

## Molecular Biophysics: The Cytosolic GH Loop Regulates the Phosphatidylinositol 4,5-Bisphosphate-induced Gating Kinetics of Kir2 Channels



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# The Cytosolic GH Loop Regulates the Phosphatidylinositol 4,5-Bisphosphate-induced Gating Kinetics of Kir2 Channels\*<sup>S</sup>

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Hai-Long An<sup>±§</sup>, Shou-Qin Lü<sup>1</sup>, Jun-Wei Li<sup>§</sup>, Xuan-Yu Meng<sup>‡</sup>, Yong Zhan<sup>§</sup>, Meng Cui<sup>‡</sup>, Mian Long<sup>11</sup>, Hai-Lin Zhang<sup>||2</sup>, and Diomedes E. Logothetis<sup>±3</sup>

From the <sup>‡</sup>Department of Physiology and Biophysics, School of Medicine, Virginia Commonwealth University, Richmond, Virginia 23298, the <sup>§</sup>Institute of Biophysics, School of Sciences, Hebei University of Technology, Tianjin 300401, China, the <sup>II</sup>Key Laboratory of Neural and Vascular Biology, Ministry of Education, The Key Laboratory of Pharmacology and Toxicology for New Drug, Hebei Province, Department of Pharmacology, Hebei Medical University, Shijiazhuang 050017, China, and the <sup>II</sup>Key Laboratory of Microgravity (National Microgravity Laboratory) and Center of Biomechanics and Bioengineering, Institute of Mechanics, Chinese Academy of Sciences, Beijing 100190, China

**Background:** The detailed mechanism of  $\text{PIP}_2\text{-}\text{induced}$  Kir channel gating remains elusive.

**Results:** Specific mutations increase the flexibility of the cytosolic GH loop and accelerate the PIP<sub>2</sub>-induced gating kinetics of Kir2 channels.

**Conclusion:** Interactions of the GH loop with the N terminus regulate the PIP<sub>2</sub>-induced gating kinetics of Kir2 channels. **Significance:** We identify a novel region in Kir channels involved in the control of PIP<sub>2</sub>-induced gating.

Inwardly rectifying  $K^+$  (Kir) channels set the resting membrane potential and regulate cellular excitability. The activity of Kir channels depends critically on the phospholipid PIP<sub>2</sub>. The molecular mechanism by which PIP<sub>2</sub> regulates Kir channel gating is poorly understood. Here, we utilized a combination of computational and electrophysiological approaches to discern structural elements involved in regulating the PIP<sub>2</sub>-induced gating kinetics of Kir2 channels. We identify a novel role for the cytosolic GH loop. Mutations that directly or indirectly affect GH loop flexibility (*e.g.* V223L, E272G, D292G) increase both the on- and especially the off-gating kinetics. These effects are consistent with a model in which competing interactions between the CD and GH loops for the N terminus regulate the gating of the intracellular G loop gate.

Inwardly rectifying  $K^+$  (Kir) channels play a vital role in many physiological processes, such as control of heart rate,

maintenance of the resting membrane potential, hormonal secretion, extracellular K<sup>+</sup> buffering in the brain, and K<sup>+</sup> secretion in the kidney. These physiological processes are regulated by specific stimuli that control the gating of Kir channels. Examples of stimuli gating Kir channels include intracellular  $H^+$  for Kir1 (1), the  $\beta\gamma$  subunits of G proteins for Kir3 (2), and phosphatidylinositol 4,5-bisphosphate  $(PIP_2)^4$  for all Kir channels (3, 4). The inwardly rectifying ATP-inhibited  $K^+$  ( $K_{ATP}$ ) (Kir6.2) channel was the first reported to depend on  $PIP_2$  (5, 6). Subsequently, all known Kir channels were shown to depend directly on phosphoinositides for maintenance of their activity (3, 5, 7-10). In general, Kir channels, as is the case with most ion channels tested, run down when PIP<sub>2</sub> is depleted (dephosphorylated by lipid phosphatases) and reactivate when PIP<sub>2</sub> is applied from the intracellular side (11). The gating kinetics of Kir channels are quite distinct between different Kir channels, but the underlying reasons are not understood.

