It is known that rectification of currents through the inward rectifier K⁺ channel (Kir) is mainly due to blockade of the outward current by cytoplasmic Mg²⁺ and polyamines. Analyses of the crystal structure of the cytoplasmic region of Kir2.1 have revealed the presence of both negatively (E224, D255, D259, and E299) and positively (R228 and R260) charged residues on the wall of the cytoplasmic pore of Kir2.1, but the detail is not known about the contribution of these charged residues, the positive charges in particular, to the inward rectification. We therefore analyzed the functional significance of these charged amino acids using single/double point mutants in order to better understand the structure-based mechanism underlying inward rectification of Kir2.1 currents. As a first step, we used two-electrode voltage clamp to examine inward rectification in systematically prepared mutants in which one or two negatively or positively charged amino acids were neutralized by substitution. We found that the intensity of the inward rectification tended to be determined by the net negative charge within the cytoplasmic pore. We then used inside-out excised patch clamp recording to analyze the effect of the mutations on blockade by intracellular blockers and on K⁺ permeation. We observed that a decrease in the net negative charge within the cytoplasmic pore reduced both the susceptibility of the channel to blockade by Mg²⁺ or spermine and the voltage dependence of the blockade. It also reduced K⁺ permeation; i.e., it decreased single channel conductance, increased open-channel noise, and strengthened the intrinsic inward rectification in the total absence of cytoplasmic blockers. Taken together, these data suggest that the negatively charged cytoplasmic pore of Kir electrostatically gathers cations such as Mg²⁺, spermine, and K⁺ so that the transmembrane pore is sufficiently filled with K⁺ ions, which enables strong voltage-dependent blockade with adequate outward K⁺ conductance.

INTRODUCTION

Inward rectification of the current through the inwardly rectifying K⁺ channel (Kir) is reportedly due to blockade by cytoplasmic Mg²⁺ (Matsuda et al., 1987; Vandenberg, 1987; Matsuda, 1988) and polyamines (Ficker et al., 1994; Lopatin et al., 1994; Fakler et al., 1995; Ishihara et al., 1996; Nichols and Lopatin, 1997). The strong voltage dependence and high susceptibility to blockade of Kir plays a key role in organizing I_Ki within the cardiac action potential (Luo and Rudy, 1994; Matsuoka et al., 2003). After the isolation of Kir1.1 (Ho et al., 1993) and Kir2.1 cDNA (Kubo et al., 1993a), the structural elements that determine the properties of inward rectification were identified using mutagenesis. It was first found that D172, located in the second transmembrane region of Kir2.1, was an amino acid residue shown to contribute to inward rectification: a D172N mutant had reduced susceptibility to blockade by both cytoplasmic polyamines and Mg²⁺ (Lu and MacKinnon, 1994; Stanford et al., 1994; Wible et al., 1994), and D172 was thought to make a strong energetic contribution to the binding of the blockers. Another amino acid residue, S165, also in the second transmembrane region, just below the GYG selectivity filter, has been shown to be important for blockade by Mg²⁺, but not by polyamines (Fujii and Kubo, 2002). Thus, it appears that the cytoplasmic blockers plug the permeation pathway at sites deep within the transmembrane pore, below the selectivity filter. On another aspect, E224 and E299, located outside the transmembrane region, appear to play different roles in inward rectification. Both E224G and E299S mutants reportedly show reduced susceptibility to blockade by cytoplasmic blockers (Taglialetela et al., 1995; Yang et al., 1995; Kubo and Murata, 2001), and based on analysis of the electrophysiological properties of E224G, E299S, and E224G/E299S, it was suggested that these residues foster blockade by mediating increases in the local spermine concentration (Kubo and Murata, 2001). Xie et al. (2002, 2003) also analyzed the

Correspondence to Yuichiro Fujiwara or Yoshihiro Kubo:
yuichiro@nips.ac.jp or ykubo@nips.ac.jp

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Abbreviations used in this paper: EGFP, enhanced green fluorescent protein; Kir, inwardly rectifying K⁺ channel.
functional significances of E224 and E299 using synthesized polyamines of various length and suggested that spermine binds to these residues, thereby contributing to a surface charge screening effect. This suggests that E224 and E299 coordinate the intermediate binding of spermine without actually occluding K+ permeation, which in turn increases the susceptibility to spermine blockade at a deeper site, near D172. On the other hand, Guo and Lu (2003) reported that polyamines block the permeation pathway at the level of E224 and E299. Thus the functional significance of these amino acid residues within the cytoplasmic pore has not yet been conclusively defined.

In recent years, the crystal structures of several Kir channels have been solved. The structure of the cytoplasmic region of Kir3.1 was determined by Nishida and MacKinnon (2002), and the structure of bacterial Kir channel was determined by Kuo et al. (2003). In addition, Pegan et al. (2005) recently succeeded in resolving the structure of the cytoplasmic region of Kir2.1. These crystal structures have revealed that Kir has a long permeation pathway comprised of transmembrane and cytoplasmic pore regions. As expected, E224 and E299 are located on the wall of the cytoplasmic pore along with two other negatively charged residues, D259 and D255 (Fig. 1), and Pegan et al. (2005) reported that this di-aspartate cluster (D259 and D255) is also important in determining the extent of inward rectification. Not expected was the finding that there were also positively charged residues (R228 and R260) on the wall, the functional significance of which is unknown.

With that as background, our aim in the present study was to clarify the functional significance of the charged amino acid residues on the wall of the cytoplasmic pore of Kir2.1 and to better understand the structure-based mechanism underlying inward rectification of Kir2.1 currents. To that end, we made single and double point mutants by substituting the aforementioned charged amino acids and systematically analyzed the inward rectification properties of the resultant channels. Using two-electrode voltage clamp, we analyzed the intensity of inward rectification of currents through the mutant channels, as well as the activation of the inward and the decay of the outward currents. Using inside-out patch clamp recording, we analyzed the mutants’ susceptibility to cytoplasmic blockers and their voltage dependence. As the characteristic properties of the inward rectification of Kir channel currents are reportedly linked to the permeant K+ ions themselves (Hagiwara and Takahashi, 1974; Hagiwara and Yoshii, 1979; Oliver et al., 1998; Pearson and Nichols, 1998; Spassova and Lu, 1998), we also analyzed the permeation properties of the mutants; the single channel properties, and the intrinsic inward rectification in the total absence of the blockers.

**MATERIALS AND METHODS**

**In Vitro Mutagenesis**
The single point mutants were made with a QuickChange site-directed mutagenesis kit (Stratagene) using mutated oligonucleotide DNA primers and wild-type (WT) Kir2.1 cDNA (Fujisawa and Kubo, 2002). The introduction of each mutation was confirmed by sequencing with a BigDye Terminator Sequencing Kit (Applied Biosystems) and an automatic DNA sequencer (ABI type 310). The electrophysiological properties of two independent mutant clones were confirmed to be identical. Double-point mutants were made using single-point mutants as templates; triple mutants were prepared using double-point 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, type 1, Sigma-Aldrich) and then injected with ~50 nl of cRNA solution prepared from linearized plasmid DNA using an RNA transcription kit (Stratagene), after which they were incubated for 2–4 d at 17°C in frog Ringer solution supplemented with 20 mM KCl (Kubo et al., 1995a). All experiments were conducted in accordance with the guidelines of the Animal Care Committees of National Institute for Physiological Sciences.

