Mechanism of IRK1 Channel Block by Intracellular Polyamines

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Abstract Intracellular polyamines inhibit the strongly rectifying IRK1 potassium channel by a mechanism different from that of a typical ionic pore blocker such as tetraethylammonium. As in other K+ channels, in the presence of intracellular TEA, the IRK1 channel current decreases with increasing membrane voltage and eventually approaches zero. However, in the presence of intracellular polyamines, the channel current varies with membrane voltage in a complex manner: when membrane voltage is increased, the current decreases in two phases separated by a hump. Furthermore, contrary to the expectation for a nonpermeant ionic pore blocker, a significant residual IRK1 current persists at very positive membrane voltages; the amplitude of the residual current decreases with increasing polyamine concentration. This complex blocking behavior of polyamines can be accounted for by a minimal model whereby intracellular polyamines inhibit the IRK1 channel by inducing two blocked channel states. In each of the blocked states, a polyamine is bound with characteristic affinity and probability of traversing the pore. The proposal that polyamines traverse the pore at finite rates is supported by the observation that philanthotoxin-343 (spermine with a bulky chemical group attached to one end) acts as a nonpermeant ionic blocker in the IRK1 channel.

Keywords: inward-rectifier K+ channel • ion permeation • protonation • polyamine • diamine

Introduction Inward-rectifier K+ channels are blocked by intracellular Mg2+ and polyamines (Vandenberg, 1987; Matsuda et al., 1987; Logothetis et al., 1987; Horie et al., 1987; Lopatin et al., 1994; Ficker et al., 1994; Fakler et al., 1995). This block depends strongly on membrane voltage, causing the channel to conduct in an inwardly rectifying manner, enabling the inward-rectifier K+ channels to accomplish various important biological tasks (Hille, 1992). The voltage dependence of channel block is not exclusively an intrinsic property of the blocking ion because it also depends strongly on the nature and concentration of permeant ions present in the extracellular solution (Oliver et al., 1998; Spassova and Lu, 1998, 1999). Quantitative analyses show that the voltage dependence of channel block results at least in part from the concomitant movement of permeant ions (energetically coupled to the blocking ions) in the transmembrane electrical field (Spassova and Lu, 1998, 1999).

Previous studies of polyamine block of strongly rectifying inward-rectifier K+ channels have revealed complex phenomena. For example, the relation between the extent of channel block and membrane voltage exhibits a shallower phase and a steep phase, and thus requires an equation with two Boltzmann terms. It has, therefore, been suggested that more than one polyamine molecule can bind to the pore (Lopatin et al., 1995) and may be required for complete block of the channel (Yang et al., 1995; Lu et al., 1999). In the present study, we investigated the mechanism underlying the complex blocking behavior of polyamines in the strongly rectifying IRK1 channel.

Methods Molecular Biology and Oocyte Preparation

IRK1 (Kir2.1) cDNA cloned into the pcDNA1/AMP plasmid (In Vitrogen Corp.) was kindly provided by Dr. Lily Y. Jan (Kubo et al., 1993). RNA was synthesized using T7 polymerase (Promega Corp.) from Not1-linearized IRK1 cDNA. Oocytes harvested from Xenopus laevis (Xenopus One) were incubated in a solution containing (mM): 82.5 NaCl, 2.5 KCl, 1.0 MgCl2, 5.0 HEPES, pH 7.6, and 2–4 mg/ml collagenase. The oocyte preparation was agitated using a platform shaker (80 rpm) for 60–90 min. It was then rinsed thoroughly with and stored in a solution containing (mM): 96 NaCl, 2.5 KCl, 1.8 CaCl2, 1.0 MgCl2, 5 HEPES, pH 7.6, and 50 μg/ml gentamicin. Defolliculated oocytes were selected and injected with RNA at least 2 and 16 h, respectively, after collagenase treatment. All oocytes were stored in an incubator at 18°C.

