Properties of an Inwardly Rectifying ATP-sensitive K⁺ Channel in the Basolateral Membrane of Renal Proximal Tubule

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ABSTRACT The potassium conductance of the basolateral membrane (BLM) of proximal tubule cells is a critical regulator of transport since it is the major determinant of the negative cell membrane potential and is necessary for pump-leak coupling to the Na⁺,K⁺-ATPase pump. Despite this pivotal physiological role, the properties of this conductance have been incompletely characterized, in part due to difficulty gaining access to the BLM. We have investigated the properties of this BLM K⁺ conductance in dissociated, polarized Ambystoma proximal tubule cells. Nearly all seals made on Ambystoma cells contained inward rectifier K⁺ channels (γ_in = 24.5 ± 0.6 pS, γ_out = 3.7 ± 0.4 pS). The rectification is mediated in part by internal Mg²⁺. The open probability of the channel increases modestly with hyperpolarization. The inward conducting properties are described by a saturating binding–unbinding model. The channel conducts Ti⁺ and K⁺, but there is no significant conductance for Na⁺, Rb⁺, Cs⁺, Li⁺, NH₄⁺, or Cl⁻. The channel is inhibited by barium and the sulfonylurea agent glibenclamide, but not by tetraethylammonium. Channel rundown typically occurs in the absence of ATP, but cytosolic addition of 0.2 mM ATP (or any hydrolyzable nucleoside triphosphate) sustains channel activity indefinitely. Phosphorylation processes alone fail to sustain channel activity. Higher doses of ATP (or other nucleoside triphosphates) reversibly inhibit the channel. The K⁺ channel opener diazoxide opens the channel in the presence of 0.2 mM ATP, but does not alleviate the inhibition of millimolar doses of ATP. We conclude that this K⁺ channel is the major ATP-sensitive basolateral K⁺ conductance in the proximal tubule.

KEY WORDS: ion channel • kidney • patch-clamp • sulfonylurea • epithelia

INTRODUCTION

The ability of the renal proximal tubule to maintain homeostatic electrolyte and water reabsorption in the face of drastic changes in dietary solute and water intake and renal hemodynamics implies that transtubular ion transport is tightly regulated. Proximal tubule potassium channels, particularly in the basolateral membrane (BLM), play pivotal physiologic roles in the regulation of membrane voltage, potassium recycling, and ultimately in transepithelial solute and water reabsorption. That the basolateral membrane potential of a proximal tubule cell is dominated by the BLM K⁺ conductance is well established (Sackin and Boulpaep, 1983). The Na⁺,K⁺-ATPase pump in the BLM provides the energy that makes ion transport thermodynamically favorable, but continuous operation of the pump requires there be a K⁺ exit pathway. The BLM K⁺ conductance provides such a pathway and thus steady state intracellular K⁺ activity can be maintained in the face of large transcellular fluxes of salt and water (Sackin and Boulpaep, 1983). Since a major portion of the transcellular Na⁺ flux is reabsorbed across the BLM by the action of the Na⁺,K⁺-ATPase, a population of BLM K⁺ channels working in concert with the pump would allow K⁺ to recycle in a regulated fashion. Moreover, hyperpolarization secondary to the opening of BLM K⁺ channels enhances the driving force for electrogenic apical Na⁺ entry and basolateral Cl⁻ efflux, resulting in net NaCl reabsorption.

The Ambystoma proximal tubule exhibits a large K⁺ conductance in the basolateral membrane (Siebens and Boron, 1987; Sackin and Boulpaep, 1981) that has been shown macroscopically to be sensitive to barium and pH. However, previous studies of the “macroscopic” BLM K⁺ conductance (Boulpaep, 1976) lack the resolution to distinguish whether there is one population of imperfectly selective K⁺ channels or a set of highly selective K⁺ channels coexisting with a population of nonselective cation channels. More recent “microscopic” single-channel patch-clamp studies of BLM K⁺ channels have shown diversity in both experimental design and findings (Tsuchiya et al., 1992; Hunter, 1991; Parent et al., 1988; Kawahara et al., 1987; Sackin and Palmer, 1987; Gögelein and Greger, 1987a, 1987b).
so a clear consensus has been elusive and details of the properties and regulation are lacking.

We have now characterized the properties and regulation (see Mauerer et al., 1998) of the principal K\(^+\) channel in the BLM in a preparation of dissociated yet polarized Ambystoma proximal tubule cells (Segal et al., 1996). Inwardly rectifying, ATP-sensitive K\(^+\) channels were present in $\geq 95\%$ of recordings from the BLM, each containing from 2 to $\leq 25$ K\(_{ATP}\) channels/patch. Although the regulation of this proximal tubule BLM K\(^+\) channel is similar to that of recently cloned K\(_{ATP}\) channels in the apical membrane of the distal nephron (ROMK1 and ROMK2), there are important differences, and ROMK has not been found in the proximal tubule (Chepilko et al., 1995; Zhou et al., 1994; Lee and Hebert, 1995; Boim et al., 1995; Ho et al., 1993). The studies reported in this paper and the companion paper elucidate the properties and the regulation, respectively, of the major K\(^+\) channel underlying the BLM K conductance that is coupled to transport in the proximal tubule.

**Materials and Methods**

**Solutions and Drugs**

The compositions of the solutions used are summarized in Table I. After titration to pH 7.5 (710A; Orion Research, Boston, MA), sucrose was added to adjust the osmolality of the solutions (3MO; Advanced Instruments Inc., Needham Heights, MA). To determine channel conductance as a function of [K\(^+\)], sucrose was added to the standard pipette KCl to maintain osmolality. Chemicals used were of the highest quality and obtained from Sigma Chemical Co. (St. Louis, MO), except thallium acetate (Aldrich Chemical Co., Milwaukee, WI), diazoxide (Calbiochem Corp., La Jolla, CA), ATP\(_S\), and ADP (Boehringer-Mannheim Biochemicals, Indianapolis, IN). Nucleotides were prepared fresh daily as 20–50-nM stocks in bath solution. Glibenclamide was dissolved in DMSO (100 nM stock).

**Cell Preparation**

Dissociated proximal tubule cells were isolated from amphibian kidneys as previously described (Segal et al., 1996). Briefly, aquatic phase *Ambystoma tigrinum* kept at 4°C were killed by submersion in 0.2% tricaine methanesulfonate. The kidneys were rapidly removed and placed in iced HEPES-buffered NaCl at pH 7.5 (solution \(a\)). The adventitial tissue was removed by hand dissection, and the renal tissue was cut into 1–2 mm\(^3\) pieces and incubated in collagenase-dispase (0.2 U/ml of collagenase; Boehringer-Mannheim Biochemicals) on a gyratory shaker for 60 min at 22°C. The enzyme reaction was stopped by washing with Ca\(^2+\)- and Mg\(^2+\)-free NaCl (solution \(b\)). The cells were then mechanically dispersed into suspension by repeated trituration, and a pellet was obtained by centrifugation at $\sim 1,600$ rpm for 3 min. Finally, the cells were resuspended in 2.5 ml NaCl (solution \(a\)) in a 35-mm culture dish, and stored at 4°C until use. The dissociated proximal tubule cells can retain their epithelial polarity for up to 14 d (Segal et al., 1996). Cells were used for experiments from 2 to 12 d after dissociation. Representative cells as seen under light microscopy (Fig. 1 \(A\)) and scanning electron microscopy (Fig. 1 \(B\)) are shown (for details of methods see Segal et al., 1996).

**Electrophysiology**

A 5-µl aliquot of cell suspension in NaCl storage solution (solution \(a\)) was placed on a Cell-Tak™-coated glass coverslip in a re-

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**Table I**

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Fulfil the following criteria (Segal et al., 1996): (a) characteristic morphology (see Fig. 1; Segal et al., 1996). An individual proximal tubule cell was selected for an experiment only if it exhibited the following characteristics: (1) absence of large vacuoles. (2) Relatively smooth appearing basolateral surface, (3) distinct apical and basolateral membrane domains and the de-}

Figure 1. Distilled Ambystoma proximal tubule cells retain epithelial cell polarity. (A) Light photomicrograph of a single dissociated Ambystoma proximal tubule cell shows distinct apical and basolateral membrane surfaces. The apical surface of these bilobated cells is the smaller lobe with a microvillar brush border. The robust cytoskeleton of these cells includes an actin-rich "waist-band" (arrowheads) between the two membrane domains that is important in retention of epithelial polarity. Giga-ohm seals can be made on both surfaces. Scale bar, 10 μm. (B and C) Scanning electron micrographs showing the sharp transition between the membrane domains and the detailed topology of the apical surface invested with its microvillar brush border, and the basolateral surface with its folds and projections. Scale bars: 10 μm in B, 1 μm in C.

cording chamber of our design (RC-5/25; Warner Instruments, Hamden, CT) mounted on an inverted microscope (Olympus IM; Olympus America, Inc., Melville, NY). The chamber has a bath volume of 500 μl, and solutions are perfused directly into an input multiplexer on the chamber at a gravity-driven flow rate of ~10 ml/min.

Nonadherent cells were washed off the coverslip with the recording solution (either Ca²⁺-free NaCl, solution C, or KCl, solution D; unless otherwise noted) and the cells were visualized under Hoffman modulation optics (Modulation Optics, Greenval, NY). Proximal tubule cells were readily recognized by their characteristic morphology (see Fig. 1; Segal et al., 1996). An individual proximal tubule cell was selected for an experiment only if it fulfilled the following criteria (Segal et al., 1996): (a) distinctly bilobated structure, (b) clearly defined brush border sharply delimited on the apical surface, (c) relatively smooth appearing basolateral membrane, and (d) absence of large vacuoles.

Patch clamp. The standard configurations for single-channel and whole-cell tight seal patch-clamp technique (Hamill et al., 1981) were used to record channel currents from the BLM. Patch pipettes were fabricated from borosilicate glass capillaries (Warner Instruments, Hamden, CT) on a two-step puller (PP-83; Narishige Corp., Midland, MI) to within 200 μm of the tip, and fire-polished just before use. When filled with KCl, the open tip pipette resistance was 3–8 MΩ when placed in the initial bath solution. A hydraulic micromanipulator (Narishige) was used to guide the patch microelectrode to the BLM of the cell. High resistance giga-ohm seals (up to 50 GΩ) were obtained on the BLM in ~75% of attempts by applying gentle suction to the pipette just after it touched the cell membrane. To achieve the whole-cell configuration, further suction was applied to rupture the cell-attached patch. Data have not been corrected for liquid-junction potentials since for most solutions they were <4 mV when measured as follows: the bath Ag-AgCl ground electrode was connected to the control KCl bath through a 3% agar bridge made of KCl pipette solution. A low resistance (<1 MΩ) pipette filled with 3 M KCl was placed in a KCl bath and the DC offset was adjusted to 0 mV in zero current clamp. The liquid-junction potentials were measured as the voltage offset resulting when the control KCl bath was replaced by the test solution. Low [Cl] solutions in which 90% of Cl⁻ was replaced by aspartate had a liquid-junction potential of 13.3 mV.

