Effects of External Rb⁺ on Inward Rectifier K⁺ Channels of Bovine Pulmonary Artery Endothelial Cells

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ABSTRACT Inward rectifier (IR) K⁺ channels of bovine pulmonary artery endothelial cells were studied using the whole-cell, cell-attached, and outside-out patch-clamp configurations. The effects of Rb⁺ on the voltage dependence and kinetics of IR gating were explored, with [Rb⁺]₀ + [K⁺]₀ = 160 mM. Partial substitution of Rb⁺ for K⁺ resulted in voltage-dependent reduction of inward currents, consistent with Rb⁺ being a weakly permeant blocker of the IR. In cells studied with a K⁺-free pipette solution, external Rb⁺ reduced inward IR currents to a similar extent at large negative potentials but block at more positive potentials was enhanced. In outside-out patches, the single-channel i-V relationship was approximately linear in symmetrical K⁺, but rectified strongly outwardly in high [Rb⁺]₀ due to a reduced conductance for inward current. The permeability of Rb⁺ based on reversal potential, \( V_{rev} \), was 0.45 that of K⁺, whereas the Rb⁺ conductance was much lower, 0.034 that of K⁺, measured at \( V_{rev} \approx 80 \text{ mV} \). The steady state voltage-dependence of IR gating was determined in Rb⁺-containing solutions by applying variable prepulses, followed by a test pulse to a potential at which outward current deactivation was observed. As [Rb⁺]₀ was increased, the half-activation potential, \( V_{1/2} \), changed less than \( V_{rev} \). In high [K⁺]₀ solutions \( V_{1/2} \) was \( V_{rev} - 6 \text{ mV} \), while in high [Rb⁺]₀, \( V_{1/2} \) was \( V_{rev} + 7 \text{ mV} \). This behavior contrasts with the classical parallel shift of \( V_{1/2} \) with \( V_{rev} \) in K⁺ solutions. Steady state IR gating was less steeply voltage-dependent in high [Rb⁺]₀ than in K⁺ solutions, with Boltzmann slope factors of 6.4 and 4.4 mV, respectively. Rb⁺ decreased (slowed) both activation and deactivation rate constants defined at \( V_{1/2} \), and decreased the steepness of the voltage dependence of the activation rate constant by 42%. Deactivation of IR channels in outside-out patches was also slowed by Rb⁺. In summary, Rb⁺ can replace K⁺ in setting the voltage-dependence of IR gating, but in doing so alters the kinetics.

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INTRODUCTION

The mechanism of gating of inward rectifier (IR) channels appears to differ qualitatively from that of other voltage-dependent ion channels. Whereas most voltage-gated ion channels are envisioned as opening and closing as a direct result of voltage-induced alterations in the structural configuration of the channel protein, the voltage-dependence of IR channel gating is profoundly affected by the concentration of permeant ions, and thus most likely arises from a distinct mechanism. Major structural differences between a recently cloned IR channel (Kubo, Baldwin, Jan, and Jan, 1993) and a large group of structurally similar voltage-gated K⁺ channels are consistent with the possibility that these two classes of channels may have distinct gating mechanisms. The voltage-dependence of the IR was first described as a strict function of the extracellular K⁺ concentration, [K+]o, such that rectification depended on the reversal potential for K⁺, E_K (Hodgkin and Horowicz, 1959; Almers, 1971; Hagiwara and Takahashi, 1974). A distinction was made between instantaneous rectification of the IR and a voltage- and time-dependent gating mechanism (Almers, 1971; Hagiwara, Miyazaki, and Rosenthal, 1976; Leech and Stanfield, 1981; Hestrin, 1981; DeCoursey, Dempster, and Hutter, 1984). Changes in intracellular K⁺ concentration, [K+]i, were shown to affect the position of the voltage-activation curve of the IR, although less profoundly than changes in [K+]o (Hagiwara and Yoshii, 1979; Hestrin, 1981; Saigusa and Matsuda, 1988; Cohen, DiFrancesco, Mulrine, and Pennefather, 1989; Pennefather, Oliva, and Mulrine, 1992). It has thus been possible to dissociate shifts in the position of the voltage-activation curve from shifts in V_rev.

Pennefather et al. (1992) proposed that inward rectification is not an intrinsic property of the single IR channel conductance, but rather that rectification results from a voltage-dependent isomerization that is intrinsic to the channel protein, and that is modulated by allosteric regulatory binding sites for K⁺ on the outside of the channel.

In the past decade, permeant ion effects on gating have been demonstrated in a number of ion channels (references cited in Shapiro and DeCoursey, 1991b). The effects of permeant ions on the gating of other channels are more subtle than the effects of [K+]o on the IR, and generally have been viewed as modulation of a gating mechanism which is intrinsic to the channel protein, rather than as evidence that gating itself might be due to interaction of permeant ions with the channel (Chen and Eisenberg, 1992). In addition to the profound effects of [K+]o on IR gating, when Tl⁺ replaces K⁺ in the external solution, activation of IR currents becomes faster (Hagiwara, Miyazaki, Krasne, and Giani, 1977) and an inactivation process occurs upon hyperpolarization (Stanfield, Ashcroft, and Plant, 1981). The permeant blockers of cardiac IK₁ channels, Rb⁺ and Cs⁺, slow activation (Tournier, Mitra, Morad, and Rougier, 1987; Mitra and Morad, 1991) and result in IK₁ current "inactivation," which appears to represent block by external divalent cations, in cells dialyzed with K⁺-free solution (Mitra and Morad, 1991).

We have studied the relationship between [Rb⁺]o and the IR gating mechanism in bovine pulmonary artery endothelial cells. Under the ionic conditions employed, the IR is the predominant ionic conductance present, and this combined with their small size and favorable geometry (compared with myocytes, for example), makes endothe-
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lial cells a nearly ideal preparation for these studies. Rb+ blocks inward K+ currents in a voltage-dependent manner, but is also weakly conductive. At the single channel level, Rb+ reduces the apparent unitary current amplitude, presumably due to block with rapid kinetics. A major result of the present study is that Rb+, like K+, can "set" the position on the voltage axis of the voltage-activation curve of the IR channel. We demonstrate that as the Rb+ mole-fraction is increased, the midpoint of the macroscopically observable gating process does not shift to the same extent as $V_{\text{rev}}$. As a result, in high $[\text{Rb+}]_o$ most of the channels are open at $V_{\text{rev}}$, whereas in high $[\text{K+}]_o$ most of the channels are closed at $V_{\text{rev}}$. In addition, Rb+ slows the kinetics of gating. Thus, in high $[\text{Rb+}]_o$, it is possible to observe large and relatively slowly decaying transient outward IR currents during depolarizing voltage pulses from potentials at which the IR conductance was activated. This phenomenon is exploited to quantify the voltage-dependence of the IR gating mechanism over the full range of Rb+ and K+ mole fractions. External Rb+ slows IR gating. The steady state activation curve is less steep in high $[\text{Rb+}]_o$, which in terms of a simple two-state model can be ascribed to a decrease in the steepness of the voltage-dependence of the activation rate. These effects of Rb+ are interpreted with respect to a quantitative model of gating in a companion paper (Pennefather and DeCoursey, 1994).

MATERIALS AND METHODS

Cells

The techniques used for isolation, culture, and identification of bovine pulmonary artery endothelial cells were described previously (Silver and DeCoursey, 1990). Briefly, endothelial cells were isolated from bovine pulmonary arteries obtained from a slaughterhouse, by enzymatic digestion: 0.25% collagenase (Worthington CL-II, Cooper Biomedical, Malvern, PA) incubated for 30 rain at 37°C. Cells were studied from passage 3 through 9. Cells were plated onto fragments of glass cover slips before electrophysiological study, in order to facilitate transfer into the recording chamber.

Solutions

The composition of the solutions used are given in Table I A and B. Solutions are indicated in the text according to their monovalent cation constituents in mM: e.g., 40 K + 120 Rb. The pipette solution used in most whole-cell experiments was KCH$_3$SO$_4$, and was nominally Mg$^{2+}$-free in order to minimize possible block of outward current (Vandenberg, 1987; Matsuda, Saigusa, and Irisawa, 1987) and interaction with the gating mechanism (Matsuda, 1988; Oliva, Cohen, and Pennefather, 1990; Silver and DeCoursey, 1990; Pennefather et al., 1992). Liquid junction potentials arising between the pipette solution and the bath solution and between the reference electrode and the bath solution were calculated as described previously (Silver and DeCoursey, 1990; Shapiro and DeCoursey, 1991a), and were <5.2 mV in all experiments. Data presented in tables and in Figs. 6-12 have been corrected, except Fig. 10, while other raw data are illustrated without correction. Most chemicals were purchased from Sigma Chemical Company (St. Louis, MO) or Aldrich Chemical Company (Milwaukee, WI).

Electrophysiology

Standard gigohm-seal recording techniques were used (Hamill, Marty, Neher, Sakmann, and Sigworth, 1981). Cells selected for recording were approximately spherical, and had input
**TABLE IA**

**Composition of Solutions**

| Name          | Na⁺ | K⁺ | Rb⁺ | Ca²⁺ | Mg²⁺ | Cl⁻ | HEPES |
|---------------|-----|----|-----|------|------|-----|-------|
| Ringer’s      | 160 | 4.5| 0   | 2    | 1    | 170.5| 5     |
| 160 Na        | 160 | 0  | 0   | 2    | 1    | 166  | 10    |
| 160 K         | 0   | 160| 0   | 2    | 1    | 166  | 10    |
| 160 Rb        | 0   | 0  | 160 | 2    | 1    | 166  | 10    |
| 4.5 K + 160 Rb| 0   | 4.5| 160 | 2    | 1    | 170.5| 5     |
| 160 K + 4.5 Rb| 0   | 160| 4.5 | 2    | 1    | 170.5| 5     |

Mixtures of K⁺ and Rb⁺ are referred to as, e.g., 40 K + 120 Rb, meaning 40 mM K⁺ and 120 mM Rb⁺ and were made by mixing appropriate amounts of 160 K and 160 Rb. Solutions with varying [K⁺]o in which K⁺ is the only permeant ion were made by combining appropriate amounts of 160 K and 160 Na. Solutions were titrated to pH 7.4 with the hydroxide of the predominant cation. HEPES is [4-(2-hydroxyethyl)-1-piperazineethane-sulfonic acid].