Patch Recording and Data Analysis

IRK1 currents were recorded in the inside-out configuration from the membrane of Xenopus oocytes (injected with IRK1 cRNA) with an Axophot 200B amplifier (Axon Instruments, Inc.). The recorded signal was filtered at 1 kHz and sampled at 5 kHz using an analogue-to-digital converter (DigiData 1200; Axon Instruments, Inc.) interfaced with a personal computer. pClamp6 software (Axon Instruments, Inc.) was used to control the amplifier and acquire the data. During the current recording, the voltage across the membrane patch was first hyperpolarized from the 0 mV holding potential to ~100 mV for 50 ms (to relieve channel block by the tested intracellular blocker), and then stepped...
to a test voltage between -100 and +100 mV, in 5- or 10-mV increments, for 0.3-1 s. Background leak current correction was carried out as previously described (Lu and MacKinnon, 1994; Spassova and Lu, 1998). All curve fittings were carried out using Origin software version 5 (Microcal Software, Inc.).

Recording Solutions

Pipette solution contained (mM): 100 KCl, 0.3 CaCl$_2$, 1.0 MgCl$_2$, and 10 HEPES, pH 7.6. The bath solutions contained (mM): 90 KCl, 5 K$_2$EDTA, 10 HEPES, pH 7.6 or 6.6, as specified. The bath solutions containing diamines, polyamines, and philanthotoxin-343 (PhTx)$^\ddagger$ were prepared daily.

RESULTS

Channel Block by Intracellular Tetraethylammonium

Tetraethylammonium and all other intracellular IRK1 channel blockers that we studied block the channel in a voltage-dependent manner. To examine the IRK1 current at various membrane voltages, we used a voltage-pulse protocol rather than a ramp, because the current through the IRK1 channel does not instantaneously reach a steady state after a change in membrane voltage (Fig. 1A). To quantify channel block by a given blocker, we analyzed its effect only on the steady state current. Although the nature of the time-dependent current decline after a step to positive membrane voltages is unknown, intrinsic gating has been suggested as the cause because the current decline persists even after the intracellular side of the membrane patch is exhaustively perfused with recording solution nominally devoid of blocking ions (Aleksandrov et al., 1996; Shieh et al., 1996). Fig. 1B shows both the “instantaneous” and the “steady state” I-V curves recorded 2 and 300 ms, respectively, after the voltage step. The instantaneous I-V curve exhibits slight inward rectification, while the steady state I-V curve shows stronger rectification. The ratio of the two I-V curves in Fig. 1B is plotted against membrane voltage in C. The valence factor $Z^*$, which quantifies the voltage dependence of the current decline, is 0.55 ± 0.01 (mean ± SEM), much smaller than those of high-affinity polyamines such as spermidine and spermine. This finding argues that the current decline results from causes other than incomplete removal of high-affinity polyamines.

For the purpose of comparison, we first examined channel block by the classic intracellular ionic pore blocker TEA (Armstrong and Binstock, 1965). Fig. 2A shows the IRK1 currents recorded without and with various concentrations of intracellular TEA under symmetric K$^+$ conditions. The membrane was first hyperpolarized to -100 mV for 50 ms, and then depolarized for 300 ms to a test membrane voltage ranging from -100 to +100 mV in 10-mV increments. Outward currents in the presence of intracellular TEA became progressively smaller with membrane depolarization. Consequently,

$^\ddagger$Abbreviations used in this paper: CNG channel, cyclic nucleotide–gated channel; PhTx, philanthotoxin-343.
the I-V curves (Fig. 2 B) at positive membrane voltages display significant downward curvature in the presence of TEA. In Fig. 2 C, we plotted the fractions of current not blocked by various concentrations of TEA as a function of membrane voltage. The curves superimposed on the data are fits of the Woodhull equation (a variation of the Boltzmann equation) (Woodhull, 1973). As expected for an ionic pore blocker, the extent of IRK1 channel block by intracellular TEA progressively increases as membrane voltage is made more positive.