Voltage-clamped membrane currents were amplified (and zero current-clamp membrane potentials were measured) with an EPC-7 patch-clamp amplifier (Medical Systems, Greenval, NY) controlled by a PDP 11/23 computer (Digital Equipment Corp., Maynard, MA). The current (filtered at 10 kHz) and voltage outputs of the EPC-7 were digitized at 44.1 kHz using a modified pulse code modulator (501ES; Sony Corp., Tokyo, Japan), and stored on videocassette (SL-HF300; Sony Corp.). Whole-cell membrane currents were also digitized at 250 samples/s (Cheshire A/D). Indec, Sunnyvale, CA) and stored directly on computer disk. Signals were monitored on an oscilloscope (205; Hameg Instruments, Inc., Frankfurt, Germany) and strip-chart recorder (220; Brush, Cleveland, OH). All experiments were carried out at room temperature (20–22°C).

Data Analysis

Current data were played back and low pass filtered at 400 Hz (902LPF eight-pole Bessel filter; Frequency Devices Inc., Haver-
hill, MA), digitized at 1,000 samples/s, and stored on the PDP-11/23. In some cases, currents were filtered at 40 Hz and digitized at 100 samples/s for current binning and averaging analysis. Datafiles were transferred to a Pentium computer (Gateway 2000, North Sioux City, SD) via Kermit (Columbia University, NY) for analysis. Custom software for data acquisition and analysis was written in our laboratory using BASIC-23, AsxBASIC 1.0 (Axon Instruments, Foster City, CA), and Matlab 4.0 (The Mathworks, Natick, MA).

Channel activity (nP) was calculated over periods of 60–500 s as follows. The closed current level (Ic) was taken as the mode of the distribution around closed events. This current was subtracted from the current of a given bin, and the difference was multiplied by the number of events in that bin. The sum of these products yields the open channel area of the histogram. nP is given by dividing the open channel area by the single-channel current, Io. That is,

\[ nP_o = \frac{\sum_{i} (I_{bin} - I_c) \cdot \text{No. events}_{bin}}{I_o}. \]  

For kinetic analysis, currents were filtered at a corner frequency (fC) of 2 kHz and sampled at 5,000 s^{-1}. An event (transitions) was counted each time a data point crossed the 50% level of the unitary channel current. For our recording system, the patch-clamp has a 5-kHz step response, a 5-kHz tape bandwidth, and a 2-kHz eight-pole Bessel filter, yielding an effective bandwidth (~3 dB point, fcorner) of 1.74 kHz. With these settings, the “dead time” of the recording system is given by Colquhoun and Sigworth (1983), T_{dead} = 0.170/ f_{eff}. The “50% delay time” of the Bessel filter is T_{50} = 0.596/ f_{C}. For f_{corner} = 1.74 kHz and f_{C} = 2 kHz, T_{dead} = 102.8 μs and T_{50} = 253 μs. Therefore, events in time histogram bins <500 s were cut off. Since an event lasting 253 μs would be the margin of detection, all dwell lifetimes <253 μs would be missed events.

Open and closed dwell-time kinetics were fit to a probability density function expressed as a sum of exponentials,

\[ f(t) = \sum_{i=1}^{n} A_i e^{-t/\tau_i}, \]  

where n is the number of open or closed states, the A_i are the relative amplitudes, and the \( t \) are time constants. This function was transformed according to \( x = \ln(t) \) and logarithmically binned (Sigworth and Sine, 1987), such that

\[ f(x) = \sum_{i=1}^{n} A_i e^{(\ln(x) - \ln(x_i))/\tau_i}. \]  

This function was used to fit the data using the Levenberg-Marquardt nonlinear least-squares fitting algorithm (Origin 4.0; Microcal Software, Inc., Northampton, MA) to find the appropriate set of \( \tau \)s. Note that if the errors follow a Gaussian distribution, this nonlinear least-squares method is equivalent to the method of maximum likelihood (Colquhoun and Sigworth, 1983).

Dose–response relations for a drug (D) were fitted to the Hill equation as

\[ I_{max} = I_0 + \left( \frac{K}{[D]} \right)^n \tau, \]  

where \( I_{max} \) is the fractional inhibition, \( K \) is the concentration of drug giving 50% inhibition, \( [D] \) is the concentration of the drug, and \( n \) is the Hill coefficient. When \( n = 1 \), this equation reduces to the Langmuir adsorption isotherm.

In the text, the number of observations or experiments is reported, whereas n in the analysis denotes either the whole data set or the subset of total experiments in which precise quantitation could be reliably applied. In some figures, a running average (using a specified window width) of current versus time is displayed. Statistical values for the n elements are given as mean ± SEM. Student’s t-test was applied where appropriate.

**Results**

**Properties of the BLM K⁺ Channel**

**Overview.** With [K⁺] = 95 mM in the patch pipette, we found the K⁺ channel in 551 of 559 seals made on the BLM (98.6%). This K⁺ channel was never detected in seals made on the apical membrane (0 of 16, 0%), consistent with our previous finding that dissociated Ambystoma proximal tubule cells retain epithelial cell polarity (Segal et al., 1996). Seals on the BLM typically contained from 2 to >25 K⁺ channels. Despite numerous attempts to minimize patch area using small-tipped pipettes (resistance ≥ 30 MΩ), patches appearing to have one and only one channel were very infrequent (n = 4, only 0.7%).

**Cell-attached patches.** When cell-attached (c/a) patches were made in NaCl bath, spontaneous inward K⁺ currents were usually observed (95%) at 0 mV (−V_{pip}) command potential. By briefly switching to zero current clamp mode, the resting membrane potential (V_m) of the cell can sometimes be estimated if R_{seal} >> R_{patch}. Using this method under these conditions, V_m averaged −37.2 ± 2.1 mV (n = 16), in good agreement with V_m = −40 mV as measured by conventional impalement (Segal et al., 1996). The V_m of the dissociated cells is ~15–20 mV less than that for a cell in the intact tubule (Sackin and Boulpaep, 1983), suggesting an anion conductance exists at either the apical or basolateral membrane. Indeed, we have characterized a cAMP-activated Cl⁻ channel in the BLM of these cells, which often appears in the same membrane patch as the K⁺ channel (not shown). Alternatively, the isolated cells may have acquired a nonspecific leak pathway that shunts the normally high K diffusion potential. Since the K⁺ and Cl⁻ activity of the pipette solution (aK = 0.80*[95] = 76 mM and aCl = 0.78*[92] = 71.8 mM) is greater than the intracellular K⁺ and Cl⁻ activity (aK = 0.80*[68] = 54.4 mM and aCl = 0.78*[20.5] = 16 mM (Sackin and Boulpaep, 1983), yielding a reversal potential (E_{rev}) of −28.7 mV. Thus, at V_{pip} = 0 mV, the inward current must be carried by K⁺ moving down its electrochemical gradient from the pipette into the cell.

Representative c/a current records at various command potentials are shown in Fig. 2 A, and the current–voltage (I-V) characteristic is shown in Fig. 2 B. The inward slope conductance is 22.2 ± 1.4 pS (taken from −40 to −80 mV, n = 8), and the outward slope conductance (from +20 to +80 mV) is 3.5 ± 0.1 pS (n = 5), and channel activity appears to increase with hyperpolarization (Fig. 2 A). Although the c/a I-V char-
Figure 2. The BLM K⁺ channel appears to be an inward rectifier. (A) Representative current records at various command potentials (−V_{cm}) from a cell-attached basolateral membrane patch containing at least two K⁺ channels. Note the increase in channel activity with hyperpolarization. The patch pipette contains 95 mM K⁺ (solution d) and the bath is NaCl (solution c). The dashed line represents the all channels closed (leak) current at each potential. Each open channel level is denoted by a dotted line. (B) Current–voltage relation for the BLM K⁺ channel for the conditions described in A. The limiting inward slope conductance is 22.2 ± 1.4 pS (n = 8) and the outward chord conductance 3.5 ± 0.1 pS (between 120 and 180 mV, n = 5). Symbols represent mean (●) ± SEM (bars).

A characteristic displays inward rectification, suggesting that the channel is an inward rectifier; under these conditions, the I-V relation would be expected to "inwardly rectify" due to Goldman-Hodgkin-Katz rectification.

Excised patches. When BLM membrane patches were excised in the inside-out (i/o) configuration into the standard bath solutions (solution c or d), channel activity typically began to decline and then disappear. Addition of 0.2 mM ATP to the bath before or just after patch excision prevented rundown and maintained channel activity indefinitely.

Measurements of single-channel current with [K⁺] = 95 mM on both sides of the membrane patch (plus 0.2 mM ATP on the cytosolic side) demonstrate that the BLM K⁺ channel is a true inward rectifier (Fig. 3, A and B). The I-V relation in symmetrical [K⁺] inwardly rectifies and reverses very close to E_K = 0 mV. The channel has an inward slope conductance of γ_{slope,in} = 24.5 ± 0.6 pS (n = 8, measured between −60 and −100 mV), and an inward chord conductance of γ_{chord,in} = 20.5 ± 0.4 pS (n = 8, measured between E_{rev} = 0 and −100 mV). The outward chord conductance measured at +80 mV is γ_{chord, out} = 3.7 ± 0.4 pS (n = 2). The outward slope conductance between +20 and +80 mV is clearly smaller.

Voltage dependence of nP_o. Since BLM membrane patches almost always contained more than one K⁺ channel, nP_o (channel activity) was used to assess the relative voltage dependence of P_o. This assumes that the number of active channel proteins in an excised patch remains constant as voltage varies. (We take n as the maximum number of simultaneously open channels observed, which places a minimum on the actual number of channel proteins in the patch.) As was the case for cell-attached patches, channel activity with KCl on both sides of the excised membrane patch increases with increasing hyperpolarization (Fig. 3 C). Although the absolute values of nP_o differed significantly among patches, relative nP_o increased e-fold per ~83-mV hyperpolarization between −40 and −120 mV, reflecting a 27% increase of nP_o for every 20 mV of hyperpolarization.

