Inward-rectifier Potassium Channels in Basolateral Membranes of Frog Skin Epithelium

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ABSTRACT Inward-rectifier K channel: using macroscopic voltage clamp and single-channel patch clamp techniques we have identified the K+ channel responsible for potassium recycling across basolateral membranes (BLM) of principal cells in intact epithelia isolated from frog skin. The spontaneously active K+ channel is an inward rectifier (Kw) and is the major component of macroscopic conductance of intact cells. The current–voltage relationship of BLM in intact cells of isolated epithelia, mounted in miniature Ussing chambers (bathed on apical and basolateral sides in normal amphibian Ringer solution), showed pronounced inward rectification which was K+-dependent and inhibited by Ba2+, H+, and quinidine. A 15-pS Kw channel was the only type of K+-selective channel found in BLM in cell-attached membrane patches bathed in physiological solutions. Although the channel behaves as an inward rectifier, it conducts outward current (K+ exit from the cell) with a very high open probability (Po = 0.74–1.0) at membrane potentials less negative than the Nernst potential for K+. The Kw channel was transformed to a pure inward rectifier (no outward current) in cell-attached membranes when the patch pipette contained 120 mM KCl Ringer solution (normal NaCl Ringer in bath). Inward rectification is caused by Mg2+ block of outward current and the single-channel current–voltage relation was linear when Mg2+ was removed from the cytosolic side. Whole-cell current–voltage relations of isolated principal cells were also inwardly rectified. Power density spectra of ensemble current noise could be fit by a single Lorentzian function, which displayed a K dependence indicative of spontaneously fluctuating Kw channels. Conclusions: under physiological ionic gradients, a 15-pS inward-rectifier K+ channel generates the resting BLM conductance in principal cells and recycles potassium in parallel with the Na+/K+ ATPase pump.

INTRODUCTION

Sodium absorption across tight-junction epithelia occurs by passive entry into the cell at the apical membrane through amiloride-blockable channels and active secretion...
across the basolateral membrane via an Na⁺/K⁺ ATPase pump. Potassium is recycled across this membrane through a channel. In the Ussing model the basolateral membrane behaves like a pure potassium electrode in that the variations in transepithelial potential in response to changes in extracellular K⁺ concentration can be predicted from the Nernst equation (Koefoed-Johnsen and Ussing, 1958). This could imply that one type of K⁺ channel, possibly showing Goldman rectification, is involved in recycling potassium. In absorptive epithelia, however, where K⁺-dependent current−voltage relations have been reported, it seems that the K⁺-conductive pathway at the basolateral membrane cannot be described by Goldman-Hodgkin-Katz (GHK) formalism but instead shows a higher conductance for inward than for outward current (inward rectification) (Nagel, 1985; Horisberger and Giebisch, 1988). Similar conclusions that an inward-rectifier K⁺ (Kᵢ) channel dominates the membrane conductance was reached in recent patch clamp studies in isolated cells from frog skin (García-Díaz, 1991).

The isolated frog skin is the classical model for sodium-absorbing tight epithelia, but little is known about the physiology of the potassium channel(s) in the basolateral membrane of this tissue. Our specific aim in this study was to identify the type of K⁺ channel in basolateral membranes responsible for steady-state K⁺ recycling and compare the single-channel and macroscopic K⁺ conductance properties in the intact epithelium.

This is the first investigation of single K⁺ channels in basolateral membranes of an intact sodium-transporting epithelium. The combination of the patch clamp recording of single-channel activity and conventional microelectrode measurements of membrane potential and conductance in the intact epithelium permits a detailed analysis of K⁺ channel physiology. The data reported here show that the macroscopic inward rectification of basolateral membrane current is generated by inward-rectifier K⁺ channels which share similar voltage-gated properties with the anomalous rectifier of excitable cells (Hagiwara, 1983).

METHODS

Epithelium Isolation

Intact epithelial sheets were isolated from the ventral skin of Rana esculenta by means of a combined enzymatic and mechanical separation technique. The isolated skin (11 cm²) was exposed on the corial side for 60 min to 1 mg/ml collagenase (125 IU activity; Worthington Biochemical Corp., Freehold, NJ) under 10 cm of hydrostatic pressure at 30°C. This treatment was terminated when blisters first formed on the epithelial side. A fine polyethylene catheter (0.25 mm o.d.) was introduced into a blister and Ringer solution was injected into the subepithelial space. This resulted in detachment of a large sheet of epithelium (7 cm²). The basement membrane could easily be detached by flushing the epithelium in calcium-free Ringer buffered with 2 mM EGTA for 3–10 min. Epithelia isolated in this way had transepithelial sodium transport rates and transepithelial resistances comparable with whole skins. The epithelium was mounted basolateral side upwards in a miniature Ussing chamber which allowed conventional short-circuit current measurements, intracellular microelectrode recording, and single-channel patch clamp studies to be performed on the same epithelium (Fig. 1).

The frog skin epithelium is a complex three- to five-cell layered structure composed of several cell types that include granular cells and mitochondria-rich cells in the layers
immediately below the outermost layer of dead cells (stratum corneum), and germinal cells in the deeper layers. There is a great deal of evidence from studies of cell volume and membrane potential responses to amiloride, ouabain and changing external Na and K concentrations that the multilayered epithelium acts as a syncytial Na transport compartment (Kristensen and Ussing, 1985; Rick, Dorge, von Arnim, and Thurau, 1978). Only the first granular cell layer has a polarized apical membrane containing Na+ channels and are termed principal cells. The cells in deeper layers are coupled to the first reactive cell layer by gap junctions (Farquhar and Palade, 1965; Sharin and Blankemeyer, 1989) so that they share the sodium transport pool. The mitochondria-rich cells, which are the site of active acid secretion, are not considered to be part of this syncytium. In our isolated epithelial preparation, principal and mitochondria-rich cells could easily be distinguished and although we also found inward-rectifier K+ channels in the basolateral membranes of mitochondria-rich cells, only the properties of Kᵢ channel in principal cells are reported here.