Capacities of typically 10–15 pF. Standard conventions are used in all configurations, i.e., outward current flowing from the intracellular to the extracellular side of the membrane is positive and is plotted as upward; applied clamp potentials which depolarize the intracellular side of the membrane are positive, and in the cell-attached patch configuration, potentials are stated relative to the cell’s resting potential (RP) which is unknown, e.g., RP + 80 mV means the membrane patch was depolarized by 80 mV above RP. Micropipettes were pulled in several stages using a Flaming Brown automatic pipette puller (Sutter Instruments, San Rafael, CA) from EG-6 glass (or infrequently from 0010 or KG-12 glass) obtained from Garner Glass Co. (Claremont, CA). Pipettes were coated with Sylgard 184 (Dow Corning Corp., Midland, MI), and heat polished to a tip resistance measured in Ringer’s solution ranging typically between 0.9–1.8 MΩ. Electrical contact with the pipette solution was achieved by a thin sintered Ag-AgCl pellet (In Vivo Metric Systems, Healdsburg, CA) attached to a silver wire covered by a Teflon tube. Pipette and the initial bath solutions were filtered at 0.1–0.2 μm (Millipore Corp., Bedford, MA). A reference electrode made from a Ag-AgCl pellet was connected to the bath through an agar bridge saturated with Ringer’s solution. The current signal from the patch

**TABLE IB**

**Composition of Solutions**

| Name          | K⁺  | Cl⁻ | F⁻  | CH₃SO₄⁻ | Mg²⁺ | Ca²⁺ | EGTA | HEPES |
|---------------|-----|-----|-----|---------|------|------|------|-------|
| *KCH₃SO₄      | 175 | 4   | 0   | 120     | 0    | 2    | 10   | 10    |
| KCH₃SO₄+2 Mg  | 160 | 6   | 0   | 140     | 2    | 1    | 10   | 10    |
| KCH₃SO₄/F     | 162 | 6   | 40  | 100     | 2    | 1    | 10   | 10    |
| NMGCH₃SO₄     | 2   | 4   | 0   | 160     | 2    | 0    | 5    | 5     |

*The KCH₃SO₄ solution also had 10 mM EDTA (ethylene-diamine-tetraacetic acid).

Contains nominally 160 mM NMG⁺ (N-methyl-D-glucamine).

K⁺ solutions were titrated to pH 7.2–7.4 with KOH, NMGCH₃SO₄ solution was titrated to pH 7.0 with NMG⁺. The dipotassium salt of EGTA, ethylene glycol bis-(β-aminoethyl ether) N,N,N',N'-tetraacetic acid, was used in K⁺ solutions. Monovalent cation concentrations are approximate and include estimates of the KOH or NMG⁺ added to titrate the buffers present. Free [Mg²⁺] is ~1.9 mM in KCH₃SO₄+2 Mg and NMGCH₃SO₄, and <50 nM in the other pipette solutions (cf Silver and DeCoursey, 1990).
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clamp (Axopatch-1A, Axon Instruments, Inc., Burlingame, CA) was recorded and analyzed using an Indec Laboratory Data Acquisition and Display System (Indec Corporation, Sunnyvale, CA). Series resistance compensation with the Axopatch-1A is prone to oscillation, and therefore was not used routinely. After whole-cell configuration was achieved by suction, the series resistance read from the dial was typically about double the pipette resistance in the bath (2–4 MΩ). The voltage error for a 1 nA current would be 2–4 mV, but most measurements of gating kinetics and voltage-dependence were done near \( V_{rev} \), where currents generally were smaller than this. Experiments were done at room temperature, with the bath temperature monitored continuously by a thinfilm platinum RTD (resistance temperature detector) element (Omega Engineering, Stamford, CT) immersed in the bath solution, and the temperature stored along with each record. Recorded bath temperature ranged 19–23°C, but was typically 21°C with a standard deviation of <1°C for each data set in Table III. When extracted rate constants were plotted against temperature, the scatter in the data obscured any temperature effects, so the data were not corrected for temperature.

Data Analysis

Data were digitized and recorded in computer files for subsequent analysis. Analysis programs were written in BASIC-23 or in FORTRAN-77. The procedures for generating null-subtracted, averaged single-channel relaxations (Figs. 11, A and B) use programs written by Dr. Richard S. Lewis (Stanford University, CA) and are described fully elsewhere (Shapiro and DeCoursey, 1991a). Current relaxations were fitted by eye to exponential functions, while Boltzmann fits were by nonlinear least squares. Significance of differences were calculated by Student’s two-tailed \( t \) test.

RESULTS

Inwardly rectifying K + (IR) channels have been observed in endothelial cells isolated from bovine pulmonary artery (Johns, Lategan, Lodge, Ryan, Van Breeman, and Adams, 1987; Cannell and Sage, 1989; Silver and DeCoursey, 1990), bovine aorta (Colden-Stanfield, Schilling, Ritchie, Eskin, Navarro, and Kunze, 1987; Takeda, Schini, and Stoeckel, 1987; Olesen, Davies, and Clapham, 1988), and human umbilical vein (Nilius and Reimann, 1990). Under the conditions of the present study of bovine pulmonary artery endothelial cells, the IR predominated whole-cell currents except in the positive potential range, and comprised the only obviously time-dependent conductance. Whole-cell currents during voltage ramps are plotted in Fig. 1. These ramp currents are equivalent to steady state current-voltage curves, except near \( V_{rev} \), because of the rapidity of gating of the IR (Fig. 1, legend). In symmetrical K + -containing solutions the IR gives rise to large inward currents with an approximately constant slope conductance, and a small hump of outward current positive to \( V_{rev} \) which vanishes 40–60 mV positive to \( V_{rev} \) (Adrian and Freygang, 1962). At more positive potentials there are outward currents unrelated to the IR which in the cell in Fig. 1 A are approximately linear and in the cell in Fig. 1 B increase progressively at large positive potentials. Some endothelial cells have a Cl − conductance which under these ionic conditions would carry some outward current, and which is reduced but not eliminated when extracellular Cl − is replaced by methanesulfonate − (Shapiro and DeCoursey, 1991c). In other cells there are increasing outward currents at quite positive potentials (as in Fig. 1 B) which are not reduced by replacement of Cl − with methanesulfonate −. Both of these conductances
are largely time-independent and are not affected when the IR is abolished by Ba\(^{2+}\) (Silver and DeCoursey, 1990), or when all external permeant cations are removed (160 Na trace in Fig. 1 B).

Rb\(^{+}\) Block

Fig. 1 A illustrates superimposed whole-cell currents recorded during voltage ramps from -100 to +60 mV in a cell exposed to a wide range of \([\text{Rb}^{+}]_o\). The presence of even small amounts of Rb\(^{+}\) in the external solution reduces inward currents through the IR. At each \([\text{Rb}^{+}]_o\) the fractional block at first increases at more negative potentials, indicating that Rb\(^{+}\) block is voltage-dependent, but at large negative potentials the blockade decreases again, with a weaker voltage dependence. Quite similar voltage-dependence of block was described by Standen and Stanfield (1980, their Fig. 2 B). That block does not continue to increase at large negative potentials suggests that Rb\(^{+}\) can permeate IR channels, and that in this voltage range the blocking and unblocking rates have a similar voltage-dependence. The voltage-dependence of Rb\(^{+}\) block can be estimated from the ratio of the current in the presence of Rb\(^{+}\) to that in 160 K. Neglecting the gradual increase at large negative potentials (Standen and Stanfield, 1980), this ratio could be fitted by a Boltzmann function, with a midpoint shifting to more positive potentials as \([\text{Rb}^{+}]_o\) increased, and which limited to a minimum that decreased as \([\text{Rb}^{+}]_o\) increased. The slope factor

\[ I = \frac{I_{\text{Rb}}} {I_{\text{K}}} = \frac{1}{1 + \exp \left( \frac{V - V_0}{\Delta V} \right) } \]

where \(V_0\) is the midpoint, \(\Delta V\) is the slope factor, and \(I_{\text{Rb}}/I_{\text{K}}\) is the ratio of currents in the presence of Rb\(^{+}\) and K\(^{+}\), respectively.

The figure shows how the current decreases as \([\text{Rb}^{+}]_o\) increases, with a clear voltage-dependence.

**Figure 1.** Whole-cell currents during voltage ramps in the indicated solutions, labeled according to the \([\text{Rb}^{+}]_o\) in mM with the remainder K\(^{+}\), except for the trace in B labeled Na\(^{+}\) which was permeant-free 160 Na. The voltage was ramped from -100 to 60 mV at 0.52 mV/ms. The pipette solution was KCH\(_2\)SO\(_4\), bath temperature was 20.5°C (A) and 21.7°C (B). At both large negative and large positive potentials, these ramp currents are equivalent to steady state current-voltage curves, because gating of the IR is very fast (\(\tau < 1 \text{ ms}\)) more than 30 mV from \(V_{1/2}\) in 160 K (Silver and DeCoursey, 1990). In the voltage range near \(V_{1/2}\) where IR channel gating is slowest, some distortion will occur, especially in Rb\(^{+}\)-containing solutions in which gating is slower, because IR channels will open or close depending on the direction of the ramp. Even in 160 Rb, the maximum value for \(\tau\) in most experiments is <30 ms, which is the time calibration bar in this figure.
averaged 12.4 ± 2.7 mV (mean ± SD, n = 7) in 140 K + 20 Rb, and the potential at which half-block occurred was -2.7 ± 8.7 mV (n = 7).

**Rb⁺ Permeation**

In Fig. 1 B ramp currents are shown at higher gain in another cell to illustrate the situation at high [Rb⁺]o. There are small inward currents in 80 K + 80 Rb, and smaller but qualitatively similar currents in 160 Rb. The small inward currents are much reduced when Rb⁺ is replaced by Na⁺. Evidently, Rb⁺ can permeate IR channels. When only Rb⁺ is present externally there is inward rectification (a superlinear I–V relationship) at large negative potentials which is eliminated in 160 Na. Inward rectification of pure Rb⁺ current through the IR also occurs in other tissues (Hagiwara and Takahashi, 1974; Standen and Stanfield, 1980).

