Cation Selectivity of and Cation Binding to the cGMP-dependent Channel in Bovine Rod Outer Segment Membranes

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ABSTRACT The properties of the cGMP-dependent channel present in membrane vesicles prepared from intact isolated bovine rod outer segments (ROS) were investigated with the optical probe neutral red. The binding of neutral red is sensitive to transport of cations across vesicular membranes by the effect of the translocated cations on the surface potential at the intravesicular membrane/water interface (Schnetkamp, P. P. M. J. Membr. Biol. 88: 249-262). Only 20-25% of ROS membrane vesicles exhibited cGMP-dependent cation fluxes. The cGMP-dependent channel in bovine ROS carried currents of alkali and earth alkali cations, but not of organic cations such as choline and tetramethylammonium; little discrimination among alkali cations (K > Na = Li > Cs) or among earth alkali cations (Ca > Mn > Sr > Ba = Mg) was observed. The cation dependence of cGMP-induced cation fluxes could be reasonably well described by a Michaelis-Menten equation with a dissociation constant for alkali cations of about 100 mM, and a dissociation constant for Ca²⁺ of 2 mM. cGMP-induced Na⁺ fluxes were blocked by Mg²⁺, but not by Ca²⁺, when the cations were applied to the cytoplasmic side of the channel. cGMP-dependent cation fluxes showed a sigmoidal dependence on the cGMP concentration with a Hill coefficient of 2.1 and a dissociation constant for cGMP of 92 μM. cGMP-induced cation fluxes showed two pharmacologically distinct components; one component was blocked by both tetracaine and L-α-diltiazem, whereas the other component was only blocked by tetracaine.

INTRODUCTION

Excised patches of plasma membrane of the outer segment of amphibian and mammalian rod photoreceptors (ROS) contain a cGMP-dependent conductance that in the intact cell appears to carry the light-sensitive current (Fesenko et al., 1985; Yau and Nakatani, 1985; Zimmerman et al., 1985; Matthews, 1987; Hanke et al., 1988; Quandt et al., 1988). cGMP-induced cation fluxes (mostly Ca²⁺) have also been described in preparations of both amphibian and bovine ROS membrane vesicles (Caretta et al., 1979, Caretta and Cavaggioni, 1983; Caretta, 1985; Koch and Kaupp, 1985; Schnetkamp and Bownds, 1987; Bauer, 1988). The cGMP-dependent channel has been purified and functionally reconstituted from bovine rods (Cook et al., 1987). The preparation of bovine ROS membrane vesicles offers an opportunity...
to study the properties of the cGMP-dependent channel from mammalian ROS, a preparation less accessible to electrical recordings due to their small size. One point of distinction between the bovine and amphibian cGMP-dependent channels has been reported: cGMP-dependent Ca$^{2+}$ fluxes in bovine ROS show two pharmacologically distinct components (Koch et al., 1987; Schnetkamp, 1987), while only a single component is observed in frog ROS (Nicol et al., 1987). In this study I have used a recently developed optical technique to measure electrogenic cation fluxes in a suspension of membrane vesicles (Schnetkamp, 1985a, b). The cation selectivity of and cation binding to the cGMP-dependent channel in bovine ROS membranes were measured, and the results are compared with those obtained for the light-sensitive and cGMP-dependent conductance observed in the amphibian rod. In addition, the cGMP-dependent channel offers a case to illustrate an optical probe for measuring small ionic currents not in single cells, but in a suspension of cells or cellular organelles.

**Materials and Methods**

**Preparation of ROS Membrane Vesicles**

Bovine eyeballs were purchased from a local abattoir and collected as fresh as possible in a light-tight container. ROS with a sealed plasma membrane were isolated and purified as Ca$^{2+}$-rich or Ca$^{2+}$-depleted intact ROS as described before (Schnetkamp, 1986). In the Ca$^{2+}$-depleted preparation dithiothreitol (0.5 mM) was present throughout the procedure. After purification and washing, intact ROS were resuspended to a final concentration of 150–250 $\mu$M rhodopsin in a medium containing 600 mM sucrose, 5% wt/vol Ficoll 400, 0.5 mM dithiothreitol, and 20 mM HEPES (adjusted to pH 7.4 with arginine). The suspension was immediately frozen by putting its container in an ethanol/dry ice bath. After thawing, the ROS suspension was subjected to a hypotonic shock by a 20-fold dilution in 10 mM HEPES (adjusted to pH 7.4 with arginine); after 2 min, 25 mM tetramethylammonium chloride (TMAC$^+$) was added from a 2.5 M stock solution to facilitate sedimentation of ROS membranes by centrifugation (20 min at 10,000 rpm or 12,000 g in a J-20 rotor [Beckman Instruments, Inc., Palo Alto, CA]). The supernatant was discarded and the pellet resuspended to a final rhodopsin concentration of 200–500 $\mu$M in a medium containing 300 mM sucrose, 2.5% Ficoll 400, 0.5 mM dithiothreitol, and 20 mM HEPES (adjusted to pH 7.4 with arginine); this preparation will be referred to as ROS membrane vesicles. Measurements of cGMP-induced cation fluxes were completed within 60 min. All the above and the following procedures were carried out under dim red light illumination.