![Ussing Model](image)

**Figure 1.** The Ussing model of sodium reabsorption through principal cells and schema of the experimental arrangement for recording short-circuit current, intracellular potential, and single-channel activity in an isolated epithelium mounted in a miniature Ussing chamber on the stage of an inverted Nikon Diavert microscope.

The experiments were carried out at room temperature. The apical side was always bathed in standard NaCl Ringer solution and the basolateral side was continually superfused in either Na or K Ringer solutions of composition shown in Table I. In solutions containing EGTA, free concentrations of calcium were calculated using a computer program (Chang, Hsieh, and Dawson, 1988). All chemicals were purchased from Sigma Chemical Co. (St. Louis, MO).

**Dissociated Cells**

Single cells were isolated for whole-cell patch clamp recording. The basolateral side of the epithelium mounted in an Ussing chamber was superfused in Ca-free Ringer solution containing 10 mM EGTA for 2–5 min. A small area of epithelium was then scraped with a broken tip patch electrode to detach single cells from the epithelial sheet. In this way cells could be isolated from every layer of the epithelium. The combination of scraping and suction of cells was also used to expose basolateral membranes of the cells in the outermost layer ("first
reactive principal cell layer") for single-channel recording. These cells have polarized apical and basolateral membranes separated by tight junctions whereas cells from deeper layers are nonpolar and behave electrically as a single basolateral membrane.

**Basolateral Membrane Current-Voltage Analysis**

Conventional microelectrodes were prepared from borosilicate glass capillaries (250F Clark Electromedical, Pangbourne, UK) using a programmable puller (DMZ Universal Puller, Werner Zeitz, Augsburg, Germany). Electrode tip resistance was between 40 and 60 MΩ when filled with 1 M KCl solution. The electrical arrangement for recording basolateral membrane current-voltage (Ib - Vb) relations has been previously described in detail (Harvey, Thomas, and Ehrenfeld, 1988). After microelectrode impalement of principal cells from the basolateral side a transepithelial bipolar pulse train (pulse duration 210 ms, interval 100 ms) incrementing in 5-mV steps to ±200 mV was generated under computer command via a voltage clamp amplifier (model VCC600, Physiologic Devices, Houston, TX). The basolateral membrane potential was measured with a high input impedance differential amplifier (model 750, World Precision Instruments Inc., Hew Haven, CT). Voltage and current outputs were digitized and stored on computer for analysis. Barium-sensitive current-voltage relations were obtained by subtracting Ib - Vb relations recorded in the absence and presence of 5 mM BaCl₂. The K⁺ selectivity of the basolateral membrane conductance was examined by running voltage clamp ramps in standard Ringer (3.7 mM K⁺) solution and in Ringer with 37 mM K⁺ added as gluconate salt. Transference numbers for potassium (T_k) were calculated from the magnitude of Vb changes in response to varying E_k by 58 mV (increasing external [K] from 3.7 to 37 mM).

**Single-Channel Recording**

The cell-attached and inside-out configurations of the patch-clamp technique (Hamill, Marty, Neher, Sakmann, and Sigworth, 1981) were used for single channel studies. Patch pipettes (tip
resistance 4–10 MΩ) were prepared from borosilicate glass capillaries (Vitrex, Modulohm, Herlev, Denmark), pulled, and polished on a programmable puller (DMZ, Zeitz Instrumente, Augsburg, Germany). The patch pipettes were filled with either standard NaCl Ringer or an intracellular-like K⁺ solution ("Kᵢₘ⁺" composition given in Table I). All solutions were filtered through 0.2-µm cellulose disks (Millipore Corp., Bedford, MA). Current and voltage recordings from patch-clamp electrodes were amplified (Biologic RK 300 patch-clamp amplifier, Biologic, Claise, France), passed through a 30-kHz filter and stored on video tape (Betamax, Sony, Tokyo, Japan) after pulse code modulation (PCM model 501ES, Sony). The single-channel current records were digitized and displayed in real time using a digital oscilloscope (model 310, Nicolet Instrument Corp., Madison, WI). Single-channel current records for illustrations were filtered at 1 kHz using a Tchebycheff filter and replayed on a chart recorder (model RS3200, Gould Instruments, Valley View, OH).

Current and voltage signals were digitized using a 16-bit A/D converter (CED 1401, Cambridge Electronic Design, Cambridge, UK) after low-pass (−3 dB) filtering at 5 kHz (8 pole Bessel filter, model 902, Frequency Devices Inc., Haverhill, MA) and analysed with a model 80486 50-MHz Dell Corp. (Limerick, Ireland). Steady-state current–voltage relations were run and analyzed using VGEN and PAT programs (J. Dempster, Strathclyde Electrophysiology Software version 6.1, University of Strathclyde, Scotland) and instantaneous current–voltage trials were generated using PULSESTIM and VCLAMP programs (Cambridge Electronic Design, release EPC 5.0). Single-channel current amplitude histograms, open probability, and open and closed dwell times were analyzed from data segments of 20–60-s duration using the eAT program. Single-channel K⁺ permeability was calculated from the general form of the GHK equation:

\[
I = \frac{P_K V^2 F}{RT} \left\{ \frac{[K]_0 \exp(VF/RT) - [K]_i}{\exp(VF/RT) - 1} \right\}
\]

where \(R\), \(T\), and \(F\) have their usual meanings, \([K]_0\) and \([K]_i\) are the external and internal concentrations of potassium, and \(P_K\) is the permeability to K⁺. Cation movement across the membrane from the external to the cytoplasmic side is defined as inward current and shown as downward deflections in single-channel records.