Macroscopic currents were recorded using two-electrode voltage clamp with a bath-clamp amplifier (OC-725C, Warner Co.). Stimulation and data acquisition and analysis were done on a Pentium-based computer using Digidata 1322A and pCLAMP software (Axon Instruments, Inc.). Intracellular glass microelectrodes were filled with 3 M potassium acetate with 10 mM KCl (pH 7.2). The microelectrode resistances ranged from 0.1 to 0.3 MΩ. Two Ag–AgCl pellets (Warner Co.) were used to pass the bath current and sense the bath voltage. The voltage-sensing electrode was placed near the oocyte (~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 ~200 Ω (Sabirov et al., 1997). As the measured current at the most hyperpolarized potential was 25 µA in the largest case, and was mainly <20 µA, the voltage-clamp error due to this series resistance was estimated to be no more than 5 mV and mostly <4 mV. This error, which was not compensated in the experiments, did not change the conclusions drawn from the comparison of WT and mutant channels.

For the data in Figs. 2–4 and Tables I and II, the recording bath solution contained 8 mM KCl, 80 mM NMDG, 70 mM HCl, 3 mM MgCl2, 5 mM HEPES, and 2 mM KOH (pH 7.4). All recordings in this work were performed while the preparation was superfused with bath solution (7 ml/min) at room temperature (20–23°C). By assuming an intracellular K+ concentration ([K+]i) of 80 mM, the EK was calculated to be −52 mV; however, the exact values of [K]i and EK for each oocyte are unknown. The EKs used for calculation of the chord conductance (Fig. 2) were therefore adjusted to yield continuous G-V plots. The adjusted EKs ranged from −54 to −49 mV.

Data from the same batch of oocytes were used for comparison of phenotypes (Figs. 2–4, Fig. 9 A, and Tables I and II) because properties such as inward rectification and blocking speed differ significantly from batch to batch. Similar tendencies were reproducibly observed in five batches of oocytes.

**Expression in HEK293T Cells**
The cDNAs for WT and mutant Kir2.1 were subcloned into the pCIXN2 expression vector (Niwa et al., 1991). The plasmid DNA
was then cotransfected into HEK293T cells (human embryonic kidney cell line) with a transfection marker, enhanced green fluorescent protein (CLONTECH Laboratories, Inc.; 1/15 the amount of plasmid DNA), using Lipofectamine Plus (GIBCO BRL), as instructed by the manufacturer. The cells were then cultured for 24 h in Dulbecco’s modified Eagle’s medium with 10% bovine calf serum. The transfectants were dissociated 12–15 h later by treatment with 0.025% trypsin in Ca2+- and Mg2+-free PBS and reseded on coverslips at a relatively low density. The reseding was done to obtain well-isolated cells and to facilitate successful G1 seal formation. We confirmed previously that the trypsin treatment did not change the electrophysiological properties of the expressed channels (Kubo and Murata, 2001; Fujiwara and Kubo, 2002). Electrophysiological recordings were performed 7–24 h after reseding, which corresponds to 18–35 h after transfection.

Macroscopic Current Recordings in HEK293T Cells

A coverslip with HEK293T cells was placed in a recording chamber containing bath solution (see below) on the stage of an inverted fluorescence microscope (IX70, Olympus), and the transfected cells were identified by the fluorescent signal from the cotransfected green fluorescent protein. Macroscopic currents were then recorded in the excised inside-out patch configuration using an Axopatch-1D amplifier. The resistance of the patch pipettes ranged from 1.0 to 1.5 MΩ, 60–80% of the voltage error due to the series resistance was compensated by a circuit in the amplifier. The high expression level achieved by the combination of HEK293T cells and the pCWN2 expression enabled macroscopic current recording using standard-sized patch pipettes. The recorded currents were low-pass filtered at 1 kHz by a Bessel filter built into the amplifier and digitized at 5 kHz. In all experiments, the recording chamber was perfused with the bath solution at a rate of 12–15 h later by treatment with 0.025% trypsin in Ca2+- and Mg2+-free PBS and reseded on coverslips at a relatively low density. The degree of rundown that occurred during the recordings used for determining I-V relationships was monitored by reapplying step pulses, and data with apparent rundown were discarded. For polyamine-free experiments, recordings were started after intensive perfusion of the bath (intracellular side) solution for 10–15 min to completely wash out endogenous polyamines.

Single-channel Recordings in Xenopus Oocytes

For single-channel recordings, the vitelline membrane was peeled off by bathing the oocytes in a hyperosmolar solution for 5–10 min (Kubo et al., 1993a). The patch pipettes were prepared from borosilicate glass (Warner Instruments) using a P97 horizontal puller (Sutter) and a fire polisher (Narishige). Pipette resistance ranged from 1 to 2 MΩ. Recordings were made in the cell-attached configuration using an Axopatch 1D amplifier (Axon Instruments, Inc.). The recorded currents were low pass filtered at 1 kHz using a Bessel filter built into the amplifier and digitized at 5 kHz. The pipette (extracellular) and bath solutions contained 116.88 mM KCl, 2 mM EDTA, 2.83 mM KH2PO4, 7.17 mM KHPO4, and 5.95 mM KOH (pH 7.2).

Leak Subtraction

Leak subtraction was not done for the data summarized in Figs. 2–6 and Fig. 9 (A and B). But when assessing the intensity of inward rectification, it is critical to avoid contamination of the leak component. We therefore avoided setting EK = 0 mV because that would be difficult to discriminate between the outward Kir currents and the leak component. We instead used mixture of K+ and NMDG in the extracellular solution to set EK at, for example, −50 mV. We then carefully monitored the reversal potential during the recordings and judged whether the leak level was ignorable by confirming that the reversal potential remained at the expected level. Only data that satisfied these criteria were used for analysis. Because we recorded intrinsic rectification using symmetrical recording solutions (EK ≈ 0 mV) in the experiments summarized in Fig. 8 and Fig. 9 C, leak subtraction was necessary. To do so, we added 20 mM Mg2+ to the bath intracellular solution to cause the Kir2.1 channels to rundown; and then after 5 min, a background current was recorded as a template for leak subtraction.

Data Analysis

Data were analyzed using Clampfit 8 and 9 (Axon Instruments, Inc.), Igor Pro (WaveMetrics, Inc.), and KyPlot (KyensLab, Inc.) software. The means of two groups were compared using Student’s t test, while pairs of means among three or more groups were compared using Tukey’s test, and the difference among the means of multiple groups was tested using one-way ANOVA.