After the above protocol cGMP-independent cation fluxes in ROS membrane vesicles were minimized, and no such permeabilities appeared to be present in membranes derived from Ca$^{2+}$-depleted ROS. In membranes derived from Ca$^{2+}$-rich ROS that were not immediately frozen, a significant K$^+$ permeability was frequently observed in the absence of cGMP. No experiments on the cGMP-independent K$^+$ permeability are reported here.

**Measurement of Cation Fluxes across ROS Membrane Vesicles**

Cation fluxes across ROS membrane vesicles were measured with the optical probe neutral red as described before (Schnetkamp et al., 1981; Schnetkamp, 1985a, b; Schnetkamp and Szerencsei, 1989). The physicochemical basis of this method is illustrated schematically in Fig. 1. Neutral red adsorbs to phospholipid bilayer membranes carrying a net negative charge (for example due to the presence of acidic phospholipids such as phosphatidylycerine and
SCHNETKAMP  cGMP-dependent Channels in Bovine Rods

FIGURE 1. Neutral red as a probe for surface potentials. A, Molecular structure of neutral red; the arrow indicates the site of protonation. B, Neutral red readily crosses biological membranes in the uncharged basic form, but not in the positively charged acidic form. C, Absorption spectrum of equal concentrations of neutral red at pH 7.4 in aqueous solution (broken line) or adsorbed to ROS membranes (solid line). D, Effect of cation concentration on amount of neutral red adsorbed to membrane surface.
phosphatidylinositol); at pH 7.4 the neutral red adsorbed to the membrane is fully protonated, whereas neutral red in aqueous solution is predominantly unprotonated (Schnetkamp et al., 1981). As a consequence, membrane-bound neutral red can be distinguished easily from its aqueous counterpart in a spectrophotometer. In a dual wavelength spectrophotometer, a change in the difference in absorption $A_{540} - A_{650}$ is a quantitative measure for a change in the amount of membrane-bound dye. Two forces control the binding of a positively charged amphiphile such as neutral red to phospholipid bilayers: first, the electrostatic surface potential as illustrated in Fig. 1; second, the transmembrane potential in the case that the charged form of neutral red can cross the bilayer membrane. The binding of neutral red to ROS membranes is not affected by changes in membrane potential (Schnetkamp and Szerencsei, 1989), leaving the electrostatic surface potential as the dominant parameter controlling changes in binding of neutral red to ROS membranes. Thus, the binding of the positively charged protonated form of neutral red is a function of the electrostatic surface potential at the membrane/water interface, which in turn is a simple function of the concentration and valency of the cations in the aqueous solution (as illustrated in Fig. 1 by potassium ions). In the next section I will demonstrate that changes in the binding of neutral red to the intravesicular membrane surface (measured by the change in the difference of absorption $A_{540} - A_{650}$) can provide a quantitative measure for cation fluxes across the vesicle membrane (see Fig. 2).

The suspension of ROS membranes was diluted to final concentration of 20 μM rhodopsin in 300 mM sucrose, 0.5 mM dithiothreitol, 50 μM neutral red, 20 mM HEPES (adjusted to pH 7.4 with arginine), and 1 mM FCCP when indicated. Aliquots of 2 ml were placed in a cuvette, kept at a constant temperature by means of a circulating waterbath, and mixed with a magnetic stirrer. The difference in light absorption ($A_{540} - A_{650}$) upon addition of the various cations, cGMP, and ionophores was monitored in a DW2C dual wavelength spectrophotometer (SLM-Aminco, Urbana, IL).

Four control experiments were done to ensure that the cGMP-induced changes in light absorption in the presence of alkali cations were due to changes in the binding of neutral red, and not to other causes. First, when the compartment containing the cGMP-dependent channel was equilibrated with alkali cations by a first addition of either cGMP (500 μM) or gramicidin, a second addition of cGMP (500 μM) did not elicit any absorption change other than that due to dilution (0.002 absorbance units). Second, addition of cGMP in the absence of the dye neutral red did not elicit any absorption change. Third, addition of cGMP in the presence of impermeable cations such as choline and tetramethylammonium did not elicit any absorption change. Finally, cGMP-induced changes in the binding of neutral red were not due to an acidification of the medium caused by hydrolysis of cGMP; no hydrolysis of cGMP occurred as tested with the pH-indicating dye phenol red (activation of the rod phosphodiesterase required the presence of GTP). In all experiments cGMP was added from concentrated stock solutions in a volume of 5 μl to a total volume in the cuvette of 2 ml. The starting value for $A_{540} - A_{650}$ amounted to ~1.5.