Channel Block by Intracellular Putrescine

Fig. 3 A shows current traces recorded in the absence or presence of various concentrations of intracellular putrescine. The recordings were carried out between −100 and +100 mV in 5-mV increments, but, for clarity, we plotted only the current traces recorded at 10-mV intervals. Fig. 3 B plots the I-V curves in the absence and presence of the three concentrations of putrescine. Unlike that in the presence of TEA (compare Fig. 2 B), the IRK1 channel current in the presence of putrescine, after reaching a peak at some positive voltages, did not continue to decline towards zero in the manner expected for a typical ionic pore blocker. Rather, a significant current persists at very positive membrane voltages. The amplitude of the residual current is smaller at higher putrescine concentrations. In Fig. 3 C, we plotted the fractions of unblocked current as a function of membrane voltage. Contrary to the expectation for voltage-dependent block by a typical ionic pore blocker, the extent of channel block for a given putrescine concentration does not follow the Woodhull equation, but approaches a non-zero level at positive membrane voltages. Putrescine clearly does not simply act as a typical ionic pore blocker.
Figure 3. Channel block by intracellular putrescine. (A) Current traces recorded in the absence or presence of various concentrations of putrescine. For each experiment, the membrane voltage was first hyperpolarized to −100 mV for 50 ms, and then stepped to various test voltages from −100 to +100 mV in 5-mV increments for 300 ms. However, for clarity, we only plotted the current traces recorded at 10-mV intervals. (B) I-V curves in the absence or presence of various concentrations of putrescine. (C) The fractions of current not blocked by putrescine are plotted against membrane voltage. The curves were drawn according to the Woodhull equation (Fig. 2).

Channel Block by Intracellular Diamines

To better understand block of the IRK1 channel by putrescine (a primary diamine) described here, we systematically tested a series of primary diamines of varying methylene chain length. Fig. 4 shows the current traces of the IRK1 channel between −100 and +100 mV, recorded in the absence and presence of nine diamines containing from 2 to 10 methylene groups (DM$_{C2}$ through DM$_{C10}$; putrescine is DM$_{C4}$). Fig. 5 shows I-V curves obtained without and with the nine diamines at various concentrations including those of Fig 4. Except for DM$_{C2}$, which blocked the channel slightly more effectively than DM$_{C3}$, blocking efficacy generally increased with methylene chain length. The effect of each diamine on the I-V curves is qualitatively similar to that of putrescine (DM$_{C4}$). However, the currents appear to reach a nonzero “plateau” at progressively lower membrane voltages as the methylene chain length increases. In Fig. 6, we plotted the fractions of unblocked current in the presence of the various diamines as a function of membrane voltage. As in the case of putrescine (DM$_{C4}$), the extent of channel block by each of the diamines tends to a nonzero level at positive membrane voltages, a behavior reminiscent of block of the retinal cGMP-gated (CNG) channel by certain diamines (e.g., DM$_{C8}$, DM$_{C9}$, and DM$_{C10}$), where these diamines act as permeant blockers (Guo and Lu, 2000).

Channel Block by Intracellular Polyamines

Fig. 7 A shows the current traces recorded between −100 and +100 mV in the absence or presence of various concentrations of the triamine spermidine. Practi-
cally, 30 nM is the lowest concentration of spermidine at which the outward IRK1 current reaches a steady state within 300 ms after a voltage step. In Fig. 7 B, we plotted I-V curves recorded in the absence or presence of spermidine at three concentrations. The I-V curve in the presence of spermidine consists of multiple phases. Fig. 7 C shows the fractions of unblocked current at the three spermidine concentrations as a function of membrane voltage. As with diamines, the extent of channel block by spermidine tends to nonzero levels at positive membrane voltages, but a noticeable hump appears in spermidine-blocking curves that was absent from those of diamines.

Fig. 8 A shows channel block by the intracellular tetramine spermine. Although spermine appears to block the channel more effectively, it displays a complex blocking behavior generally similar to that of spermidine. The I-V curves in the presence of spermine also consist of multiple phases (Fig. 8 B). The spermine-blocking curves also have a hump, albeit less prominent than in the case of spermidine, preceding the plateau phase at positive voltages (Fig. 8 C). Over a comparable voltage range, the IRK1 channel blocking curves for spermine and spermidine shown here resemble somewhat those in the CNG channel (Guo and Lu, 2000), except that the humps are much less prominent in the IRK1 channel.