**Whole-Cell Current–Voltage Analysis in Isolated Cells**

The whole-cell configuration was obtained from the cell-attached mode after breaking the membrane patch by applying a brief negative pressure pulse (−200 mbar) in the pipette. Whole-cell currents were amplified (RK300 Biologic), digitized at 20 kHz (CED 1401) and sampled in real time on hard disk at an acquisition rate of 5 kHz. Current–voltage relations were analyzed using CED VCLAMP software.

**Whole-Cell Current Noise Analysis**

The application of current noise analysis to epithelial cells has been reviewed in detail by Van Driessche and Zeiske (1985) and we give only the essentials of the technique as applied in this study. Current noise was recorded from isolated dispersed single principal cells perfused through the patch pipette with an intracellular-like K⁺ solutions ("whole-cell" solution in Table I). Whole-cell current fluctuations were amplified by the patch clamp amplifier and split into low and high pass components using Bessel filters with cutoff frequencies at 1 kHz and 0.1 Hz, respectively, and stored on video tape. The current noise signal was replayed through the CED1401 A/D converter, digitized at 5 kHz, and analyzed by a fast Fourier transform using customized software. The variance of the fluctuations in current noise is presented as a function of the alternating current frequencies contained in the current noise (power density spectra).
An average power spectrum was compiled from 200 record segments (each of 250 ms in duration) and channel noise was recorded at pipette holding potentials between ±100 mV relative to $E_K$. A Lorentzian curve was fitted to the power spectrum using a Levenberg-Marquadt iterative fitting algorithm (K. Brown, University of Cincinnati). For a channel that fluctuates between distinct open and closed states, the power spectrum will contain a Lorentzian defined by the equation:

$$L(f) = S_o/(1 + (f/f_c)^2)$$

where $S_o$ is the low frequency asymptote or plateau value and $f_c$ the corner or cutoff frequency at which the spectral power is $S_o/2$. The cutoff frequency is related to the time constant ($\tau$) of channel open/closure fluctuations by

$$\tau = (2\Pi f_c)^{-1}$$

where $\tau = (\alpha + \beta)^{-1}$ and $\alpha$ and $\beta$ are on- and off-rates for channel transitions from open to closed and closed to open states, respectively, based on a two-state model, OPEN $\xrightarrow{\alpha} \text{CLOSED}$. The time constant is related to open probability by

$$P_o = \beta/(\alpha + \beta)$$

Closed and open transition rates were calculated from

$$\beta = P_o'2\Pi f_c, \quad \alpha = (2\Pi f_c) - \beta$$

using values for open probabilities ($P_o$) determined in excised inside-out recordings under identical bathing conditions and voltage. The open and closed rate constants $\alpha$ and $\beta$ were compared with time constants $t_o$ and $t_c$ measured from the exponential distribution of open and closed times, respectively. The variance in channel current-noise ($\sigma^2$) was calculated by integrating the power spectrum

$$\sigma^2 = (\Pi f_c S_o)/2$$

and the single-channel conductance ($G_o$) was then determined from

$$G_o = \sigma^2/|I_m(V_p - E_K)|$$

where $I_m$ is the mean direct current recorded in the power spectrum and $V_p$ is the pipette holding voltage. Single-channel current ($I_s$) can then be obtained from

$$I_s = G_o(V_p - E_K)$$

The density of channels in the cell membrane ($N$) was estimated from

$$N = \Pi f_c S_o/2(G_o')^2 P_o(1 - P_o)$$

Data are presented as mean values ± SEM.

RESULTS

Macroscopic Properties of Basolateral Membrane Conductance

The basolateral membrane current–voltage relationship ($I_b - V_b$) shows steep inward rectification under spontaneous Na$^+$ transport conditions (Fig. 2A). The membrane is therefore more conductive for inwardly directed current (cation entering or anion leaving the cell) than for outward current (cation leaving or anion entering the cell). The degree of rectification and the potential at which the current changes sign
(reversal potential, $V_r$) were a function of the serosal K$^+$ concentration. Raising [K$^+$] in the serosal bath from 3.7 to 37 mM produced an increase in conductance for inward current which was 10-fold greater than for outward current and shifted $V_r$ by +48 mV. These effects are expected if the basolateral membrane current is carried mainly by potassium. Practically all types of K$^+$ channels are blocked by barium ions and the experiment in Fig. 2 A shows that addition of 5 mM BaCl$_2$ to the serosal bath completely eliminated the inward current rectification. Ba$^{2+}$ reduced the potassium transference from 0.82 ± 0.05 to 0.09 ± 0.03 (n = 10 cells) and the membrane potential no longer responded to changes in external K$^+$. The K$^+$-specific current-voltage relations of the basolateral cell membranes was extracted by taking the difference between $I_b - V_b$ curves measured in control and in the presence of barium (Fig. 2 B). Ba$^{2+}$-sensitive $I_b - V_b$ relations show strong inward rectification and the increased membrane conductance for inward current is evident at hyperpolarized membrane potentials. Acidification of the epithelium at pH 6.5 produced a similar decrease in basolateral membrane conductance as in Ba$^{2+}$ (Fig. 3 A). Quinidine at high concentrations (100 μM) also inhibited the inward rectification of basolateral membrane current (Fig. 3 B). Quinidine has been used to block Ca$^{2+}$-activated K$^+$ currents in turtle colon (Chang and Dawson, 1988) but other inhibitors of these classes of K$^+$ channels were without effect on membrane conductance. For example, $I_b - V_b$ relations were insensitive to quartenary ammonium ions such as (5 mM) tetraethyl ammonium (Fig. 3 C), tetrapentyl ammonium, or choline. The honey bee venom, apamin, which is considered specific for calcium-dependent "small" conductance K$^+$ channels, had no effect (at 10$^{-9}$ M) on basolateral membrane current. The maxi-K channel blocker, charybotoxin (10$^{-8}$ M) was also without effect, making it unlikely that the basolateral membrane K$^+$ conductance is generated by calcium-activated K$^+$ channels. Basolateral membrane current was, however, sensitive to removal of extracellular Ca$^{2+}$ which enhanced inward rectification (Fig. 3 D).