**Calibration of cGMP-induced Cation Fluxes**

In all experiments reported in this study, cGMP-induced cation influx into ROS membrane vesicles was electrically compensated for by an efflux of protons via the electrogenic protonophore FCCP. Therefore, it is possible to obtain a quantitative calibration of cation flux as indicated by the cGMP- or gramicidin-induced changes in light absorption due to the unbinding of neutral red by measuring the proton efflux with the pH-indicating dye phenol red. Phenol red does not bind to ROS membranes and reports on the pH in the suspension medium (Schnetkamp and Kaupp, 1985); under our experimental conditions pH-indicating absorption changes were linearly related to the amount of HCl added. In this experiment,
FIGURE 2. Calibration of cation flux across ROS membrane vesicles as a function of cation-induced changes in light absorption due to unbinding of neutral red at the intravesicular membrane/water interface. A, Na\(^+\)-induced release of neutral red from the intravesicular surface (solid circles) was compared with (a) Na\(^+\)-induced release of neutral red from the extravesicular surface (open circles) and (b) Na\(^+\)-induced proton release from the intravesicular surface (triangles). The suspension medium of ROS membrane vesicles in the neutral red experiments contained 300 mM sucrose, 50 \(\mu\)M neutral red, and 20 mM HEPES (brought to pH 7.4 with arginine); the suspension medium of ROS membrane vesicles in the proton release measurements contained 300 mM sucrose, 20 mM tetrathylammonium chloride, 40 \(\mu\)M phenol red, and 0.5 mM HEPES (brought to pH 7.4 with arginine). All measurements were done in the dual wavelength mode as \(A_{405} - A_{650}\) (neutral red) or \(A_{570} - A_{650}\) (phenol red). Absorption changes were normalized by dividing the value observed at each given Na\(^+\) concentration by the value observed for the highest Na\(^+\) concentration of 100 mM. Neutral red release from the extravesicular membrane surface was measured by the instantaneous absorption changes upon addition of NaCl; release of neutral red from the intravesicular surface was measured by the absorption changes in the presence of extravesicular Na\(^+\) observed upon addition of gramicidin. Absorption changes originating from the intravesicular membrane surface were typically twice as large as those originating from the extravesicular membrane surface. Proton release from the intravesicular space was measured by the \((A_{570} - A_{650})\) upon addition of gramicidin 10 s after addition of NaCl. B, Na\(^+\)-induced proton release from the intravesicular space (measured with the pH-indicating dye phenol red) was compared with Na\(^+\)-induced changes in the binding of neutral red to the intravesicular membrane surface (measured as \(A_{440} - A_{660}\)). Absorption changes by the dye phenol red were calibrated into moles H\(^+\) release/mole rhodopsin by adding known amounts of HCl to the suspension.

ROS membrane vesicles were made permeable to Na\(^+\) and protons by addition of the channel ionophore gramicidin. Fig. 2 A compares the dependence on Na\(^+\) concentration of (a) the amplitude of Na\(^+\)-induced unbinding of neutral red from the extradiskal membrane surface, (b) the amplitude of the Na\(^+\)-induced unbinding of neutral red from the intradiskal membrane surface, and (c) the Na\(^+\)-induced release of protons from the intradiskal space; the amplitudes of all three parameters depended on the external Na\(^+\) concentration in a very similar manner. The relatively large changes observed between 1 and 10 mM are typical for Na\(^+\)-induced...
screening of surface potentials (Schnetkamp, 1985a). Na⁺-induced changes in light absorption due to unbinding of neutral red were linearly related to Na⁺-induced proton release (Fig. 2 B) and can be converted in a quantitative measurement of Na⁺ flux. The above results could be interpreted to indicate that neutral red bound to the surface of ROS membranes acts as a pH indicator at the membrane surface rather than as an indicator of surface potential; these two mechanisms are indistinguishable since the surface potential controls the surface pH, and hence the degree of protonation of acidic and basic groups on the membrane surface. Sodium ions (or any other cation) entering the intravesicular space replace protons as counterions for the fixed negatively charged groups on the membrane. Neutral red and protons both carry a single positive charge and their binding to the membrane depends on an identical way on the surface potential and cation-induced changes thereof. The observation that changes in neutral red binding on the external and internal membrane surface, respectively, and proton release from the intravesicular space were all very similar with respect to their dependence on the Na⁺ concentration, supports the conclusion that Na⁺-induced proton release is mediated by Na⁺-induced changes in surface potential and a concomitant degree of protonation of membrane components. The issue of intravesicular pH changes is discussed in more detail in Schnetlomp (1985a).