Role of magnesium. It has been shown that [Mg^{2+}], mediates at least part of the inward rectification in other inwardly rectifying K⁺ channels (Matsuda et al., 1998).
Figure 3. The BLM K⁺ channel is a true inward rectifier. Representative current records at various command potentials (−Vpp) from an inside-out basolateral membrane patch in symmetrical [K] containing at least three K⁺ channels. The patch pipette and bath each contain 95 mM K⁺ (solution d) with 0.2 mM ATP added to the bath. The dashed line represents the all channels closed (leak) current at each po-
1987; Horie et al., 1987; Ficker et al., 1994) by blocking outward currents in a voltage-dependent manner. When Mg$^{2+}$, was removed from the “cytosolic” side of i/o patches (solution e plus 0.2 mM Na$_2$ATP), we observed flickering, and then rundown of the channel. This is in sharp contrast to the behavior of ROMK1 channels, in which channel rundown is slowed in a Mg$^{2+}$-free bath (McNicholas et al., 1994). Since rundown occurs in the absence of free Mg$^{2+}$, and the presence of 0.2 mM Na$_2$ATP ($n = 4$), it appears that at least the complex of Mg-ATP is required to prevent rundown. Interestingly, it has been shown that both Mg-ATP and free Mg$^{2+}$ are required to sustain channel activity for an ATP-insensitive inward rectifier K$^+$ channel (Fakler et al., 1994). However, since higher levels of ATP block the BLM K$^+$ channel (see below), it is not possible for us to dissociate the role of free Mg$^{2+}$, from that of the Mg-ATP complex. Millimolar concentrations of ATP (or indeed, any nucleotide) block the BLM K$^+$ channel, and this effect is independent of both free Mg$^{2+}$, and Mg-ATP.

In a solution containing 368 nM Mg-ATP and 200 nM free [Mg$^{2+}$], (solution f) (Fabiato and Fabiato, 1979), channel rundown did not occur. Although Fakler et al. (1994) showed that >10 µM free [Mg$^{2+}$], is required to prevent rundown from occurring in the K$_0$2.1 channel, just 200 nM free [Mg$^{2+}$], is sufficient for the BLM K$^+$ channel. Under this condition, the outward unitary conductance of the latter increases, but the inward conductance is essentially unaffected. The enhanced outward current is ATP sensitive, as 5 mM ATP blocked 89.3 ± 6.3% ($n = 3$) of the current. The outward chord conductance with [Mg$^{2+}$] = 200 nM increased from 4.25 ± 0.59 pS to 14.4 ± 1.2 pS ($n = 4$, measured at +80 mV); inward currents were not affected by the change in [Mg$^{2+}$], (Fig. 3 B). Inward rectification was reestablished by returning bath [Mg$^{2+}$], to 1 mM.

Assuming a single-binding site, the Mg$^{2+}$ block may be described using a one-site model according to Woodhull (1973):

$$K(V) = K(0\ \text{mV}) \cdot \exp(-\delta V zF/RT), \quad (5)$$

where $V$ is the command potential, $\delta$ is the valence of 2, and $K(V)$ and $K(0\ \text{mV})$ are the concentrations of Mg$^{2+}$, causing half-maximal block at $V$ and 0 mV, respectively. The factor $\delta$ may be the electrical distance of the binding site from the outside of the channel pore. Alternatively, $\delta$ can be considered together with another term to yield either the equivalent valence ($z$) of, or the effective potential ($V_e$) sensed by, the Mg$^{2+}$ (Hille, 1992).

Current–voltage relations from inside-out patches in symmetrical K$^+$ for several [Mg$^{2+}$], were constructed, from which the current was normalized to that observed in a Mg$^{2+}$-free bath, and the relative current was plotted against command potential for each [Mg$^{2+}$], (not shown). These data were then fitted (nonlinear least squares fitting routine) by the following function that incorporates Eq. 5:

$$\frac{I}{I_{\text{Mg}}^{2+}} = \frac{K(V)}{K(V) + [\text{Mg}^{2+}]} . \quad (6)$$

The fit according to Eq. 6 yields the following results: $\delta = 0.57, K(0\ \text{mV}) = 7.7\ \text{mM}, K(60\ \text{mV}) = 0.72\ \text{mM}$.

Concentration and voltage dependence of $i_c$. To further investigate the biophysical properties of this channel, we asked the question of how varying extracellular [K$^+$] would affect the current carried by the channel. To isolate the change in [K$^+$] from any change in driving force across the patch, the same [KCl] was used in the pipette and bath, thus clamping $E_{\text{rev}}$ to 0 mV. This approach allowed us to measure the change in absolute conductance while maintaining a constant relative permeability (i.e., we varied the Goldman-Hodgkin-Katz current equation while holding the result of the Goldman-Hodgkin-Katz voltage equation constant). Under these conditions, the command potential is the only driving force for net K$^+$ movement across the membrane patch. Pipette and bath [KCl] ranged from 5 to 205 mM while osmolality was kept at 400 mosm/kg using sucrose as necessary. Since $i_{sc}$ for [KCl] = 95 mM was the same in both standard KCl solution (solution d, 200 mosm/kg) or 400 mosm/kg KCl solution, the toxicity change itself does not significantly alter the conducting properties of the channel.

Channel events (inward current) were analyzed at command potentials of −20, −40, −60, −80, and −100 mV for [K$^+$] = 5, 10, 20, 25, 55, 95, 155, and 205 mM, and the I-V relationship was determined for each of these KCl concentrations. Channel events for [K$^+$] = 5 mM could not be adequately resolved and were not included in the further analysis. For each voltage, the...
single channel current increases with [K+] and approaches a limiting current (\(i_{\text{max,slope}}\)). Similarly, the single channel chord conductance (\(\gamma_{\text{chord}}\)) also increases with [K+] and saturates (as shown for −100 mV in Fig. 4 A). Fitting the \(\gamma_{\text{chord}}\) versus [K+] data at −100 mV with the Hill equation:

\[
\gamma = \gamma_{\text{max}} \cdot \left[1 + \left(\frac{[K^+]}{K_d}\right)^n\right]^{-1}
\]

yielded a \(K_d\) of 77 mM, a Hill coefficient of 0.95, and a maximum value for \(\gamma_{\text{chord}}\) of 36.6 pS (Fig. 4 A). Plots of \(\gamma_{\text{chord}}\) versus [K+] at different voltages yielded similar results for \(\gamma_{\text{max}}\). The average value for \(\gamma_{\text{max}}\) determined at −60, −80, and −100 mV was 34.3 pS. \(\gamma_{\text{max,sc}}\) was the same for all command potentials fitted. Indeed, the intersection of the channel’s operating surface with the I-V plane as K shows a linear conductance. This mean value for \(\gamma_{\text{max}}\) was applied to obtain \(K_d\) values and Hill coefficients for all the voltages (Table II).

Comparison of the dissociation constants \(K_d\) for the different command potentials shows that \(K_d\) decreases with hyperpolarization (Fig. 4 B and Table II). The relationship between \(K_d\) (mM) and voltage (mV) can be described by a single exponential decay

\[
K_d(V) = K_{d,\infty} + K_{d,0} \cdot \exp\left(-\frac{V - \psi}{\delta}\right)
\]

where \(K_d\) is the asymptotic value for \(K_d\) as \(V_{\text{pip}} \to \infty\), \(K_{d,0}\) is the difference of \(K_d\) at the reversal potential (0 mV) and \(K_{d,\infty}\). Finally, \(\psi\) is the voltage change required to effect an e-fold change in \((K_d - K_{d,\infty})\).

Cation selectivity. The cation to anion preference of the total BLM conductance was determined by salt dilution experiments using i/o membrane patches. Starting in symmetrical 95 mM K+ and 94.5 mM Cl− (\(E_K = E_{Cl} = 0\) mV) and holding at −Vpip = 0 mV, the bath was changed to a 14 mM K+ and 13.5 mM Cl− solution (solution i, \(E_K = +48.6\) mV, \(E_{Cl} = -49.4\) mV) plus sucrose to maintain isoosmolality. This maneuver resulted in large inward currents (\(n = 3\)), reflecting the cation (K+) moving down its chemical gradient. The reversal potential for this membrane current (including leak) was at least +40 mV (\(n = 3\)). While holding at this \(E_{\text{rev}}\), outward current developed when the 10% KCl bath was replaced with 100% KCl, due to K+ moving along its electrical gradient. Thus, the BLM conductance is cation selective.

Using the Goldman-Hodgkin-Katz voltage equation (Hodgkin and Katz, 1949; Goldman, 1943) for K+ and Cl−, the permeability ratio \(p_{K}/p_{Cl}\) can be estimated (see MATERIALS AND METHODS):

\[
E_{\text{rev}} = \frac{RT}{F} \ln \frac{p_K [K]_o + p_{Cl} [Cl]_i}{p_K [K]_i + p_{Cl} [Cl]_o}
\]

\[
\frac{p_K}{p_{Cl}} = \exp\left(\frac{E_{\text{rev}}}{RT/F}\right) \cdot \frac{[K]_o - [Cl]_i}{[K]_i - \exp\left(\frac{E_{\text{rev}}}{RT/F}\right) \cdot [K]_i}. \tag{9}
\]

\[\text{T A B L E ~ I I}
\]

<table>
<thead>
<tr>
<th>Command potential (mV)</th>
<th>(K_d) (mM)</th>
<th>Hill coefficient</th>
</tr>
</thead>
<tbody>
<tr>
<td>−20</td>
<td>146</td>
<td>0.94</td>
</tr>
<tr>
<td>−40</td>
<td>110</td>
<td>1.02</td>
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<td>92</td>
<td>0.96</td>
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<td>−80</td>
<td>85</td>
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</tr>
<tr>
<td>−100</td>
<td>77</td>
<td>0.95</td>
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</table>
where \( R = 8.315 \text{ J K}^{-1} \text{ mol}^{-1}, F = 9.648 \cdot 10^4 \text{ C mol}^{-1}, T = 293 \text{ K}, E_{\text{rev}} \geq +40 \text{ mV}, [\text{Cl}]_o = 94.5 \text{ mM}, [\text{K}]_o = 95 \text{ mM}, [\text{Cl}]_i = 13.5 \text{ mM}, \) and \([\text{K}]_i = 14 \text{ mM}, \) yielding \( p_\text{K}:\ p_\text{Cl} \) of at least 17.1. Note that even a small change in \( E_{\text{rev}} \) to +44 mV would double the selectivity ratio. It is emphasized that this value represents the minimum cation to anion preference of the K\(^+\) channel since this is the selectivity ratio of the whole basolateral membrane patch including chloride and leak conductances.