In high [Rb⁺]o solutions in Fig. 1 B, there is a hump of outward current just positive to Vrₑ. In fact the slope conductance is greater for outward currents just positive to Vrₑ than for inward currents just negative to Vrₑ. This outward current hump is due to outward K⁺ current (from the cell containing K⁺) flowing through IR channels, because the hump disappears when Rb⁺ is replaced by Na⁺ (Fig. 1 B), and the behavior of the hump is consistent with the gating kinetics of the IR. Thus, when the potential is ramped more slowly the hump is reduced, because the channels have more time to close at a given potential. Ramping from positive to negative potentials eliminates the hump (not shown) because the IR channels are closed at positive potentials, and by the time they open the voltage is negative to Vrₑ.

**Rb⁺ Conductance with K⁺-free Pipette Solution**

The possibility that intracellular K⁺ might interact with Rb⁺ permeation led us to examine the Rb⁺ conductance in endothelial cells using K⁺-free pipette solutions. Whole-cell currents are illustrated in Fig. 2 in a cell dialyzed with NMGCH₃SO₃ (Table I B), during voltage ramps from +80 mV to −160 mV. When the bath solution was changed from Ringer’s with 4.5 mM K⁺ (not shown) to 160 K, the inward current increased ~20-fold, and outward current remained small. This behavior, together with the ubiquitous presence of large IR currents in endothelial cells, and the absence of other K⁺-selective conductances at negative voltages with low intracellular free Ca²⁺, supports the conclusion that these currents are due to IR channels. Changing the bath to 160 Rb greatly decreased the inward current (Fig. 2). Addition of 5 mM BaCl₂, an IR blocker, nearly abolished the inward current in 160 Rb, providing further support for the identity of the conductance as IR. The inward currents in 160 Rb were also greatly reduced by replacement with Na⁺, TMA⁺ (tetramethylammonium), or TEA⁺ (tetraethylammonium). In conclusion, the Rb⁺ conductance of the IR in endothelial cells is low regardless of the presence or absence of intracellular free Ca²⁺, supports the conclusion that these currents are due to IR channels. Changing the bath to 160 Rb greatly decreased the inward current (Fig. 2). Addition of 5 mM BaCl₂, an IR blocker, nearly abolished the inward current in 160 Rb, providing further support for the identity of the conductance as IR. The inward currents in 160 Rb were also greatly reduced by replacement with Na⁺, TMA⁺ (tetramethylammonium), or TEA⁺ (tetraethylammonium). In conclusion, the Rb⁺ conductance of the IR in endothelial cells is low regardless of the presence or absence of intracellular K⁺.

Whole-cell currents in several Rb⁺ and K⁺ mixtures are also illustrated in Fig. 2. As in cells studied with K⁺ in the pipette (Fig. 1), small additions of Rb⁺ greatly reduced inward K⁺ currents. One intriguing difference is that the enhancement of Rb⁺ block by hyperpolarization was not observed in cells studied with K⁺-free pipette solutions; instead the slope conductance increased monotonically with hyperpolarization at all [Rb⁺]o. The ratio of inward current in Rb⁺-containing solutions to that in 160 K increased at more negative potentials. Block was only weakly voltage-dependent at
160 mM Rb solution with 5 mM BaCl₂ added, "+5 BaCl₂" is nearly superimposed on the voltage axis. The changes observed were rapidly reversible.

large negative potentials and similar to that with normal K⁺ in the cell, but increased at more positive potentials. This is in contrast with the situation in normal [K⁺], where the opposite voltage-dependence was observed in the voltage range positive to ~80 mV, namely relief of block by depolarization. This difference was not due to ramping "down" from positive to negative potentials, because ramping down with normal K⁺ in the cell produced results indistinguishable from those when ramping up (except near V_{rev} where gating is slowest). Evidently intracellular K⁺ relieves block at potentials near V_{rev}. Perhaps the relief of Rb⁺ block in cells with normal [K⁺], reflects interactions of Rb⁺ and internal K⁺ within the channel.

Relative Conductance and Permeability

In Fig. 3 the ratio of current at -90 mV in the presence of various [Rb⁺]₀ to the current in 160 K is plotted, both for cells with K⁺ internally (●) and for cells perfused
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Regardless of the internal solution, the reduction of inward current by Rb+ is much greater than would result from simply replacing external K+ with an inert impermeant ion. The current at -90 mV was reduced by 50% at <10 mM Rb+. Evidently, Rb+ interferes with K+ permeation. In three cells studied with high [K+]i, corrected for "leak" currents measured in permeant-free 160 Na, the conductance in 160 Rb relative to that in 160 K was 0.023 measured at Vm = -20 mV and increased to 0.034 at Vm = 80 mV. In cells perfused with K+-free pipette solutions, the ratio of the slope conductance at -100 mV in 160 Rb to that in 160 K was 0.025 ± 0.009 (mean ± SD, n = 8), assuming that 5 mM BaCl2 completely blocked the IR and can be used to estimate the leak. Thus, the relative conductance of Rb+ at large negative potentials is the same whether or not K+ is present internally. In contrast to the low conductance of Rb+, when 160 K is replaced by 160 Rb (with K+ present internally) Vm shifted on average by -20.1 mV, indicating a permeability ratio PRb/PK of 0.45 according to the Goldman-Hodgkin-Katz voltage equation.

Instantaneous Current-Voltage Relationships in Rb+ and K+

Macroscopic instantaneous I–V relationships could be determined up to a few tens of mV above Vm in K+-containing solutions. At 80 mM (Fig. 4A) and 160 mM [K+]o (data not shown), the I–V relation was approximately linear over the voltage range studied. The instantaneous I–V relation was somewhat easier to resolve in high [Rb+]o because of the effects of Rb+ on IR gating. In 80 K + 80 Rb (Fig. 4A) inward currents were greatly reduced, with a smaller but substantial reduction of outward currents. The result is that the I–V relation rectifies strongly in the outward direction. The strong outward rectification at high [Rb+]o is more obvious in Fig. 4B in which the same data are plotted at higher gain, along with the I–V relation in the same experiment in 4.5 K + 160 Rb. In other experiments rectification in 160 Rb solutions was comparable to that in 4.5 K + 160 Rb, and in 160 K was similar to that in 80 K + 80 Na. In high [Rb+]o, the slope conductance for inward currents is 10–20 times smaller than that for outward currents. This ratio may slightly underestimate the extent of rectification, because the outward currents were corrected for time-independent leak current, whereas the inward currents were not corrected (see Fig. 4, legend), and the voltage range was limited. In intermediate [Rb+]o solutions, I–V relations fell between these extremes.

K+ and Rb+ Permeation in Single IR Channels

IR currents in symmetrical K+ in the cell-attached patch configuration. Inward rectifier currents were often observed in cell-attached patches after seals were formed in preparation for whole-cell experiments, using K+-containing “internal” solutions (Table I B). Although several other types of channels were sometimes seen, IR channels have a characteristic behavior which makes their identification straightforward. One signature is that the open probability is high at negative potentials, whereas at positive potentials no outward current can be resolved. Fig. 5 illustrates the open-channel i–V relationship from an experiment in which conditions were maximized for detecting outward currents. The patch membrane was ramped rapidly from negative to positive potentials in order to increase the likelihood that the
Figure 4. Instantaneous current-voltage relationships in 80 K + 80 Na (80 K, open symbols) and in 80 K + 80 Rb (80 Rb, filled symbols) (A), and at higher gain in the same cell in 4.5 K + 160 Rb (160 Rb, open symbols) and in 80 K + 80 Rb (80 Rb, filled symbols) (B). In all parts, triangles indicate peak currents during pulses to negative potentials at which the IR is fully activated, diamonds indicate directly measured instantaneous currents following negative prepulses which maximally activated the IR conductance, and squares indicate the amplitude of a single exponential fitted to the transient outward current following a prepulse. These last data are thus effectively corrected for any time-independent current unrelated to the IR, but for small depolarizations may underestimate the IR component if there are maintained outward currents. Pipette KCH$_3$SO$_3$, 22.4°C.

Channel, open at negative potentials, would remain open during the ramp beyond $V_{rev}$. The single open-channel current was constructed by sorting and averaging sections of data in which the channel was open and sections in which the channel was closed, and subtracting the latter from the former. The slope conductance of the net open-channel current was 30 pS within ~60 mV of $V_{rev}$ and increased to 42 pS at

Figure 5. Averaged, leak-subtracted single open-channel current-voltage ($i-V$) relationship in a cell-attached patch, with KCH$_3$SO$_3$ in the pipette and Ringer's in the bath. Slope conductance at large negative potentials 42 pS. The patch membrane potential was ramped repeatedly from RP - 100 mV to RP + 160 mV at 5.08 mV/ms, sample interval 100 μs, ramp duration 50.8 ms, filter 2 kHz, 18.9°C. See text for details.
more negative potentials. In this experiment, and in all other cell-attached patch experiments, it was never possible to identify positively any instance of outward current through IR channels, either in voltage ramps or pulses. We attempted to enhance the signal-to-noise ratio by defining as “open” 50 ramp records in which the channel was open up to $V_{rev}$. The average obtained still showed no hint of outward current. In the cell-attached patch configuration, inward rectification of the IR channel appears to be absolute, within the resolution of our techniques.

**IR currents in symmetrical K\(^+\) in the outside-out patch configuration.** In contrast to their behavior in cell-attached patches, IR channels in outside-out patches excised from endothelial cells can exhibit distinct outward current. In the experiment shown in Fig. 6A, the voltage was ramped from negative to positive, and the average leak-subtracted open-channel $i-V$ relation was found to be approximately linear with a conductance of 27 pS. The IR channel remained open up to +80 mV in only one record, so the data are noisy at positive potentials. During all other ramps the channel closed, as is evident from the averaged current from 60 sweeps, which is superimposed in this figure. The behavior of the averaged single-channel current is qualitatively similar to that of whole-cell macroscopic currents (e.g. Fig. 1), with a hump of outward current which vanishes 40–50 mV positive to $V_{rev}$. The outward current hump appears to be more pronounced in single channels in excised patches than at the whole-cell level.