Channel Block by Spermidine at Two Different Intracellular pH

To test whether the multiphasic polyamine blocking curve is due to nonuniform protonation of the polyamine and/or the channel, we examined how intracellular pH affects channel block by spermidine. Fig. 9 A shows IRK1 channel current traces obtained at membrane voltages between −100 and +100 mV. The two sets of current traces Fig. 9 A, top, were recorded without and those below with intracellular spermidine. Fig. 9 A, left, were obtained at intracellular pH 7.6, those on the right at pH 6.6. As shown previously (Shieh et al., 1996), lowering intracellular pH from 7.6 to 6.6 reduced the outward current through the IRK1 channel at positive voltages. In Fig. 9 B, the fractions of unblocked current, obtained at intracellular pH 7.6 and 6.6, are plotted as a function of membrane voltage. Lowering pH from 7.6 to 6.6 results in noticeable, if
small, changes in the spermidine-blocking curve: the hump decreases but the residual current at positive voltages increases.

Channel Block by a Spermine Derivative

To test whether the residual current at positive voltages is due to spermine being a permeant blocker, we examined block of the IRK1 channel by PhTx, essentially spermine with a bulky chemical group attached to one end. Fig. 10 A shows the current traces recorded at membrane voltages between −100 and +100 mV in the absence or presence of 10 μM PhTx. The corresponding I-V curves are shown in Fig. 10 B. In Fig. 10 C, we plotted the fractions of unblocked current as a function of membrane voltage. As in the case of TEA, the extent of channel block by PhTx increased monotonically with membrane voltage. Both the hump and the trailing “plateau” phase seen in the spermine-blocking curve are absent. The curve superimposed on the data points in Fig. 10 C is a fit of the Woodhull equation. Thus, PhTx blocks the IRK1 channel very much like a typical (nonpermeant) ionic pore blocker.

DISCUSSION

Block of the IRK1 channel by intracellular polyamines appears to be complex. To understand its mechanism, we examined channel blocking behaviors of di- and polyamines as a function of concentration and membrane voltage. Some of these blocking behaviors are reminiscent of those occurring in the CNG channel (Guo and Lu, 2000).

Channel Block by Diamines

As observed previously by Pearson and Nichols (1998), channel block by all primary diamines tested remains incomplete at positive membrane voltages (Fig. 6). To analyze the voltage dependence of channel block, Pearson and Nichols (1998) modified the Woodhull equation by adding a constant to account for the current remaining at positive voltages. The residual current is not likely to result from (a) diamine-induced discrete subconductance state(s) because we found that the amplitude of the residual current decreases when diamine concentration is raised (Fig. 6). We observed a similar phenome-
non in block of CNG channels by some longer diamines that appear to permeate very slowly through the channel (Guo and Lu, 2000). Thus, the persisting current observed here at very positive voltages likely results also from diamines being permeant blockers, as illustrated in Fig. 11 A, for which the fraction of unblocked channels and thus unblocked current is given by:

\[
\frac{I}{I_0} = \frac{1}{1 + [DM] \left( \frac{k_1 e^{-\frac{z_1 F V_m}{RT}}}{k_2 e^{-\frac{z_2 F V_m}{RT}}} + \frac{k_2 e^{-\frac{z_2 F V_m}{RT}}}{k_1 e^{-\frac{z_1 F V_m}{RT}}} \right)} \quad (1)
\]

or

\[
\frac{I}{I_0} = \frac{1}{1 + \left[ \frac{[DM]}{k_1 e^{-\frac{z_1 F V_m}{RT}}} \right]^{\left( \frac{k_1}{k_2} \frac{z_2 F V_m}{RT} \right)}} \quad (2)
\]