**Selectivity among cations.** Two approaches were used to determine the selectivity of this inwardly rectifying K\(^+\) conductance on the BLM. (a) Using c/a and i/o patches, the K\(^+\) in the patch pipette was replaced with the chloride salt of Na\(^+\) \((n = 48),\) Rb\(^+\) \((n = 4),\) Li\(^+\) \((n = 4),\) Cs\(^+\) \((n = 2),\) or NH\(_4\)\(^+\) \((n = 7).\) Each solution was adjusted to pH 7.5 with the respective hydroxide salt. In all cases, c/a and i/o patches failed to show inward channel currents. These results strongly suggest that the BLM K\(^+\) channel is highly selective for K\(^+\) and excludes these cations, since channel activity is seen in >98% of seals made on the BLM when K\(^+\) is in the pipette. (b) Outside-out patches were made to exclude the remote possibility that the c/a and i/o patches used above did not contain any channels. The pipette was filled with KCl, and 0.2 mM Mg-ATP was included to prevent channel rundown. After recording in a KCl bath, the test cation was introduced into the bath (as the chloride salt) and the voltage protocol was repeated. KCl bath exchanges were interposed between test cations.

The cation selectivity as determined from outside-out patches is exemplified in Fig. 5. For Na\(^+\) and Li\(^+\), the current at −40 mV is negligible, but a modest inward current is carried by Rb\(^+\), Cs\(^+\), and NH\(_4\)\(^+\). Exchanging the test cation with KCl returned the ensemble currents to the control level in all cases. That the outward current carried by K\(^+\) remains essentially unchanged during each bath cation substitution indicates that the test cations do not act as channel blockers from the outside. The ensemble current obtained at \( V_{\text{pip}} = -40 \) mV shows that the selectivity of the channel for these cations is K\(^+\) >> Rb\(^+\) ≈ Cs\(^+\) ≈ NH\(_4\)\(^+\) ≈ Na\(^+\) ≈ Li\(^+\).

**Thallium.** Many types of K\(^+\) channels have been shown not only to conduct Tl\(^+\), but often better than they conduct K\(^+\) (Hille, 1992). To assess the Tl\(^+\) conductance of the BLM K\(^+\) channel, the patch pipette was filled with 90 mM Tl-acetate (solution g; the Cl salt of Tl\(^+\) was not used due to its low aqueous permeability) and the bath was filled with K-acetate (solution h). Under these conditions, inward Tl\(^+\) currents were observed in both c/a and i/o patches. The kinetic behavior of these channel events was notably different from those seen when the channel conducts K\(^+\). When conducting Tl\(^+\), the channel openings displayed more bursting, with each opening interrupted by fast flickery closures. The probability that this is actually a different channel is low since (a) the frequency of finding the K\(^+\) channel exceeds 98%, (b) other cation-selective channels were rarely observed, (c) the disparate kinetics

**Figure 5.** The BLM K\(^+\) conductance is highly selective for K\(^+\). The cationic selectivity of the BLM is demonstrated in this outside-out patch with KCl (solution d) + 0.2 mM ATP in the pipette and Cl-salt of the test cation in the bath. The running average of current at −40 mV shows that cationic selectivity is K\(^+\) >> Rb\(^+\) = Cs\(^+\) = NH\(_4\)\(^+\) > Na\(^+\) = Li\(^+\). The dashed line is the zero-current line, and the dotted line is the all channels closed (leak) current at a command potential of −40 mV. The voltage protocol is indicated below the current data.
In conclusion, these findings indicate that the permeability sequence of this K⁺ channel is Tl⁺ > K⁺ >> Cs⁺ = Rb⁺ = NH₄⁺ > Li⁺ = Na⁺ = Cl⁻.

Kinetics. Due to the high density of this K⁺ channel in the BLM, a patch apparently containing only one channel is extremely rare. In over 550 seals, only four membrane patches appeared to contain only one channel (0.7%). Since the open probability (Pₒ) of the channel is only 0.05 ± 0.01 (n = 4), long recordings were required to accumulate enough transitions for meaningful analysis of the long closed state. Kinetic analyses from such patches show that under resting state conditions at −Vpip = −60 mV, the BLM K⁺ channel has two apparent open states and two apparent closed states. Parameters from one c/a patch and one i/o patch show that the open dwell lifetimes are (ms): τₒ₁ = 0.78 (c/a), 1.21 (i/o), and τₒ₂ = 4.7 (c/a), 6.6 (i/o). The closed dwell lifetimes are (ms): τₙ₁ = 1.27 (c/a), 0.72 (i/o), and τₙ₂ = 397 (c/a), 502 (i/o). Fig. 6 shows the open and closed time histograms for a c/a patch.

Inhibitor Profile

The BLM K⁺ channel is inhibited by barium and glibenclamide. The channel is insensitive to tetraethylammonium (up to 10 mM) applied either to the extracellular or cytoplasmic side.

Barium. We have previously shown by recording single-cell membrane potential that the whole cell conductance is dominated by a barium-sensitive K⁺ conductance (Segal et al., 1996). Perforated patch whole-cell recordings show that the barium-sensitive whole cell conductance inwardly rectifies. When 2 mM Ba²⁺ was included in the KCl pipette solution (unpaired experiments), channel openings were rare and a flickery state was noted. In contrast to the voltage dependence of channel activity without Ba²⁺ in the pipette (see Fig. 3 C), steady depolarization now has the effect of increasing nPₒ, and subsequent hyperpolarization reduced activity. This is presumably due to the voltage dependence of the Ba²⁺ block of the channel (data not shown).

To quantify the Ba²⁺ block and use each patch as its own control, outside-out patches were made (n = 3). In outside-out membrane patches at −40 mV, Ba²⁺ inhibits the channel with a Kᵦ = 460 μM (Fig. 7 A).

Glibenclamide. This sulfonamide is known to inhibit ATP-sensitive K⁺ channels in a number of epithelial tissues by binding to the sulfonamide receptor (SUR). SUR1 has recently been cloned (Aguilar-Bryan et al., 1995) and is thought to associate with the Kᵦ channels, thereby conferring sulfonamide sensitivity. However, SUR1 may not be present in the kidney (Inagaki et al., 1995), which may explain in part the much higher dose of glibenclamide required to inhibit renal Kᵦ channels (Hebert and Ho, 1994).
the BLM K⁺ channel is glibenclamide sensitive, albeit at “renal doses.” We treated 22 patches with glibenclamide; 16 excised inside-out patches were exposed to 500 μM, while 6 cell-attached patches were exposed to low (0.01–10 μM) concentrations. In 7 of 16 inside-out patches, 500 μM glibenclamide inhibited activity by 42.3 ± 5.6% (Fig. 7 B). Remarkably, the inhibition was much more potent in the cell-attached patches: 10 μM exerted an 83 ± 2% inhibition in three patches, and 100 nM exerted a 70 ± 3% inhibition in three other patches (data not shown).

ATP Sensitivity

Similar to other Kᵦ ATP channels, low doses of ATP are required to prevent Ambystoma BLM K⁺ channel rundown, whereas millimolar doses inhibit channel activity. At a dose of 5 mM ATP, >90% of channel activity is inhibited. This effect is reversible as nPₒ returns to baseline when the bath [ATP] is returned to 0.2 mM (Fig. 8 A). The dose–response curve for ATP has a Kᵦ = 2.4 mM (Fig. 8 B). The Hill coefficient of ~4 may suggest that the channel has a tetrameric structure, with each subunit possessing an ATP binding site. Thus, in the Ambystoma proximal tubule, the BLM K⁺ channel that appears to be the major K⁺ conductance of the cell is ATP sensitive.

Among the nucleoside diphosphates, ADP is less potent than ATP (ADP inhibits by 65.6 ± 8.1%, n = 4, P < 0.01), but more potent than CDP, GDP, IDP, TDP, or UDP (data not shown). This suggests that the putative nucleotide binding site(s) recognize NDPS as well as NTPs, and that nucleotide hydrolysis is probably not occurring at this site. Indeed, even nucleoside monophosphates have a moderate inhibitory effect, although nucleosides themselves are without effect. The relative potency of the adenosine nucleosides (at 5 mM) in inhibiting the BLM K⁺ ATP channel is ATP (93.3 ± 1.9%) > ADP (65.6 ± 8.1%) > AMP (38.7 ± 3.7%) > adenosine (1 ± 2%) (Fig. 8 C, n = 3–9).

Other nucleotides. The effect of nucleotides was tested in excised i/o patches. All the NTPs tested reversibly inhibited BLM K⁺ channel activity at a dose of 5 mM (ATP 93.3 ± 1.9%, n = 9; CTP 70.3 ± 10.0%, n = 5; GTP 62.3 ± 7.5%, n = 3; ITP 61.1 ± 0.8%, n = 9; TTP 53.7 ± 6.7%, n = 2; UTP 71.7 ± 5.7%, n = 2) (Fig. 8 C). These results suggest that each compound probably interacts with common cytoplasmic nucleotide binding site(s). Note that ATP is significantly more potent than the other nucleoside triphosphates (P < 0.02), but there is no significant difference among the nonadenosine nucleotides.

Rundown of the BLM K⁺ Channel

One characteristic of Kᵦ ATP channels is “rundown,” a gradual loss of activity when the membrane patch is deprived of cytosolic ATP (Findlay and Dunne, 1986). Typically, both Mg²⁺ and ATP are required to prevent rundown in Kᵦ ATP channels (Ashcroft and Ashcroft, 1990). Likewise, the BLM K⁺ channel runs down in the absence of either Mg²⁺ or ATP (or both). Lower concentrations of ATP (100–200 μM) will prevent or “rescue” channel rundown. The experiment shown in Fig. 9 A summarizes the characteristics of BLM K⁺ channel rundown. Channel activity typically begins to decrease
Figure 8. Nucleotides reversibly inhibit the BLM K⁺ channel. (A) ATP, reversibly inhibits the BLM K⁺ channel in inside-out patches. In the experiment depicted, up to 13 channel open levels are seen under control conditions (top) at −60 mV with KCl (solution d) in the pipette and NaCl (solution c) plus 0.2 mM ATP in the bath. (middle) Addition of 5 mM ATP to the cytoplasmic side almost completely blocks channel activity (98% decrease) with only rare openings to one open level. (bottom) The inhibition is readily reversed upon returning to 0.2 mM ATP. (B) Dose–response curve for inhibition by ATP. The inhibitory effect of ATP was determined in inside-out patches under the conditions described in A. Relative nPo (nPo/nPo,control) is plotted versus [ATP]. The data (●) was fitted with the Hill equation (solid line) yielding a Ki = 2.4 mM and nH = 3.95. Adenosine nucleotides and nucleoside triphosphates (each at 5 mM) inhibit the BLM K⁺ channel. The inhibitory effect of ATP, ADP, AMP, and adenosine was determined under the conditions described in A. Values plotted are average (nPo,test/ nPo,control) SEM (n = 3–9). The rank order of inhibition is ATP (93.3%) > ADP (65.6%) > AMP (38.7%). Adenosine has no inhibitory effect on nPo of the BLM K⁺ channel. Although all the nucleoside triphosphates inhibit the channel, ATP exerts a significantly stronger block than the other NTPs tested (P < 0.02). There is no significant difference among the other NTPs.