RESULTS

cGMP-induced Na⁺ Transport in Membrane Vesicles Derived from Intact Bovine ROS

In the first experiment I describe the basic observation on cGMP-induced changes in Na⁺ flux in membrane vesicles derived from intact ROS after freeze-thawing and hypotonic shock; this preparation consists of pieces or vesicles of inverted plasma membrane with the disks attached to them via filamentary structures (Molday et al., 1987). Closed vesicles could be vesicles of inverted plasma membrane, intact disks, or perhaps vesicles of fused disk and plasma membrane. In the following sections I will use the term ROS membrane vesicles to indicate the ensemble of closed membrane vesicles derived from intact bovine ROS. In order to be detected with the method used in this study, cGMP-dependent channels have to reside in sealed vesicles with the cGMP binding sites of the channel on the extravesicular membrane surface.

Addition of 100 mM NaCl to a suspension of ROS membrane vesicles in a buffered sucrose medium containing 50 μM neutral red and 1 μM FCCP caused an instantaneous change in absorption (A₄₅₀ - A₆₅₀) due to dilution and to Na⁺-induced unbinding of neutral red from the external surface of the membrane vesicles (Fig. 2). The instantaneous change in light absorption was followed by time-resolved changes indicating unbinding of neutral red from the intravesicular membrane surface; a typical example is depicted in Fig. 3. As discussed in the Methods, the unbinding of neutral red is most likely caused by Na⁺-induced changes in the surface potential at the intravesicular membrane/water interface and therefore reflects Na⁺ transport into the vesicles. Addition of the ionophore gramicidin, a nonselective channel for alkali cations, caused a rapid unbinding of neutral red from the intravesicular membrane surface, consistent with a rapid increase in the intravesicular Na⁺ concentration due to Na⁺ influx via gramicidin. The calibration plot illustrated in Fig. 2 can be used to convert changes in light absorption due to
unbinding of neutral red into quantitative cation fluxes independent of the molecular mechanism underlying the unbinding of neutral red.

In the absence of gramicidin or cGMP, Na⁺ influx into ROS membrane vesicles ranged from 0 to 0.1 pA when expressed as a current through the ensemble of vesicles derived from a single bovine ROS. In this and subsequent calculations I assumed that the total rhodopsin concentration in bovine ROS amounts to 3 mM and that a single bovine ROS is a cylinder of 1 x 20 μm and contains 2.9 x 10⁷ rhodopsin molecules.

Addition of 500 μM cGMP before addition of gramicidin caused a rapid increase in the rate of absorption change in the same direction as that observed for gramicidin. However, the amplitude of the cGMP-induced absorption change amounted to only 20% of that observed for gramicidin; a value of 20–25% was observed consistently in many preparations of ROS membrane vesicles. This result suggests that cGMP opens an Na⁺-permeable channel. Adding gramicidin after addition of cGMP caused a rapid change in light absorption to the same value as that observed when only gramicidin was added, whereas addition of cGMP after addition of gramicidin did not cause any absorption change (not shown). The effects of gramicidin and cGMP were not additive, i.e., membrane vesicles equilibrated with Na⁺ via the cGMP-dependent channel did not take up any further Na⁺ after addition of gramicidin. The average cGMP-induced Na⁺ flux in ROS membrane vesicles (derived from a single ROS) at 100 mM Na⁺ amounted to 5 x 10⁶ Na⁺/outer segment per s or a current of 0.8 pA; flux rates were obtained by measuring the initial rate of cGMP-induced change in light absorption and converting this into flux with the calibration curve shown in Fig. 2.
cGMP-induced Na⁺ Fluxes Reflect a Conductance Mechanism

In the above experiment I used the electrogenic protonophore FCCP with the objective of providing an outward proton current to compensate for the putative cGMP-induced inward Na⁺ current; this aspect is investigated in more detail in the experiments illustrated in Fig. 4A. Addition of cGMP in the presence of an inward gradient of Na⁺ did not cause a rapid change in light absorption in the absence of FCCP. For comparison, traces are included for the absorption changes when no cGMP was added, or when Na⁺ was replaced with the impermeable cation tetramethylammonium. In a parallel experiment I measured cGMP-induced proton release from ROS membrane vesicles with the pH-indicating dye phenol red (with a protocol similar to that used for the experiment illustrated in Fig. 2); addition of cGMP in the presence of an inward gradient of Na⁺ caused very little proton release unless FCCP was present (not illustrated). A likely explanation for the above observations is that in the absence of FCCP little mass influx of Na⁺ via the cGMP-dependent channel occurred, but instead an inside-positive membrane potential developed. In the presence of FCCP, the cGMP-induced influx of Na⁺ appeared

