Control of rectification and permeation by two distinct sites after the second transmembrane region in Kir2.1 K⁺ channel

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1. The rectification property of the inward rectifier K⁺ channel is chiefly due to the block of outward current by cytoplasmic Mg²⁺ and polyamines. In the cloned inward rectifier K⁺ channel Kir2.1 (IRK1), Asp172 in the second transmembrane region (M2) and Glu224 in the putative cytoplasmic region after M2 are reported to be critical for the sensitivity to these blockers. However, the difference in the inward rectification properties between Kir2.1 and a very weak inward rectifier sWIRK could not be explained by differences at these two sites.

2. Following sequence comparison of Kir2.1 and sWIRK, we focused this study on Glu299 located in the centre of the putative cytoplasmic region after M2. Single-point mutants of Kir2.1 (Glu224Gly and Glu299Ser) and a double-point mutant (Glu224Gly–Glu299Ser) were made and expressed in Xenopus oocytes or in HEK293T cells.

3. Their electrophysiological properties were compared with those of wild-type (WT) Kir2.1 and the following observations were made. (a) Glu299Ser showed a weaker inward rectification, a slower activation upon hyperpolarization, a slower decay of the outward current upon depolarization, a lower sensitivity to block by cytoplasmic spermine and a smaller single-channel conductance than WT. (b) The features of Glu224Gly were similar to those of Glu299Ser. (c) In the double mutant (Glu224Gly–Glu299Ser), the differences from WT described above were more prominent.

4. These results demonstrate that Glu299 as well as Glu224 control rectification and permeation, and suggest the possibility that the two sites contribute to the inner vestibule of the channel pore. The slowing down of the on- and off-blocking processes by mutation of these sites implies that Glu224 and Glu299 function to facilitate the entry (and exit) of spermine to (and from) the blocking site.
M2 region, which corresponds to D172 of Kir2.1, sWIRK showed a very weak rectification (Kubo et al. 1996). The amino acid residue that corresponds to E224 of Kir2.1 is G231, as for Kir1.1 (G223). By mutating E179 of sWIRK to Gln (Q) (E179Q, G231), the inward rectification property was almost completely lost and it showed outward rectification even in the presence of cytoplasmic blockers (Kubo et al. 1996). As this feature clearly differed from the double mutant of Kir2.1 (D172N–E224G) (Yang et al., 1998) and from this study have been published in abstract form compared with those of WT Kir2.1. Preliminary results were made, and their electrophysiological properties were almost half-way through the putative cytoplasmic region in Kir7.1 (Fig. 1B) (N171D) (Lu & MacKinnon, 1994). In contrast, the charged amino acid was introduced into the M2 region which also showed strong rectification when a negatively charged in all strong inward rectifiers and in Kir1.1, critical for the strong inward rectification was speculated. Therefore, the amino acid sequence of sWIRK was compared in this study with that of other strong inward rectifiers such as Kir2.1, and E299 of Kir2.1 almost half-way through the putative cytoplasmic region following the M2 region was focused upon (Fig. 1A and B). The corresponding amino acid residues were negatively charged in all strong inward rectifiers and in Kir1.1, which also showed strong rectification when a negatively charged amino acid was introduced into the M2 region (N171D) (Lu & MacKinnon, 1994). In contrast, the corresponding residue was Ser in sWIRK, in Kir4.1 and in Kir7.1 (Fig. 1B). Single-, double- and triple-point mutants of Kir2.1 at the three sites D172, E224 and E299 were made, and their electrophysiological properties were compared with those of WT Kir2.1. Preliminary results from this study have been published in abstract form (Kubo, 1999).

**METHODS**

*In vitro mutagenesis*

The single-point mutants were made using the Sculptor kit (Amersham), using mutated oligonucleotide DNA primers and single-stranded (ss)DNA of WT mouse Kir2.1. The introduction of a mutation was confirmed by sequencing with a PRISM kit (Applied Biosystems) using ABI 377–18 DNA sequencer (Applied Biosystems). The electrophysiological properties of two independent clones of the mutant were confirmed to be identical. The double-point mutants were made using the ssDNA of the single-point mutants, and the triple mutants were prepared based on the ssDNA of the double mutants.