The susceptibility of WT and mutant channels to blockers was evaluated based on Kd values, which were calculated by fitting the data with Hill’s equation. The voltage dependence of the blockade was analyzed using Kd values obtained at various voltages using the following equation (Hille, 2001):

$$\log_{\sigma} \left( \frac{Kd_{(A)}}{Kd_{(B)}} \right) = \frac{2.303RT}{F}\alpha \left( E_{(A-B)} \right),$$

where R is the Gas constant, T is the absolute temperature, and F is Faraday’s constant. In this study, the z values were 2 and 4 for the Mg2+ and spermine blockades, respectively, and σ was calculated by fitting voltage-Kd plots with a linear line (Figs. 5, 6, and 9).

Single-channel conductance and the open-close event histogram were analyzed using Clampfit 9 (Axon Instruments, Inc.) and Igor Pro.
∆ and negative (functional mutants were compared with images of positive (WT) camera (Carl Zeiss MicroImaging, Inc.). The images of the non-Fluorescent images were acquired using an AxioCam cooled CCD slide glasses using PermaFluor mounting medium (Shandon). After 24 h, they were fixed by 4% paraformaldehyde and then rinsed three times with PBS. The coverslips were mounted on Stockklausner et al., 2001), and we used it as a negative control of infected with the EGFP-tagged constructs using the Lipofectamine Plus (GIBCO BRL) according to the manufacturer’s instructions.

Detection of Cell Surface Expression
We analyzed the surface expression of nonfunctional mutants of Kir2.1 by detecting fluorescent signals attached to the channels under fluorescent microscope. Enhanced green fluorescent protein (EGFP)–tagged constructs of Kir2.1 WT, nonfunctional mutants, and ΔFCYENEV were made by PCR using primers that include EcoRI(5′) and BamHI(3′) sites. ΔFCYENEV is a deletion mutant of the trafficking motif (F374CYENEV380) (Ma et al., 2001; Stockklausner et al., 2001; Kuo et al., 2003; Pegan et al., 2005), but the detail has not been analyzed intensively regarding their contributions to the susceptibility to intracellular blockers, the single channel properties, and the intrinsic inward rectification. To address this issue, we systematically substituted these residues such that they were nearly unchanged in size, but were made electrostatically neutral (i.e., E→Q, D→N, and R→Q). As an initial step, we analyzed the electrophysiological properties of mutants using two-electrode voltage clamp with a *Xenopus* expression system.

**RESULTS**

**Charged Amino Acids on the Wall of the Inner Vestibule of Kir2.1**

X-ray crystallographic analysis has shown that Kir has a long permeation pathway that extends through a cytoplasmic region and a transmembrane region; the scheme drawn in Fig. 1 A refers to the structures of Kir2.1, Kir3.1, and KirBac1.1 (Nishida and MacKinnon, 2002; Kuo et al., 2003; Pegan et al., 2005). Multiple alignment of the amino acid sequences of Kir family proteins are shown in Fig. 1 B; the amino acid residues facing the aqueous cavity of the cytoplasmic pore are colored yellow, negatively charged ones are colored red, and positively charged ones are colored blue. The surface of the permeation path through the cytoplasmic pore of one Kir2.1 subunit is illustrated in Fig. 1 C; the amino acid residues facing the aqueous pore are shown as sticks, the others as cartoons. Note that within the structure of Kir2.1, a cluster of hydrophilic residues lays between the hydrophobic residues F274 and M301. With the channel oriented as in the figure, the negatively charged residues E224 and E299 are located on the top of the cluster, and the positively charged residues R228 and R260 are below them, as is negatively charged D259. Negatively charged D255 is located at the bottom of the cytoplasmic pore, far below the transmembrane region.

E224 and E299 have been investigated previously using E224G and E299S mutants (Kubo and Murata, 2001; Guo and Lu, 2003; Xie et al., 2004). The function of the other charged residues (D255, D259, R228, and R260) have been also studied (Nishida and MacKinnon, 2002; Kuo et al., 2003; Pegan et al., 2005), but the detail has not been analyzed intensively regarding their contributions to the susceptibility to intracellular blockers, the single channel properties, and the intrinsic inward rectification. To address this issue, we systematically substituted these residues such that they were nearly unchanged in size, but were made electrostatically neutral (i.e., E→Q, D→N, and R→Q). As an initial step, we analyzed the electrophysiological properties of mutants using two-electrode voltage clamp with a *Xenopus* expression system.

**Contribution of Charged Amino Acids to Inward Rectification**

We examined the electrophysiological effects of neutralizing the charged residues within the cytoplasmic pore of Kir2.1 (Fig. 2 and Table I). Macroscopic
currents recorded in 10 mM K\(^+\) are shown in Fig. 2 A, while the I-V relationships are shown in Fig. 2 B. As an index of the intensity of the inward rectification, the ratios of the current amplitudes measured 100 ms after the onset of step pulses to +50 and −100 mV were calculated (Fig. 2, C and D).

We first describe the effects of neutralizing the negative charges. Inward rectification was weakened in all of the single point mutants (E224Q, D259N, and E299Q) (Fig. 2 C and Table I, group 1), especially in E224Q, which showed a large outward current (Fig. 2 A). Addition of E299Q or D259N mutation to E224Q did not affect inward rectification remarkably. We also analyzed the functional significance of D255, located at the bottom in the cytoplasmic aqueous pore (Fig. 1 C) and found that D255N mutation did not affect inward rectification remarkably. We also analyzed the functional significance of D255, located at the bottom in the cytoplasmic aqueous pore (Fig. 1 C) and found that D255N mutation had little effect on the intensity of inward rectification (Table I, group 2). Pegan et al. (2005) reported that both D259 and D255 were important for inward rectification. This discrepancy might reflect the difference in the mutants used: they used D255R, whereas we used D255N. These results suggest that negative charges (E224, E299, and to a minor extent D259) on the wall of the cytoplasmic pore are key determinants of the strength of the inward rectification of the currents through Kir2.1.

We next analyzed the effects of neutralizing the positive charges within the cytoplasmic pore of Kir2.1. Data obtained with R228Q, R260Q, and R228Q/R260Q using two-electrode voltage clamp are summarized in Table I, group 3. Although both R228Q and R260Q were highly expressible, no macroscopic currents through the R228Q/R260Q double mutant were observed. R228Q and R260Q currents showed strong inward rectification, and the difference from WT was subtle. We suspected that the significance of these positively charged residues was masked by the presence of adjacent negative charges, so we made the E224Q/R228Q and E224Q/R260Q double mutants and compared them with E224Q, E224Q/R260K, and E224Q/R260Q (Fig. 2 and Table I, group 4). Introduction of the additional R228Q mutation substantially increased the intensity of the inward rectification of E224Q, whereas the R228K mutation had no clear effect (Fig. 2 C and Table I, group 4). Similar tendencies were observed with E224Q, E224Q/R260Q, and E224Q/R260K. To further examine the electrostatic effect of these residues, we also made two inversely charged single point mutations, R228E and R260E, but the latter carried no detectable current. R228E mutation strengthened the
intensity of the inward rectification of E224Q, and the effect was stronger than that seen with E224Q/R228Q (Table I, group 5).