**Effects of [Rb\(^+\)]\(_o\) on IR currents in the outside-out patch configuration.** Single-channel IR currents in the presence of different [Rb\(^+\)]\(_o\) were compared directly using outside-out patches. Three consecutive ramp records in 160 K are superimposed in Fig. 6B (darker lines). About six IR channels were active, giving rise to the equivalent of scaled-down macroscopic IR currents. In 150 K + 10 Rb solution, the inward currents exhibited voltage-dependent block and were reduced by >50% at large negative potentials compared with 160 K (three consecutive traces are superimposed in Fig. 6B), without any obvious change in the hump of outward current. The lack of change in outward current suggests that the same number of channels was open at negative potentials as in 160 K, that is, Rb\(^+\) did not decrease the number or availability of functional IR channels. The reduction of inward current by 10 mM Rb\(^+\) is comparable to its effect on whole-cell currents (Figs. 1A and 3). Although single-channel current fluctuations are evident in Fig. 6B in 160 K, in 150 K + 10 Rb unitary transitions were smaller and barely resolvable, the largest of which corresponded roughly with a conductance of 8–12 pS. Evidently, Rb\(^+\) reduces the amplitude of single IR channel currents; if Rb\(^+\) simply reduced the number of channels open at negative potentials without affecting the unitary conductance single-channel current levels would be the same size and as distinct as in 160 K. The lower conductance could be explained if each Rb\(^+\) ion which permeates the channel transiently “blocks” the channel, i.e., impedes the flow of K\(^+\) through the channel, before passing through to the intracellular solution, with blocking kinetics too fast to permit resolution of discrete blocking events.

The number of functional IR channels in this patch declined with time. The unitary open-channel $i-V$ relationships in Fig. 6A, C, and D were recorded late in the experiment when only one or two channels remained active. When the solution bathing the outside-out patch was 80 K + 80 Rb or 160 Rb, unitary inward currents

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**SILVER ET AL. Effects of Rb\(^+\) on Inward Rectifier Gating**
FIGURE 6. IR channel currents during voltage ramps in the same outside-out membrane patch, with nominally Mg²⁺-free KCH₃SO₄/F in the pipette, and various solutions in the bath. After the patch was excised, the membrane resistance was high with < 1 pA of leak current at ±80 mV in permeant-free 160 Na.

(A) Unitary IR channel i-V relation with 160 K in the bath, after leak subtraction as described in Fig. 5. Superimposed is the leak-subtracted average current in 60 ramp records. The positive value of \( V_{m} \), +14 mV, in the presence of approximately symmetrical K⁺ concentrations presumably reflects uncorrected liquid junction potentials and/or drift. Data collected 60 min after formation of the outside-out patch, when only one IR channel was active in the patch. Ramp rate 2.9 mV/ms, filter 2 kHz, holding potential +10 mV, 18.5°C.

(B) Currents during voltage ramps in 160 K (dark lines) and in 150 K + 10 Rb (light lines). Three records in each solution are superimposed, without leak correction. Note that, unlike the other parts of this figure, there are 4–6 IR channels open at the start of each ramp. Incomplete capacity compensation probably caused the current to reverse at more negative potentials (due to a small constant vertical offset) than in the other parts of this figure in which leak correction was applied. Recorded 20 min after patch excision. Ramp rate 1.8 mV/ms, filter 2 kHz, 21.4°C.

(C) Unitary IR channel current with 80 K + 80 Rb in the bath, leak-corrected as described in the text. No distinct inward currents were seen, but outward current events were observed...
through the IR could not be detected, but outward currents could still be resolved. The net open-channel $i-V$ relations are plotted in Figs. 6, C and D. Because IR channels close with depolarization the $i-V$ relation could be determined only up to the most positive potential at which a channel remained open. Although inward unitary currents were not resolvable, it is clear that the open channel $i-V$ relations exhibited strong outward rectification at high [Rb+]o. We attempted to estimate the unitary conductance for inward currents by sorting the ramp records according to whether the IR channel in the patch was open or closed above $V_{re}$. If no opening was detectable the channels were considered "closed" in the negative range but "open" if a channel was open above $V_{re}$. In 80 K + 80 Rb the estimated conductance was 1.1 pS, or 0.037 of the ~30 pS for the unitary IR conductance in 160 K. For comparison, the macroscopic current at −90 mV in 80 K + 80 Rb is 0.046 that in 160 K (Fig. 3). The procedure just described is obviously crude, and would underestimate the true conductance if there are records in which a channel was open at negative potentials but closed before the potential was enough beyond $V_{re}$ for outward current to be detectable. In 160 Rb (Fig. 6D) the $i-V$ relation rectifies strongly outwardly, consistent with the macroscopic instantaneous $I-V$ relation in Fig. 4B. Using the relative current estimate in 160 Rb in Fig. 3 of 0.027, the scaled unitary conductance would be only 0.8 pS in 160 Rb, a value below our resolution at the single channel level. In summary, in all solutions studied the rectification of single-channel currents qualitatively resembles the rectification of the macroscopic IR currents.

**Effects of [Rb+]o on the Voltage-dependence of IR Gating**

*Steady state voltage-dependence.* The left inset in Fig. 7 illustrates the measurement of the steady state activation curve of the IR in 160 K (cf Silver and DeCoursey, 1990). Prepulses to a range of potentials were applied, which were long enough for the gating process to equilibrate, followed by a test pulse, in this case to −19 mV. The instantaneous current, obtained by extrapolating back to the start of the test pulse a single exponential curve fitted by eye to the test current, reflects the fraction of channels activated at the end of the prepulse. For large negative prepulses which fully activate the IR, the instantaneous test current is large, whereas after large positive prepulses which close IR channels, the test current is small initially and then rises to the same steady-state value. The instantaneous test current, plotted in Fig. 7 (■) after normalization to its fitted minimum and maximum values, reflects the which tended to close at positive potentials. The voltage was ramped from −100 to +80 mV, but no channels stayed open above ~65 mV. The slope conductance by least-squares of the data from −100 to 0 mV is 1.1 pS, a value which should be regarded as very approximate; the slope conductance of the current positive to +37 mV is 26 pS. Recorded 53 min after patch excision, ramp speed 1.6 mV/ms, data sampled at 2 kHz and later Gaussian-filtered at 1 kHz, 18.3°C.

(D) Unitary IR channel current with 160 Rb in the bath. As in C, only outward currents were resolved and the data are leak-corrected. No channels remained open above ~50 mV. The slope conductance at positive potentials, indicated by the line, is 31 pS; that for inward currents is too small to calculate meaningfully. Recorded 43 min after patch excision, ramp speed 0.89 mV/ms, filter 1 kHz, holding potential 0 mV, 18.4°C.
voltage-dependence at steady-state of the gating mechanism of the IR. The original data were fitted to a Boltzmann function:

\[
I_0(V) = \frac{(I_{0,\text{max}} - I_{0,\text{min}})}{1 + \exp(V - V_{1/2}/k)} + I_{0,\text{min}}
\]

where \(I_0\) is the extrapolated instantaneous current at potential \(V\), \(I_{0,\text{max}}\) is the fitted maximum value of \(I_0\), \(I_{0,\text{min}}\) is the fitted minimum value of \(I_0\), \(V_{1/2}\) is the potential at which the IR conductance is half-activated, and \(k\) is a slope factor. Measured in 160 K (\(\bullet\)), \(V_{1/2}\) was 6.5 mV negative to \(V_{\text{rev}}\). Note that the voltage axis is labelled relative to the reversal potential observed in each solution. The voltage dependence of gating is quite steep, with a slope factor, \(k\) of 3.7 mV. Mean values for these parameters in many cells are given in Table II. While the focus of the present study was on effects of \(Rb^+\), we interspersed several measurements of the same cells in \(K^+\)-containing

\[\text{FIGURE 7. Steady state voltage-dependence of IR gating in a cell in 160 K (\(\bullet\)) and in 160 Rb (\(\bullet\)). The amplitude of the activating (K\(^+\)) or deactivating (Rb\(^+\)) transients during the test pulses are plotted as a function of the prepulse potential. Data are normalized according to the fitted minimum and maximum for each data set, and the voltages on the abscissae are relative to the observed } V_{\text{rev}} \text{ in each solution (} -3.2 \text{ mV in 160 K and } -27.1 \text{ mV in 160 Rb). Curves show fits to a Boltzmann function (Eq. 1), with midpoint } V_{1/2} \text{ and slope factor } k \text{ in 160 Rb } -16.5 \text{ mV (absolute potential) and 6.46 mV, respectively, and in 160 K, } -9.7 \text{ and } 3.72 \text{ mV. Pipette } \text{KCH}_3\text{SO}_4. (Left) Measurement of activation curve in 160 K. Prepulses 20 ms long were applied from a holding potential of +1 mV, which was also the zero-current potential, followed by a test pulse to } -19 \text{ mV. Illustrated prepulses are for } -29 \text{ to } +6 \text{ mV in 5 mV increments. Filter 2 kHz, sample interval 0.1 ms, 20.7°C. (Right) Measurement of activation curve in 160 Rb. Prepulses were 140 ms long, to allow full equilibration of gating, followed by a test pulse to } +2 \text{ mV. Illustrated prepulses are to } -43 \text{ to } +7 \text{ mV in 10 mV increments. Arrow indicates zero current. Filter 2 kHz, sample interval 0.2 ms, 20.5°C.}\]
solutions. As in our previous study (Silver and DeCoursey, 1990), $V_{1/2}$ was always negative to $V_{re}$ in K+-containing, Rb+-free solutions, and within the error of the measurement was displaced from $V_{re}$ by the same amount at all [K+]o studied. Combining data at all [K+]o, $V_{1/2}$ was 6.9 mV negative to $V_{re}$ in the previous study (Table II, top line) and 6.0 mV in the present study (Table II, third line). The slope factor, $k$, was also similar in both data sets, 4.3 and 4.5 mV, respectively.

