ATP with continuous illumination (Fig. 7). A third possible control lies in regulation of the time delay before PDE activity reaches its peak after
a flash. The delay is clearly longer in Ca++ solutions lacking added ATP (Fig. 11 a). One speculation is that ATP might be acting through its role in rhodopsin phosphorylation (Hubbell and Bownds, 1979; Liebman and Pugh, 1979 and 1980), but recent experiments (Hermolin and Bownds1) have shown that the ATP effect on PDE can be observed under conditions that suppress rhodopsin phosphorylation. Further experiments will be required to determine the locus of the calcium and ATP effects.

It is important to note that controls illustrated by the responses to single flashes of light do not offer a complete explanation of the response of the enzyme complex to continuous illumination. The data of Fig. 12 reveal that a single flash diminishes the effectiveness of a second, similar flash. Thus, if a series of flashes is presented, the response to each grows smaller, and finally a constant rate is set for a given level of rhodopsin bleaching. Cyclic GMP levels and the permeability of isolated outer segments also assume constant values at intermediate levels of illumination, and the present data suggest that this behavior may be determined, at least in part, by the behavior shown in Fig. 12. The process responsible for attenuating responsiveness to successive flashes is short lived, for a flash delivered 1 min after the first flash elicits an equally large response. This suggests that a flash is followed by the transient accumulation of a substance that either inhibits further activation or accelerates inactivation of the PDE response to successive flashes of light. The data do not permit a clear distinction between these possibilities. The lifetime of the inhibitory process is on the same order as that of metarhodopsin II in these preparations, but no effect of ATP and low calcium on the decay kinetics of this intermediate were observed. Further work on the inhibitory process will require that its kinetics be measured as calcium and ATP concentrations are varied.

In studying the PDE complex that apparently is central in the regulation of visual transduction, one faces an array of potential controls. In addition to the GTPase, and, more recently, its “helper” protein (Shinozawa et al., 1980), there is an inhibitor, a light-activated $K_m$ shifter, a Ca++-dependent regulator (Robinson et al., 1980), and now an ATP-dependent regulator (Liebman and Pugh, 1979 and 1980). Current work in our laboratory is directed toward specifying the locus of the Ca++ and ATP controls and probing further analogies between PDE activity and in vivo physiology.

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