Kir channels, like other K<sup>+</sup> channels, are homo- or heterotetramers of four subunits. Each subunit shares a common structure consisting of a cytosolic N-terminal domain, an outer transmembrane helix (M1), an ion-selective P loop (selectivity filter (SF)), an inner transmembrane helix (M2) lining most of the permeation pathway, and a cytosolic C terminus controlling ion flow (12, 13) (Fig. 1*A*). The SF and bundle crossing of the M2 helix are thought to serve as gates, constriction points along the permeation pathway limiting ion flow (12). Mechanisms of gating (*i.e.* transitions between the closed and open states) for the SF and M2 gates have been studied and proposed (14). The Nand C termini, which are located on the intracellular side of the membrane, form a comparable cytosolic extension to the transmembrane portion of the pore. In addition to the SF and M2 transmembrane gates, a cytosolic constriction (HI or G loop) of



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<sup>&</sup>lt;sup>1</sup> To whom correspondence may be addressed: Key Laboratory of Microgravity (National Microgravity Laboratory) and Center of Biomechanics and Bioengineering, Institute of Mechanics, Chinese Academy of Sciences, Beijing 100190, China. Tel.: 86-10-82544131; E-mail: mlong@imech.ac.cn.

<sup>&</sup>lt;sup>2</sup> To whom correspondence may be addressed: Key Laboratory of Neural and Vascular Biology, Ministry of Education; The Key Laboratory of Pharmacology and Toxicology for New Drug, Hebei Province; Department of Pharmacology, Hebei Medical University, Shijiazhuang 050017, China. Tel.: 86-311-86265562; Fax: 86-311-86265562; E-mail: z.hailin@yahoo.com.

<sup>&</sup>lt;sup>3</sup> To whom correspondence may be addressed: Department of Physiology and Biophysics, School of Medicine, Virginia Commonwealth University, Richmond, VA 23298. Tel.: 804-828-5878; Fax: 804-828-7382; E-mail: delogothetis@vcu.edu.

<sup>&</sup>lt;sup>4</sup> The abbreviations used are: PIP<sub>2</sub>, phosphatidylinositol 4,5-bisphosphate; M2, inner transmembrane helix; SF, selectivity filter; MD, molecular dynamics; RMSF, root mean square fluctuation; TEVC, two-electrode voltage clamp; Ci-VSP, *Ciona intestinalis* voltage-sensitive phosphatase.

the permeation pathway has been proposed as a third gate (9) (Fig. 1A). Specific acidic residues localized in the cytosolic pore are thought to give rise to inward rectification (15). Functional studies and mutagenesis data have identified several residues as potential PIP<sub>2</sub>-interacting sites in both the N- and C termini of the cytoplasmic domain of the Kir2.1 channel. Many of these residues are basic and thus capable of forming salt bridge interactions with the negatively charged head group of  $PIP_2$  (3, 8, 10, 16). Recent crystallographic evidence has confirmed this view (17, 18). Moreover, crystal structures of a Kir3.1 chimera caught the intracellular G loop gate in dilated (open) and constricted (closed) conformations. The movement in the G loop observed in different structures of the cytoplasmic domains is also notable (9, 19). However, it is not known how the G loop gate integrates into channel gating and how it affects the gating kinetics of Kir channels.

Although recent crystallographic studies have greatly enhanced our understanding of the three-dimensional structure of Kir channels (13, 17, 20), it has not been possible yet to understand the dynamic gating mechanism of these channels induced by molecules such as  $PIP_2$ . A recent study of multiple KirBac3.1 structures in either the open or closed conformations has highlighted the importance of intersubunit interactions between the N and C termini and the G loops of adjacent subunits in Kir channel gating (21).