**Single-Channel Recording**

*Cell-attached recordings in NaCl Ringer solution.* We found it relatively easy to obtain patch-clamp seal resistances $>$ 10 GΩ on basolateral cell membranes of principal

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Figure 2. Macroscopic current-voltage relations of basolateral cell membranes of isolated frog skin epithelium. (A) Control situation: standard Na Ringer solution (Ⅰ) and after addition of 5 mM BaCl$_2$ (Ⅱ). (B) “Difference” or Ba$^{2+}$-sensitive current-voltage relation for A.
cells, but almost 50% of basolateral membrane patches showed no channel activity. When the experiments were made with sodium Ringer solution in the bath and patch pipette, only one type of K⁺-selective channel was found in over 500 cell-attached patches in normal Na-transporting epithelia. The spontaneously active K⁺ channels showed properties of inward rectification which was similar to the voltage dependence of the macroscopic basolateral membrane current. The single-channel current–voltage relationship (Fig. 4) is opposite to that expected from GHK formalism.

**FIGURE 3.** Current–voltage relations for basolateral cell membranes. (A) Effect of intracellular acidification from a control pH 7.4 (●) to pH 6.4 (■) and comparison with effects of 5 mM BaCl₂ (★). (B) Effect of quinidine 100 μM: control (●); quinidine (■). (C) Effect of tetraethylammonium (TEA) 5 mM: control (●); TEA (○). (D) Effect of removing Ca²⁺ from basolateral Ringer solution, control (▲), calcium-free (▲).
Despite the favorable chemical gradient for K⁺ exit from the cell into the bath (and patch pipette), the channel behaves as an inward rectifier, being more conductive for inward current than for outward current. The unit current amplitude was small (0.5 pA) at the resting membrane potential and a slope conductance of 3pS was calculated from the current–voltage relation at zero holding potential. When the potential across the patch was hyperpolarized by > 40 mV the single-channel current reversed direction. As expected for a K⁺-selective channel, the reversal potential \( V_r = E_m - V_p \) at which the current changes sign agrees with reasonable measurements of the Nernst potential for K⁺ (\( [K^+] = 120 \text{ mM}, \ E_K = -88 \text{ mV}, \ E_m = -50 \text{ mV}, \) and \( V_p = 40 \text{ mV} \)) (Harvey and Kernan, 1984). At \( V_p > 40 \text{ mV} \) the electrochemical driving force favors K⁺ entry from the bath (and pipette) into the cell. A mean single-channel conductance (Eq. 8) of 15 ± 2 pS (\( n = 8 \) cells) was calculated between the reversal potential and \( V_p = 100 \text{ mV} \).

The experiment in Fig. 5A shows single-channel current recordings from an inward-rectifier K⁺ channel (Kir) in the basolateral membrane of a principal cell in the cell-attached mode when both the bath and pipette contained normal NaCl Ringer solution. At the depolarized membrane potential \( (V_p = -30 \text{ mV}) \), the channel was open and passed outward current for practically the whole duration of the recording. When the membrane potential was hyperpolarized \( (V_p > 40 \text{ mV}) \), the channel conducted inward current and openings occurred in bursts with rapid transitions to the closed state during the burst. Channel activity often reflected the openings of multiple ion channels in the patch and was measured as the product of the number of active channels and open probability \( (N'P_o) \). Channel activity was calculated from

\[
N'P_o = \sum (n't_n)
\]

where \( n \) represents the channel state, \( 0 = \text{closed}, 1 = \text{one channel open}, 2 = \text{two channels open}, \) etc., and \( t_n \) is the duration of state \( n \). Channel activity is shown as a function of applied voltage in Fig. 5B, and was near-maximal at the resting membrane potential and decreased for inward K⁺ current when the membrane voltage was hyperpolarized beyond \( E_K \). These results show that the Kir channel can conduct K⁺ out of the cell with a very high open probability. An evaluation of how the Kir channel could influence the macroscopic basolateral membrane conductance can be obtained from the voltage dependence of the product of single-channel current and channel activity \( (I'N'P_o) \) as shown in Fig. 5C. Kir channel activity will confer inward rectification properties on the basolateral membrane, and behaves as a near-constant current source for outward current. This would appear to be an ideal property for a channel recycling K⁺ in parallel with a constant current Na/K pump.