### Table II

| [K+]o | [Rb+]o | $V_{1/2}$ | n | $k$ | n |
|-------|--------|-----------|---|-----|---|
| all*  | 0      | −6.9 ± 0.7| 39| 4.3 ± 0.1 | 43 |
| 160   | 0      | −4.2 ± 1.4| 5 | 4.4 ± 0.4 | 4  |
| all   | 0      | −6.0 ± 0.9| 25| 4.53 ± 0.19| 25 |
| 160   | 0      | −3.8 ± 2.0| 3 | 4.17 ± 0.80| 3  |
| 160   | 4.5    | −6.4 ± 1.5| 2 | 4.49 ± 0.68| 2  |
| 150   | 10     | −4.7 ± 1.4| 5 | 4.50 ± 0.30| 5  |
| 140   | 20     | −5.8 ± 0.7| 11| 4.68 ± 0.40| 11 |
| 120   | 40     | −3.8 ± 1.1| 13| 4.92 ± 0.16| 13 |
| 80    | 80     | 11.5 ± 1.4| 12| 5.32 ± 0.10| 12 |
| 40    | 120    | 16.8 ± 1.2| 4 | 5.68 ± 0.43| 4  |
| 20    | 140    | 10.0 ± 2.1| 5 | 5.87 ± 0.17| 7  |
| 10    | 150    | 18.3 ± 2.6| 6 | 6.19 ± 0.35| 7  |
| 4.5  | 160    | 16.0 ± 0.6| 19| 6.48 ± 0.19| 19 |
| 0    | 160    | 17.3 ± 2.2| 6 | 6.16 ± 0.27| 6  |

*Data from a previous study (Silver and DeCoursey, 1990) including only experiments with Mg²⁺-free pipette solutions, as in the present study. Because both parameters were found to be identical for all [K+]o, values from [K+]o ranging 4.5–160 mM are combined (all). Parameters defined in Eq. 1. Superscripted symbol pairs indicate significance levels in t tests vs control data in the top and third rows, respectively: * = $p < 0.05$; ** = $p < 0.01$ for both controls; † = $p < 0.01$; ‡ = $p > 0.05$). Means ± SE are given for $n = $ number of experiments.

Determination of the activation curve of the IR using the variable prepulse method in solutions containing high [Rb⁺]o differed from that in solutions containing only K⁺ in that the optimal test potential was positive to $V_{re}$. The right-hand inset in Fig. 7 illustrates the measurement in 160 Rb in which the amplitude of the deactivating outward current was evaluated. Following negative prepulses the outward "tail current" is large, while after large depolarizing prepulses the amplitude of the test current is small. The extrapolated instantaneous current during the test pulse in 160 Rb is plotted in Fig. 7 (●). The midpoint, $V_{1/2}$, was 10.6 mV positive to $V_{re}$. When K⁺ was replaced by Rb⁺ in this experiment, $V_{re}$ changed by −24 mV while $V_{1/2}$ shifted only −7 mV. Table II summarizes the data at various [Rb⁺]o. As the Rb⁺ mole-fraction increased and $V_{re}$ shifted to more negative potentials, $V_{1/2}$ changed more gradually, with the result that $V_{1/2}$ was well positive to $V_{re}$ in 160 Rb. As shown in Table II, this shift was significant at all [Rb⁺]o > 40 mM.

The voltage-dependence of IR gating was less steep in 160 Rb, with a slope factor of 6.5 mV in the cell in Fig. 7 compared with 3.7 mV in 160 K. Table II shows that, like $V_{1/2}$, $k$ was significantly increased at [Rb⁺]o > 40 mM.
Effects of [Rb+]o on macroscopic IR currents. Rb+ greatly alters the appearance of IR current relaxations. The time-dependent activation of inward currents which is a hallmark of macroscopic IR currents in K+-containing solutions is harder to resolve in high [Rb+]o. The cell in Fig. 8A was held at -3 mV, well positive to $V_{re}$ at -26 mV. The current during the family of voltage steps exhibits small time-dependent relaxations. For the voltage steps negative to the holding potential, activating currents can be seen, which are outward at -8 and -18 mV and inward at more negative potentials because $V_{re}$ (indicated by an arrow) was -26 mV. Activation is slower in 160 Rb than in K+-containing solutions, and in both solutions becomes more rapid at more negative potentials. The decay of current during the most positive pulse (+2 mV) reflects deactivation of the IR conductance which was partially activated even in this cell held 23 mV positive to $V_{re}$.

Fig. 8B illustrates the kinetics of IR current deactivation in 160 Rb in the same cell as in A. A hyperpolarizing prepulse activated the IR conductance, and then a range of test pulses were applied. Deactivation first becomes evident positive to $V_{re}$, in contrast with K+-containing solutions in which the IR conductance is already
largely deactivated at $V_{rev}$. The time constant of deactivation quantified by fitting the transient outward current with a single exponential becomes faster rapidly with further depolarization. Also observed in some experiments were smaller slowly-decaying outward currents, which might be due to the IR or to some other conductance.

$V_{rev}$ in high $[Rb^+]_o$. A critical measurement in this study is that of $V_{rev}$, the reversal potential of the IR conductance. In high $[K^+]_o$ solutions in which $V_{rev}$ is near 0 mV and the IR conductance is large compared with other conductances, the zero current potential corresponds closely with $V_{rev}$. However, in high $[Rb^+]_o$, where the IR conductance is greatly reduced and $V_{rev}$ is more negative, leak or other extraneous conductances may introduce errors into the measurement. To determine $V_{rev}$ as accurately as possible, we used the procedure illustrated in Fig. 8 C, in which a depolarizing prepulse which closed the IR channels (off-scale) was followed by a range of test pulses, and the current relaxation was observed. From the direction of the increase in IR current as channels open, $V_{rev}$ can be determined. In the experiment in Fig. 8 C in 80 K + 80 Rb, the activating currents reversed at -13 mV. For this illustration 3 mV increments are shown, but for final determination of $V_{rev}$ voltage increments of 1 mV were used. It is noteworthy that in Fig. 8 C distinct activation of outward current is evident, that is, there is significant steady state outward IR current in high $[Rb^+]_o$ solutions.

One of the classical behaviors of the IR, illustrated in Fig. 9 A, is that the voltage-dependence of activation shifts with $V_{rev}$ when $[K^+]_o$ is varied, provided that $[K^+]_i$ is constant. At all $[K^+]_o$, $V_{1/2}$ was a few mV negative to $V_{eff}$. The same parameters are plotted in Fig. 9 B for various mole-fractions of $Rb^+$ and $K^+$. While $V_{rev}$ was 20 mV more negative in 160 mM $Rb^+$ than in 160 mM $K^+$, $V_{1/2}$ barely changed. Thus, the two parameters became dissociated in the presence of $Rb^+$, and at $[Rb^+]_o > 80$ mM $V_{1/2}$ was positive to $V_{rev}$.

Effects of $[Rb^+]_o$ on the Gating Kinetics of the IR

$Rb^+$ slows IR gating. Macroscopic IR current relaxations, such as those in Fig. 8, were fitted with single exponentials over a range of $Rb^+$ mole fractions. Although small deviations from single exponential behavior were observed, no consistent pattern emerged. As was found previously for solutions containing various $[K^+]_o$, the $\tau-V$ relationship is bell-shaped, with a maximum near the midpoint of the activation curve. The relationship between $1/\tau$ and voltage is shown in Fig. 10 for a cell in 20 K + 140 Rb solution. Within the accuracy of the data, IR gating in all $[Rb^+]_o$ and $[K^+]_o$ mixtures could be expressed in the same form as in $K^+$-containing solutions, as a simple two-state system in which the opening rate, $\alpha$, and the closing rate, $\beta$, are both exponentially dependent on voltage. The solid curve in Fig. 10 shows the fitted $\tau-V$ relationship, while the dotted lines show the components $\alpha(V)$ and $\beta(V)$, where

$$\alpha(V) = \tilde{\alpha} \exp \left[ \frac{(V_{1/2} - V)}{k_a} \right]$$

and

$$\beta(V) = \tilde{\beta} \exp \left[ \frac{(V - V_{1/2})}{k_B} \right].$$

The experimental values for $\alpha$ and $\beta$ (shown as + and ×, respectively) were
Figure 9. Comparison of the reversal potential, $V_{\text{rev}}$, and midpoint of activation of the IR, $V_{1/2}$, when $[K^+]_o$ was increased by substituting it for Na$^+$ (A) or Rb$^+$ (B). All values have been corrected for liquid junction potentials, and are plotted as mean ± 1 SE.

(A) The sum of $[K^+]_o$ and $[Na^+]_o$ for all solutions is 160 mM, except for Ringer's (Table I). Data from cells studied also in Rb$^+$ solutions were pooled with data from an earlier study (Silver and DeCoursey, 1990). Numbers of experiments, for increasing $[K^+]_o$, are: $V_{\text{rev}}$: 23, 6, 5, 8, 7; $V_{1/2}$: 25, 6, 5, 14, 8, 7.

(B) The sum of $[K^+]_o$ and $[Rb^+]_o$ is 160 mM, except for 4.5 K + 160 Rb and 160 K + 4.5 Rb (second data points from either end). Note that the ordinates encompass a smaller voltage range in B. Numbers of experiments, for decreasing $[Rb^+]_o$, starting with 160 mM $[Rb^+]_o$ are: $V_{\text{rev}}$: 7, 21, 6, 5, 15, 10, 4, 2, 7; $V_{1/2}$: 6, 19, 7, 7, 4, 12, 15, 10, 4, 2, 7). The dependence of $V_{\text{rev}}$ on $[K^+]_o$ and $[Rb^+]_o$ has an apparent minimum in the region of high $[Rb^+]_o$ and low $[K^+]_o$. Similar effects were noted by Hagiwara and Takahashi (1974). Based on $P_{Rb}$ measured in 160 Rb, the Goldman-Hodgkin-Katz voltage equation predicts a monotonic relationship between Rb$^+$ mole-fraction and $V_{\text{rev}}$, with the calculated value for $V_{\text{rev}}$ positive to the observed values in 80 mM and higher $[Rb^+]_o$. In six of nine cells in which $V_{\text{rev}}$ was measured in 160 mM Rb$^+$ and at least one other solution in this range, $V_{\text{rev}}$ was (anomalously) more negative at the lower $[Rb^+]_o$. 
calculated from \( \tau \) and \( P_{\text{open}} \), the open probability derived from the fitted parameters of the activation curve, given that \( \tau = 1/(\alpha + \beta) \) and \( P_{\text{open}} = \alpha/(\alpha + \beta) \). We used \( V_{1/2} \) as the reference potential for the rate constants, that is, \( \alpha \) and \( \beta \) are defined at \( V_{1/2} \) in Eqs. 2 and 3. This procedure was adopted, as opposed to fixing the rate constants at their values at 0 mV for example, because when defined in this way their values were found previously to be independent of \([K^+]_o\) (Silver and DeCoursey, 1990). Table III gives the values for the rate constants and their slope factors, \( k_\alpha \) and \( k_\beta \), for the whole range of \([Rb^+]_o\) studied. The separation of the two rate constants, however, is not straightforward, because due to the nature of IR gating their extraction from the data is model-dependent (see Discussion). As defined here, \( Rb^+ \) reduced \( \alpha \) by a factor of 8, and \( \beta \) by a factor of 4. The voltage dependence of the opening rate, reflected in \( k_\alpha \), was significantly less steep in high \([Rb^+]_o\), whereas \( k_\beta \) was not altered.