Here we set out to study Kir channel gating kinetics, focusing on the cytosolic channel regions using a voltage-gated phosphatase to control PIP<sub>2</sub> levels (22) and computational, mutagenesis, and electrophysiological approaches to probe the role of specific secondary structural elements and specific residues within them. We observed highly correlated movements of the Kir2.1 (Val-223) (of the CD loop) and Glu-272 (of the GH loop) with Ala-306 at the G loop gate (see sequence alignment in supplemental Fig. S1). Mutations, at Val-223 in Kir2.1 or Val-224 in Kir2.2, accelerated both the opening and closing gating kinetics of the channel and increased the flexibility of the GH loop. Similar results were obtained with mutations of key residues within the GH loop. Our results are explained by a model in which PIP<sub>2</sub> induces the switching of the N terminus from the CD loop to the GH loop (around the short  $\alpha$ G helix) (Fig. 1C). Mutations that decrease the GH loop interactions with the N terminus (e.g. V223L, E272G, D292G) increase the flexibility of the GH loop and allow the N terminus to gate the G loop with faster kinetics.

#### **EXPERIMENTAL PROCEDURES**

Chemicals—Phosphatidylinositol-4,5-bisphosphate diC8  $PIP_2$  was purchased from Avanti Lipids and was prepared as described previously (8, 10). All other chemicals were purchased from Sigma.

Homology Modeling—Modeler V9.5 (23) was used to add missing residues to the crystal structures of Kir2.1 (PDB code 1U4F), and the full-length crystal structure of Kir2.2 (PDB code 3JYC). Modeler was also used to create a homology model of the cytosolic domain of Kir2.2. The mutant channels were constructed by substituting the WT side chain with the specified side chains. The models were then subjected to at least 3000 steps of a steepest descent minimization using the CHARMM

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program with the implicit membrane/solvent Generalized Born (GB) model (24).

Molecular Docking-AUTODOCK (25) was used for the docking studies. We replaced  $PIP_2$  with its analog diC1  $PIP_2$ , which has two methyl groups. The atomic charges of the PIP<sub>2</sub> head group were taken from the *ab initio* calculations by Lupyan and colleagues (26). A grid map was generated for the Kir2.2 full-length structure using CHNOP (i.e. carbon, hydrogen, nitrogen, oxygen, and phosphor) elements sampled on a uniform grid containing  $120 \times 120 \times 120$  points 0.375 Å apart. The center of the grid box was set to the center of known critical PIP<sub>2</sub>-sensitive residues, *i.e.* Gln-51, Arg-65, Lys-183, Arg-186, Lys-188, Lys-189, Arg-190, Arg-219, Lys-220, Arg-229, and Arg-313. The Lamarckian Genetic Algorithm (LGA) was selected to identify the binding conformations of the ligands. 100 docking simulations were performed, and the final docked PIP<sub>2</sub> analog configurations were selected on the basis of docked binding energies and cluster analysis. The PIP<sub>2</sub>-Kir2.2 complex was constructed on the basis of the docked PIP<sub>2</sub> analog-Kir2.2 complex structure and refined by CHARMM using the same protocol as described above.

Molecular Dynamics (MD) Simulations-The crystal structure of the Kir2.1 cytosolic domain was initialized as follows, solvating the molecule in a rectangular water box of 82 imes 104 imes103  $\text{\AA}^3$  and neutralizing the water box by adding Na<sup>+</sup> and Cl<sup>-</sup> of  $\sim 100$  mm. MD simulations were performed using NAMD with CHARMM27 all-atom force field parameters (27). An integration time step of 1 fs, a uniform dielectric constant of 1.0, a scaling factor for 1-4 interactions of 1.0, and periodic boundary conditions were applied in all simulations. A smooth (12-16)Å) cutoff and the Particle Mesh Ewald (PME) (28) were employed to calculate van der Waals forces and full electrostatics, respectively. Prior to the equilibration process, energy minimization (5000 steps with backbone atoms of C-terminal fixed and then another 5000 steps with all atoms free), followed by a heating-up process from 0 to 300 K over 35 ps, were performed. Then, two 5-ns equilibration processes with either all atoms free or partly constrained were performed with the temperature held at 300 K using Langevin dynamics while the pressure was held at 1 atm using the Langevin piston method. An RMSF (root mean squared fluctuation) analysis was performed on the basis of the two equilibration processes.