**Cell-attached recordings with KCl in the patch pipette.** In cell-attached patches bathed with 125 mM KCl solution in the pipette and NaCl Ringer solution in the bath, >75% of patches showed spontaneous K⁺-selective channel activity. Similar Kir channel activity was recorded in Kext (1 mM Ca²⁺) or in Kint pipette solutions (100 mM Ca²⁺) but the latter solution was used preferentially to allow transfer from cell-attached to whole-cell recording configurations in the same cell. The experiment in Fig. 6A shows single-channel currents recorded when pipette voltage was varied between ±100 mV in 10-mV steps with Kint in the pipette and normal Ringer in the
The channel conducted $K^+$ from pipette into the cell at hyperpolarized membrane potentials but outward currents were barely detectable even when the membrane was depolarized by 100 mV. The corresponding current–voltage relationship is included in Fig. 6 B. Single-channel conductance measured between $V_p -20$ and +100 mV was 35 ± 4 pS ($n = 12$). In the presence of high external $K^+$ the channel is transformed to an almost pure inward rectifier and only movement of $K^+$ into the cell is possible. The voltage dependence of channel activity ($N\cdot P_o$) recorded from a multichannel patch clearly shows this diodelike property (Fig. 6 C). Channel activity increased with membrane hyperpolarization and this is in contrast to channel behavior observed with standard Na Ringer solution in the pipette where $P_o$...
decreased with membrane hyperpolarization. The open probability for inward current was higher in low calcium pipette solutions compared to high calcium solutions, with Ca\(^{2+}\) producing a flicker-type block of inward current openings (not shown). The blocking effect of extracellular calcium was also noted for macroscopic inward current (Fig. 3 D). The single-channel current–voltage relation was practically linear for inward current in high K pipette solutions. Fitting the GHK equation to the current–voltage data between \(V_p\) 0 and 100 mV yielded a mean value for the potassium permeability = 0.38 (±0.08) \(10^{-13}\) cm/s (\(n = 12\)).

![Figure 6](image_url)
Single-Channel Biophysics in Excised Inside-out Membranes

Na Ringer in the pipette. In excised inside-out membrane patches, the Kir channel conserved the basic properties of inward rectification observed in the cell-attached configuration. The patches were normally excised into intracellular-like KCl Ringer solution (Kint at pCa 7, pH 7.2). Single Kir channel currents recorded over a wide voltage range are shown in Fig. 7A from an excised patch exposed to a Kint solution on the cytosolic side and standard Na Ringer in the pipette. The membrane was voltage clamped for 800 ms to the potentials (Vc) indicated on the right of the trace.

Figure 7. Current-voltage relationship of Kir channels in excised inside-out basolateral membranes exposed to standard Na Ringer in the patch pipette and Kint solution in the bath. (A) Single-channel recording at imposed membrane potentials. The patch potential is shown on the left of each current trace and the closed level is indicated by a dashed line. Cutoff frequency 1 kHz. (B) Mean single-channel current-voltage relationship (n = 8) showing inward rectification and reversal potential of -85 mV, which is close to Ek (-88 mV).

The calculated Ek across the patch is -85 mV and it can be seen in Fig. 7A that the channel passes outward current (from bath into pipette) for Vc less negative than Ek and inward current for Vc more negative than Ek. Under these physiological ionic gradients, the reversal potential is close to Ek, indicating a high selectivity for potassium. The mean single-channel current-voltage relationship is shown for eight inside-out patches in Fig. 7B and presents clear inward rectification. The slope conductance for inward current (17 ± 2 pS, n = 18 cells) and for outward current (5 ± 2 pS, n = 18 cells) correspond to the conductances measured in the cell-attached mode. The high open probability for outward current and the flickery-type
openings during inward current are also comparable with the cell-attached observations. The relative permeability of Na\(^+\) to K\(^+\) \((P_{Na}/P_{K})\) was obtained in mixed solutions of Na and K from

\[ V_r = \frac{RT}{F} \ln \left( \frac{[K]_o + (P_{Na}/P_{K})[Na]_o}{[K]_i + (P_{Na}/P_{K})[Na]_i} \right) \]

where \([K]_o, [Na]_o\) are the concentrations (in mM) of external (patch pipette) K\(^+\) (3.7) and Na\(^+\) (120), and \([K]_i, [Na]_i\) are the internal (bath) K\(^+\) (120) and Na\(^+\) (12) concentrations, respectively. \(P_{Na}\) and \(P_{K}\) are the permeability to Na\(^+\) and K\(^+\), respectively. The reversal potential \((V_r) = -79 \pm 0.6\) mV \((n = 7\) patches\) and the

![Figure 8](image)

**Figure 8.** Voltage dependence of Kir channel current in excised inside-out basolateral membranes exposed to K\(_{int}\) solution on both sides. (A) Single-channel recording as a function of imposed membrane voltage. Values of \(V_c\) are shown on right of trace, and the closed level is indicated by a dashed line. Note absence of channel opening at depolarized membrane potentials and multiple-channel activity during hyperpolarization. Cutoff frequency 1 kHz. (B) Single Kir channel current-voltage relationship showing diode-like behavior. Line through data points fitted by eye (mean values for eight cells).

Mean calculated \(P_{Na}/P_{K} = 0.013\), indicating the channel is 77 times more selective for potassium.

**Channel activity in symmetrical KCl solutions.** In inside-out patches bathed in symmetrical \(K_{int}\) Ringer solutions, no outward currents were recorded between 0 and +100 mV relative to \(E_K\) (0 mV), whereas the channel was activated at \(V_c\) more negative than \(E_K\). Representative single-channel recordings are shown as a function of membrane potential (±100 mV) in Fig. 8A and the current–voltage relationship from a mean of eight cells is given in Fig. 8B. The channel was inactive at depolarized potentials whereas channel activity \((N-P_o)\) was greatly increased by membrane hyperpolarization. Since the current–voltage relation presents an ohmic
conductance at hyperpolarized membrane potentials, an estimate of the single-channel K⁺ permeability for inward current was calculated from the general form of the GHK equation. The potassium permeability was 0.41\cdot10^{-13} \text{ cm/s}, which is in excellent agreement with $P_K$ calculated in cell-attached patches.

**Cytosolic Mg²⁺ and inward rectification.** Removing magnesium ions from the internal side of inside-out patches, resulted in a loss of inward rectification (Fig. 9 A). Single $K_r$ channels displayed near linear current–voltage relations when exposed to Mg-free (2 mM EDTA) symmetrical $K_{\text{int}}$ solutions (Fig. 9 B). The mean single-channel conductance was $35 \pm 2 \text{ pS}$ ($n = 6$ patches) over the membrane potential range ±100 mV.