![Graph A](image1)

**Figure 4.** cGMP-induced Na⁺ fluxes reflect cGMP-induced Na⁺ currents. Experimental conditions were as described in the legend of Fig. 3; 100 mM NaCl was added 10 s before the start of the recordings and 500 µM cGMP was added at time zero. The calibration bar represents a change in light absorption by 0.01 absorbance unit. Temperature, 25°C. A, Recordings are labeled to indicate the omission of FCCP (no FCCP), omission of cGMP (no GMP), replacement of Na⁺ by tetramethylammonium ion (TMA⁺), and control (FCCP). B, The recordings were started by addition of 500 µM cGMP 10 s after addition of 100 mM NaCl. The arrows indicate the time of addition of FCCP. In the recording that showed an immediate response to addition of cGMP, FCCP was present at the start.
to be compensated for by an efflux of protons. FCCP does not appear to have a pharmacological effect on the cGMP-dependent channel as judged from the observation that FCCP had no effect on cGMP-induced Ca$^{2+}$ release measured with arsenazo III (not illustrated). cGMP-induced Ca$^{2+}$ release was observed in the absence of any ionophores (Caretta and Cavaggioni, 1983; Koch and Kaupp, 1985; Schnetkamp, 1987; Bauer, 1988). In the above studies cGMP-induced Ca$^{2+}$ release was probably electrically compensated for by cGMP-induced alkali cation influx and no net current presumably occurred (discussed in Koch and Kaupp, 1985; Schnetkamp, 1987).

**Do cGMP-induced Na$^+$ Fluxes Inactivate?**

The amplitude of the cGMP-induced changes in light absorption was much smaller compared with that induced by gramicidin (Fig. 3). This could be interpreted in two ways: it could reflect inactivation of the cGMP-dependent channels a few seconds after opening, or it could suggest that only 20% of the membrane vesicles contained a functional cGMP-dependent channel. The use of FCCP offers an opportunity to investigate the possibility of inactivation of the cGMP-dependent channel. In this experiment cGMP was added to ROS membrane vesicles immediately after establishment of an inward Na$^+$ gradient, but before addition of FCCP; after some time interval FCCP was added to initiate a net inward Na$^+$ flux and outward proton flux. Increasing the time interval between additions of cGMP and FCCP, respectively, should give rise to progressively smaller signals if the cGMP-dependent channels inactivate in this time interval (Fig. 4 B). This is not observed, demonstrating that the cGMP-dependent channel does not inactivate on a time scale of 2 min.

**Cation Selectivity of the cGMP-dependent Channel**

The cation selectivity of the cGMP-dependent channel was investigated in the experiments illustrated in Fig. 5 and summarized in Table I. Most of the experiments with alkali cations were conducted at a temperature of 5°C, which decreased the rate of cGMP-induced cation fluxes by about fivefold compared with those observed at 25°C (Fig. 5; note that the traces are separated by 0.01 absorption units). The first thing that can be noticed in Fig. 5 is that the amplitude of the cGMP-induced absorption changes was identical for the different alkali cations tested; this is consistent with the notion that electrostatic screening of surface potentials by cations is independent of the type of cation; it suggests that screening rather than charge compensation (binding) is the dominant mechanism by which alkali cations reduce the surface potential. The amplitude of the cGMP-induced signals observed for divalent cations (tested at 20 mM) was somewhat more variable, suggesting that charge compensation (binding) contributes to the reduction of surface potential by divalent cations (not illustrated); this result can be anticipated from similar effects of divalent cations on the surface potential of simple phospholipid bilayer membranes (McLaughlin et al., 1978, 1981). The ion selectivity of cation fluxes through the cGMP-dependent channel was obtained by measuring the initial rates of cGMP-induced light absorption changes in the presence of different cations; the measurements are equivalent to electrically recorded conductance ratios and the results are
Figure 5. Ion selectivity of cGMP-induced alkali cation fluxes. Experimental conditions were as in the legend of Fig. 3; 100 mM of the indicated cation chlorides was added 10 s before addition of 500 μM cGMP at time zero. TMA is tetramethylammonium ion. Note that the recordings are separated by 0.01 absorbance units. Temperature, 5°C.

summarized in Table I. No cGMP-induced fluxes were observed for the organic cations tetramethylammonium and choline.