**Two-electrode voltage-clamp recordings in Xenopus oocytes**

*Xenopus* oocytes were collected from frogs anaesthetized in water containing 0.15% tricaine; after the final collection the frogs were killed by decapitation. Isolated oocytes were treated with collagenase (2 mg ml⁻¹, type 1, Sigma), and injected with approximately 50 nl of cRNA solution, prepared from the linearized plasmid DNA by RNA transcription kit (Stratagene). The injected oocytes were incubated for 2–4 days in frog Ringer solution (Kubo et al. 1993a) supplemented with 20 mM KCl at 17°C. Macroscopic current was recorded under two-electrode voltage clamp using a ‘bath-clamp’ amplifier (OC-725-BHV, Warner Co.). Stimulation, data acquisition and analysis were done on a Pentium-based computer using Digidata 1200A and pCLAMP software (Axon Instruments). Intracellular glass microelectrodes were filled with 3 M potassium acetate with 10 mM KCl (pH 7.2). The microelectrode resistance ranged from 0.1 to 0.4 MΩ. Two Ag–AgCl pellets (Warner Co.) were used to pass the bath current and to sense the bath voltage. The voltage-sensing electrode was placed near the oocyte (approximately 2 mm away) on the same side as the voltage-recording microelectrode. The bath current-passing pellet and the current injection microelectrode were placed on the other side. Under these conditions, the series resistance between the oocyte surface and the bath voltage-sensing pellet was approximately 200 Ω (Sabirov et al. 1997). As the measured current at the most hyperpolarized potential was 50 μA in the largest case, and mostly less than 20 μA, the voltage-clamp error due to this series resistance was estimated to be no more than 10 mV and mostly less than 4 mV. The error, which was not compensated in the experiments, did not change the conclusions of the comparison of WT and mutant channels in the present study. The recording bath solution contained 10 mM KCl, 80 mM N-methyl-glucamine, 70 mM

![Figure 1](Image)

**Figure 1.** The position of E299 of Kir2.1, and comparison of the corresponding amino acid residues among the inwardly rectifying K⁺ channel family

A, schematic drawing of the structure of the Kir2.1 channel based on the initial model by Kubo et al. (1993a). E299 is located at the centre of the putative cytoplasmic chain after the M2 region. D172 in the M2 region and E224 just after M2 are also depicted. B, alignment of the amino acid sequences of the inwardly rectifying K⁺ channel family at E299 of Kir2.1 and the surrounding region. The original reports used for the alignment are as follows. Kir1.1, Ho et al. (1993); Kir2.1, Kubo et al. (1993a); Kir2.2, Koyama et al. (1994); Kir2.4, Topert et al. (1998); Kir3.1, Kubo et al. (1993b) and Dascal et al. (1993); Kir3.2, Lesage et al. (1994); Kir3.4, Krapivinsky et al. (1995) and Iizuka et al. (1995); Kir6.1, Inagaki et al. (1995a); Kir6.2, Inagaki et al. (1995b); Kir7.1, Krapivinsky et al. (1998) and Doring et al. (1998); Kir4.1, Bond et al. (1994) and Takumi et al. (1995); sWIRK, Kubo et al. (1996).
HCl, 3 mM MgCl₂ and 5 mM Hepes (pH 7.4). All recordings were carried out at room temperature (23 ± 2°C).

By assuming the intracellular potassium concentration ([K⁺]) of the oocytes to be 80 mM, the equilibrium potential of the potassium ion (E_K) was calculated to be −52 mV. However, the exact values of [K⁺] and E_K of each oocyte are unknown. Therefore, E_K, used for the calculation of the chord conductance (Figs 2, 3 and 10) was adjusted to yield a continuous conductance–voltage (g–V) plot. The adjusted E_K ranged from −54 to −49 mV.

Data from the same batch of oocytes were used for comparison of the phenotypes in Figs 2–6 and 10, because properties such as inward rectification and blocking speed are influenced by the cytoplasmic polyamine concentration, and because they differ significantly depending on the batch. Similar tendencies of the data in Figs 2–6 and 10 were reproducibly observed in five batches of oocytes.

**Single-channel recordings in Xenopus oocytes**

For single-channel recording, the vitelline membrane was peeled off by bathing oocytes in a high osmolarity solution (Kubo et al. 1993a) for 5–10 min. The patch pipettes were prepared from borosilicate glass (Warner Instruments) using a P-97 horizontal puller (Sutter) and a fire polisher (Narishige, Japan). Recordings were obtained under the cell-attached configuration using an Axopatch-1D amplifier (Axon Instruments). The current was low-pass filtered at 1 kHz by a Bessel filter built into the amplifier, and was digitized at 5 kHz. The recordings shown in Fig. 9 were digitally filtered at 200 Hz. The pipette (extracellular) and the bath solution contained 130 mM KCl, 3 mM MgCl₂, 10 mM Hepes and 4 mM KOH (pH 7.4). The patch pipette resistance ranged from 3 to 10 MΩ.

**Expression in HEK293T cells**

The cDNAs of WT and mutant Kir2.1 were subcloned into the expression vector pCXN2 (Niwa et al. 1991). The plasmid DNA was transfected into HEK293T cells (human embryonic kidney cell line) using Lipofectamine Plus (Gibco BRL) following the manufacturer’s protocol. Enhanced green fluorescent protein (GFP; Clontech; 1/10 the amount of plasmid DNA) was co-transfected as a transfection marker. The cells were cultured in Dulbecco’s modified Eagle’s medium with 10% bovine calf serum for 24 h. The cells were then dissociated by treatment with 0.2% trypsin in Ca²⁺-, Mg²⁺-free PBS, medium with 10% bovine calf serum for 24 h. The cells were then seeded on coverslips at a relatively low density. The re-seeding was done to obtain well-isolated cells and to facilitate successful giga-ohm seal formation. It was confirmed that the trypsin treatment did not change the electrophysiological properties of the expressed channels. Electrophysiological recordings were carried out 4–30 h after re-seeding, which corresponds to 28–54 h after transfection.