At hyperpolarized potentials, these mutations also affected the G-V relationship (Fig. 2 E). The macroscopic channel conductance recorded using two-electrode voltage clamp was clearly influenced by introducing the E224Q mutation: it did not reach saturation, even at the most hyperpolarized potential recorded (−160 mV). The macroscopic conductance of E224Q/R228Q saturated at membrane potentials more hyperpolarized than −130 mV, while the G-V relationship of E224Q/R228K did not saturate at −160 mV, as was observed with E224Q (Fig. 2 E). These changes in the macroscopic conductance suggest that K⁺ permeation and the unbinding of cytoplasmic blockers at hyperpolarized potentials were influenced by the neutralization of charged residues within the cytoplasmic pore of Kir2.1.

To further confirm the electrostatic effect of these residues, we also analyzed the effect of R→Q neutralized mutation on other background mutants: E299Q, D259N, E224G, or E229S (Table I, groups 6 and 7). Addition of an R228Q or an R260Q mutation also strengthened inward rectification of the backgrounds (Table I, groups 6 and 7). In some cases, the mutant channels carried no detectable current, even when

| TABLE I |

| Summarized Data on the Contribution of Charged Residues to the Intensity of Inward Rectification Obtained using Two-electrode Voltage Clamp |

| Group | Mutant | Inward rectification intensity | Statistical analysis |
|-------|--------|-------------------------------|----------------------|
| 1     | WT     | 0.034 ± 0.003 (n = 6)         | control             |
|       | E224Q  | 0.560 ± 0.026 (n = 5)         | ***                 |
|       | D259N  | 0.047 ± 0.003 (n = 6)         | *                   |
|       | E299Q  | 0.082 ± 0.013 (n = 5)         | *                   |
|       | E224Q/E299Q | 0.534 ± 0.025 (n = 8) | ***                 |
|       | E224Q/D259N | 0.606 ± 0.029 (n = 5) | ***                 |
|       | E299Q/D259N | ND                         |                     |
|       | E224Q/E299Q/D259N | ND                     |                     |
| 2     | WT     | 0.044 ± 0.005 (n = 5)         | control             |
|       | D255N  | 0.047 ± 0.007 (n = 5)         | P > 0.05            |
|       | E224Q  | 0.668 ± 0.013 (n = 4)         | control             |
|       | E224Q/D255N | 0.604 ± 0.027 (n = 5) | P > 0.05            |
| 3     | WT     | 0.040 ± 0.007 (n = 5)         | control             |
|       | R228Q  | 0.082 ± 0.009 (n = 7)         | *                   |
|       | R260Q  | 0.074 ± 0.010 (n = 7)         | P > 0.05            |
|       | R228Q/R260Q | ND                         |                     |
| 4     | E224Q  | 0.56 ± 0.03 (n = 4)           | ***                 |
|       | E224Q/R228K | 0.48 ± 0.02 (n = 5)         |                     |
|       | E224Q/R228Q | 0.18 ± 0.01 (n = 4)         | control             |
|       | E224Q/R260K | 0.36 ± 0.02 (n = 3)         | **                  |
|       | E224Q/R260Q | 0.18 ± 0.02 (n = 3)         | control             |
| 5     | E224Q  | 0.46 ± 0.02 (n = 5)           |                     |
|       | E224Q/R228Q | 0.20 ± 0.01 (n = 4)         | control             |
|       | E224Q/R228E | 0.15 ± 0.01 (n = 4)         | **                  |
| 6     | E299Q  | 0.096 ± 0.016 (n = 6)         | control             |
|       | E299Q/R228Q | 0.046 ± 0.006 (n = 6) | **                  |
|       | E299Q/R260Q | ND                         |                     |
|       | D259N  | 0.065 ± 0.008 (n = 5)         | control             |
|       | D259N/R228Q | ND                         |                     |
| 7     | E224G  | 0.30 ± 0.02 (n = 5)           | control             |
|       | E224G/R228Q | 0.14 ± 0.01 (n = 6)         | ***                 |
|       | E224G/R260Q | 0.23 ± 0.03 (n = 5)         | *                   |
|       | E299S  | 0.40 ± 0.05 (n = 5)           | control             |
|       | E299S/R228Q | 0.12 ± 0.02 (n = 6)         | ***                 |
|       | E299S/R260Q | 0.36 ± 0.05 (n = 5)         | P > 0.05            |
| 8     | WT     | 0.036 ± 0.004 (n = 5)         | control             |
|       | H229Q  | 0.046 ± 0.006 (n = 4)         | P > 0.05            |
|       | H229K  | 0.099 ± 0.020 (n = 4)         | **                  |

For comparison of the phenotypes, mean values for WT and mutant channels were analyzed statistically using Tukey’s test or Student’s t-test (***, P < 0.001; **, P < 0.01; *, P < 0.05). ND denotes channels that were expressed only poorly or not at all, and so could not be analyzed. Data from the same batch of oocytes were used for comparison of phenotypes.
the oocytes were injected with highly concentrated (>20×) cRNA. There is a possibility that they are not expressed on the membrane surface. We therefore attached a tag of EGFP to the mutants and analyzed fluorescence signal on the membrane surface of transfected L929 cells. Fluorescent signals were clearly detected on the cell surface for WT-EGFP (positive control), while they were detected only from the cytoplasm for ΔFCYENEV-EGFP (negative control) (see Fig. S1, available at http://www.jgp.org/cgi/content/full/jgp.200509434/DC1). The signals of all nonfunctional mutants were clearly detected on the cell surface similarly to those of WT-EGFP (Fig. S1). These results suggest that the lack of function is not due to the problem of surface expression, implying that the role of the charged amino acid residues on K⁺ permeation is critical. As all double mutants on the background of E224Q showed sufficient functional expression, we used, in this study, E224Q as a representative background mutant that showed weak inward rectification.

Contribution of Charged Amino Acids to Activation

The activation phases upon hyperpolarization of WT Kir2.1 and the mutants with neutralized charges were recorded in 10 mM K⁺. The current traces obtained with WT at −70 to −150 mV, with E224Q, E224Q/R228K, E224Q/R228Q, and E224Q/R260Q at −100 to −160 mV are shown in Fig. 3 A. The time constants obtained by fitting the data with a single exponential function are plotted in Fig. 3 (B and C); note that the inactivating component of the WT, E224Q/R228Q, and E224Q/R260Q currents (Fig. 3 A) was ignored in the fitting. At strongly hyperpolarized potentials, the activation phases of WT, D259N, and E299Q were too rapid to be separated precisely from the capacitative current, so they were not used for analysis. Upon introduction of E224Q mutation, the activation phases were noticeably slower (Fig. 3 A). We also analyzed neutralized mutants of the negative charges (E224, E299, and D259) and observed that they slow, with different extents, the activation of the inward current (Fig. 3 B and Table II, group 1). Addition of R228Q mutation to E224Q markedly accelerated activation of the inward current, whereas addition of R228K mutation had no significant effect (Fig. 3, A and C). Likewise, addition of R260Q mutation also markedly accelerated activation of the inward current, whereas addition of R260K mutation also had a significant but much smaller effect. It was also noteworthy that the voltage dependences of the time constants were strengthened by R228Q or R260Q mutation, contrary to the changes by neutralization of negative charges (Fig. 3, B and C). Taken together, these findings indicate that neutralization of positive charges on the wall of the