**Effect of \( Rb^+ \) on deactivation kinetics in outside-out patches.** The slowing of deactivation by \( Rb^+ \) was observable in isolated patches as well as in macroscopic currents. It is difficult to resolve IR gating at the single-channel level even in \( K^+ \) solutions (Kurachi, 1985; Burton and Hutter, 1990), because gating occurs near \( V_{re} \), where unitary currents are small. Here we consider only the strongly voltage- and \([K^+]_o\)-dependent gating which corresponds with macroscopic IR activation/deactivation kinetics, and not the voltage-dependent inactivation of some IR channels at large negative potentials (Sakmann and Trube, 1984; Rae, 1986; Josephson and Brown, 1986; McKinney and Gallin, 1988). We could not resolve convincingly the activation kinetics of the IR at the single-channel level in high \([Rb^+]_o\). However, it was possible to resolve deactivation kinetics in isolated patches. The outside-out patch illustrated in Fig. 11 had many IR channels, with a steady state \( I-V \) relation measured in 160 K shown in Fig. 11 C. Deactivation was studied by applying depolarizing pulses from \(-20\) mV, where the IR channels were maximally activated, to more positive potentials...
at which the channels closed. Outward current transients composed of ensembles of many capacity-corrected records at +30 mV are shown in Fig. 11 A measured in 160 K, and in Fig. 11 B in 160 Rb. Both transients were well-fitted by a single-exponential decay, and the time constant, $\tau$, was 3.1 ms in 160 K and 6.7 ms in Rb+. Deactivation was slower in 160 Rb up to +60 mV at which point it was not clearly distinguishable from the capacity transient.

The voltage-dependence of IR gating kinetics in whole-cell experiments in 160 K and 160 Rb are summarized in Fig. 12. The curves show $1/\tau$ calculated from mean values for the rate constants, slope factors, and $V_{1/2}$ (indicated as tic marks). The

| [K+]o | [Rb+]o | $\alpha$ | $k_a$ | $\beta$ | $k_b$ | $n$ |
|-------|--------|--------|-------|--------|-------|-----|
| mM    | mV     | s⁻¹    | mV    | s⁻¹    | mV    |     |
| all*  | 203 ± 14 | 17.7 ± 0.7 | 97. ± 6 | 9.0 ± 0.3 | 34    |
| all   | 0      | 191 ± 13.5 | 19.0 ± 0.9 | 93. ± 15 | 7.5 ± 0.4 | 20, 17 |
| 160   | 0      | 211      | 16.9    | 61      | 5.5    | 2       |
| 160   | 4.5    | 205      | 22.3    | 138     | 7.8    | 1       |
| 150   | 10     | 195. ± 7 | 24.4 ± 1.3 | 87. ± 14 | 7.4 ± 0.9 | 4       |
| 140   | 20     | 164. ± 12 | 28.1 ± 2.3 | 74. ± 13 | 7.5 ± 0.5 | 10, 5   |
| 120   | 40     | 1104. ± 14 | 29.3 ± 1.2 | 64.3 ± 6.5 | 6.2 ± 0.4 | 13      |
| 80    | 80     | 151. ± 4.0 | 28.3 ± 1.5 | 61.4 ± 2.7 | 7.5 ± 0.4 | 14      |
| 40    | 120    | 222.8 ± 3.9 | 322.7 ± 2.5 | 340.6 ± 11 | 9.2 ± 0.7 | 3, 5    |
| 20    | 140    | 1218. ± 3.9 | 28.1 ± 0.8 | 242 ± 2.6 | 3.9 ± 0.6 | 7       |
| 10    | 150    | 226.3 ± 3.4 | 261 ± 0.5 | 241 ± 3.0 | 3.9 ± 0.5 | 6       |
| 4.5   | 160    | 341. ± 2.4 | 255 ± 1.0 | 316 ± 2.6 | 9.4 ± 0.3 | 20      |
| 0     | 160    | 57.5 ± 3.7 | 254. ± 2.0 | 235 ± 3.0 | 10.0 ± 0.8 | 7       |

Parameters are defined as Eqs. 2 and 3. *Data from Silver and DeCoursey (1990) including only experiments with Mg2+-free pipette solutions, as in the present study. Because all four kinetic parameters were found to be identical for all [K+]o values for [K+]o ranging 4.5-160 mM are combined. Symbol pairs indicate significance of differences vs control data in the first and second rows, respectively ($p < 0.05$; $\dagger$ = $p < 0.01$; $\ddagger$ = $p < 0.01$ for both controls; $\n$ = $p > 0.05$). Means ± SE are given for $n$ = number of experiments; when two values for $n$ are given the first value refers to $\alpha$ and $k_a$ and the second indicates $\beta$ and $k_b$. Parameters related to activation ($\alpha$, $k_a$) are more reliable than those for deactivation ($\beta$, $k_b$) at low [Rb+]o, whereas the reverse is true at high [Rb+]o. Deactivation was particularly poorly-resolved at low [Rb+]o, and $1/\tau$ sometimes exhibited sublinear voltage-dependence at positive potentials, so $k_b$ was poorly defined in these solutions. In most cases, the deactivation $\tau$ was determined directly from current transients (e.g., Fig. 11 B), whereas in the earlier study, deactivation was generally determined from envelopes, which were considered more reliable in K+-containing (i.e., Rb+-free) solutions.

Voltage-axis is labeled in terms of absolute potential. Plotted in this way, deactivation (positive limb) is slowed in 160 Rb about twofold, in agreement with the single-channel deactivation ensembles in Fig. 11. Activation (negative limb) is slowed in 160 Rb by more than an order of magnitude, with the slowing effect increasing at more negative potentials. If the data are expressed relative to $V_{1/2}$, then the slowing of deactivation by Rb+ is increased slightly, while that of activation is less; corresponding with the values given in Table III. Quantifying IR gating with reference to $V_{res}$, because classically IR gating has been expressed as a function of $V-V_{res}$, Rb+ slows both $\alpha$ and $\beta$ by about the same amount.
Figure 11. Ensemble IR channel tail currents in an excised outside-out patch in 160 K (A) and in 160 Rb (B) with KCH3SO3 + 2 Mg in the pipette. There were many IR channels in this patch. For both measurements, the patch was held at -20 mV (at which potential Pop/ was maximal) and repeated pulses to +30 mV were applied. To subtract capacity currents, the membrane was pulsed to -10 mV at which potential the gK was still maximally activated, and the averaged data were then subtracted from the raw records at +30 mV after appropriate scaling. Superimposed on the data points are smooth curves showing the fit (by eye) to a single exponential, with r = 6.7 ms in 160 Rb and 3.1 ms in 160 K. The amplitude of each transient was ~ 15 pA. In A and B, respectively, sample interval was 50 μs and 60 μs, 22.6°C and 22.0°C, 50 and 30 pulses averaged, filter 5 kHz in both. (C) Steady state I-V relationship of IR channels in the same outside-out patch with 160 K in the bath. Currents were measured at the end of 30 ms test pulses. Leak subtraction was based on scaled average currents during four repeated pulses, each 1/6 the amplitude of the test pulses, starting from a holding potential of +30 mV at which the channels appeared to be closed.

Figure 12. Voltage dependence of 1/τ in 160 K (upper curve) and in 160 Rb (lower curve), calculated from mean parameter values for rate constants and slope factors. The rate constants given in Table III (top and bottom rows) are expressed relative to V1/2, but this figure is plotted in terms of absolute potential; mean V1/2 values in 160 K and 160 Rb are indicated on the x-axis by tic marks at -7.7 and -16.3 mV, respectively.
The mole-fraction dependence of the slowing of the rate constants by Rb\(^+\) is plotted in Fig. 13. The effect of Rb\(^+\) on the opening rate, \(\alpha\) (●), is greater than that on \(\beta\) (□), at least as these parameters were defined. Both parameters were reduced by small additions of Rb\(^+\), approaching their limiting values in 160 Rb after ~ 50% of the K\(^+\) was replaced by Rb\(^+\).

**Figure 13.** Effect of [Rb\(^+\)]\(_o\) on the rate constants of IR gating, \(\overline{\alpha}\) (●) and \(\overline{\beta}\) (□). Data from Table III are plotted ±1 SE. The data at 0 mM [Rb\(^+\)]\(_o\) include all [K\(^+\)]\(_o\) (4.5 to 160 mM) from Silver and DeCoursey (1990); for clarity new control data, also including all [K\(^+\)]\(_o\), are plotted arbitrarily at 3 mM [Rb\(^+\)]\(_o\).