**Macrosopic current recordings in HEK293T cells**

A coverslip of HEK293T cells was placed in a recording chamber containing bath solution (see below) on an inverted fluorescence microscope (IX70, Olympus, Japan), and transfected cells were identified by the fluorescence signal of the co-transfected GFP. The macroscopic current was recorded under the excised inside-out patch condition using an Axopatch-1D amplifier. The resistance of the patch pipettes ranged from 2 to 4 MΩ. Fifty to eighty per cent of the voltage error due to the series resistance was compensated by a circuit in the amplifier. The combination of HEK293T cells and the pCXN2 expression vector enabled high expression sufficient for macroscopic current recording using standard-sized patch pipettes. The current was low-pass filtered at 1 kHz by a built-in circuit in the amplifier and digitized at 5 kHz.

The pipette (extracellular side) solution contained 160 mM KCl, 120 mM N-methyl-glucamine, 103 mM HCl, 3 mM MgCl₂, 10 mM Hepes and 4 mM KOH (pH 7.4). The bath (intracellular side) solution was low-pass filtered at 1 kHz by a Bessel filter built into the amplifier, and was digitized at 5 kHz. The recordings shown in Fig. 9 were digitally filtered at 200 Hz. The pipette (extracellular) and the bath solution contained 130 mM KCl, 3 mM MgCl₂, 10 mM Hepes and 4 mM KOH (pH 7.4). The bath solution was measured to be approximately 10 mM and membrane potentials were corrected for this value.

The extent of rundown during a set of I–V relationship recordings was monitored by applying step pulses again, and data with apparent rundown were discarded. Due to this limitation, it was not possible to obtain a series of data for all concentrations of blockers from one patch, and the data shown in Figs 7 and 8 were obtained from multiple patches.

**RESULTS**

**Macrosopic current recordings of WT and mutant Kir2.1**

The electrophysiological properties of a point mutant of Kir2.1 in which Glu at position 299 was mutated to Ser (E299S) were compared with those of WT Kir2.1 and the D172N and E224G mutants (Fig. 2). The properties of the double mutants of the three sites, D172N–E224G, D172N–E299S and E224G–E299S, were also analysed (Fig. 3). The macroscopic currents in 10 mM K⁺ under two-electrode voltage clamp in Xenopus oocytes are shown in Figs 2A and 2B. The current–voltage (I–V) relationships (Figs 2B and 3B) and the conductance–voltage (g–V) relationships (Figs 2C and 3C) are also shown.

The observed features were as follows. (1) The D172N mutation weakened the inward rectification slightly, and removed the activation phase upon hyperpolarization (Figs 2 and 3). (2) Both the E224G and E299S mutations remarkably weakened the inward rectification of WT Kir2.1 and the D172N mutant. In the E224G and E299S mutants, significant outward currents were observed even 100 ms after the beginning of the depolarized pulses. Both of the mutations slowed down the activation upon hyperpolarization and the decay of the outward current upon depolarization (Figs 2 and 3). (3) The properties of the E224G and E299S mutants were highly similar to each other (Figs 2 and 3). (4) The double mutation E224G–E299S enhanced the changes observed in the E224G and E299S mutants (Fig. 3).

In the following sections, the properties of WT and mutant channels were compared with respect to inward
rectification, activation at hyperpolarized potentials and decay of the outward current at depolarized potentials.

Comparison of the inward rectification properties of WT and mutant Kir2.1

The intensity of the inward rectification of WT Kir2.1 and the D172N, E224G, E299S and E224G–E299S mutants is compared in Fig. 4. As the conductance of the E224G–E299S double mutant did not reach a plateau level even at the most hyperpolarized potential (Fig. 3C), it was not possible to fit the $g$–$V$ plot reliably with the Boltzmann equation and to compare the fitting parameters quantitatively. Instead, the intensity of the inward rectification was compared semi-quantitatively by normalizing the current amplitudes to the values at $-100$ mV (Fig. 4A and B). As an index of the intensity of the inward rectification, the ratio of the current amplitudes at $+50$ and $-100$ mV was calculated, and the accumulated data ($n = 3–4$) are shown (Fig. 4C and D). It is clear that the D172N mutant showed slightly weaker inward rectification than WT, and both E224G and E299S mutants showed much less intense inward rectification. The inward rectification of E224G–E299S, especially at 5 ms from the beginning of the step pulses, was even weaker than that of the E224G or E299S mutants (Fig. 4).