The cGMP-induced Ca^{2+} currents (at 20 mM Ca^{2+}) were on average 3.5-fold smaller (SD = 0.9, 13 preparations) as compared with cGMP-induced Na^+ currents (at 100 mM Na^+); i.e., the cGMP-induced Ca^{2+} flux was 7-fold smaller than the cGMP-induced Na^+ flux.

| Cation | Relative fluxes | Standard deviation | No. of observations |
|--------|-----------------|--------------------|---------------------|
| Li^+   | 0.83            | 0.12               | 8                   |
| Na^+   | 0.88            | 0.10               | 8                   |
| K^+    | 1.00            | 0.10               | 7                   |
| Ca^{2+}| 0.59            | 0.10               | 11                  |
| Mg^{2+}| 0.50            | 0.10               | 11                  |
| Mn^{2+}| 0.86            | 0.26               | 5                   |
| Ca^{2+}| 1.00            | 0.26               | 5                   |
| Sr^{2+}| 0.64            | 0.09               | 10                  |
| Ba^{2+}| 0.47            | 0.07               | 10                  |

cGMP-induced alkali cation fluxes were measured at 500 μM cGMP and 100 mM of the cation chloride salts; fluxes were normalized with respect to the fluxes of the most conductive cation K^+. cGMP-induced divalent cation fluxes were measured at 500 μM cGMP and 20 mM of the divalent cation chloride salts; fluxes were normalized with respect to the fluxes of the most conductive divalent cation Ca^{2+}. Results from experiments conducted at 5 and 25°C were pooled; no difference in ion selectivity was apparent when experiments conducted at 5°C were compared with those conducted at 25°C.
Cation Binding to the cGMP-dependent Channel

The cGMP-induced cation fluxes were dependent on the external cation concentration; both for Ca\(^{2+}\) and alkali cations the dependence of cGMP-induced flux rate on the external cation concentration was reasonably well described by a single-site Michaelis-Menten equation (Figs. 6 and 7). This suggests that the cGMP-dependent channel has a binding site for Ca\(^{2+}\) with a dissociation constant of 2 mM (Fig. 6), and a binding site for alkali cations with dissociation constants between 75 and 150 mM (Fig. 7).

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**Figure 6.** Ca\(^{2+}\)-dependence of cGMP-induced Ca\(^{2+}\) fluxes. cGMP-induced Ca\(^{2+}\) fluxes were measured as a function of Ca\(^{2+}\) concentration following the protocol described in Figs. 3 and 5. The flux rate or current was normalized with respect to the maximal current extrapolated from the data; maximal current and dissociation constant were obtained from a linear regression of a Scatchard plot of the data. The dissociation constant obtained was 2.2 mM. The computer-drawn curves represent Michaelis-Menten equations with, from top to bottom, Ca\(^{2+}\) dissociation constants of 1.5, 2.2, and 3.0 mM. Temperature, 25°C.

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**Figure 7.** Alkali cation dependence of cGMP-induced alkali cation fluxes. cGMP-induced alkali cation fluxes were measured as a function of cation concentration following the protocol described in Figs. 3 and 5. The flux rate or current was normalized with respect to the maximal current extrapolated from the data; maximal current and dissociation constant were obtained separately for each cation from a linear regression of a Scatchard plot of the data. The dissociation constants obtained were 80 mM (Li\(^+\), circles), 144 mM (Na\(^+\), triangles), 120 mM (K\(^+\), squares), and 101 mM (Cs\(^+\), inverted triangles). The computer-drawn curves represent Michaelis-Menten equations with, from top to bottom, dissociation constants of 80, 100, 120, and 140 mM. Temperature, 5°C.
Interaction between Cations in the cGMP-dependent Channel

The experiments illustrated in Figs. 6 and 7 show that the cGMP-dependent channel has a higher affinity for Ca\(^{2+}\) as compared with Na\(^{+}\) or other alkali cations; consistent with this, cGMP-induced Ca\(^{2+}\) fluxes at saturating Ca\(^{2+}\) concentration were smaller than cGMP-induced Na\(^{+}\) fluxes at saturating Na\(^{+}\) concentration, since Ca\(^{2+}\) probably resided longer in the channel than Na\(^{+}\). Therefore, the combined cGMP-induced cation flux in the presence of both Ca\(^{2+}\) and Na\(^{+}\) is expected to be less than the cGMP-induced cation flux in the presence of Na\(^{+}\) alone; i.e., Ca\(^{2+}\) is expected to block Na\(^{+}\) fluxes. The experiment illustrated in Fig. 8 shows that this is not the case. cGMP-induced fluxes for the individual ionic species Na\(^{+}\), Ca\(^{2+}\), and Mg\(^{2+}\), respectively, are shown in Fig. 8 A; when applied to the cytoplasmic side of the channel Mg\(^{2+}\), but not Ca\(^{2+}\), inhibited cGMP-induced Na\(^{+}\) fluxes (Fig. 8 B) in agreement with observations on the cGMP-dependent channel in toad rods (Yau et al., 1986). The cGMP-induced Na\(^{+}\) and Ca\(^{2+}\) fluxes were additive at concentrations below their respective dissociation constants (not illustrated).
Dependence of cGMP-dependent Fluxes on cGMP Concentration