where \([DM]\) is the intracellular diamine concentration (the extracellular diamine concentration is zero) and all rate constants are assumed to vary exponentially with membrane voltage. The four independent parameters are: \(K_1\), the equilibrium dissociation constant for channel-diamine interaction; \(\frac{k_2}{k_1}\), the relative probability of a diamine bound in the pore permeating the pore versus returning to the intracellular solution; and the corresponding apparent valences \(Z_1\) and \(Z_2\). The curves superimposed on the data in Fig. 6 are fits of Eq. 2. Values for these four adjustable parameters, determined from the fits, are plotted against diamine methylene chain length in Fig. 12. The value of \(K_1\) generally decreases with increasing methylene chain length (Fig. 12 A), consistent with the observation that longer diamines are more potent blockers (Fig. 5). Thus, as previously suggested (Pearson and Nichols, 1998), hydrophobic interactions are critical in mediating interactions between diamines and the pore. 

Fig. 12 C shows that \(\frac{k_2}{k_1}\) is rather small and decreases (from 0.05 to 0.01) with increasing methylene chain length. Since \(\frac{k_2}{k_1}\) is the relative probability of a molecule bound in the pore traversing the pore versus returning to the intracellular solution, the result shows that diamines traverse the pore very infrequently.
However, a few percent of bound diamine molecules popping through the pore allow sufficient $K^+$ permeation to account for the residual current at positive membrane voltages. Furthermore, $k_2/k_1$ decreases with increasing methylene chain length, consistent with the expectation that more hydrophobic diamines are less likely to traverse the pore.

The valence factors $Z_1$ and "$z_1 + z_2$" also increase with methylene chain length (Fig. 12, B and D). Empirically, $Z_1$ and "$z_1 + z_2$" for a given diamine appear to be numerically similar. If, for the sake of discussion, one assumes that for a given diamine, quantities $Z_1$ and "$z_1 + z_2$" are equal, then Eq. 1 simplifies to:

$$\frac{I}{I_0} = \frac{1}{1 + \frac{[DM]}{K_1e^{\frac{-Z_1F\nu_m}{RT}} + \frac{k_{-2}}{k_1}}}.$$  

At a sufficiently high membrane voltage, Eq. 3 approaches a voltage-independent constant value determined solely by diamine concentration:

$$\frac{I}{I_0} = \frac{1}{1 + \frac{k_1}{k_{-2}}[DM]},$$  

which explains why at a given diamine concentration the voltage-dependent inhibition curve can be well fitted using the Woodhull equation, provided an extra constant is included to represent residual current at positive membrane voltages (Pearson and Nichols, 1998).

Eq. 4 can be appreciated in an intuitive way. When a lodged diamine molecule pops through the pore, a brief window of opportunity is created for $K^+$ ions to "leak" through the pore; the window closes when another diamine molecule becomes lodged in the pore.
Either a higher diamine binding rate ($k_1[\text{DM}]$) or a lower rate of bound diamine popping through the pore ($k_{-2}$), or both, will reduce the time available for $K^+$ to leak and thus leave a smaller residual current. This explains why the residual current is inversely related to diamine concentration (Fig. 6). Although diamines may permeate the channel, their intrinsic rate of permeation is too slow to account for the residual current. Nearly all the observed residual current must be carried by $K^+$ ions because it vanishes at sufficiently high blocker concentrations (Fig. 6).

Channel Block by Polyamines

Fig. 7 C shows the voltage-dependent blocking curve of the IRK1 channel in the presence of three concentrations of the triamine spermidine. Above 0 mV, the curve descends rapidly as membrane voltage is increased, and it then forms a hump before approaching a nonzero level at extreme positive voltages. Clearly, the curve cannot be fitted by a Woodhull equation. These blocking curves are reminiscent of those of the CNG channel in the presence of intracellular polyamines, although the hump observed here is much less prominent (Guo and Lu, 2000). To account for the multiphasic spermidine-blocking curve, we used a model, shown in Fig. 11 B, which assumes that binding of spermidine to the channel can lead to two blocked states. In each of the two blocked states, spermidine is bound with different affinity and has a different probability of traversing the pore. Interconvertibility between the two blocked states cannot be assessed based on the steady
states. The fraction of unblocked channel and thus un-
state data presented here. For simplicity, our model as-
sumes no direct transition between the two blocked
states. The fraction of unblocked channel and thus un-
blocked current is given by Eq. 5:
\[
\frac{1}{I_0} = \frac{1}{1 + [\text{PM}]},
\]