It was additionally observed that E224G–E299S and E224G–E299Q had similar inward rectification properties, and that D172N–E299S and D172N–E299Q were also similar (Fig. 4C and D). These results imply that the negative charge of E299 contributes to inward rectification and that the changed phenotype of E299S is not due to Ser itself, and similarly in the case of the E224G mutant (Yang et al. 1995). However, as there was a difference in the inward rectification properties between E299S and E299Q (Fig. 4C and D), supplementary effects of the size and the shape of the amino acid residue at position 299

Figure 2. Macroscopic current recordings under two-electrode voltage clamp of WT Kir2.1 (IRK1) and D172N, E224G and E299S mutants expressed in *Xenopus* oocytes

A, representative current recordings in 10 mM K$_G$. The holding potential was $-50$ mV, and step pulses from $+50$ to $-140$ mV were applied by 10 mV decrements. B, $I$–$V$ relationships of the data in A. The current amplitudes at 5 ms (●) and at 100 ms (○) of the step pulses are plotted. C, chord $g$–$V$ relationship of the data in B.
were suggested, in addition to the negative charge. To examine further the effect of electrostatic charge at this residue, E299R and E299K mutants were also prepared. Neither of them, however, expressed functional current, showing the importance of the amino acid residue at position 299 for channel function. For the following analysis, E299S was used as a representative mutation of this site.

Comparison of the activation kinetics of WT and mutant Kir2.1

The activation phase upon hyperpolarization was not clearly observed in the D172N (Fig. 2A), D172N–E224G or D172N–E299S mutants (Fig. 3A). The activation phases at −120 to −160 mV in 10 mM KCl of WT Kir2.1 and the E224G, E299S and E224G–E299S mutants are shown in Fig. 5A. The time constants of the fits with a single-exponential function are plotted in Fig. 5B. The inactivating component of WT Kir2.1 (Fig. 5A) was ignored in the fitting. Fitting with a double-exponential function was also done for E224G–E299S to improve the fit, and the two time constants were also plotted (Fig. 5B). The activation of E224G and E299S is clearly slower than that of WT. In E224G–E299S, an additional very slow component was observed. The voltage dependency of the time constants did not differ remarkably between WT and the mutants. These features were confirmed in three oocytes of the same batch, and a similar tendency was reproducibly observed in other batches.

Comparison of the outward current of WT and mutant Kir2.1

The time course of the tail currents at +50 mV, when depolarized from various prepulse potentials, was analysed. In Fig. 6A, recordings of the outward current are shown on a magnification scale normalized to the amplitude of the inward current at −100 mV. The decay of the outward current of WT Kir2.1 was too fast to be analysed separately from the capacitative current. In contrast, the tail currents of the mutants were larger in amplitude and declined much more slowly. They could be fitted not with a single- but with a double-exponential

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**Figure 3.** Macroscopic current recordings under two-electrode voltage clamp of double mutants of Kir2.1 (D172N–E224G, D172N–E299S and E224G–E299S) expressed in *Xenopus* oocytes

A detailed explanation is given in Fig. 2.
function. A triple-exponential function did not improve the fit. The two time constants obtained are plotted in Fig. 6B. The two time constants of E224G and E299S were in the same range, and those of E224G–E299S were much larger. These features were confirmed in three oocytes from the same batch, and also in other batches. The slight voltage dependency of the time constants (Fig. 6B) might be, at least partly, due to the artifact of the current measurements. The error due to the series resistance between the oocyte surface and the bath potential electrode was negligible for the conclusions drawn here.

**Fraction of the slowly decaying component of the outward current of E224G–E299S**

The slowly declining component of E224G–E299S did not increase proportionally as the amplitude of the preceding hyperpolarizing pulse increased. The amplitudes of the two components calculated in the fit of Fig. 6B are plotted in Fig. 6C. It is clear that the fast component increased monotonically and proportionally as the preceding step pulses were more hyperpolarized. In contrast, the increase in the amplitude of the slow component was not monotonic; the increase started to be augmented at −100 mV. These results were reproducibly observed in three oocytes from the same batch, and also in other batches.

If there is only a single open state (and no inactivated or blocked state) at the preceding hyperpolarized potentials, the ratio of the amplitudes of the fast and slow components would be expected to be independent of the preceding voltage steps. Therefore, the results in Fig. 6 imply that there are multiple states at hyperpolarized potentials in E224G–E299S and that the ratio of the probabilities of these states changes depending on the

![Image](image_url)
prepulse potential. A non-monotonic increase in the slow component was also noticed in E224G and in E299S (Fig. 6A).