Figure 3. Comparison of the hyperpolarization-evoked activation kinetics in WT Kir2.1 and mutants. (A) Inward currents recorded using two-electrode voltage clamp with Xenopus oocytes at the indicated membrane voltages in 10 mM K⁺. (B) The activation phases were fitted with a single exponential function, and the time constants of the fittings are plotted. The symbols used are as shown in the figure. Bars depict means ± SEM (n = 6–8). (C) Same as in B; in this case, data obtained from E224Q, E224Q/R228K, E224Q/R228Q, E224Q/R260K, or E224Q/R260Q were compared. Bars depict means ± SEM (n = 3–4).
cytoplasmic pore of Kir2.1 accelerated the activation of the inward current.

Contribution of Charged Amino Acids to the Decay of the Outward Current

We next analyzed the time courses of the tail currents at +50 mV after depolarization from various prepotentials. In Fig. 4 A, recordings of the outward currents are shown using a magnification scale normalized by the amplitude of the inward current at −120 mV. None of the currents could be fitted with a single exponential function. With the exception of the current through E224Q/D259N, they could be fitted with a double-exponential function, and representative time constants are plotted in Fig. 4 (B and C). The decay of the outward current through WT Kir2.1 was too fast to be analyzed separately from the capacitative current. By contrast, much larger in amplitude and slowly declining tail currents were observed with E224Q (Fig. 4, A and B). We also analyzed neutralized mutants of other negative charges (E2299 and D259) and observed that they all slow the decay of the outward current (Fig. 4 B and Table II, group 1). The tail currents carried by E224Q/R228Q and E224Q/R260Q were smaller in amplitude and declined faster than those through E224Q (Fig. 4, A and C), while the currents through E224Q/R228K and E224Q/R260K were similar to those through E224Q (Fig. 4, A and C). It was also noteworthy that the tail current amplitudes increased after prepulses of strong hyperpolarized potentials in some mutants, apparently in E224Q, E224Q/R228K, and E224Q/R260K (Fig. 4 A). One possible interpretation about these phenomena is that these may be caused by the increase of the release of polyamines from the intermediate binding site in the cytoplasmic pore level when strong hyperpolarization is applied to these mutants (Kubo and Murata, 2001); and therefore current amplitudes would increase extra.

Contribution of Charged Residues to Blockade by Mg²⁺

Given that inward rectification of Kir is caused by blockade of the permeation pathway by cytoplasmic Mg²⁺ or polyamines, we next determined how the susceptibility to blockade by these inhibitors is affected by mutation. The analysis summarized below was performed using WT Kir2.1 and R228Q, D259N, E224Q, E224Q/R228Q, and E224Q/D259N as representative mutants. Macromolecular currents were recorded from inside-out membrane patches excised from HEK293T cells expressing WT or mutant Kir2.1 (Fig. 5); cells were held at −50 mV, and 500-ms test pulses from +70 to −120 mV were applied in 10-mV decrements. Representative

### Table II

| Group | Mutant | Activation speed | Decaying speed of the outward current |
|-------|--------|------------------|--------------------------------------|
|       |        | Single exp fitting (test −100 mV) | Control | Fast component (pre −150 mV) | Slow component (pre −150 mV) |
|       |        | Statistic analysis |          | Statistical analysis | Statistical analysis |
| 1     | WT     | 2.23 ± 0.10 ms (n = 4) | control | ND (too fast) | ND (too fast) |
|       | E224Q  | 15.78 ± 0.49 ms (n = 6) | *** control | 37.2 ± 2.2 ms (n = 6) | 4526 ± 207 ms (n = 6) |
|       | D259N  | 1.63 ± 0.14 ms (n = 4) | P > 0.05 | 16.3 ± 1.1 ms (n = 6) | * 93 ± 3 ms (n = 6) *** |
|       | E299Q  | 5.29 ± 0.92 ms (n = 4) | ** | 1.8 ± 0.6 ms (n = 6) | ** 28 ± 1 ms (n = 6) *** |
|       | E224Q/D259N | 17.82 ± 0.97 ms (n = 6) | *** P > 0.05 | 72.9 ± 2.0 ms (n = 6) | ** 6009 ± 175 ms (n = 6) *** |
|       | E224Q/D259N | 42.39 ± 2.99 ms (n = 6) | *** *** | 508.2 ± 16.6 ms (n = 5) | *** (−) |

Mean values were compared statistically between WT and mutant channels using Tukey’s test or Student’s t-test (***, P < 0.001; **, P < 0.01; *, P < 0.05). ND denotes phenotypes that were too fast to be analyzed; (−), not detectable.
series of recordings made in the nominal absence of polyamines and in the presence of various concentrations of free Mg\(^{2+}\) (calculated as described in MATERIALS AND METHODS) are shown in Fig. 5. In the absence of polyamines and Mg\(^{2+}\), the I-V relationships of WT and the mutants were not linear under these experimental conditions; for example, the I-V relationship for WT showed outward rectification consistent with the GHK equation, and that of E224Q showed intrinsic inward rectification.

At depolarized potentials, the outward current through WT was blocked almost instantaneously, even by low concentrations of Mg\(^{2+}\) (3 μM), while D259N currents were blocked more slowly by 3 μM Mg\(^{2+}\) (Fig. 5 A), suggesting that D259N was slightly less susceptible to blockade by Mg\(^{2+}\) than WT. This slow blockade of outward currents by Mg\(^{2+}\) was also observed with the other mutants. With E224Q, for example, a very slow blockade was observed in the presence of 300 μM Mg\(^{2+}\); this blockade was accelerated by the introduction of an R228Q mutation (Fig. 5 A), and was slowed by introduction of a D259N mutation (current traces not shown).

We then analyzed the dose–block relationship by measuring the macroscopic conductances recorded from a single patch in a series of Mg\(^{2+}\) concentrations; data showing apparent rundown were discarded. The normalized dose–block relationships for currents measured 500 ms after the onset of the step pulses are plotted in Fig. 5 B and fitted with Hill’s equation; the Kd values (n = 3) obtained at each voltage were well fitted by a linear function as shown in Fig. 5 C. The voltage dependence of the Kd values varied among mutants (Fig. 5, B and C), but were always linearly correlated. The means calculated from the fitted lines in Fig. 5 C are shown in Fig. 5 D, and the σ values (calculated as described in MATERIALS AND METHODS) are shown in Fig. 5 E.