\[
\frac{1}{I_0} = \frac{1}{1 + [\text{PM}]} \left( \frac{\frac{k_1^b e}{\text{RT}} + \frac{k_2 e}{\text{RT}}}{\frac{k_1^a e}{\text{RT}} + \frac{k_2 e}{\text{RT}}} \right)
\]

or, upon rearranging:
\[
\frac{1}{I_0} = \frac{1}{1 + [\text{PM}]} \left( \frac{\frac{k_1^b e}{\text{RT}} + \frac{k_2 e}{\text{RT}}}{\frac{k_1^a e}{\text{RT}} + \frac{k_2 e}{\text{RT}}} \right)
\]

where \([\text{PM}]\) is the intracellular polyamine concentra-
tion (the extracellular polyamine concentration is zero)
and all rate constants are assumed to vary exponentially
with membrane voltage. As shown in Fig. 7, the spermi-
dine-blocking curves can be well fitted by Eq. 6.

Since the IRK1 channel is inhibited by intracellular
protons, and polyamines are not fully protonated near
neutral pH (Palmer and Powell, 1974), we wonder
whether the different blocked channel states result
from nonuniform protonation of either the polyamines
or the channel. Therefore, we examined how altering
intracellular pH affects the spermidine-blocking curve.
Lowering intracellular pH from 7.6 to 6.6 reduces the
hump in the curve, but enhances the residual current
at positive voltages (Fig. 9 B). Consequently, the block-
ing curves for the two pH values cross, a phenomenon
qualitatively similar to that seen with spermine in the
CNG channel. To analyze the data, we used a model
that assumes that the two blocked states result from
nonuniform protonation of a site in spermidine (or in
the channel) (Guo and Lu, 2000). According to the
model:

\[
\frac{1}{I_0} = \frac{1}{1 + [\text{PM}]} \left( \frac{\frac{k_1^b e}{\text{RT}} + \frac{k_2 e}{\text{RT}}}{\frac{k_1^a e}{\text{RT}} + \frac{k_2 e}{\text{RT}}} \right)
\]

\[
\frac{1}{I_0} = \frac{1}{1 + [\text{PM}]} \left( \frac{\frac{k_1^b e}{\text{RT}} + \frac{k_2 e}{\text{RT}}}{\frac{k_1^a e}{\text{RT}} + \frac{k_2 e}{\text{RT}}} \right)
\]

where \(\Theta = 10^{-\text{pH}}/(10^{-\text{pH}} + 10^{-\text{pKa}})\). The smooth traces
superimposed on the blocking curves in Fig. 9 B are si-
multaneous fits of Eq. 7 to the two curves. From the fit,
we deduced a pKa for the titrated site of 8.1 ± 0.1
(mean ± SEM). Given this value, much larger changes
in the blocking curve should be observed if the intra-
cellular pH were raised from 7.6 rather than lowered.
Unfortunately, channel currents run down very rapidly
above pH 8.0.