Comparison of the sensitivity to the block by $\text{Mg}^{2+}$ of WT and mutant Kir2.1

Kir2.1 WT and mutant cDNA was subcloned into the expression vector pCXN2 and transfected to HEK293T cells by the lipofection method. The macroscopic current was recorded from excised inside-out membrane patches. The recordings, in the nominal absence of polyamines and in the presence of various concentrations of free $\text{Mg}^{2+}$, are shown in Fig. 7A. In the absence of polyamines and $\text{Mg}^{2+}$, the $I-V$ relationships of WT and mutant Kir2.1 were almost linear over the time range studied. By increasing $[\text{Mg}^{2+}]$, the outward current of WT Kir2.1 at depolarized potentials was blocked almost instantaneously, but the outward current of the mutants was blocked much more slowly. The normalized $I-V$ relationships at 500 ms from the beginning of the step pulses are plotted and the accumulated data of the inward rectification index in various $[\text{Mg}^{2+}]$ ($n = 3-6$) are shown in Fig. 7B and C, respectively. The normalized current amplitudes at 500 ms did not differ significantly between WT and the mutants.

The outward current amplitude and the inward rectification index of WT Kir2.1 in the absence of $\text{Mg}^{2+}$ and polyamines showed high variation, as judged by the large standard deviation (Fig. 7C). As the sensitivity of the WT channels to block by spermine is extremely high as shown later, this variation is thought to be due to residual cytoplasmic polyamines, even after intensive perfusion of the bath (intracellular) solution with polyamine-free solution. The WT channel data in Fig. 7A and B in $\text{Mg}^{2+}$-free solution are from a case in which prominent outward current was observed.

Comparison of the sensitivity to block by spermine of WT and mutant Kir2.1

The current recordings in Fig. 8A were obtained in the absence of $\text{Mg}^{2+}$ and in the presence of various concentrations of spermine, and the normalized $I-V$ plots at 3 s from the beginning of the step pulses are shown in Fig. 8B. WT Kir2.1 was highly sensitive to block. E224G and E299S were much more resistant to block by spermine, and the decrease in sensitivity was even more prominent in E224G–E299S. These observations were confirmed in the accumulated data of the rectification index in Fig. 8C and D ($n = 3-6$). The sensitivities of E224G and E299S were approximately 100–1000 times lower.
than that of WT, and that of E224G–E299S was 10 000–100 000 times lower than that of WT (Fig. 8C and D). By fitting the data of Fig. 8B, the $K_d$ values at +30 mV were estimated to be approximately 0.0005 µM for WT, 0.2 µM for E224G, 0.1 µM for E299S and 30 µM for E224G–E299S. The $K_d$ value for the double mutant E224G–E299S assuming a simple additivity of changes in binding energy of E224G and E299S is predicted by the following equation: $K_d(E224G–E299S) = (K_d(E224G)K_d(E299S))/K_d(WT)$ (Yang et al. 1995). As the predicted value (40 µM) is close to the experimental value (30 µM), it is suggested that E224 and E299 contribute independently to spermine binding.

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**Figure 6. Comparison of the outward current of WT Kir2.1 and the E224G, E299S and E224G–E299S mutants expressed in *Xenopus* oocytes under two-electrode voltage clamp (A and B) and the dependency of the fraction of the slowly decaying component of E224G–E299S on the preceding step pulses**

$A$, outward current recordings at +50 mV when depolarized from various potentials ranging from +50 to −160 mV. [K+]o was 10 mM. $B$, the recordings of E224G, E299S and E224G–E299S were fitted with a double-exponential function, and the two time constants are plotted. The bottom axis indicates the voltage of the preceding step pulses. The symbols used are as follows: E224G, ■; E299S, □; E299S–E224G, ●. The filled symbols represent the values of the fast component and the open symbols are for the slow one. Data from two oocytes from the same batch are plotted for each channel. $C$, the amplitudes of the fast (top) and the slow (bottom) components used for the fitting of E224G–E299S in $B$. 
Inward rectification property of Kir2.1 E299S mutant

Figure 7. The sensitivity of the macroscopic currents of WT Kir2.1 and the E224G, E299S and E224G–E299S mutants to block by Mg$^{2+}$.

A, macroscopic current recordings from an excised patch of HEK293T cells transfected with Kir2.1 WT or mutant cDNA. The concentration of free Mg$^{2+}$ was determined by adding MgCl$_2$ and EDTA following the equation of Fabiato & Fabiato (1979). [K$^+$]$_i$ in the pipette was 20 mM, and [K$^+$]$_o$ in the bath was 140 mM. The calculated $E_K$ was −49 mV. The holding potential was −50 mV and step pulses from +30 down to −90 mV were applied by 10 mV decrements. B, normalized $I$–$V$ relationships at 500 ms from the beginning of the step pulses. The current amplitudes were normalized by the values at −90 mV. The symbols used are as follows: 0 µM Mg$^{2+}$, ○; 3 µM, △; 30 µM, ■; 300 µM, ◦; 3 mM, ▲. C, comparison of the rectification indexes at 500 ms from the beginning of the step pulses. Rectification index was calculated as the ratio of the current amplitudes at +30 and −90 mV. The mean and the standard deviation of the accumulated data ($n = 3$–6) are plotted. The five bars indicate the values in 0, 3, 30, 300 and 3000 µM Mg$^{2+}$, from left to right.
Figure 8. The sensitivity of the macroscopic current of WT Kir2.1 and the E224G, E299S and E224G–E299S mutants to block by spermine