**DISCUSSION**

**Rb\(^+\) Permeation Through IR Channels**

Although Rb\(^+\) has long been known to interfere with K\(^+\) permeation through IR channels (Bolingbroke, Harris, and Sjodin, 1961; Adrian, 1964), it is not completely impermeant. Replacement of external K\(^+\) with Rb\(^+\) greatly reduces but does not abolish inward rectification, which can be further reduced by formaldehyde (Hutter and Williams, 1979), TEA\(^+\) (Standen and Stanfield, 1980), or Ba\(^2+\) (this study). The relative permeability of Rb\(^+\) in IR channels based on the Goldman-Hodgkin-Katz voltage equation is 0.3–0.4 in starfish egg cells, and the relative limiting slope conductance is 0.05 (Hagiwara and Takahashi, 1974). These values agree well with those obtained in the present study for macroscopic currents, with \(P_{\text{Rb}}/P_{\text{K}} = 0.45\) and the relative Rb\(^+\) conductance 0.034 at \(V_{\text{rev}} \approx 80\) mV. Our results reconfirm that IR channels are more selective than most other voltage-dependent K\(^+\) channels, such as various delayed rectifiers, whose \(P_{\text{Rb}}/P_{\text{K}}\) is > 0.7 (references in Hille, 1992) and \(g_{\text{Rb}}/g_{\text{K}}\) under biionic conditions is ~ 0.5 (references in Shapiro and DeCoursey,
Consistent with the macroscopic instantaneous \( I-V \) relation, single channel rectification was strongly outward in outside-out patches with high \([Rb^+]_o\), that is, external \( Rb^+ \) greatly reduces inward currents but permits relatively large outward \( K^+ \) current through the IR. Lower \( Rb^+ \) mole fractions produce voltage-dependent reduction of the single-channel current consistent with effects on macroscopic current-voltage relationships.

In mixtures of \( Rb^+ \) and \( K^+ \) the inward IR current amplitude was reduced by small additions of \( Rb^+ \) more than expected if \( Rb^+ \) were simply an inert impermeant ion. \( Rb^+ \) in the external solution, while measurably permeant, evidently inhibits \( K^+ \) permeation through IR channels. In cells containing \( K^+ \), block by \( Rb^+ \) is enhanced by hyperpolarization at small negative potentials, reaching a limiting level of block which is relieved by extreme hyperpolarization. Similar voltage-dependent block by \( Rb^+ \) of IR channels in other preparations has been taken as evidence that \( Rb^+ \) blocks at a site within the pore (Hagiwara and Takahashi, 1974; Standen and Stanfield, 1980). The slope factor derived from the voltage dependence of the relative conductance in \( Rb^+ \)-containing solutions, 10–16 mV, corresponds to an electrical distance, \( \delta \), in a one-site channel model (Woodhull, 1973) of 1.6–2.5, which suggests that the IR channel is multiply occupied by ions and that the block site is within the channel. When \( K^+ \)-free pipette solutions were used, block at large negative potentials was similar in magnitude and voltage dependence to that with normal \([K^+]_i\). However, at less negative potentials block increased with depolarization when \([K^+]_i\) was low, but decreased with depolarization when \([K^+]_i\) was normal. Internal \( K^+ \) thus appears to relieve block by external \( Rb^+ \) at potentials approaching \( V_{rev} \) even though the net current is inward. Similarly, Standen and Stanfield (1980) found that increasing \([K^+]_i\) reduced the affinity of \( Rb^+ \) for its block site in the IR of skeletal muscle. Voltage-dependent block by \( Rb^+ \) may be more accurately described as voltage-dependent interaction between external \( Rb^+ \) and internal \( K^+ \).

No time-dependence of block of macroscopic currents was detected, furthermore, the reduction of unitary current by \( Rb^+ \) occurred without detectable blocking events. Evidently \( Rb^+ \) block occurs at frequencies above the bandwidth of our recording. If the block site in the channel is occupied during permeation, then the minimum block rate can be estimated from the \( Rb^+ \) conductance in 160 Rb, using the scheme of Standen and Stanfield (1980):

\[
\text{Rb}_o^+ + R \xrightleftharpoons[k_{i[Rb^+]}]{k_{-i[Rb^+],i}} \text{Rb}^+R \xrightarrow[k_{i[Rb^+]}]{k_{-i[Rb^+],i}} \text{Rb}_i^+ + R,
\]

where \( \text{Rb}_o^+ \) and \( \text{Rb}_i^+ \) represent a \( Rb^+ \) ion in the external or internal solution, \( R \) is the receptor site at which block occurs, and \( k_{i}, k_{2}, k_{-1}, \) and \( k_{-2} \) are rate constants. Because \([Rb^+]_i\) is assumed to be negligible in these experiments, \( k_{-i[Rb^+],i} \) is neglected. The unitary conductance at \( V_{rev} \) = 90 mV, scaled according to Fig. 3, gives a net \( Rb^+ \) flux of \( 4.7 \times 10^5 \) ions/s. If we take this as the rate \( Rb^+ \) exits the block site to the interior, \( k_{2} \), then from the half-blocking concentration of \( ~7 \text{mM} \) \([Rb^+]_o\) measured at \(-90 \text{mV} \) (Fig. 3), the forward block rate constant, \( k_{1} \), is \( 6.7 \times 10^7/M \cdot s \). This value represents the minimum rate of \( Rb^+ \) permeation/blocking events, which could be even higher if \( Rb^+ \) sometimes blocks a channel without permeating, if \( Rb^+ \) re-enters the channel from the inside, or if the block site is not continuously occupied in 160 Rb. That \( Rb^+ \)
block is only weakly voltage-dependent at large negative potentials indicates that $k_1$ and $k_2$ either are both practically voltage-independent or have similar voltage-dependencies. For this analysis we assume the former possibility, and ascribe most of the voltage-dependence of block at potentials near $V_{1/2}$ to the rate that $\text{Rb}^+$ exits the block site to the external solution, $k_{-1}$. At large negative potentials $k_{-1}$ is thus assumed to be negligible, so that $K_i = (k_{-1} + k_2)/k_1$ simplifies to $K_i = k_2/k_1$. The block rate constant for $\text{Rb}^+$ here, $6.7 \times 10^7/M/s$, is much higher than that reported by Matsuda, Matsuura, and Noma (1989), $1.75 \times 10^6/M/s$, for block of $I_{K1}$ channels by $\text{Rb}^+$ at large negative potentials at 25°C. In that study, low concentrations of $\text{Rb}^+$ were used, 20–100 $\mu$M, in order to resolve single blocking events, which usually were all-or-none but sometimes produced subconductance levels. The higher block rate constant for endothelial cell IR channels and the higher $[\text{Rb}^+]_o$ used in the present study probably explain why we could not resolve discrete blocking events.

In contrast with the low $\text{Rb}^+$ conductance of the IR measured with high $[\text{K}^+]_i$ here and in other tissues (Hagiwara and Takahashi, 1974; Hutter and Williams, 1979; Standen and Stanfield, 1980), Mitra and Morad (1991) observed inward $\text{Rb}^+$ currents 51% as large as $\text{K}^+$ currents through the $I_{K1}$ in cardiac myocytes studied with permeant ion-free pipette solutions. However, IR channels in endothelial cells exhibited the same low $\text{Rb}^+$ conductance of $\sim 3\%$ the $\text{K}^+$ conductance whether the cell was dialyzed with $\text{K}^+$ or impermeant NMG$^+$ pipette solutions. Evidently, the permeation properties of $I_{K1}$ channels differ from those of IR channels of endothelial cells.

**Effects of $[\text{Rb}^+]_o$ on the voltage-dependence and kinetics of IR gating.** A major result of this study is that external $\text{Rb}^+$ shares with external $\text{K}^+$ the ability to establish the voltage-dependence of IR gating. Because of the well-known block of IR currents by $\text{Rb}^+$, we were at first surprised that it was possible to resolve IR gating at high $[\text{Rb}^+]_o$. As the $\text{Rb}^+$ mole-fraction increases and $V_{rev}$ shifts to more negative potentials, $V_{1/2}$ changes more gradually, with the result that $V_{1/2}$ is well positive to $V_{rev}$ in 160 $\text{Rb}^+$. The practical result is that in high $[\text{Rb}^+]_o$, most of the channels are open at $V_{rev}$ while in high $[\text{K}^+]_o$, most of the channels are closed at $V_{rev}$. This fact, combined with the slowing effect of $\text{Rb}^+$ on IR gating and the strong outward rectification when $[\text{Rb}^+]_o$ is large (Figs. 4 and 6), facilitated measuring the voltage-dependence of macroscopic IR gating in high $[\text{Rb}^+]_o$. The variable prepulse technique requires clear resolution of channel gating during the test pulse. In both $\text{Rb}^+$ and $\text{K}^+$ solutions, gating kinetics are slowest and therefore easiest to resolve near the midpoint of the activation curve; which in $\text{K}^+$-containing solutions is just negative to $V_{rev}$ (e.g., Hagiwara et al., 1976; Leech and Stanfield, 1981; Tourneur et al., 1987; Cohen et al., 1989; Silver and DeCoursey, 1990), and in high $[\text{Rb}^+]_o$ is just positive to $V_{rev}$.

Because the voltage-dependence of IR gating depends strongly on $[\text{K}^+]_o$, we assume that the effects of $\text{Rb}^+$ are ascribable to binding of $\text{Rb}^+$ at the same site(s) at which $\text{K}^+$ affects gating. This site might be within the pore or at some external location remote from the permeation pathway. In cardiac myocytes external $\text{Cs}^+$ appears to share the ability of $\text{K}^+$ and $\text{Rb}^+$ in being able to activate $I_{K1}$ channels (Mitra and Morad, 1991; Carmeliet, 1992). That external $\text{Rb}^+$ and $\text{K}^+$ strongly influence channel opening is consistent with the idea that the relevant gating sites are accessible from the external solution while the channel is closed. Spalding, Swift,
Senyk, and Horowicz (1982) reached similar conclusions from K\(^+\) efflux studies in skeletal muscle. They found that in addition to its ability to inhibit K\(^+\) efflux, presumably through IR channels, Rb\(^+\) can also activate the IR like K\(^+\) does by binding to externally accessible sites.