The cGMP-induced, cation-selective fluxes illustrated so far were all recorded at a saturating concentration of 500 μM cGMP. Fig. 9A illustrates the dependence of Na⁺ fluxes on the concentration of cGMP. The cGMP-induced Na⁺ flux depended in a sigmoidal manner on the cGMP concentration; the data were transformed into a Hill plot which yielded a straight line with a Hill coefficient of 2.0 and a dissociation constant for cGMP of 75 μM (Fig. 9B). The cGMP-induced fluxes of the different alkali cations tested did not differ significantly in their dependence on the concen-
tration of cGMP; in six different preparations the average dissociation constant for cGMP was 92 μM (SD = 9) and the average Hill coefficient was 2.12 (SD = 0.11).

**Evidence for the Existence of Two Distinct Forms of the cGMP-dependent Channel**

Previous studies with bovine ROS membrane vesicles showed that the cGMP-induced Ca²⁺ flux measured with arsenazo III had two components with different kinetics, with different dependence on cGMP concentration, and with a different pharmacology; similar experiments in frog ROS revealed only a single component (Koch et al., 1987; Nicol et al., 1987; Schnetkamp, 1987). The cGMP-induced fluxes observed in this study showed little evidence for two components with different kinetics (e.g., Fig. 5). In the experiment illustrated in Fig. 10, I have used the pharmacology of the cGMP-dependent channel to test whether cGMP-induced Na⁺ fluxes have two distinct components. One component of cGMP-induced Ca²⁺ release is fully activated by 50 μM cGMP and can be blocked by tetracaine, but not by L-cis diltiazem; the other component of cGMP-induced Ca²⁺ release requires a much higher concentration of 500 μM cGMP to be fully activated and is blocked by both L-cis

![Figure 10. Pharmacology of cGMP-induced fluxes. Experimental conditions were as described in the legend of Fig. 3. 100 mM NaCl was added 10 s before addition at time zero of 50 μM cGMP (A) or 500 μM cGMP (B). No cGMP was added in the recordings labeled “no cGMP.” Tetracaine and the two stereoisomers of diltiazem were present at a final concentration of 20 μM as indicated; d-cis diltiazem reduced the amplitude of cGMP-induced changes in light absorption by 25%. Temperature, 25°C.](image-url)
diltiazem and (less effectively) tetracaine (Koch et al., 1987; Schnetkamp, 1987). Drugs such as tetracaine and diltiazem are positively charged amphiphiles and adsorb strongly to negatively charged phospholipid bilayer membranes (much like neutral red); for this reason tetracaine and diltiazem have a strong effect on the surface potential and in our assay their use is limited to concentrations not exceeding 20 μM. I used d-cis diltiazem, the stereoisomer of L-cis diltiazem which does not block the cGMP-dependent channel, as a control to cancel out the effect of the drugs on the surface potential.

If cGMP-induced Ca^{2+} fluxes observed with arsenazo III (Koch and Kaupp, 1985; Schnetkamp, 1987) are carried by the same channel responsible for cGMP-induced Na^{+} fluxes observed in this study, the latter should show two components with the following characteristics: cGMP-induced Na^{+} fluxes at 50 μM cGMP should be blocked by tetracaine, but not by L-cis diltiazem, whereas cGMP-induced Na^{+} fluxes at 500 μM cGMP should be blocked by both L-cis diltiazem and (less effectively) by tetracaine. The experiment illustrated in Fig. 10 shows exactly this pattern (similar results were obtained in eight other preparations for cGMP-induced Ca^{2+} and Mg^{2+} fluxes as well as for cGMP-induced fluxes of the other alkali cations tested). Note that in the presence of L-cis diltiazem the cGMP-induced Na^{+} fluxes at 50 and 500 μM cGMP, respectively, were nearly identical as observed before for cGMP-induced Ca^{2+} fluxes measured with arsenazo III (Schnetkamp, 1987).