A, macroscopic current recordings from an excised patch of HEK293T cells transfected with the channel cDNA. The concentrations of spermine (Spm) are indicated in each panel. Note that data for 0.01, 0.1 and 1 µM spermine are shown for WT Kir2.1, and data for 1, 10 and 100 µM spermine are shown for the mutants. \([K^+]_o\) and \([K^+]_i\) were 20 and 140 mM, respectively. The holding potential was −50 mV, and step pulses from +30 down to −90 mV were applied by 10 mV decrements. B, normalized \(I–V\) relationships at 3 s from the beginning of the step pulses. The current amplitudes were normalized to the values at −90 mV. The symbols used for WT are as follows: 0 µM, ◦; 0.01 µM, Δ; 0.1 µM, ■; 1 µM, ○. The symbols for the mutants are as follows: 0 µM, ◦; 1 µM, Δ; 10 µM, ■; 100 µM, ○. C and D, comparison of the rectification indexes at 3 s from the beginning of the step pulses. Rectification index was calculated as the ratio of the current amplitudes at +30 and −90 mV. The mean and the standard deviation (n = 3–6) are plotted. The four bars indicate the data in 0, 0.01, 0.1 and 1 µM spermine for WT (C), and 0, 1, 10 and 100 µM spermine for the mutants (D) from left to right.
Comparison of the single-channel conductance of WT and mutant Kir2.1

The single-channel recordings in Fig. 9A were obtained from Xenopus oocytes under the cell-attached patch configuration in 140 mM K⁺ at −120 mV. The single-channel current–voltage relationships and the accumulated data for the single-channel conductance are shown in Fig. 9B and C. The values were (means ± S.D.) 19.7 ± 0.49 pS for WT, 16.7 ± 0.46 pS for E224G, 16.5 ± 0.35 pS for E299S and 12.7 ± 0.45 pS for E224G–E299S (n = 3). The open probability, the mean open time and the mean closed time did not show any obvious differences. Yang et al. (1995) reported that open channel noise is increased in the E224G mutant. A slight increase was also observed in the mutants of this study as shown in the expanded traces of Fig. 9A (right). The variance of the current fluctuation of the open channel at −120 mV was 0.024 pA² for WT, 0.050 pA² for E224G, 0.054 pA² for E299S and 0.032 pA² for E224G–E299S, when that of closed level was in the same range as follows: 0.014 pA² (WT), 0.012 pA² (E224G), 0.011 pA² (E299S) and 0.014 pA² (E224G–E299S). Taken together, these results suggest that E299 as well as E224 contribute to the permeation properties.

Phenotype of the D172N–E224G–E299S mutant and comparison with that of sWIRK

To examine whether the three sites focused on in the present study determine the inward rectification property completely, two triple mutants of Kir2.1, D172N–E224G–E299S and D172N–E224G–E299Q, were made. The E179Q mutant of sWIRK, which has no negative charges at the three sites in question (E179Q–G231–S311) were also made. The mutants were expressed in Xenopus oocytes and recorded in 10 mM K⁺ under two-electrode voltage clamp. The current recordings

![Figure 9. Comparison of the single-channel recordings of WT Kir2.1 and the E224G, E299S and E224G–E299S mutants](image-url)

A, representative single-channel recordings in Xenopus oocytes under the cell-attached patch configuration. The [K⁺] in the pipette and in the bath was 140 mM. The holding potential of the displayed recordings was −120 mV. The dashed lines indicate the zero current level after subtracting the leak current. The traces are shown on an expanded time scale on the right to demonstrate the fluctuation of the open channel current. B, plot of the relationship of the single-channel current amplitude and the holding potential. The symbols used are as follows: WT Kir2.1, ●; E224G, ▲; E299S, □; E224G–E299S, ▼. C, the single-channel conductance calculated from the slope of the plot in B. The mean and the standard deviation (n = 3) are plotted.
(Fig. 10A), the $I-V$ plots (Fig. 10B) and the $g-V$ plots (Fig. 10C) are shown. As we reported previously (Kubo et al. 1996), E299Q of sWIRK showed a slight outward rectification, and no clear sign of inward rectification. The conductance at hyperpolarized potentials decreased. A kind of gating mechanism that leads to channel closure at negative voltages might explain this property. In contrast, a weak but clear inward rectification remained in the triple mutants of Kir2.1 (D172N–E224G–E299S and D172N–E224G–E299Q). The accumulated data ($n = 3$) of the rectification index support these observations (Fig. 10D and E).

Figure 10. Macroscopic current recordings under two-electrode voltage clamp of the sWIRK mutant E179Q and two triple mutants of Kir2.1, D172N–E224G–E299S and D172N–E224G–E299Q, expressed in Xenopus oocytes.