Contribution of Charged Residues to Blockade by Spermine

We next determined how the susceptibility to blockade by intracellular spermine was affected by the mutations. Macroscopic currents were recorded from inside-out membrane patches excised from HEK293T cells expressing WT or mutant Kir2.1 using 3-s step pulses. Representative series of recordings showing spermine blockade in the absence of Mg\(^{2+}\) are shown in Fig. 6. Outward currents through WT and D259N were blocked.
Figure 5. Comparison of the susceptibility of macroscopic currents through WT Kir2.1 and mutants to blockade by intracellular Mg$^{2+}$. (A) Macroscopic currents recorded in the presence of the indicated concentrations of Mg$^{2+}$, from excised patches of HEK293T cells expressing WT Kir2.1 or mutants. The concentrations of Mg$^{2+}$, are shown on the top; note the difference in the concentration ranges used for WT and E224Q. K$^+$, in the pipette was 20 mM, and K$^+$, in the bath was 140 mM. The calculated $E_K$ was $-49$ mV. The holding potential was $-50$ mV, and step pulses from $+70$ to $-120$ mV were applied in 10-mV decrements. (B) Normalized dose–block relationships derived from data in A; values were measured 500 ms after the onset of each step pulse. (C) Relationships between the $K_d$ and voltage for Mg$^{2+}$ blockade of outward K$^+$ currents. Data obtained from three patches were plotted for each channel, and each set was fitted with a line. (D) Relationships between the voltage and the mean of the fitted lines in C, and the dashed lines indicate SEMs. (E) $Z_d$ values of each channel calculated from the slopes of the lines in C. Mean values were compared statistically between WT and the mutants using Tukey’s test (***, P < 0.001; * P < 0.05; NS, P > 0.05); statistical comparison of the data obtained from E224Q and E224Q/R228Q is also shown. (F) Values of the Hill coefficient for each channel. Bars depict means ± SEM ($n = 3$). The symbols used are as indicated in the figure.
Figure 6. Comparison of the susceptibility of macroscopic currents through WT Kir2.1 and mutants to blockade by intracellular spermine.

(A) Macroscopic currents recorded in the presence of the indicated concentrations of spermine from excised patches of HEK293 T cells expressing WT Kir2.1 or the indicated mutant. Note the difference in the concentration ranges used for WT and mutants, as well as the difference in the time scales. K\textsuperscript{\textit{o}} and K\textsuperscript{\textit{i}} were 20 mM and 140 mM, respectively. The holding potential was −50 mV, and step pulses from +70 to −120 mV were applied in 10-mV decrements. (B) Normalized dose-block relationships; values were measured 800 ms (in WT and D259N) or 3,000 ms (in E224Q and E224Q/R228Q) after the onset of each step pulse. (C) Relationships between K\textit{d} and voltage for spermine blockade of outward K\textsuperscript{+} currents. Data obtained from three patches were plotted for each channel. The entire dataset could not be fitted with a straight line, but data from each patch was fitted with a straight line for the voltage range from −70 to −10 mV. (D) Relationships between voltage and the mean K\textit{d} value in C. (E) Zd values of each channel calculated from the slopes of the lines in C. Mean values were compared statistically between WT and the mutants using Tukey's test (****, P < 0.001; *, P < 0.05; NS, P > 0.05); statistical comparison of the data obtained from E224Q and E224Q/R228Q is also shown. (F) Values of the Hill coefficient of each channel. Bars depict means ± SEM (n = 3). The symbols used are as indicated in the figure.
at depolarized potentials, even by 0.01 μM spermine. With E224Q, on the other hand, the slow blocking phase was not clearly observed, even with 1 μM spermine, but observed in the presence of 100 μM. Moreover, the blockade was almost completely eliminated by addition of a D259N mutation to E224Q. In the presence of 100 μM spermine, only a 7.4 ± 0.4% (n = 4) reduction in the amplitude in the outward current was observed at +70 mV, so that the analyses summarized in Fig. 6 could not be performed with E224Q/D259N. By contrast, with addition of an R228Q mutation to E224Q, outward currents were blocked by 0.1 μM spermine (Fig. 6).

The normalized steady-state dose–block relationships fitted with Hill’s equation is shown in Fig. 6 B; Kd values (n = 5) at each voltage and the lines fitted to them are shown in Fig. 6 C. The Kd values all showed voltage dependence, the extent of which varied, but using all of the data, none could be fitted by a line due to the presence of extra conductances at highly depolarized potentials (Fig. 6 C). Two possible reported explanations for these extra conductances are (1) that spermine permeates through Kir2.1 at a highly depolarized potential (Guo and Lu, 2000a) and (2) that a second Kir2.1 form is present with a low Kd for spermine blockade and a weak voltage dependence (Ishihara and Ehara, 2004). We did not try to clarify which of these models is correct, and, for the sake of convenience, the voltage dependences of spermine blockade were calculated by linear fitting of the data obtained at voltages ranging from −70 to −10 mV (Fig. 6 C). The mean Kd values and zr values are shown in Fig. 6 (D and E). The Kd value for the spermine blockade of D259N at +40 mV was slightly larger than that for WT, and that for E224Q was much larger. The voltage dependence of the blockade of D259N was slightly weaker than in WT, and that in E224Q was much weaker. The diminished voltage dependency seen with E224Q was partially restored by addition of a R228Q mutation. Hill coefficients did not significantly correlate with voltage (Fig. 6 F).

Contribution of Charged Residues to Intrinsic Inward Rectification

The properties of Kir2.1 rectification are determined not only by cytoplasmic blockers but also by the characteristics of permeation in the absence of those blockers. For that reason, we next determined how the properties of the intrinsic rectification of the pore were affected by neutralization of the charged residues. Inward rectification in the absence of intracellular and extracellular blockers was analyzed using inside-out excised patch clamp recordings with a HEK293 cell expression system. For this analysis, step pulses from +100 to −100 mV were applied in 10-mV decrements, and two holding potentials were used: 0 and −100 mV. That the data obtained with the two holding potentials did not differ confirmed the accuracy of the recordings. Representative macroscopic currents recorded using symmetrical 140 mM K+ solutions and the mean normalized I-V relationships (n = 3–4) are shown in Fig. 8 (A and B). As an index of the intensity of the inward rectification, the ratios of the current amplitudes at +100 and −100 mV were calculated for each of the two holding potentials (Fig. 8 C). The intrinsic inward rectification of WT was weak, while that of D259N was slightly stronger than that of WT, and that of E224Q was much stronger. The intensity of the intrinsic rectification of E224Q was weakened by addition of an R228Q mutation. These results suggest that the intensity of intrinsic inward rectification was also controlled by the electrostatic circumstances within the cytoplasmic pore.

We also focused on the negatively charged amino acid residue E125, which is located within the extracellular loop of the channel protein (Fig. 10 A). Neutralization of E125 (E125N) was reported to reduce the susceptibility of the channel to blockade, the voltage dependence of the blockade by extracellular
Ba\textsuperscript{2+} and Mg\textsuperscript{2+}, and the single channel conductance of K\textsuperscript{+} permeation (Alagem et al., 2001; Murata et al., 2002). Based on those reports, we made E125Q and E125K mutants and analyzed their intrinsic rectification. Representative macroscopic currents and the mean I-V relationships are shown in Fig. 8 (D and E). The intrinsic inward rectification was weakened by the introduction of the E125Q mutation, and it was further weakened, rather outwardly rectified, by the E125K mutation. These changes caused by mutation of E125 were also observed in the presence of other background mutations (E224G and E224Q; unpublished data), which implies that E125 influences the intrinsic rectification from the extracellular side in a manner similar to the negative charges within the cytoplasmic pore.