**Inhibition of $K_r$ Channels**

The effects of barium were tested on excised inside-out patches bathed in normal Ringer and $K_{\text{int}}$ solutions on extra- and intracellular sides, respectively. Barium (5 mM) added to the cytosolic side reduced $K_r$ channel activity instantaneously and reversibly without affecting the open-state conductance. The effects of $\text{Ba}^{2+}$ on channel kinetics were voltage-dependent and the reduction in $P_o$ was greater for outward current. Barium reduced $P_o$ for inward current from 0.74 ± 0.04 to 0.28 ± 0.03 and outward current $P_o$ from 0.85 ± 0.04 to 0.02 ± 0.01 ($n = 8$ cells). Inhibitors of calcium-dependent K⁺ channels were without effect when applied to the cytosolic side of excised patches, such as internal tetraethylammonium or tetrapotassiumammonium (5 mM), or external charybdotoxin (10 nM) or apamin (10 nM), whereas quinidine in high concentrations (>100 \text{ µM}) produced a flicker-type block. In excised outside-out patches exposed to symmetrical $K_{\text{int}}$ solutions, external Cs⁺ caused a flicker-type block which increased with membrane hyperpolarization. Maximal inhibition was produced at 5 mM CsCl with an apparent $K_i = 300 \text{ µM}$.

The $K_r$ channel is less permeant to Rb⁺ and a permselectivity ratio $P_{\text{Rb}}/P_K =$
0.67 ± 0.04 (n = 6 patches) was calculated from changes in the reversal potential after equimolar substitution of KCl with RbCl.

**Whole-Cell Recordings of Kᵦ Current**

Whole-cell current was recorded from isolated single principal cells separated from neighboring cells by mechanical suction while incubated in calcium-free Ringer solutions. Under physiological ionic gradients (Na Ringer in bath, Kᵦ pCa 8 solution in pipette), the whole-cell current displayed marked inward rectification. In kᵦ solutions, the whole-cell currents exhibited time-dependent inactivation which was prevented when the pipette filling solution contained guanosine 5'-O-(3-thiotriphosphate) GTPγS (10 μM). We, therefore, routinely used GTPγS in whole-cell pipette filling solutions for the investigation of Kᵦ currents.

Fig. 10 A shows whole-cell currents generated at holding potentials between ±100 mV for cells bathed externally in 120 mM K⁺, 54 mM K⁺, and 3.7 mM K⁺. The amplitude of inward current was increased at high external K⁺ concentrations as expected for a K⁺-selective membrane. The whole cell current records and the corresponding current–voltage relations in Fig. 10 B, illustrate the loss of outward K⁺ current in high external K solutions which is a characteristic property of single Kᵦ channels. The whole-cell slope conductance for inward current greatly exceeds the conductance for outward current and changes in Vᵦ produced by raising bath K⁺ concentration agree with the predicted shift in E_K (Fig. 10 C). A slope of 56 mV/decade change in [Kᵦ]ᵦ was calculated for the relationship Vᵦ versus log [Kᵦ]ᵦ. Whole-cell currents are generated essentially by potassium flux through inward rectifier K⁺ channels. The dependence of whole-cell conductance (for inward current) on external [Kᵦ]ᵦ is shown in Fig. 11. The data were fit by a form of the Michaelis-Menten equation:

\[
G_{cell} = \frac{G_{max}[K^+]_0}{[K^+]_0 + [K^+]_{50}}
\]

This analysis gave an upper limit for cell membrane conductance of 5,294 pS and a [Kᵦ]₅₀ = 32 mM at which cellular conductance was half-maximal. A double logarithmic plot of Gcell versus [Kᵦ]₀ had a slope of 0.59 which is close to the value expected (0.5) if conductance is proportional to the square root of [Kᵦ]₀, as reported for inward rectifiers in excitable cells (Hagiwara, 1983; Harvey and Ten Eick, 1988).

**Whole-Cell Current Noise Analysis**

The kinetics of Kᵦ channels and their contribution to the cell membrane conductance was further investigated by fluctuation noise analysis of whole-cell currents. Whole-cell membrane current noise was recorded at different external [K⁺] (3.7, 54, and 120 mM). The pipette solution contained Kᵦ solution (pCa 8) with GTPγS (whole-cell solution in Table 1). The power density spectra presented in Fig. 12 A were recorded at the Nernst potential for K and at holding potentials which generated electrochemical driving forces ΔµK = +90 or +100 mV favoring K⁺ efflux and ΔµK = −90 or −100 mV driving K⁺ influx. The power density spectra could be fit by a single Lorentzian function (solid line through the data points) which is indicative of
Figure 10. (A) Whole-cell current recordings as a function of external bath K⁺ concentration (Kᵢₒ solution in the pipette) in 120 mM K, 54 mM K, and 4 mM K. Negative (outward) currents were recorded at pipette potentials imposed in 10-mV steps to +100 mV. Positive (inward) currents were generated in increasing amplitude at pipette voltages clamped in 10-mV steps to −100 mV. Current signals were filtered at 5 kHz. Note disappearance of outward (negative) currents and increased amplitude of inward currents at raised external [K⁺]. (B) Current–voltage relationship for the conditions 120 mM K (■), 54 mM K (○), and 3.7 mM K (●). In each case the current–voltage relation shows inward rectification and deviates from Goldman formalism. (C) The dependence of reversal potentials of whole-cell current on extracellular K⁺ concentration (mean values for 12 cells at each different extracellular K concentration. The cell membrane behaves as a K⁺ electrode.