A detailed explanation of A–C is given in Fig. 2. D and E, the ratio of the current amplitudes at +50 and at -100 mV was calculated as an index of the rectification intensity. The mean and the standard deviation ($n = 3$) of the indexes at 5 ms (D) and at 100 ms (E) from the beginning of the step pulses are shown.
DISCUSSION

The difference between the intensity of inward rectification of Kir2.1 and that of sWIRK (Kubo et al. 1996) could not be explained by differences at the two sites already reported, D172 and E224. Therefore, the presence of an additional site(s) was postulated that contributes to the extent of the inward rectification. E299 of Kir2.1 in the region after M2 was newly identified in this study. The phenotypes of E224G and E299S were highly similar each other, and differed from that of WT Kir2.1 in terms of the rectification and permeation properties as follows. (1) The inward rectification was less intense (Figs 2–4). (2) The activation upon hyperpolarization was slower (Fig. 5). (3) The decay of the outward current at depolarized potentials was slower (Fig. 6). (4) The sensitivity to block of the outward current by spermine was severely reduced (Fig. 8). (5) The single-channel conductance was smaller (Fig. 9). Qualitatively similar but more enhanced changes were observed in the double mutant E224G–E299S.

The structure of the inner vestibule of Kir2.1

The observations above are surprising because E299 is located almost half-way through the long cytoplasmic chain after M2, and is very distant from E224 on the linear sequence (Fig. 1). In mutagenesis studies including Cys-scanning mutagenesis, there is always an argument that the mutagenesis effect might be due to drastic changes of the overall structure. Although this possibility remains in the present study, the relatively intact channel function of E299S and E299Q suggests that it is more straightforward to assume that the negative charges of E299 and E224 contribute to the permeation pathway. As no clear change of phenotype from WT could be observed in the E244Q mutant, the specificity of the effects of mutations at E299 and E224 was supported. Taking together the present results and the previous reports on E224 (Yang et al. 1995; Taglialatela et al. 1995; Nichols & Lopatin, 1997), it is likely that the cytoplasmic chain after M2 is infolded to comprise the inner vestibule, which is continuous with the membrane-spanning pore region, and that both E224 and E299 contribute to the permeation pathway (Fig. 11).

This speculation was supported by two recent reports by Lu et al. (1999a,b). They demonstrated by a stoichiometric covalent modification study that the inner vestibule of Kir2.1 is formed by both cytoplasmic amino- and carboxyl-terminal domains, and that the vestibule is spacious enough to fit multiple methanthiosulfonate reagents. They estimated the diameter to be larger than 12 Å even at the depth of D172. This image is quite different from the resolved pore structure of the bacterial two-transmembrane-type K⁺ channel KcsA (Doyle et al. 1998). A difference between Kir2.1 and KcsA was also reported in the arrangement of the transmembrane segments (Minor et al. 1999) and in the pore structure (Thompson et al. 2000). Thompson et al. (2000) showed by mutagenesis study that residues beyond the selective filter of Kir2.1 channels contribute to permeation and blockage by extracellular cations, contrary to the expectations from the crystal structure of KcsA channels.

On the other hand, structural similarity between another inward rectifier channel Kir6.2 and KcsA was reported. Loussouarn et al. (2000) showed by Cys-scanning mutagenesis that at least three subunits are required to fit Cd²⁺. Thus, they proposed that the inner vestibule of the pore of the Kir6.2 channel is relatively narrow, similar to that of the KcsA channel, and that it has a state-dependent flexibility. A high structural similarity of the KcsA to the Kir6.2 channel was also reported by modelling based on the KcsA structure (Capener et al. 2000). Taken together, there is still some controversy on the structure of Kir channels. This discrepancy might be partly due to the differences between Kir2.1 and Kir6.2.

To have a conclusive understanding, we await the resolution of the structure of the Kir channel including the amino- and carboxyl-terminal regions in the crystal, because there is a possibility that the inner vestibule could be formed in a different way in the absence of these regions.

![Figure 11. Scheme to explain the location and the functional role of E224G and E299S, and the possible presence of two open states](image)
Differences between the functional role of D172 and those of E224 and E299

In the D172N mutant, the sensitivity to block by spermine was decreased and the activation phase at hyperpolarized potentials disappeared (Stanfield et al. 1994; Wible et al. 1994), suggesting that D172 has a strong energetic contribution to the binding of spermine in the channel pore. D172 was also reported to contribute to the binding of Mg2+ in the pore (Oishi et al. 1998). Thus, it is thought that D172 is a binding site or locates close to a binding site for these cytoplasmic blockers.