Contribution of the Histidine Residue to Inward Rectification Properties

We next examined the functional significance of H226 located at the center of other charged residues (Fig. 1 C), whose electrostatic charge changes in accordance with the environmental pH (pKa value \textasciitilde6.0). This H226 residue was reported to contribute to the intracellular pH (pH\textsubscript{i})-sensitive flickery gating of the single channel in the E224G and E224A mutants (Xie et al., 2004). We made some H226 mutants and compared their inward rectification intensities with that of WT under the two-electrode voltage clamp (Fig. 9 A and Table I, group 8). H226K showed weaker inward rectification than WT, and a remarkable outward tail current was observed (Fig. 9 A). The macroscopic current of H226Q was mostly very small, and a recording with exceptionally high expression obtained by injecting \times20 concentrated cRNA to oocytes is shown in Fig. 9 A. Its rectification intensity was not different from that of WT (Fig. 9 A). The macroscopic currents of the H226A, R, E, and D mutants were also too small to be analyzed precisely. We next compared current through WT under pH\textsubscript{i} 7.2 and pH\textsubscript{i} 5.8 and the H226K mutant under pH\textsubscript{i} 7.2, to know the contribution of a positive charge at position 226 (Fig. 9 B). An analysis of H226K under pH 5.8 could not be done because the current disappeared in all trials. The Kd value for the Mg\textsuperscript{2+}, blockade of H226K (pH 7.2) at +70 mV was larger than that of WT (pH 7.2), and it was closer to that of WT (pH 5.8). The voltage dependence of the blockade of H226K (pH 7.2) was weaker.
than that of WT (pH_i 7.2), and it was similar to that of WT (pH_i 5.8) (Fig. 9 B, right). We also analyzed the intrinsic inward rectification of WT and H226K from the identical patch. The intrinsic inward rectification of H226K (pH_i 7.2) was slightly stronger than that of WT (pH_i 7.2). The intrinsic inward rectification of WT became stronger with the decrease in pH_i (Fig. 9 C), and a decrease in the amplitude was also observed. The mean ± SEM values of the intrinsic inward rectification indexes and their statistical analyses using Tukey’s test in WT were 0.78 ± 0.02 (pH_i 7.2, control) and 0.52 ± 0.03 (pH_i 5.8, P < 0.001), and the values in H226K were 0.65 ± 0.02 (under pH_i 7.2, P < 0.05), respectively (n = 4). As the values in H226K under pH_i 6.6 showed a slight but nonsignificant decrease (0.56 ± 0.03, P > 0.05 (in comparison with H226K [pH_i 7.2], n = 4), it seems the effect of pH_i on intrinsic rectification is mediated also by mechanisms other than the protonation at 226. In summary, a positive charge at 226 provided by H226K mutation or by intracellular acidification of WT was shown to weaken inward rectification, to decrease the sensitivity to the block by Mg^{2+}, and to increase intrinsic inward rectification in the absence of blockers. These data show that a positive charge at 226 also plays significant roles as other charged amino acid residues do.

**DISCUSSION**

**Correlation of Location of Charged Amino Acids and their Contribution to Channel Properties**

We observed that mutations that neutralized the negative charges on the wall of the cytoplasmic pore (pH_i 7.2, control) and 3.70 ± 0.12 (pH_i 5.8, P < 0.001), and the values in H226K were 1.40 ± 0.06 (under pH_i 7.2, P < 0.05), respectively (n = 4). As the values in H226K under pH_i 6.6 showed a slight but nonsignificant decrease (1.20 ± 0.06, P > 0.05 (in comparison with H226K [pH_i 7.2], n = 4), it seems the effect of pH_i on intrinsic rectification is mediated also by mechanisms other than the protonation at 226. In summary, a positive charge at 226 provided by H226K mutation or by intracellular acidification of WT was shown to weaken inward rectification, to decrease the sensitivity to the block by Mg^{2+}, and to increase intrinsic inward rectification in the absence of blockers. These data show that a positive charge at 226 also plays significant roles as other charged amino acid residues do.

![Figure 8](image-url)
Figure 9. Comparison of the inward rectification properties of macroscopic currents through WT Kir2.1 and the indicated mutants of the H226 residue. (A) Macroscopic currents recorded using two-electrode voltage clamp with *Xenopus* oocytes and the intensities of the inward rectification of them were analyzed as in Fig 2. Bars depict means ± SEM (n = 4–5), and the values were compared statistically between WT and the mutants using Tukey’s test (**, P < 0.01; P > 0.05). (B) Comparison of the susceptibility of macroscopic currents through WT Kir2.1 and H226K under the indicated intracellular pH to blockade by intracellular Mg^{2+}. Macroscopic currents recorded in the presence of the indicated concentrations of Mg^{2+}, from excised patches of HEK293T cells as in Fig. 5. The concentrations of Mg^{2+} are shown in the figure. The relationships between the Kd and voltage for Mg^{2+} blockade of outward K^{+} currents are shown on the right. The solid lines with symbols indicate the means of the fitted lines with data obtained from four patches for H226K (pH 7.2) and WT (pH 7.2), and the dashed lines indicate SEMs. For reference, the data of WT (pH 7.2) shown in Fig. 5 were also plotted with SEM. (C) Comparison of the intrinsic inward rectification of WT and H226K as in Fig. 8. Bars depict means ± SEM (n = 4) of the normalized current at each voltage. The symbols used are as indicated in the figure.
(E224Q, E299Q, and D259N) reduced the intensity of inward rectification and slowed the activation of inward currents and the decay of outward currents, and that additional mutations neutralizing the positive charges (R228Q and R260Q) had the opposite effect. We also observed protonation at H226 by intracellular acidification or the H226K mutation changed these channel properties. It thus appears that increases in the net negativity within the cytoplasmic pore tends to enhance inward rectification properties (Figs. 2–4, Fig. 9, and Tables I and II), which is consistent with the interpretation proposed by Pegan et al. (2005).

As a next step, we selected several point mutants that reflect this tendency and, using inside-out patch clamp recordings, systematically analyzed the contributions of the affected residues to the blockade by intracellular blockers and to K⁺ permeation (Figs. 5–9). The analyzed parameters were all influenced by the electrostatic negativity within the cytoplasmic pore. With a decrease in the number of negative charges, (1) Kd values for both the Mg²⁺ and spermine blockades were increased, (2) the voltage dependencies were reduced, (3) the single-channel conductance was reduced, (4) open-channel noise was increased, and (5) intrinsic inward rectification was enhanced. These values were partially restored by additional neutralization of a positive charge. Thus, both channel blockade and K⁺ permeation were influenced by the net charge within the cytoplasmic pore.