Figure 11. Whole-cell conductance for inward current plotted as a function of external potassium concentration (n = 4 cells). The solid line through the data points was drawn to Eq. 13 using values for $G_{\text{max}} = 5.294$ nS and $[\text{K}⁺]_{\text{out}} = 32$ mmol/liter.
an homogeneous population of spontaneously fluctuating channels. The results of the Lorenzian analysis for outward currents and inward currents at various extracellular K concentrations are summarized in Table II. The noise power decreased for outward currents and increased for inward current at high external [K]. This finding is in agreement with the K dependence of single channel and whole-cell currents. For example, cells exposed to physiological ion gradients at a holding potential of 15 mV ($\Delta\mu K = +100$ mV), the power density spectra of outward current fluctuations had a

![Figure 12](image)

**Figure 12.** (A) Power density spectra (PDS) of fluctuations in whole-cell outward and inward currents as a function of applied voltage and external K+ concentration ($K_o$) (mean values for six cells at each $K_o$). The pipette was filled with $K_{ion}$ solution (Table I). The bath concentrations of K+ and the electrochemical driving forces are given by the following symbols (•) $K_o = 120$ mM, $\Delta\mu K = -100$ mV. (★) $K_o = 54$ mM, $\Delta\mu K = -100$ mV. (●) $K_o = 3.7$ mM, $\Delta\mu K = -100$ mV. (○) $K_o = 3.7$ mM $\Delta\mu K = +100$ mV. The PDS were fit by a single Lorentzian function (solid line through data points) and a summary of data analysis is given in Table II. Raising external K+ increased the Lorentzian plateau and decreased the corner frequency for inward current fluctuations. Outward current noise power was greatly reduced in high K+ solutions (not shown). No distinct Lorentzian could be discerned in the PDS when the patch membrane potential was clamped to the Nernst potential for K+. (B). Voltage dependence of the plateau Lorentzian in the PDS of whole-cell current noise recorded under different external K+ concentrations (mean values for six cells at each $K_o$). The pipette filling solution was $K_{ion}$ (120 mM K) throughout. The extracellular K+ concentrations were as follows: (●) 120 mM $K_o$, (★) 54 mM $K_o$, (●) 3.7 mM $K_o$. In each case the minimum Lorentzian plateau value was measured at the K equilibrium potential (↓).

A single corner frequency of 9 Hz and a Lorentzian plateau of 0.21 pA^2 s^-1. When the same cells were bathed in 54 mM K+ the Lorentzian plateau was reduced 10-fold. Cells bathed in symmetrical K+ solutions of 120 mM KCl, did not contain an evident corner frequency for outward current fluctuations.

The power density spectra of inward current noise fluctuations was also affected by varying external K+ (Table II). The high corner frequency of inward current noise reflects rapid fluctuations between open and closed states which is a characteristic of
single $K_{ir}$ channels. Calculations from noise analysis of single-channel open and closed times and conductance yielded values in agreement with the patch clamp data for single $K_{ir}$ channels (Table II). For example, for inward current fluctuations recorded under physiological ionic gradients, the calculated off-rate $\alpha = 490$ s$^{-1}$ (2 ms) is in excellent agreement with measured values for single $K_{ir}$ channel mean open time $= 3$ ms. The equivalence between the mean open times obtained from single-channel recording and the off-rates determined from noise analysis demonstrates that the analyses are inherently consistent. Finally, the voltage dependence of the plateau Lorentzian ($S_o$) in the power density spectrum recorded at different extracellular K concentrations is shown in Fig. 12 B. The plateau Lorentzians show a minimum value at membrane potentials equal to the Nernst potential for K$^+$. For inward currents, the calculated number of active channels $N$ increases with mem-

**TABLE II**

| $S_o$ | $f_c$ | $\alpha$ | $\beta$ | $P_0$ | $G_o$ | $I_s$ | $N$ |
|-------|-------|----------|--------|-------|-------|-------|------|
| pA$^2$/Hz | Hz | s$^{-1}$ | pS | pA channels/cell |

**Outward currents**

| 3.7 mM K | 0.21 | 9.0 | 1.55 | 55 | 0.97 | 5.2 | 0.7 | 109 |
| 54 mM K | 0.027 | 16 | 78.5 | 22 | 0.22 | 1.4 | 0.31 | 41 |

**Inward currents**

| 3.7 mM K | 0.408 | 156 | 490 | 490 | 0.50 | 11.5 | 0.98 | 204 |
| 54 mM K | 3.11 | 88 | 216 | 357 | 0.61 | 19.7 | 1.77 | 577 |
| 120 mM K | 8.10 | 75 | 132 | 339 | 0.78 | 25 | 2.5 | 727 |

*Open probability was determined from excised inside-out patches under similar ionic gradients. Fluctuations in whole cell outward current at 120 mM extracellular K were of too low amplitude to give meaningful analysis. For comparison, the reciprocal of the measured single $K_{ir}$ channel mean open time (off-rate) is given in parentheses under each $\alpha$ value.

brane hyperpolarization and extracellular K whereas the converse is true for outward currents. This phenomenon reflects the behavior of single $K_{ir}$ channels to augment $N$ in response to increasing extracellular K and hyperpolarization (compare Fig. 5 with Fig. 6 and Fig. 7 with Fig. 8). It is reasonable to conclude, therefore, that the whole-cell current noise is generated solely by fluctuations of a single population of inward-rectifier K$^+$ channels and the macroscopic membrane conductance is dominated by $K_{ir}$ channel activity.