Palmer and Powell (1974) determined the three pKa
values of spermidine as 10.9, 9.8, and 8.3 (100 mM
ionic strength and 25°C), the latter comparable to the
value of 8.1 that we find for the titratable site influen-
cing spermidine block. If the titratable site is indeed an
amine group in spermidine, then the fully protonated
spermidine should correspond to the higher affinity
and more permeant conformation, while the partially
protonated molecule corresponds to the lower affinity
and less permeant component. These inferences are
compatible with the observation that the apparent va-
lence of the high-affinity blocking component is larger
than that of the low affinity one. However, the absolute
value of the valence factor undoubtedly does not solely
reflect the charges on spermidine, because the valence
value associated with the high-affinity blocking compo-
nent is ∼5. Although, unlike TEA or Mg\(^{2+}\) (Spaso-
va and Lu, 1998), polyamines such as spermidine and
spermine may bind in the narrow region of the pore
where most of the voltage drop occurs, the movement
of a polyamine molecule alone within the electrical
field obviously cannot account for an observed valence
of ∼5. Therefore, the rather large valence values associ-
ated with channel block by polyamines (and diamines)
most likely reflect a concerted movement of both the
blocking ion and K\(^{+}\) ions residing in the pore, as shown
for block of the ROMK1 channel by TEA and Mg\(^{2+}\)
(Spassova and Lu, 1998, 1999).
Intracellular protonation inhibits the IRK1 channel and also affects its gating, presumably by titrating a site in the channel with apparent pKa = 6.2 (Shieh et al., 1996), rendering the channel population also nonhomogeneous near neutral pH. Therefore, despite the fact that the multiphasic blocking curve can be well accounted for by a model in which spermidine exists in two protonated states with different affinities and permeabilities, we wish to emphasize that, in principle, the different blocking states may also in part result from nonhomogeneity of the channel.

As shown in Fig. 8 C, the blocking behavior of the tetramine spermine is very similar to that of the triamine spermidine and can be accounted for by the same model. However, the blocking behavior of a spermine derivative, PhTx, is quite different—no significant residual current was observed at positive voltages when the channel was blocked by PhTx (Fig. 10), as in the glutamate- and cGMP-gated channels (Bähring et al., 1997; Guo and Lu, 2000). It is as if the bulky group attached to the end of spermine effectively prevents the molecule from sliding through the channel. Compared with spermine, PhTx has a much lower affinity and apparent valence, consistent with a scenario where the bulky group hinders the interaction between spermine and the channel. A much lower affinity of the channel for a similar toxin, PhTx 433, than for spermine has also been observed by Lee et al. (1999). As shown in Fig. 10 C, the voltage-dependent block by PhTx is well described by the Woodhull equation. These findings support the idea that polyamines are permeant blockers in the IRK1 channel. Although the current is nearly voltage independent at negative potentials for a given concentration of PhTx, as well as di- and polyamines, it decreases when their concentration is increased. Thus, these polyamines must also inhibit the channel in a voltage-independent manner; e.g., by polyamine binding at the innermost part of the pore, where no signifi-
cant voltage drop occurs, or by a mechanism that does not involve the pore.

As in the case of diamines, comparable \( Z^1 \) and \( Z^2 \) values for each state (\( a \) and \( b \)) are needed to fit the polyamine-blocking curves. For the purpose of illustration, if one assumes \( Z^a_1 \) and \( Z^a_2 \) to be numerically equal (denoted \( Z^a \)) and \( Z^b_1 \) and \( Z^b_2 \) to be numerically equal (denoted \( Z^b \)), then Eq. 6 becomes Eq. 8:

\[
\frac{l}{I_o} = \frac{1}{1 + k_1^o \frac{Z}{Z^a F V / RT}}
\]

or, upon rearranging:

\[
\frac{l}{I_o} = \frac{1}{1 + \frac{k_1^o \frac{Z}{Z^a F V / RT}}{k_1^b \frac{Z}{Z^b F V / RT} + K_1^b}}
\]

Furthermore, for spermidine concentrations much greater than the sum of \( k_1^o / k_1^b \) and \( k_1^o / k_1^b \), Eq. 9 approaches Eq. 10:

\[
\frac{l}{I_o} = \frac{1}{1 + \frac{1}{k_1^o \frac{Z}{Z^a F V / RT} + K_1^b}}
\]
showing that the voltage dependence of channel block by polyamines at high concentrations can be approximated by an equation with two Boltzmann terms (compare Lopatin et al., 1995). When $K_1^a << K_1^b$ and also $Z_1^a > Z_1^b$, at some positive voltages, Eq. 9 approximates:

$$\frac{1}{I_o} = \frac{1}{1 + \frac{Z_1^a}{k_1^a + k_1^b} e^{-\frac{Z_1^b}{k_2^b}}},$$  \hspace{1cm} (11)$$

producing a hump (Fig. 7 C). At extreme positive voltages, Eq. 11 approaches Eq. 12:

$$\frac{1}{I_o} = \frac{1}{1 + \frac{Z_1^a}{k_1^a + k_1^b} [PM]} e^{-\frac{Z_1^b}{k_2^b}},$$  \hspace{1cm} (12)$$

accounting for the residual current at these voltages. As in the case of diamines, the amplitude of the residual current should decrease with increasing spermidine concentration and tend towards zero at very high concentrations, as observed (Fig. 7 C).