Both Rb\(^+\) and Cs\(^+\) slow I\(_{K1}\) activation measured at a fixed potential in cardiac myocytes (Tourneur et al., 1987; Mitra and Morad, 1991). Rb\(^+\) slows gating of the IR in endothelial cells as well, but evaluating whether opening or closing kinetics are altered is not straightforward. Because the IR has a voltage-dependence which depends strongly on external permeant ion concentrations, its conductance classically has been expressed as a function of \(V - V_{\text{rev}}\) (e.g., Hagiwara and Takahashi, 1974). The entire \(\tau-V\) relationship (including both activation and deactivation) can be superimposed at any [K\(^+\)]\(_o\) by shifting the data to superimpose \(V_{1/2}\), and consequently we defined the rate constants with reference to that potential previously (Silver and DeCoursey, 1990) and in the present study (Table III). In contrast with the parallel shift of \(V_{\text{rev}}\) and \(V_{1/2}\) when [K\(^+\)]\(_o\) is changed, when [Rb\(^+\)]\(_o\) is increased these parameters become dissociated. Thus the choice of \(V_{1/2}\) (or \(V_{\text{rev}}\)) as a reference potential for extracting kinetic parameters becomes arbitrary. Referred to \(V_{1/2}\) (Table III) \(\alpha\) was slowed (eightfold) more than \(\beta\) (fourfold) by Rb\(^+\); referred to \(V_{\text{rev}}\), both \(\alpha\) and \(\beta\) were slowed approximately equally; and expressed in terms of absolute potential (Fig. 12) the effect of Rb\(^+\) was predominantly on \(\alpha(V)\) (>10-fold slowing) with only a twofold decrease in \(\beta(V)\). If the voltage dependence of gating were set by a current-polarity-dependent mechanism, for example, then using \(V_{\text{rev}}\) as a reference point would be reasonable. Nevertheless, whether the rate constants are defined at \(V_{\text{rev}}\), \(V_{1/2}\), or at absolute potentials, both \(\alpha\) and \(\beta\) are slowed by Rb\(^+\).

Rb\(^+\) also decreased the steepness of the voltage-dependence of steady state IR gating (\(k\) in Eq. 1 and Table II). The effective valence, \(z^\theta\), decreased from 5.8 (based on the weighted average value of \(k\) from previous and new data for all [K\(^+\)]\(_o\) to 4.0 (combining data from the three highest [Rb\(^+\)]\(_o\), which were indistinguishable from each other). Decomposition of the voltage-dependence of gating into that of channel opening (\(k_\alpha\)) and channel closing (\(k_\beta\)) reveals that Rb\(^+\) mainly affects the voltagesensitivity of opening (42% increase in \(k_\alpha\), Table III) without affecting that of closing. The \(z^\theta\) derived from \(k_\alpha\) was decreased from 1.4 in K\(^+\) solutions to 1.0 at high [Rb\(^+\)]\(_o\). The \(z^\theta\) derived from the steepness of the voltage-dependence of deactivation (\(k_\beta\)) in the present study was much higher than that of activation in both K\(^+\) and high [Rb\(^+\)]\(_o\) solutions, 3.0 and 2.7, respectively. Because the sum of \(z^\theta\) derived from \(k_\alpha\) and \(k_\beta\) is somewhat less than that from the steady state activation curve (4.4 vs 5.8, respectively, for K\(^+\) solutions and 3.7 vs 4.0 for high [Rb\(^+\)]\(_o\)), the voltage-dependence of at least one of the slope factors is likely underestimated, or the two-state model is inadequate.

Is a blocking particle model indicated? The possibility that the voltage-dependence of the IR might be due to an intracellular cationic blocking molecule has been considered by numerous investigators (Armstrong, 1969; Hagiwara and Takahashi, 1974; Hille and Schwarz, 1978; Vandenberg, 1987; Matsuda et al., 1987). There are several examples in which a large molecule physically occludes the mouth of a K\(^+\) channel, and the unblocking rate is increased by adding permeant ions to the opposite side of the channel. Internal quaternary ammonium ions block open K\(^+\)
channels and adding K⁺ to the external side of the membrane promotes unblock
(Armstrong, 1969, 1971; Armstrong and Hille, 1972). External charybdotoxin
physically occludes Ca²⁺-activated K⁺ channels and is pushed off the blocking site by
internal K⁺ (MacKinnon and Miller, 1988). In a variety of K⁺ channels inactivation is
slowed while recovery from inactivation is accelerated by increasing [K⁺]₀ (references
cited in DeCoursey, 1990), which is consistent with K⁺ influx destabilizing the
NH₂-terminal region of the K⁺ channel protein which acts as an inactivation particle
(Zagotta, Hoshi, and Aldrich, 1990; Hoshi, Zagotta, and Aldrich, 1990) in a
"ball-and-chain" model of inactivation (Armstrong and BezaniUa, 1977).

Might IR gating be the result of an analogous mechanism in which gating is due to plugging
the intracellular mouth of the pore by peptide residues comprising an intracellular
portion of the IR channel protein? Occupancy of binding sites in the pore by
permeant ions might destabilize the binding of a tethered blocking peptide. IR
gating might then have the requisite dependence on [K⁺]₀, with voltage dependence
imparted by voltage-dependent occupancy of a site or sites within the pore.

The most severe difficulty with the intrinsic blocking particle model arises from the
steepness of the voltage-dependence of IR gating, with an effective zδ of 3.1–5.9
(Leech and Stanfield, 1981; Tourneur et al., 1987; Cohen et al., 1989; Burton and
Hutter, 1990; Silver and DeCoursey, 1990). Channel opening has a zδ of 1.4 in K⁺
and 1.0 in high [Rb⁺]₀ solutions. These latter values are similar to the voltage-
dependence of unblocking in the examples cited above of K⁺ displacing a blocking
particle, with zδ ≈ 1.0 (Armstrong, 1971; MacKinnon and Miller, 1988; Demo and
Yellen, 1991). However, in these examples the onset of block/inactivation is practi-
cally voltage-independent; the analogous process, IR channel closing, in contrast is
very steeply voltage-dependent. IR channel closing moves 2.7–3.0 equivalent charges
across the membrane in K⁺ and high [Rb⁺]₀ solutions. From these examples of
permeant ions ejecting blocking/gating moieties from superficial sites at the mouth
of K⁺ channels, we find no precedent to support the idea that an intrinsically
voltage-independent gating process which does not involve a conformational change
of the channel protein might gain sufficient voltage-dependence simply from the
occupancy of permeant monovalent ions in the pore to account for IR gating.

Finding an effective δ > 1 for ionic blockers classically has been taken as evidence
that a channel is multiply occupied by ions during permeation (Hille and Schwarz,
1978). The largest zδ thus far reported for IR blockers is 1.5–1.6 for block by Cs⁺
(Hagiwara et al., 1976; Shioya, Matsuda, and Noma, 1993), 1.57 for Na⁺ (Harvey and
Ten Eick, 1989), and 1.6–2.5 for Rb⁺ (present data); the largest value we are aware of
for any other K⁺ channel is 1.6 (see Hille and Schwarz, 1978). Steeper voltage-
dependent block which is sensitive to V–E_R can be produced by an impermeant
blocking particle in a model long-pore, multi-ion channel, particularly if it is
multivalent, but only at permeant ionic activities causing saturating occupancies
(Hille and Schwarz, 1978). Nevertheless, even the divalent blocker Mg²⁺ within the
IR channel itself has a zδ of only 1.14 (Matsuda, 1991). In summary, a simple
blocking particle seems an unlikely mechanism for IR channel gating, although the
gating mechanism may be modulated by occupancy of the channel. In the following
paper (Pennefather and DeCoursey, 1994) we present an "electrochemical gating"
model in which both K⁺ and Rb⁺ act as allosteric modulators of an intrinsically
voltage-dependent isomerization between open and closed states of the IR channel.
Steep voltage and concentration dependence of IR activation is the result of required binding of several K⁺ or Rb⁺ to gating sites on the channel, in conjunction with cooperative gating of multiple bores of the channel.

Dependence of rectification on patch clamp configuration. The extent of outward current through single IR channels in patches was qualitatively different in cell-attached and excised outside-out patch configurations. No outward current could be detected in cell-attached patches under conditions designed to maximize it (Fig. 5). The absence of detectable outward current through IR channels in cell-attached patches has been reported in many cell types (e.g., Sauve, Roy, and Payet, 1983; Sakmann and Trube, 1984; Rae, 1986; Vandenberg, 1987; Matsuda et al., 1987; McKinney and Gallin, 1988). In contrast, outward current through IR channels was clearly evident in the outside-out patch configuration, both using a nominally Mg²⁺-free pipette solution (Fig. 6) and one containing 1.9 mM Mg²⁺ (Fig. 11 C). Both steady-state and time-dependent outward currents were also detected in whole-cell experiments with high or low [Mg²⁺]. (Silver and DeCoursey, 1990). In symmetrical K⁺ the unitary i-V relation indeed is approximately linear, consistent with the observations of Vandenberg (1987) and Matsuda et al. (1987) in cardiac Ik₁ channels in inside-out patches. However, a gating process brings P_open to near 0 at large positive potentials, both with high and low [Mg²⁺]. The behavior of single IR channels thus differs in cell-attached patches and in outside-out patches, and this difference is not entirely attributable to the presence or absence of internal Mg²⁺ in the outside-out patch. Deactivation of outward currents was slower in excised patches than at corresponding potentials in whole-cell experiments. Matsuda (1988) reported analogous progressive slowing of deactivation of Ik₁ channels in inside-out patches. This kinetic change in IR channel behavior may account for the enhanced outward current in excised patches. There may be an intracellular cofactor other than Mg²⁺, which is lost after patch excision as suggested by Matsuda (1988), either attached to the channel protein or diffusible in the cytoplasm, that blocks outward currents or modulates the gating process.

An inward rectifier channel similar in most respects to that described here has been isolated and expressed recently in oocytes (Kubo et al., 1993). It lacks the S4 segment believed to comprise the voltage sensor of depolarization-activated K⁺ channels (Stühmer, Conti, Suzuki, Wang, Noda, Yahagi, Kubo, and Numa, 1989). Time-dependent current relaxations corresponding with the gating process described here were not detected, but may be resolved in future studies. It may be that macroscopic gating requires additional structures or post-translational modification of the channel. Site-directed mutagenesis may reveal whether the proposed gating-ion binding sites exist, and should provide additional evidence of the mechanism of IR gating.

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