**DISCUSSION**

In the original Ussing model, the basolateral cell membrane conductance was portrayed as K$^+$ diffusion from the cell. Since then many microelectrode studies have
confirmed that potassium recycling across the basolateral membrane accounts for much of the total membrane conductance. The basolateral membrane behaves almost as a pure K\(^+\) electrode in its potential response to changes in K\(^+\) chemical gradient (Nagel, 1979). In this study, we have identified the type of K\(^+\) channel normally present in the basolateral cell membrane of identified principal cells. The channel displays inward rectification under physiological ionic gradients and is the major determinant of membrane conductance. Our results from intact polarized epithelia can be compared to a recent patch clamp study of isolated frog skin cells which showed that most of the cell conductance is due to the activity of K\(^+\) channels (García-Díaz, 1991). In the latter study ion channel activity was recorded in primary cultures of trypsin-isolated cells from \textit{R. pipiens} and \textit{R. catesbiana}. Inward rectification was a great deal less pronounced and single K\(^+\) channel conductance was slightly larger (44 pS) in the cells studied by García-Díaz compared to the diodelike inward rectification and single-channel conductance of 35 pS recorded in our experiments conducted with KCl Ringer (pCa 8) in the patch pipette. In experiments on trypsin-isolated cells from frog skin we have found the cell isolation procedure by itself can activate a calcium-dependent K\(^+\) channel of 45 pS conductance (Harvey and Urbach, 1992). Inclusion of millimolar calcium in the patch pipette can also activate this channel at strong depolarizing potentials. This channel has a linear current–voltage relationship and is completely blocked by 100 μM quinidine. The presence of the ohmic K\(^+\) channel in the Garcia-Diaz study would produce a masking of the inward rectifier and confer quinidine sensitivity to whole-cell K\(^+\) currents. In our study on intact sodium-absorbing polarised epithelia, there is excellent agreement between the electrophysiological properties of the basolateral membranes of principal cells and single Kir channels.

The property of anomolous rectification would seem to limit the usefulness of the Kir channel to recycle K\(^+\). However, under physiological conditions, the open probability is high for outward current over the normal range of membrane potentials. The low conductance of the Kir channel for outward current would allow efficient voltage clamping of the membrane potential close to \(E_K\) with minimum loss of K\(^+\) from the cell.

\textit{Comparison with Other Epithelial Inward-rectifier K Channels}

Inward rectification of K\(^+\) current is not a novel feature of epithelial cells and has been reported in macroscopic current–voltage relations for basolateral membranes of toad urinary bladder (Garty, 1984), frog skin (Nagel, 1985), and \textit{Amphiuma} collecting tubule (Horisberger and Giebisch, 1988). Single-channel patch clamp data have revealed inward rectifier K channels in the basolateral membranes of dogfish rectal gland (Greger et al., 1987), and in basolateral membranes of renal tubule cells, \textit{Necturus} proximal tubule (Kawahara, Hunter, and Giebisch, 1987), rabbit proximal tubule (Gogelein, Greger, and Schlatter, 1987; Parent, Cardinal, and Sauve, 1988), cultured MDCK cells (Ponce and Cereijido, 1991), and frog early distal tubule (Wang and Giebisch, 1989; Hunter, 1991). Inward rectifier K\(^+\) channels are not confined solely to basolateral membranes but have also been reported in luminal membranes of MDCK (Friederich et al., 1989), cultured opossum (Ohno-Shosaku, Kubota, Yamaguchi, and Fujimoto, 1990), and rat collecting tubule (Frindt and Palmer, 1989;
Wang and Giebisch, 1989) and rabbit thick ascending limb of Henle's loop (Bleich, Schlatter, and Greger, 1990; Wang, White, Geibel, and Giebisch, 1990). Irrespective of their origin or localization, epithelial inward rectifier \( K^+ \) channels appear to share a common function; they are involved in moving \( K^+ \) out of the cell either as a net secretion (when present in the apical membrane) or \( K^+ \) recycling (when localized in the basolateral membrane) and as in frog skin principal cells, are the major determinant of membrane conductance.

The \( K_{ir} \) channel in frog skin epithelium presents some unusual features. Under physiological ion gradients, the single channel conductance increases with hyperpolarization but open probability increases with depolarization. This means that inward rectification cannot be ascribed to voltage dependence of channel activity \( (N\cdot P_o) \). Here we have shown that inward rectification results from block of outward current by cytosolic magnesium ions. The transformation of the \( K_{ir} \) channel to a pure inward rectifier in symmetrical \( K^+ \) solutions and the ohmic properties in Mg-free solutions under these conditions may indicate that Mg\(^{2+}\) inhibition of \( K_{ir} \) channels is potentiated by raising external \([K^+]_o\). Recently, it was shown that Mg\(^{2+}\) block of the \( K_{ir} \) channel in atrial muscle can be influenced by external \( K^+ \) (Matsuda, 1991).

The inhibition by external Ca\(^{2+}\) of inward currents in intact epithelia may explain the reduction of open probability with depolarization in calcium-containing external solutions. These results are similar to the observations of Ca\(^{2+}\) blockade of inwardly rectifying \( K^+ \) currents in guinea-pig ventricular myocytes (Biermans, Vereecke, and Carmeliet, 1987).

The anomalous rectifier properties of the \( K_{ir} \) channel may have a role in the kaliuretic response to hyperkalemia. The increased inward rectification in high K solutions would effectively reduce \( K^+ \) recycling across the basolateral membrane and thus favour net \( K^+ \) secretion. Conductance for inward whole-cell \( K_{ir} \) current increased proportionally with the square root of external \( K^+ \) concentration whereas outward conductance decreased approximately fourfold at \([K^+]_o\) 3.7 to 10 mM. It would thus be expected that physiological changes in plasma \([K^+]_o\) of 2–8 mM which occur during adaptation of frogs and toads to high external KCl media will produce a marked reduction in \( K^+ \) recycling and facilitate \( K^+ \) secretion, as is observed under these conditions (Van Driessche and Zeiske, 1985).

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