Currents through the IRK1 channel after a jump to positive membrane voltages decline with time even in patches exhaustively perfused with nominally blocking ion-free solution. It has been suggested that this decline in current results from intrinsic gating (Aleksandrov et al., 1996; Shieh et al., 1996; Lee et al., 1999), although both the extent and the rate of the current decline appear to vary considerably among various reported studies. Here, we found that the voltage dependence of the current decline is much weaker than that of block by high-affinity endogenous polyamines spermine and spermidine. This finding argues that the current decline reflects either the hypothesized intrinsic gating or channel block by some contaminating blockers other than spermine or spermidine. For presentation purposes, we will adopt the current nomenclature "intrinsic gating" when referring to the voltage-dependent current decline. The hypothesized intrinsic gating will complicate our analysis somewhat, but will not fundamentally affect our conclusions. The hump in the spermidine (or spermine)-blocking curve occurs between +20 and +50 mV, where the current decline due to intrinsic gating is minimal (Fig. 1). To include intrinsic gating (or channel block by contaminating blocking ions), at least one more non- or subconducting state (Ch*) is required (Fig. 13). For simplicity, we treat Ch* here as a nonconducting state. In principle, channels in the Ch* state may or may not be occupied by a polyamine, and channels in Ch state occupied by a polyamine may or may not undergo the gating transition. Additional non-steady state experiments are needed to evaluate these possibilities. According to Fig. 13, the reciprocals of both equilibrium constants in steady state Eq. 7 need to be multiplied by a term, as indicated by Eq. 13:
where $K^*$ is $[\text{Ch}^*]/[\text{Ch}]$, and $Z^*$ is the associated valence factor. Under our experimental conditions, the fitted values for both $K^*$ (0.03 ± 0.01; mean ± SEM) and $Z^*$ (0.55 ± 0.01) are small (Fig. 1 C), yielding multiplier values that range from 0.97 at 0 mV to 0.80 at +100 mV. Therefore, the addition of this nonconducting state would cause only minor changes in the fitted equilibrium constants ($K^*$ and $K_0$) and valence factors ($Z^*$ and $Z_0$), but has no fundamental impact on our analysis.

Models involving channel gating have previously been proposed to explain various aspects of intracellular cotransmission-induced inward rectification in the IRK1 channel. For example, Aleksandrov et al. (1996) proposed that intracellular Mg$^{2+}$ and polyamines inhibit the channel by controlling a gate located at its intracellular end. Subsequently, Lee et al. (1999) proposed a more detailed physical model where one end of the polyamine molecule is tethered to a channel gate while the other plugs the pore. The spermine-gate complex is assumed to block the channel with high affinity, while spermine itself has only low affinity. Kinetically, both their model and our model have one conducting and three nonconducting states, including the nonconducting gating state. However, the four-state model of Lee et al. (1999), without additional features, predicts that the current in the presence of polyamine will decrease with increasing membrane voltage in a multiexponential fashion and eventually vanish. In other words, their model accounts for neither the hump nor the residual current at positive voltages that we observed in the polyamine-blocking curve, because it lacks a voltage-dependent mechanism that opposes the current reduction due to channel gating or block. To account for these unusual characteristics, our model invokes two essential features: in the two blocked states, the polyamine molecules are bound not only with different affinities, but also, more critically, with different probabilities of traversing the pore. The $K^*$ leak currents resulting from the finite polyamine permeabilities can fully account for both the hump and the residual current at positive voltages. Our proposal is supported by the absence of significant humps or residual currents at positive voltages in the curves of the bulky blocker PhTx or of TEA.

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