(rundown) upon excision of the membrane patch into a nucleotide-free bath. If this process is allowed to continue, channel activity will cease, usually irreversibly. When 0.2 mM of ATP is added back, channel activity can be restored. When ATP is removed, all channels rapidly close. In the continued presence of ATP-γS, re-addition of ATP is again able to rescue rundown, and activity returns to baseline upon washout of the ATP-γS. Frequently (but not invariably), ATP-γS has an inhibitory effect on single channel activity when added in the presence of ATP, which is reversible as long as the exposure is not prolonged (n = 6), as shown in Fig. 9 B. When ATP-γS is added in the absence of ATP, channel activity runs down very quickly, usually irreversibly (n = 4, data not shown).

Thus, ATP-γS cannot substitute for ATP in sustaining channel activity, suggesting that phosphorylation itself is not sufficient to prevent rundown. Although phosphorylation may be necessary, it appears that the nucleoside triphosphate must be hydrolyzable to maintain channel activity. This hypothesis is supported by the finding that CTP, GTP, ITP, TTP, and UTP could all prevent or rescue rundown, but the corresponding NDPs could not. Rescue does not appear to require the cAMP-dependent protein kinase, since channel activity can be restored even in the presence of a high concentration of protein kinase inhibitor (PKI, 1 μg/ml, P0300; Sigma Chemical Co.).

Since it has been reported that removal of free Mg²⁺ nearly abolishes rundown of K ATP in cultured CRI-G1
ATP but not ATP-γS prevents and rescues channel rundown. Hydrolyzable nucleoside triphosphates prevent and rescue BLM K⁺ channel rundown. A running average (current versus time, window width 768 ms) of a representative experiment is shown. The pipette is KCl (solution d), the bath is NaCl (solution c), the command potential is −60 mV. Upon excision in a nucleotide-free bath, channel activity decreases (channel rundown). After the addition of 0.2 mM ATP to the bath, channel activity slowly recovers. The addition of 0.2 mM ATP-γS (a poorly hydrolyzable ATP analogue) in the continued presence of 0.2 mM ATP has no effect. However, when ATP is removed, ATP-γS is not able to support channel activity, which rapidly declines and runs down. Readdition of ATP leads to full recovery of channel activity. Single-channel traces showing that ATP-γS has an inhibitory effect on K⁺ ATP channel activity in excised inside-out BLM patches. When compared with control conditions (top), the addition of ATP-γS (middle) reduces nPₒ. This inhibition is reversible as long as the exposure to ATP-γS is not prolonged (bottom). (C) Removal of Mg²⁺ does not prevent channel rundown in an ATP-free bath. The top panel shows that rundown of the BLM K⁺ ATP channel upon excision into an ATP-free bath proceeds despite removal of bath Mg²⁺. Representative traces from the regions marked by α, β, and γ are shown at bottom.

Insulin-secreting cells (Kozlowski and Ashford, 1990) and partially inhibits rundown of ATP-regulated ROMK1 channels excised in an ATP-free bath (McNicholas et al., 1994), we assessed BLM K⁺ ATP channel activity under these conditions. The representative experiment shown in Fig. 9 C shows that rundown of the BLM K⁺ ATP channel still occurs in the absence of ATP despite excision of the patch into a Mg²⁺-free bath.

Diazoxide. The synthetic K⁺ ATP channel opener diazoxide was applied to the cytoplasmic side of i/o patches. It has been shown that this benzothiadiazine can open K⁺ ATP channels in the presence of Mg-ATP, but it may have an inhibitory effect in the absence of Mg-ATP (Kozlowski et al., 1989). Initial excision of the patch into an ATP-free bath leads to channel rundown as discussed above, and 200 μM diazoxide alone does not rescue rundown. However, addition of 0.2 mM Mg-ATP in the continued presence of, or after exposure to, diazoxide increases channel activity well in excess of that before rundown (n = 3, Fig. 10). The inhibitory effect of 5 mM ATP is not diminished in the presence of, or by previous exposure to, diazoxide (n = 5, data not shown).

Since low levels of Mg-ATP are required for diazoxide to open the BLM K⁺ channel, diazoxide probably does not interact with the rundown site. As hypothesized by others (Edwards and Weston, 1993, 1995), diazoxide may be acting at or near the nucleotide binding
site that mediates inhibition. On the other hand, the interaction at this second site is more complex than a simple competition between the K channel opener and the nucleotide, since diazoxide does not relieve the inhibition by millimolar levels of ATP.

**Discussion**

The proximal tubule (a leaky epithelium) can be considered to function within the general scheme of the epithelial transport model first proposed by Koefoed-Johnsen and Ussing (1958) (KJU), in which the apical membrane is primarily Na⁺ selective and the BLM is primarily K⁺ selective. Although this model was first applied to tight epithelia such as frog skin (Koefoed-Johnsen and Ussing, 1958) and urinary bladder (Davis and Finn, 1982), its essence holds for leaky epithelia such as small intestine (Gunter-Smith et al., 1982) and the proximal tubule (Matsumura et al., 1984). In the KJU model, maintenance of unidirectional Na transport requires that K moves in a closed circuit (recycling) across the BLM. That is, barring intracellular accumulation of K (or K⁺ secretion), the K⁺ pumped into the cell by the pump must be matched by an outward K⁺ current across the BLM.

Steady state vectorial transport in the proximal tubule thus requires continuous activity of the basolateral Na⁺,K⁺-ATPase pump, which consumes ATP and obligates intracellular accumulation of K⁺. A conductance for K⁺ is necessary both to allow this K⁺ to recycle and to maintain V_{bl}. Experiments performed by Matsumura et al. (1984) on perfused *Necturus* proximal tubules first demonstrated that the BLM G_K varies as a function of pump activity, and they suggested that the regulation of BLM G_K was linked to cellular metabolism, as had been previously proposed for red cells (Romero, 1978) and suspensions of rabbit cortical tubules (Balaban et al., 1980).

This hypothesis was bolstered when the first single-channel records of an ATP-sensitive K⁺ channel (K\_ATP) from cardiac muscle were published (Noma, 1983). Similar ATP-sensitive K⁺ (K\_ATP) channels were

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**Figure 10.** The K channel opener diazoxide activates the BLM K⁺ channel. Diazoxide activates the BLM K⁺ channel in the presence of ATP. In the experiment depicted, the pipette contains KCl (solution c), the bath contains NaCl (solution c), and the command potential is −80 mV. (top) A running average (window width 16 ms) of current versus time. (bottom) Sample current traces from the same experiment. With diazoxide alone, the channel opens infrequently (nP_o = 0.15) and no more than two channels are open at a time. Diazoxide combined with 0.2 mM ATP, however, promptly increases channel activity and within 5 min, up to 13 simultaneously open channels are evident (nP_o = 5.27). The all channels closed level (dashed line) and open channel levels (dotted lines) are indicated.
subsequently found in pancreatic β-islet cells (Cook and Hales, 1984), skeletal muscle (Spruce et al., 1985), and smooth muscle (Staden et al., 1989). The $K_i$ for ATP is 10–100 μM for all these Type I $K_{ATP}$ channels (Ashcroft and Ashcroft, 1990), and they are inhibited by sulfonylurea agents (Edwards and Weston, 1993). A $K_{ATP}$ channel with these Type I properties has not been identified in epithelial tissues. However, a related $K_{ATP}$ channel was recently demonstrated in nonperfused (Tsuchiya et al., 1992) and perfused (Hurst et al., 1993; Beck et al., 1993) rabbit proximal tubules. In preliminary experiments, we have also found this channel in the BLM of nonperfused rabbit proximal tubule, and in the present study we provide the first detailed description of the analogous BLM $K_{ATP}$ channel in an amphibian using a novel preparation of dissociated proximal tubule cells that retain epithelial polarity (see Fig. 1 and Segal et al., 1996).

Investigators have used several techniques to patch the BLM, including mechanically stripping off the basement membrane (Sackin and Palmer, 1987; Kawahara et al., 1987) and tearing tubules to access the lateral membrane (Gögelein and Greger, 1987). However, enzymatic treatment of nonperfused tubules (Tsuchiya et al., 1992; Parent et al., 1988), perfused tubules (Hurst et al., 1993; Beck et al., 1993), and single cells (Hunter, 1991) has been employed most commonly.

We have previously shown that the dissociated *Ambystoma* proximal tubule cells (see Fig. 1) retain epithelial cell polarity (Segal et al., 1996). Indeed, the $K^+$ channel described in the present study was present in >98% of seals made on the BLM, but was never detected in the apical membrane. With high $[K]$ on both sides of the membrane patch, this BLM $K^+$ channel exhibits inward rectification with $\gamma_{\text{slope, in}} = 24.5 \pm 0.6$ pS and $\gamma_{\text{chord, out}} = 3.7 \pm 0.4$ pS. This non-ohmic behavior in both cell-attached and -free membrane patches is due at least in part to a voltage-dependent open channel block by cytosolic $Mg^{2+}$ (Matsuda et al., 1987). Removal of $Mg^{2+}$ from the cytosolic side of i/o patches leads to an increase in outward conductance without affecting inward conductance. In c/a patches made on frog BLM (NaCl pipette in a KCl bath), Hunter (1991) estimated that the blocking site was 51.3% across the membrane from the cytosolic side (i.e., 0.513). This compares well to the value of 0.57 obtained for the *Ambystoma* BLM $K^+$ channel in the present study.

Inward rectification of current is not necessarily due to a decrease in the intrinsic outward conductance of the channel pore. Indeed, Matsuda et al. (1987) first described how ohmic single channel units could behave as rectifiers. They showed that inward rectification of a cardiac myocyte $K^+$ channel was mediated by two effects: (a) $Mg^{2+}$-independent voltage gating in which ohmic channels are open at hyperpolarized voltages and essentially inactivate upon depolarization, and (b) a $Mg^{2+}$-induced voltage-dependent open channel (fast) block. Our data for the BLM $K^+$ channel are in keeping with these observations since (a) the conductance is ohmic in the absence of cytosolic $Mg^{2+}$, and (b) $Mg^{2+}$ appears to cause a flickery open channel block of the outward current in the steady state (e.g., see Fig. 3 A). Naturally occurring polyamines (e.g., spermine) have also been shown to mediate inward rectification of some $K^+$ channels (Ficker et al., 1994). However, spermine (up to 5 mM) does not affect the *Ambystoma* BLM $K^+$ channel.