In all of our analyses, the magnitudes of the changes caused by D259N mutation were substantially milder than those caused by E224Q mutation. For instance, the magnitude of the effect of mutation on inward rectification was very large with E224Q, moderate with E299Q and D259N, and small with D255N (Table I, groups 1 and 2). Moreover, addition of either R228Q or R260Q mutation to E224Q significantly shifted the parameters toward WT values in all analyses, and the magnitudes of their effects were similar. The contributions of these charges can be explained by their location within the cytoplasmic pore (Fig. 10 A). With the channel oriented as in Fig. 10 A, a negatively charged ring composed of E224 and E299 is located at the top of the cytoplasmic pore, and an E224 ring is located inside of the E299 ring. A negatively charged ring composed of D259 is located below them, within the cytoplasmic pore, and D255 much farther below. Two positively charged rings composed of R228 and R260 overlap one another and are located between the E224 and D259 rings. One would expect that charged amino acids located near the mouth of the transmembrane pore would make larger contributions to inward rectification, even though the width of the cytoplasmic pore is nearly constant. Thus, the locations of the charged amino acids correlated well with the magnitudes of their contributions to the inward rectification.

Contribution of the Charged Amino Acids in Concentrating K⁺ within the Cytoplasmic Pore

We observed that the reduction of the net negative charge within the cytoplasmic pore increased the incidence of flickering subconducting states, which resulted in an increase in the open-channel noise and a decrease in the single-channel conductance. A possible interpretation about the arising of the subconducting states was proposed by Xie et al. (2004). They assumed the subconductance was caused by a fast gating, and K⁺ flux through Kir2.1 is regulated by the gate trapping spermine located at bundle crossing or below. This interpretation is based on the observation that spermine unblock is markedly slowed in E224Q, suggesting that the subconducting states may be trapping spermine in the pore (Xie et al., 2004). Another interpretation has also been proposed that the subconductance is derived from the lack of K⁺ in the selectivity filter. Lu et al. (2001) analyzed the single channel properties of Kir2.1 by introducing backbone mutations into the selectivity filter. They suggested that the subconducting states resulted from the collapse of the ion–ion interacting permeation, which was caused by the lack of specific K⁺ coordination within the selectivity filter (Lu et al., 2001). In addition, symmetric increases in [K⁺] on both sides of the channel enhance the main-state probability (Chang et al., 2005), suggesting that for efficient permeation, it is necessary to have the selectivity filter filled with permeating K⁺. In that regard, the cytoplasmic negative charge may play a role in elevating the local concentration of K⁺ in the transmembrane permeation pathway by attracting K⁺ into the cytoplasmic vestibule, thereby supporting permeation through the selectivity filter. This interpretation may also explain the effect of neutralizing mutations on the intrinsic inward rectification, which is possibly due to saturation of the outward flux of K⁺ ions through the pore. Saturation of the ion flux was also reported for permeation through the gramicidin channel when the concentration of charge carrier was low (Andersen, 1984; Koepe and Anderson, 1996), as well as in the BK channel in the absence of negative charges located on the transmembrane inner vestibule (Brelidze et al., 2003; Brelidze and Magleby, 2005). When neutralized, the cytoplasmic pore could be an obstacle to the entry of K⁺ into the permeation pathway, so that resistance within the pore, rather than the K⁺ flux through the selectivity filter, might become the rate-limiting step for K⁺ efflux. We therefore propose that the negative electrostatic potential within the cytoplasmic pore plays a key role in increasing the local concentration of K⁺ to supply sufficient K⁺ ions to the transmembrane region, which in turn enables smooth permeation of fully coordinated K⁺ ions within the selectivity filter. Considering the electrostatic effect of the charged amino acid at E125 at the extracellular mouth (Fig. 8, D and E), we also speculate that the valance of K⁺ ions pooled on the
each side of the GYG selectivity filter may determine the intrinsic rectification intensity.

The Cytoplasmic Pore Is Not the Final Plugging Site for the Blockers

In this and our earlier study (Kubo and Murata, 2001), we analyzed the speed of hyperpolarization-evoked activation (i.e., off-blocking) in various mutants. We found that activation tended to slow down with a reduction in the net negative charges within the cytoplasmic pore. These results were in clear contrast with the observation in the transmembrane D172N mutation, in which activation became almost instantaneous (Lu and MacKinnon, 1994; Stanfield et al., 1994; Wible et al., 1994; Lu et al., 1999; Fujiwara and Kubo, 2002; Chang et al., 2003). In the resolved crystal structure of Kir2.1, the vacancy created by the cytoplasmic pore is large enough to hold 25 water molecules (≈17 × 17 × 30 Å), far too large for the blocker molecules to stably plug the channel and stop the K+ flux at that point in the permeation pathway (Fig. 10 A). We therefore propose that the final plugging site is not within the cytoplasmic pore, but that the cytoplasmic pore nevertheless affects the permeation of both intracellular blockers and K+ through it.

Strong Voltage Dependence of the Blockade

Voltage dependence reflects the movement of charged particles across an electric field. Charge movements through the electric field within Kir channels are reportedly caused mainly by movements of permeant K+ ions pushed by the blockers during the blocking process rather than the movement of the blockers themselves (Oliver et al., 1998; Pearson and Nichols, 1998; Spassova...
and Lu, 1998). In the present study, we observed clear changes in the voltage dependence of the blockade with neutralization of charges on the surface of the cytoplasmic pore even using Mg$^{2+}$, which is very unlikely to occlude permeation at the wide cytoplasmic pore. So if the final plugging site(s) for intracellular blockers was unaltered by the mutations within the cytoplasmic pore, why did the voltage dependence of block change so much?

With the assumption that the blockade occurred within the transmembrane region of the permeation pathway, beyond the cytoplasmic pore, we suggest so far that the negativity within the cytoplasmic pore electrostatically concentrates not only blockers but also K$^+$, so that there is sufficient K$^+$ to be smoothly coordinated within the transmembrane pore (Fig. 10 B, left). During the blocking process, blockers can push out more K$^+$ ions from the electric field, so that a more voltage-dependent block would arise (Fig. 10 B, left). That neutralization of the negative charge within the cytoplasmic pore reduced the voltage dependence of the blockade can thus be explained by a reduction in the ability to concentrate K$^+$ ions (Fig. 10 B, right). We think this K$^+$-based explanation is compatible with various observations made with Kir2.1 and also with Kir6.2 (Kurata et al., 2004), but we have no conclusive structural information about the coordination sites of K$^+$ or intracellular blockers within the transmembrane and/or cytoplasmic pore. There is also a possibility that long-range effects in protein (Bichet et al., 2004; Gazzarrini et al., 2004) due to the cytoplasmic mutations may influence the transmembrane structure. The transmembrane structure of Kir2.1 is thought to be different from KcsA (Minor et al., 1999; Chatelain et al., 2005), so its crystal analysis will be needed to fully clarify the inward rectification mechanism in Kir2.1.

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