In the case of E224G, E299S and E224G–E299S, the sensitivity to block by spermine at depolarized potentials was much lower than that of WT (Fig. 8). In contrast with D172N, the activation speed at hyperpolarized potentials (unblocking) and the decay speed of the outward current at depolarized potentials (blocking) were slower than those of WT (Figs 5 and 6). Thus, we suggest that both the blocking and unblocking rates are decreased in these mutants, and that overall the effect of the mutation on the blocking rate is greater. If these residues were the pore-plugging binding site for spermine, the off-blocking rate in the mutants might be increased (i.e. the activation at hyperpolarized potentials might be faster). Therefore, E224 and E299 are unlikely to be the pore-plugging binding site for spermine. What then is the functional role of these sites?

One possibility is as follows. E224 and E299 are located at the inner vestibule and serve as an intermediate binding (and non-plugging) site which facilitates entry and exit of spermine to and from the final pore-plugging binding site located deeper in the pore. In the WT channel, the binding of spermine at the E224 and E299 level upon depolarization could result in an increase in the local concentration of spermine at the inner vestibule, and thus an increase in the on-blocking rate. Upon hyperpolarization, the presence of the intermediate binding (and non-plugging) site at the E224 and E299 level could facilitate the popping out of spermine from the D172 level, increasing the off-rate. Overall the effect on the blocking rate is much greater than the effect on the unblocking rate and so the blocking potency of spermine at depolarized potentials is dramatically enhanced with E224 and E299 in the WT. By mutating E224 and/or E299, the facilitating mechanisms described above for the entry and exit of spermine to and from the pore-plugging binding site may be affected, resulting in a dramatic decrease in both the blocking and the unblocking rates.

Two open states in the E224G–E299S double mutant

If the discussion on E224 and E299 in the previous section is true, there is a possibility that there exist two open states, namely one open state with no spermine in the inner vestibule and another conducting state with spermine at the E224 and/or E299 level. In the present study, data that are not contradictory with this possibility were obtained for the E224G–E299S mutant. As shown in Fig. 6, the fraction of the slowly decaying component of the outward current depended on the preceding prepulses. One possible interpretation, in which two open states were assumed, is shown schematically in Fig. 11. At less hyperpolarized potentials, spermine is released from the D172 level and the channel enters the open state. However, the released spermine still exists at the E224 and E299 levels. When depolarized back from the less hyperpolarized potential, the block of the outward current would proceed relatively fast, due to the high local concentration of spermine. In contrast, when strong hyperpolarization is applied, the cytoplasmic blocker would be released from the E224 and E299 level, and the local concentration of the blocker would be decreased. In this case, the blocking when depolarized would proceed only slowly. The same analysis could not be performed in WT Kir2.1, due to the very fast decay of the outward current. However, it is possible that a similar phenomenon to E224G–E299S occurs in WT Kir2.1.

In agreement with this discussion, Lee et al. (1999) also demonstrated the possibility that E224 functions as an intermediate binding (and non-plugging) point for spermine, by the combined use of spermine and a large spermine-related toxin which may reach the E224 level but not the D172 level. They argued that spermine binds first to the intrinsic gate close to E224, and then plugs the channel pore. Although Lee et al. (1999) assumed the intrinsic gate around E224 to be located in the cytoplasm, it is more likely that the carboxyl-terminal region including E224 (Yang et al. 1995) and E299 is folded to form the permeation pathway, because mutations at E224 and E299 affected the permeation properties (Fig. 9).

To explain the complex blocking behaviour of Kir2 channels by polyamines, a state model was proposed in which two binding sites for spermine were assumed (Lopatin et al. 1995; Guo & Lu, 2000). Lopatin et al. (1995) also produced a structural model in which two polyamines tandemly plug a long pore. It is possible that the deeper binding site is close to the D172 level and that the shallower one is close to the E224 and E299 levels. In the model by Lopatin et al. (1995), a state in which polyamine binds only to the shallower site was not postulated. It might correspond to the second open state discussed in the present study and by Lee et al. (1999).

Other amino acid residues for the control of inward rectification

There was a marked difference in the rectification property of E179Q of sWIRK (E179Q, G231, S311) and those of the triple mutants of Kir2.1 (D172N, E224G, E299S or Q) (Fig. 10), although both lack negatively charged amino acids at any of the three sites focused on in this study. It was also observed that the introduction of the S311E mutation to WT sWIRK (E179, G231, S311E)
or to the sWIRK E179Q mutant (E179Q, G231, S311E) did not significantly increase inward rectification (data not shown). The G231E mutation in WT sWIRK also did not clearly strengthen the inward rectification (data not shown). Taken together, these results suggest that the three sites are still insufficient to fully explain the intensity of the inward rectification, and that another important residue(s) for the strong inward rectification is missing in sWIRK. An amino acid residue just before M1 (M84 of Kir2.1), which was reported to be critical for the unblocking process (Ruppersberg et al. 1996), could not explain the difference between sWIRK and Kir1.1 because both of them have Lys at this residue. Further studies by systematic chimeras and mutants would provide a clue to a more conclusive understanding of this.


References


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