*Properties of $K^+$ conduction through the pore.* Constant-field theory predicts that net $K^+$ flux across the channel should increase linearly with rising $[K^+]$. Most channels do not exhibit this behavior due to “competition” for binding sites within the pore, and saturation occurs at higher concentrations of the permeant ion (Hille, 1992). In our study, conductance versus $[K^+]$ was fit with the Hill equation (Table II). Hill coefficients of $\sim$1 suggest that one $K^+$ ion interacts with the channel at a time, allowing permeation to be modeled as follows:

$$
\begin{align*}
K^+_{\text{outside}} + \text{channel} & \rightarrow [K^+\text{bound to channel}] \\
& \leftarrow k_{-1} \\
& \rightarrow K^+_{\text{inside}} + \text{channel} \\
& \leftarrow k_{-2}
\end{align*}
$$

At high $[K^+]$ concentrations, saturation occurs when the rate of ion entry approaches the maximum rate of unbinding. For the BLM $K^+$ channel, the maximum inward conductance ($\gamma_{\text{max}} = 34.3$ pS) is voltage-independent, suggesting that at high concentrations of $K^+$, the electrical driving force is the rate-limiting factor for ion flux. However, the binding-unbinding rate is voltage dependent, suggesting that the loading of the binding site for $K^+$ is dependent on the membrane potential (Hille, 1992). Kawahara et al. (1987) also noted saturation of the *Necturus* BLM $K^+$ channel conductance with external $[K^+]$ in c/a patches.

*Voltage dependence.* Hyperpolarization increases $nP_e$. Such voltage sensitivity renders the channel susceptible to changes in membrane voltage, a potential regulator of the BLM $K^+$ conductance. On the BLM in the *Necturus* proximal tubule, hyperpolarization increased $K^+$ channel activity (Sackin and Palmer, 1987; Kawahara et al., 1987). In frog kidney, one study showed that $nP_e$ increased with hyperpolarization (Kawahara, 1990), whereas
in another study, no voltage dependence of $nP_o$ was observed (Hunter, 1991). In preliminary experiments using rabbit proximal tubule, we again found that hyperpolarization increases $nP_o$, although Parent et al. (1988) reported the opposite. Note that for the cloned renal K+ channels, ROMK1 (Ho et al., 1993) and ROMK2 (Chepilko et al., 1995), which have not been detected in proximal tubule, $nP_o$ decreases with hyperpolarization.

Selectivity and blockade. The channel is highly selective for K+ over a number of cations (Na+, Rb+, Li+, NH4+, and Cs+) as well as over Cl−. When K+ in the patch pipette was replaced by these cations, $c/a$ and $i/o$ patches failed to show inward channel currents. Paired experiments in outside-out patches show that $g_{Ko}/g_{Na}$ exceeds 30:1, and that the cation selectivity of the BLM K+ channel is $K^+ >> Rb^+ > Cs^+ > NH_4^+ > Na^+ > Li^+$ (see Fig. 5). Detailed selectivity data are lacking in other studies on BLM K+ channels in the proximal tubule, but more distal K+ channels, including ROMK, conduct Rb+. With Rb+ in the patch pipette, $g_{Rb}/g_K = 0.36$ for ROMK2 (Chepilko et al., 1995). Under similar conditions, we did not observe inward Rb+ currents through the BLM K+ channel.

Like other types of K+ channels (Hille, 1992), the BLM K+ channel can conduct thallium. Indeed, the single channel conductance is higher for Ti+ than it is for $K^+$ ($g_{Ti}/g_{K} = 1.2$). Since the reversal potential under bimonic conditions (Ti+ pipette, K+ bath) is slightly positive, Ti+ is also more permeant than K+. This differs from ROMK2, in which Ti+ is also more permeant than K+ but the conductance for Ti+ is less than that for $K^+$ ($g_{Ti}/g_{K} = 0.7$) (Chepilko et al., 1995). Thus, for the BLM K+ channel, Ti+ has a higher conductance and is more permeant than $K^+$.

Similar to other renal K+ channels, this BLM K+ channel is insensitive to tetraethylammonium. However, it is well known that the proximal tubule K+ conductance can be inhibited by millimolar concentrations of Ba2+ (Sackin and Boulaap, 1981) and the whole-cell conductance of the dissociated Ambystoma cells is barium sensitive (Segal et al., 1996). Outside-out patches from the BLM show that K+ channel activity is inhibited by barium with a $K_i$ of 460 μM at a command potential of −40 mV (see Fig. 7). Complete inhibition occurs at [Ba2+]o = 10 mM.

Sensitivity to Nucleotides, Sulfonylureas, and Activation by Diazoxide

Nucleotides. The Ambystoma BLM K+ channel is sensitive to ATP, albeit at millimolar levels. Since the first demonstration of an ATP-sensitive K (KATP) channel in cardiac muscle (Noma, 1983) and pancreatic cells (Cook and Hales, 1984), potassium channels inhibited by nucleotides have been described in a wide variety of tissues (Ashcroft and Ashcroft, 1990). In pancreatic β cells, KATP channels function in the regulation of insulin secretion. Ashcroft and Ashcroft (1990) have classified such inwardly rectifying KATP channels as Type I based on their exquisite sensitivity to ATP ($K_i \sim 10–100$ μM) and sulfonylureas.

There is no doubt that KATP channels function primarily to couple cell metabolism to membrane potential (Ashcroft and Ashcroft, 1990), and perhaps epithelial transport (Mauzer et al., 1998). Curiously, epithelial KATP channels appear to lack the exquisite sensitivity of Type I KATP channels, possibly for reasons discussed below. The proximal tubule BLM K+ channel described in this study is reversibly inhibited by ATP with a $K_i$ of 2.4 mM at pH 7.5. This is a reasonable operating point given that intracellular ATP levels in proximal tubule are 4–5 mM, and ATP levels have been shown to fall to ~2 mM when transport is stimulated (Beck et al., 1991; Tsuchiya et al., 1992). The role of intracellular ATP in coupling channel activity to pump turnover is discussed in the companion paper.

It is likely that the intracellular ATP-binding site contains a motif that recognizes a spectrum of nucleotides since the channel is also reversibly inhibited by GTP, ITP, UTP, CTP, and TTP (see Fig. 8 C). This putative binding site recognizes NTPs, NDPs, and NMPs, but not the nucleosides themselves. Adenosine nucleotides exert significantly stronger inhibition than their nonadenosine counterparts, and the triphosphate of a given nucleoside is a better inhibitor than the diphosphate and the monophosphate. Since the binding of NDPs and even NMPs is effective, channel inhibition probably does not require hydrolysis. In contrast to the KATP channel on the apical membrane of principal cells (Wang and Giebisch, 1991), the inhibitory effect of the nucleotides is probably not critically dependent on Mg2+ since nucleotide sodium salts (free Mg2+ ≈ 25 μM) are as effective as the magnesium salts (free Mg2+ ≈ 1.3 mM).

Sulfonylureas. In cell-free patches, the Ambystoma and rabbit BLM K+ channels are (variably) inhibited by submillimolar levels of glibenclamide, whereas native cell Type I KATP channels are sensitive to nanomolar concentrations of glibenclamide (Ashcroft and Ashcroft, 1990). Indeed, no renal and perhaps no epithelial K+ channel has the sensitivity to sulfonylureas (or ATP) that Type I KATP channels possess. This may suggest that either Type I KATP channels are only found in specific tissues, or that associated molecules (e.g., the sulfonylurea receptor) conferring sulfonylurea sensitivity are cell specific. In this regard, it is of interest that all but one of the cloned KATP Channels fail to exhibit the degree of sulfonylurea sensitivity seen in the native β cell KATP channels. The exception is the KATP channel composed of Kir6.2 and the sulfonylurea receptor itself (Inagaki et al., 1995).
In the kidney, cloned ROMK channels are ATP regulated in the sense that they rundown in the absence of ATP, but also have a much lower (if any) sensitivity to ATP and sulfonylureas (Ho et al., 1993; Zhou et al., 1994). At least on Northern analysis, the kidney also appears to lack SUR1 (Inagaki et al., 1995), which may partially account for the decrease in glibenclamide sensitivity. Of note, the renal K+ channel ROMK2 appears to be sensitive to glibenclamide when coexpressed with the cystic fibrosis transmembrane conductance regulator, but becomes much less sensitive after PKA phosphorylation of the patch (McNicholas et al., 1996).

Since phosphorylation by PKA activates the BLM KATP channel (Mauerer et al., 1998), it may be that high current patches are already highly phosphorylated, and thus relatively insensitive to glibenclamide. That glibenclamide inhibition of the BLM KATP channel appears significantly more potent in cell-attached patches with smaller currents is consistent with this idea.

**Diazoxide.** Diazoxide is a benzothiadiazine K channel opener (KCO) that activates several types of KATP channels. In many tissues, it is thought that activation by KCOs requires Mg-ATP (Edwards and Weston, 1993; Ashcroft and Ashcroft, 1990), as illustrated for the BLM K+ channel in Fig. 10. Diazoxide also does not alter the requirement for Mg-ATP to prevent BLM K+ channel rundown (see below). Indeed, diazoxide can be inhibitory in the absence of Mg-ATP, perhaps by accelerating channel rundown (Kozlowski et al., 1989). Therefore, it is not likely that diazoxide exerts its effect on the rundown site of the BLM K+ channel.

For both the frog muscle KATP channel and the *Ambystoma* BLM K+ channel, KCOs lose their effectiveness at high (5–10 mM) intracellular [ATP]. This reciprocal effect on channel activity suggests a competitive interaction between KCOs and nucleotides. The activation by KCOs in the frog skeletal KATP channel is not Mg2+-dependent and, like the BLM K+ channel, does not occur in the absence of ATP. Further studies may thus show that diazoxide opens the BLM K+ channel by a mechanism similar to that recently proposed for KCO activation of the skeletal muscle KATP channel (Fourestier et al., 1996).

**Rundown of the BLM K+ Channel**

A hallmark of KATP channels is that they exhibit the phenomenon of rundown, a gradual loss of activity when the membrane patch is deprived of cytosolic ATP. As first noted by Findlay and Dunne (1986), rundown is “a paradoxical situation in that K+ channels that are inhibited by intracellular ATP require intracellular ATP to retain the ability to open.” In a recent study on frog proximal tubule cells (Robson and Hunter, 1997), washout of intracellular ATP reduced the whole-cell barium-sensitive conductance by ~60% over 10 min. Inclusion of 2 mM ATP in the patch pipette not only prevented this rundown, but the barium-sensitive conductance increased by 54% over 10 min.