**DISCUSSION**

*An Optical Probe to Measure Cation Fluxes in Cell Suspensions*

Membrane vesicles derived from intact bovine ROS offer a convenient preparation to study the properties of the cGMP-dependent channel from a mammalian source with the optical probe neutral red; a single measurement with neutral red averages ~4 × 10^{8} channels originating from 30 different animals. Binding of neutral red to the internal and the external surfaces of ROS membrane vesicles is sensitive to changes in intravesicular and external cation concentration, respectively; the properties of cation-induced changes in binding of neutral red are consistent with the notion that cations influence binding of neutral red via their effect on the surface potential at the membrane/water interface (Schnetkamp, 1985a; Figs. 1 and 2). The electrogenic protonophore FCCP in combination with a large proton buffer capacity of ROS membrane vesicles appeared to provide a current loop for inward cation currents through the cGMP-dependent channel; inward cation fluxes via the cGMP-dependent channel or via the added channel ionophore gramicidin were mirrored by outward proton fluxes (measured with the pH-indicating dye phenol red), and this provided a quantitative calibration of changes in light absorption (caused by unbinding of neutral red) into cation flux or current (Fig. 2). The relationship between changes in light absorption due to unbinding of neutral red and proton/cation fluxes was linear; this reflects the fact that the binding of both neutral red and protons to the membrane surface depends in an identical way on the surface potential since both carry a single positive charge.

The cGMP-dependent channel appeared to be the dominant conductive mechanism of Na^{+} transport present in ROS membrane vesicles. The average cGMP-
induced Na\(^+\) flux (at 100 mM Na\(^+\)) in ROS membrane vesicles derived from a single
ROS amounted to 5 × 10\(^6\) Na\(^+\)/outer segment per s, equivalent to a current of 0.8
pA, whereas Na\(^+\) currents in the absence of cGMP ranged between 0 and 0.1 pA;
Na\(^+\) transport in the absence of cGMP increased upon aging and probably reflected
passive leakage through the membrane. cGMP-induced fluxes were calculated from
the initial rates of changes in light absorption, converted to fluxes with the help of
the calibration plot illustrated in Fig. 2. A cGMP-induced current of 0.8 pA or an
Na\(^+\) flux of 5 × 10\(^6\) Na\(^+\)/ROS per s can be compared with cGMP-induced Ca\(^{2+}\)
release of 1.1 × 10\(^5\) Ca\(^{2+}\)/ROS per s from the same preparation of bovine ROS
membrane vesicles measured with arsenazo III (Schnetkamp, 1987) or with cGMP-
induced Na\(^+\) currents of 2 nA in excised patches of bovine ROS plasma membrane
(Quandt et al., 1988 and manuscript submitted); those patches were derived from
the same preparation of intact ROS that was used to prepare the ROS membrane
vesicles in this study. The discrepancy in the magnitude of cGMP-induced currents
observed in ROS membrane vesicles and excised patches of plasma membrane,
respectively, suggests that the cGMP-dependent channels observed in ROS mem-
brane vesicles reside in membranes with a much lower channel density than the
plasma membrane. Only 20–25% of the bovine ROS vesicles contained functional
cGMP-dependent channels (Figs. 3 and 4), similar to the fraction observed for
cGMP-induced Ca\(^{2+}\) fluxes in bovine ROS membrane vesicles (Koch and Kaupp,
1985; Schnetkamp, 1987) or in leaky but otherwise undisturbed frog ROS (Schnet-
kamp and Bownds, 1987).

**Ion Selectivity of cGMP-dependent Channel in Bovine Rods**

The cGMP-dependent channel in bovine ROS membranes passed all alkali cations
tested with a flux ratio of Li : Na : K : Cs = 0.8 : 0.8 : 1 : 0.6, and it passed all
divalent cations tested with a flux ratio of Mg : Mn : Ca : Sr : Ba = 0.5 : 0.86 : 1 :
0.64 : 0.47, whereas no fluxes were observed for choline, and cGMP-induced Ca\(^{2+}\)
fluxes (at 20 mM) were on average sevenfold smaller compared with cGMP-induced
Na\(^+\) fluxes (at 100 mM). Qualitatively this is consistent with the observation that the
apparent dissociation constant of the cGMP-dependent channel for Ca\(^{2+}\) is consid-
erably smaller than for alkali cations. The above results are in general agreement
with the ion selectivity of the light-sensitive current as measured in amphibian rods
by means of rapid ion substitution experiments (Yau and Nakatani, 1984; Hodgkin
et al., 1985; Cervetto et al., 1988; Menini et al., 1988), although the precise
sequence and flux ratios differ somewhat between the different studies; these
differences may arise from the different species or experimental protocols used (e.g.,
measuring inward currents versus measuring fluxes in the reverse direction). The
properties of the cGMP-dependent channel purified and reconstituted from bovine
ROS (Hanke et al., 1988) differ from the light-sensitive current in intact amphibian
rod cells or from the results of this study with respect to their much lower efficacy in
carrying Li\(^+\) and Ca\(^{2+}\) currents. Some of the apparent discrepancy may be resolved
by the suggestion that two forms of the cGMP-dependent channel exist differing
most noticeably in the ability to pass Ca\(^{2+}\) current (Cervetto et al., 1988).

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