Accordingly, the BLM K+ channel runs down in the absence of ATP, and low concentrations of ATP (100–200 μM) are required to prevent or rescue rundown in cell-free patches. Once rundown has begun, application of ATP will restore channel activity, but the ATP-restorable current decreases as the time between rundown and ATP application lengthens. The BLM K+ channel specifically requires Mg-ATP to support activity as rundown occurs if either ATP, Mg2+, or both are removed (see Table III). Typically, both Mg2+ and ATP are required to prevent rundown in KATP channels (Ashcroft and Ashcroft, 1990). Mg2+ is required as a cofactor, as channel rundown occurs in a Mg2+-free solution despite the presence of Na-ATP. Free Mg2+ concentrations as low as 200 nM are sufficient to prevent BLM K+ channel rundown in the presence of ATP. The possibility that free Mg2+ itself (rather than the Mg-ATP complex) plays a distinct role in the rundown of the BLM K+ channel cannot be excluded, but is difficult to test since rundown occurs if either Mg2+ or ATP is removed. Interestingly, free Mg2+ itself (≥10 μM) can prevent rundown of KATP2.1, an ATP-insensitive inward rectifier (Fakler et al., 1994).

Paradoxically, removal of free Mg2+ nearly abolishes rundown of KATP in cultured CRI-G1 insulin-secreting cells (Kozlowski and Ashford, 1990) and partially inhibits rundown of ATP-regulated ROMK1 channels excised in an ATP-free bath (McNicholas et al., 1994). The latter effect is presumed to occur via the inhibition of a Mg2+-dependent phosphatase (McNicholas et al., 1994). Thus, removal of Mg2+ inhibits rundown in ROMK1, but actually produces rundown in the BLM K+ channel (see Fig. 9C). We conclude that the Mg2+-dependent phosphatase thought to be involved in the rundown of ROMK1 (McNicholas et al., 1994) does not mediate rundown of the BLM K+ channel.

ATP can serve as a substrate in hydrolysis reactions

<table>
<thead>
<tr>
<th>Table III</th>
<th>Rundown for ROMK1 Compared with the BLM K+ Channel</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nucleotide</td>
<td>Free Mg2+</td>
</tr>
<tr>
<td>0.2 mM ATP</td>
<td>1 mM Mg2+</td>
</tr>
<tr>
<td>0.2 mM ATP-γS</td>
<td>1 mM Mg2+</td>
</tr>
<tr>
<td>ATP-free</td>
<td>1 mM Mg2+</td>
</tr>
<tr>
<td>0.2 mM ATP</td>
<td>Mg2+-free</td>
</tr>
<tr>
<td>ATP-free</td>
<td>Mg2+-free</td>
</tr>
</tbody>
</table>

*As expressed in *Xenopus* oocytes; data from McNicholas et al. (1994).*
mediated by an ATPase and in phosphorylation reactions mediated by a kinase. We asked whether phosphorylation itself (without hydrolysis) can prevent BLM K\(^+\) channel rundown, as has been proposed for other channels, including ROMK1 (McNicholas et al., 1994). To address this issue, we used ATP-γS, a poorly hydrolyzable ATP analogue that is an effective substrate for most kinases (e.g., PKA) in (thio)phosphorylation reactions (Eckstein, 1985). Similar to results reported for the cell K\(_{ATP}\) channel (Ohno-Shosaku et al., 1987), our findings (see Fig. 9 and Table III) indicate that ATP-γS cannot substitute for ATP in sustaining channel activity, whereas all hydrolyzable nucleoside triphosphates we tested do prevent or rescue rundown. The corresponding nucleoside diphosphates were also ineffective.

In summary, rundown of the BLM K\(^+\) channel in excised patches seems to be prevented by a high affinity nucleotide binding site, which hydrolyzes nucleoside triphosphates in the presence of Mg\(^{2+}\). Rundown is prevented by ATP even in the presence of high concentrations of a protein kinase inhibitor, strongly suggesting that typical protein phosphorylation processes alone are insufficient to prevent/rescue channel rundown. Fakler et al. (1994) obtained similar results in K\(_{2.1}\) channels.

**Comparison with Other Studies**

Gögelein and Greger (1987b) found a very weak inward rectifier K\(^+\) channel on the lateral membrane of rabbit proximal straight tubule. In symmetrical KCl with 1 mM Mg\(^{2+}\) in the bath, this channel had a limiting g\(_{\text{in}}\) of \(~\sim 45\) pS and a limiting g\(_{\text{out}}\) of \(~\sim 40\) pS. In contrast to the results of the present study, their channel’s open probability did not increase with hyperpolarization, and n\(_P\) was unaffected by Ca\(^{2+}\) (<1 nM–1 μM) applied to inside-out patches (Mauerer et al., 1998). They did not observe channel rundown in excised patches, the channel was tetraethylammonium sensitive, and the effect of ATP was not tested. Thus, it is not likely that their channel was the BLM K\(^+\) channel we find in *Ambystoma* proximal tubule.

Kawahara et al. (1987) found a 31-pS K\(^+\) channel in 42% of cell-attached patches made on the BLM of *Necturus* proximal tubule. They also found saturation of inward channel conductance at \(~\sim 50\) pS (compared with \(~\sim 35\) pS for *Ambystoma*) and an apparent K\(_m\) of 65.5 mM (compared with 77 mM in *Ambystoma*). Whereas we observed inward rectification for all [K], their I–V plots were restricted to inward currents. Similar to our findings in *Ambystoma*, the K\(^+\) channel in *Necturus* showed high selectivity (P\(_{K}/P_{Na}\) = 10:1 in *Necturus* versus \(~\sim 30:1\) in *Ambystoma*), channel activity increased with hyperpolarization, and external Na\(^+\) did not influence channel behavior. They also found that hyperpolarization did not affect channel open time; rather, the increase in n\(_P\) was due to shortening a closed state lifetime. Significantly, Kawahara et al. (1987) were unable to maintain channel activity in excised patches, stating that “patches became unstable and noisy after excision.” It is likely that this was channel rundown since this description is reminiscent of the channel “choking” and rundown we observe in an ATP-free bath.

Similarly, Sackin and Palmer (1987) observed two BLM K\(^+\) channels in *Necturus*, a short open-time channel that they studied in detail, and a long open-time channel that disappeared after excision, perhaps because of rundown. The short open-time channel is different from the BLM K\(^+\) channel of *Ambystoma* since it was nonrectifying and did not rundown when excited in an ATP-free bath. The long open-time channel was thought to be the same as that studied in cell-attached patches by Kawahara et al. (1987), and probably corresponds to the BLM K\(^+\) channel in *Ambystoma* except the channel in *Necturus* had a mean open time of \(~\sim 60\) ms, much longer than that seen in our study.

Parent et al. (1988) made cell-attached patches on rabbit BLM and found an inward-rectifier K\(^+\) channel that had different properties than both the *Ambystoma* BLM K\(^+\) channels in the present study. Their voltage dependence was opposite to ours, as they concluded that hyperpolarization induced a long closed state.

Hunter (1991) made cell-attached patches on the BLM of single frog proximal tubule cells and found only one kind of K\(^+\) channel, an inward rectifier with an inward g\(_{\text{dope}}\) = 32.4 pS, and an outward g\(_{\text{chord}}\) = 6.2 pS at +80 mV, similar to the one we studied on the BLM of *Ambystoma* proximal tubule. Depolarization reduces the macroscopic BLM K\(^+\) conductance and single-channel conductance in both preparations. A Boltzmann fit of the g–V curves (redrawn using Fig. 5 of Hunter, 1991, and the inset of Fig. 3 B in the present study) shows a V\(_{1/2}\) = +31.7 mV and a width of 14.5 mV for the BLM K\(^+\) channel in c/a patches of frog, compared with a V\(_{1/2}\) = −14.9 mV and a width of 21.3 mV for the BLM K\(^+\) channel in i/o patches of *Ambystoma*. Thus, the sensitivity of conductance versus voltage is such that there is an e-fold change in g per 14.5 mV in frog (c/a), and per 21.3 mV in *Ambystoma* (i/o). The discrepancy in V\(_{1/2}\) may be due to different experimental conditions and levels of cytosolic Mg\(^{2+}\). Hunter (1991) also reported that the addition of glucose and/or alanine did not affect BLM K\(^+\) channel activity in the frog cells. Although addition of alanine depolarized V\(_{m}\) > 30 mV, he was unable to explain the repolarization of the BLM under his experimental conditions. In contrast, the *Ambystoma* BLM K\(^+\) channel under discussion is activated by these substrates, which can explain the repolarization of the BLM (Mauerer et al., 1998). In a
Table IV

<table>
<thead>
<tr>
<th>Species</th>
<th>g</th>
<th>NP</th>
<th>f</th>
<th>A</th>
<th>G_K</th>
<th>G_KLM</th>
<th>G_K/G_KLM</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>µS</td>
<td>µS/cm²</td>
<td>µS/cm²</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Necturus</td>
<td>31.2*</td>
<td>0.14*</td>
<td>1.0*</td>
<td>33.6*</td>
<td>33–59</td>
<td>0.57–1.00</td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.42*</td>
<td></td>
<td></td>
<td>14.1</td>
<td></td>
<td>0.24–0.43</td>
<td></td>
</tr>
<tr>
<td>Rana</td>
<td>5.6</td>
<td>0.22</td>
<td>0.62</td>
<td>48</td>
<td>33–59</td>
<td>0.81–1.45</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>51.3 ¶</td>
<td></td>
<td>0.94</td>
<td></td>
</tr>
<tr>
<td>Rabbit</td>
<td>10**</td>
<td>0.06**</td>
<td>1.0</td>
<td>1.0**</td>
<td>100**</td>
<td>714 §§</td>
<td>0.14</td>
</tr>
<tr>
<td>Ambystoma</td>
<td>4</td>
<td>0.20</td>
<td>0.986</td>
<td>1.77</td>
<td>44.6</td>
<td>51.5 ¶</td>
<td>0.87</td>
</tr>
</tbody>
</table>

*From Kawahara et al., 1987; cell-attached at 0 mV; ‡assumed in the calculation in Kawahara et al. (1987), despite finding f = 0.42; ‡from Maunsbach and Boulpaep (1984), for Necturus; ‡from Hunter (1991), cell-attached at 0 mV; ‡from Maunsbach and Boulpaep (1984), for Ambystoma; ‡‡presumed, since data not given in Parent et al. (1988); §§from Lapointe et al. (1984); ¶from Maunsbach and Boulpaep (1984), for Ambystoma; **from Parent et al. (1988), cell-attached at −50 mV; ††present study.

Recent follow-up study, Robson and Hunter (1997) proposed that, based on differential sensitivity to barium and quinidine, the frog cells have two separate K⁺ conductances. The inwardly rectifying conductance that was inhibited by both agents appears similar to the Ambystoma BLM K_ATP channel, although direct sensitivity to ATP or glibenclamide was not tested in their study.

Tsuchiya et al. (1992) found an ATP-sensitive K⁺ channel in five patches made on the BLM of nonperfused rabbit S1 and S2 segments. With 145 mM K⁺ in the patch pipette and a bath containing 140 mM Na⁺ and 5 mM K⁺, they reported an inward g_slope = 56 pS, although an I-V plot was not given. In c/a patches at a command potential of 0 mV, they found a baseline nP_o of 0.72, much higher than that found in the present study and the two studies in which perfused tubules were patched (see below) (Hurst et al., 1993; Beck et al., 1993). Although 1 mM ATP applied to the cytoplasmic side of an inside-out patch reversibly reduced nP_o by 77% (from 0.70 to 0.16), the sensitivity of the basolateral conductance was reduced in a perfused tubule when 1 mM ATP was added to the bath.

Beck et al. (1993) were the first to patch clamp the BLM of (collagenase-treated) perfused rabbit proximal tubules, and found a K⁺ channel whose activity correlated to transport activity. This channel showed inward rectification in c/a patches (with g_in = 61 pS and g_out = 17 pS) and P_o was voltage independent. In a follow-up study (Hurst et al., 1993), the same group showed that 2 mM ATP (i/o) inhibited channel activity by ~80% and 100 μM diazoxide (c/a) effectively opened this BLM K⁺ channel.

The ROMK family of K⁺ channels is, to date, the only cloned ATP-regulated renal K⁺ channel (Ho et al., 1993). The ROMK1 channel cloned from rat kidney is an inwardly rectifying K⁺ channel with g_in = 39 pS and a high open probability (0.8–0.9) at voltages more depolarized than −60 mV, and is not activated by hyperpolarization. ROMK1 exhibits channel rundown when membrane patches are excised in ATP-free bath, although ROMK1 is not sensitive to ATP or glibenclamide (Ho et al., 1993). In situ hybridization studies show that transcripts (mRNA) for ROMK are absent.

### Basolateral membrane

**Figure 11.** Equivalent electrical cell/circuit. Adjacent proximal tubule cells are outlined as dashed lines, and the equivalent circuit discussed in the text is diagrammed on the left. The basolateral membrane potential is denoted by voltage source V_bl. The current produced by the Na,K-ATPase pump, I_Pump, is given by I_Pump = I_pump + I_pump. Basolateral ion channel currents are shown as their Thévenin equivalents with their conductances, G_K and G_Cl, represented by the equivalent resistor, in series with their Nernst potentials, E_K and E_Cl, respectively. The paracellular current, I_shunt, is represented by I_shunt = I_shunt + I_shunt. R_A is the resistance of the apical membrane. Kirchoff’s current law at Node A is given by Eq. 11.
from the proximal tubule (Lee and Hebert, 1995). Combined with the differences in mechanism of channel rundown (see above and Table III), it is highly unlikely that the BLM $K^+$ channel is ROMK1, although it may be within the same family of inwardly rectifying $K^+$ channels with two transmembrane domains (Ho et al., 1996; Kubo et al., 1993).

**Contribution of the BLM $K^+$ Channel to Total BLM Conductance**

The component of BLM conductance ($G_{BLM}$) due to the BLM $K^+$ channel ($G_K$) can be estimated as:

$$G_K = g \cdot nP_o \cdot f$$

where $g$ is the “physiologic” single channel conductance, $n$ is the average number of channels in a patch, $P_o$ is the mean open time of one channel, $f$ is the incidence of finding a channel in the patch, and $a$ is the area of the membrane patch.

In the present study, the conductance ($g$) for outward currents was $\sim 4$ pS, the average $nP_o$, extrapolated to 0 mV was 0.20, and channel activity was found in 98.6% of patches. Given that the average inner diameter of our patch pipette was 1.5 $\mu m$, the minimal membrane area of a patch is 1.77 $\mu m^2$. Using these estimates in *Ambystoma*, the value for $G_K$ is (4 pS*0.20*0.986)/1.77 $\mu m^2$ = 0.446 pS/$\mu m^2$ or 44.6 S/cm$^2$ ($R_K = 22.4$ k$\Omega$·cm$^2$). Considering the estimate of $G_{BLM} = 51.3$ S/cm$^2$ ($R_{BLM} = 19.5$ k$\Omega$·cm$^2$) in *Ambystoma* proximal tubule (Maunsbach and Boulpaep, 1984), $G_K/G_{BLM} = 44.6/51.3 = 0.87$. Thus, under the conditions used in our experiments, $G_K$ may account for 87% of $G_{BLM}$. However, to the extent that the area of the membrane patch is larger than our estimate, $G_K$ will be smaller.

The basolateral membrane area of dissociated *Ambystoma* proximal tubule cells exhibits less infoldings than these cells in situ (Segal et al., 1996). Assuming a whole-cell membrane capacitance of $\sim 120$ pF (Segal et al., 1996) and a specific capacitance of 1 pF per 100 $\mu m^2$, we can estimate that the surface of a single isolated cell is $\sim 12,000$ $\mu m^2$. Using a BLM surface area of 10,000 $\mu m^2$ ($100 \times 10^{-6}$ $cm^2$) and a $G_{BLM} = 51.3$ S/cm$^2$, the total BLM conductance is 5.13 nS. Using the transport number $G_K/G_{BLM}$ of 0.87 from Table IV, $G_K = 4.46$ nS. Despite possible changes in membrane area (e.g., due to membrane recycling) and/or channel density that could occur in the dissociated cells over time, this is a reasonable estimate since perforated-patch whole-cell studies with KCl pipette and NaCl bath ($E_K = -84$ mV) show current that reverses at $V_m = -40$ mV with $\sim 240$ pA of outward current at 0 mV command potential, and a whole-cell conductance of 6 nS. Thus, the total BLM conductance is 5.13 nS, of which 4.46 nS is $K$ conductance, and the total apical conductance is 0.87 nS (the difference of 6–5.13).

Similar comparisons between $G_K$ derived from single-channel data and $G_{BLM}$ based on cable analysis are shown in Table IV for *Necturus, Rana*, and rabbit proximal tubule, respectively. With the exception of the rabbit, the transport numbers show that the BLM conductance of all other species is dominated by $K^+$ and, specifically for *Ambystoma*, $K_{ATP}$ channels. From this information, one can estimate the $K^+$ channel density on the BLM of dissociated *Ambystoma* proximal tubule cells. If $P_o$ was unity, the number of channels would be given by $4.46$ nS/4 $pS = 1.115$ channels. However, since $P_o = 0.05$, the number of $K^+$ channels on the BLM is $\sim 22,300$ per cell; a channel density of 2.23 channels/$\mu m^2$. Using the estimate of membrane patch area of 1.77 $\mu m^2$, one would expect on average approximately four channels per patch, in good agreement with our experience.

**Physiological Role of the BLM $K_{ATP}$ Channel**

The task of the proximal tubule under physiologic conditions is to effect transport; i.e., to maintain ionic flux in the direction of reabsorption. The electrochemical difference across the BLM of proximal tubule, given by $V_{in} - E_K$, is an invariably positive driving force promoting $K^+$ efflux (Boulpaep, 1979). The presence of other conductances in the BLM with reversal potentials less negative than $E_K$ provides a nonzero difference between $E_K$ and $V_{in}$ that maintains K efflux. The BLM $Na^+, K^+-ATPase$ pump loads the cell with two $K^+$ ions to effect the transcellular transport of three $Na^+$ ions. An efflux pathway for $K^+$ is necessary to recycle $K^+$ across the BLM, thus permitting continuous operation of the pump. Pump activity and the opening of these $K^+$ channels hyperpolarize the BLM, favoring apical $Na^+$ entry. An interesting contrast is that the negative shift in membrane potential consequent to the opening of $K_{ATP}$ channels tends to depress cell function in excitable cells, but tends to promote transport in epithelia. The magnitude of BLM $K^+$ efflux must match the $K^+$ current of the pump. The regulatory effects of hyperpolarization and intracellular ATP levels on the BLM $K_{ATP}$ channels thus subserve this role.

This system of a variable BLM $K^+$ conductance coupled to changes in pump rate in a single cell has further implications when extrapolated to the level of an epithelium with a paracellular shunt pathway. Such an epithelial model should include all current pathways (see Fig. 11, adapted from Fig. 5 of Sackin and Boulpaep, 1983); namely, (a) the active $Na^+$ and $K^+$ pump currents, (b) the basolateral $K^+$ leak current, (c) the other basolateral leak currents, principally carried by $Cl^-$, and (d) the shunt current carried by $Cl^-$ and, to a
lesser extent, by Na\(^+\). In this equivalent electrical cell/circuit, Kirchhoff’s current law at node A gives

\[
I_{\text{pump}}^{\text{K}} + I_{\text{Cl}}^{\text{bl}} - I_{\text{Cl}}^{\text{shunt}} - I_{\text{Na}}^{\text{bl}} = 0 ,
\]

(11)

where \(I_{\text{pump}}^{\text{K}}\) is the transcellular component of Cl\(^-\) absorption, and \(I_{\text{Cl}}^{\text{shunt}}\) is the paracellular component of Cl\(^-\) absorption and any Na\(^+\) backleak. If the outward current through BLK K channels could be exactly matched to \(I_{\text{pump}}^{\text{K}}\), then Eq. 11 becomes

\[
I_{\text{pump}}^{\text{K}} - I_{\text{Cl}}^{\text{bl}} - (I_{\text{Cl}}^{\text{hun}} + I_{\text{Na}}^{\text{shunt}}) = 0 .
\]

(12)

When Eq. 12 is satisfied, all actively transported Na\(^+\) is accompanied by Cl\(^-\) absorption (sum of transcellular and paracellular), except for the backleak, \(I_{\text{Na}}^{\text{shunt}}\). Under these optimal conditions, the proximal tubule accomplishes NaCl reabsorption at maximum efficiency.

Note that if \(G_K^{\text{bl}}\) did not increase with pump rate, \(I_{\text{Cl}}^{\text{shunt}}\) would rise only slightly due to the small increase in driving force. Under these conditions, \(I_{\text{pump}}^{\text{K}} > I_{\text{Cl}}^{\text{bl}} + (I_{\text{Cl}}^{\text{hun}} + I_{\text{Na}}^{\text{shunt}})\) and the efficiency of NaCl reabsorption deteriorates. That is, the incremental increase in NaCl reabsorption would be less than that of the pump. Therefore, the tight linkage between the Na\(^+\),K\(^-\)-ATPase pump and the K\(^+\) leak through the BLK K\(_{\text{ATP}}\) channels is especially important in leaky epithelia that absorb large, fluctuating quantities of NaCl such as the proximal tubule. The regulation of the BLK K\(_{\text{ATP}}\) channels and how they are coupled to the pump during transport is the subject of the companion paper (Mauerer et al., 1998).

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R E F E R E N C E S