For the full-length Kir2.2 channel simulation, the channels were immersed in an explicit palmitoyloleoyl-phosphatidylcholine bilayer generated from the visual molecular dynamics membrane package. After being solvated with SPC water molecules, neutralized by Na<sup>+</sup> as the counter ions, and including K<sup>+</sup> located in the selectivity filter as obtained from the crystal structures, each system involved  $\sim$ 141,000 atoms in the MD simulations. GROMACS v4.0.5 was used to conduct the simulation with the GROMOS96 53a6 force field. The force field parameters for PIP<sub>2</sub> were generated from the Prodrg server (8), and the same atomic charges of the PIP<sub>2</sub> head group in the docking step were used in the MD simulations. The lipid parameters were obtained from Dr. Tieleman. Long range electrostatics were calculated using the PME8 method with a 12 Å cut-off. Van der Waals interactions were modeled using Lennard-Jones 6–12 potentials with a 14 Å cut-off. All simulations



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were conducted at a constant temperature of 300 K using the Berendsen thermostat. The system pressure was coupled at isotropic (X+Y, Z) directions referenced to 1 bar using the Berendsen method9. All bonds were constrained with the LINCS algorithm. The time step was 2 fs, and the neighboring list was updated every 10 time steps.

Prior to production runs, energy minimization of 3000 steps of steepest descent were carried out on each system, followed by a 0.5-ns two-step equilibration process with either all atoms free or partly constrained. An electric field of 0.06 V/nm was applied in this step as well as the production run, along the *z* axis of the box, to maintain the lower potentials in the intracellular side. The treatment of the electric field has been detailed in Refs. 29, 30. A 100-ns production run was conducted on each system, and coordinates were saved every 10 ps for analysis. Visual molecular dynamics were used for conformational visualization.

*Interaction Network Analysis*—The interactions of hydrogen bonds, salt bridges, and hydrophobic contacts analysis were calculated using the Simulaid (31–34) program after energy minimization. The Simulaid outputs for interactions were reorganized with in-house scripts for the facility of comparison among the systems.

*Molecular Biology*—All cDNA constructs were subcloned into the pGEMHE plasmid vector and used as described (10). Point mutants were produced by Pfu mutagenesis with a Quikchange kit (Stratagene). Sequences were confirmed by DNA sequencing. Recombinant Kir2.1, its mutants, and Ci-VSP were expressed in *Xenopus laevis* oocytes as described previously (10, 35). cRNA was produced with T7 RNA polymerase using a kit (Promega). cRNAs of the various Kir2.1, Kir2.2, their mutants, and of Ci-VSP were injected in the range of 0.5–10 ng/oocyte depending on the functional expression level of the given construct.

Electrophysiology-Recordings in X. laevis oocytes were performed 1 to 2 days after cRNA injection. Whole-oocyte currents were measured by conventional two-electrode voltage clamp (TEVC) using a GeneClamp 500 amplifier (Molecular Devices, CA). Electrodes were filled with 3 M KCl dissolved in 1% agarose to prevent the leakage of KCl into the oocytes. The electrodes had resistances less than 1  ${}_{M}\Omega$ . Oocytes were constantly perfused with a high-potassium solution (ND96K) containing 96 mм KCl, 1 mм NaCl, 1.8 mм CaCl<sub>2</sub>, 1 mм MgCl<sub>2</sub>, 5 mM HEPES (pH 7.4 with KOH). A low-potassium solution (ND96) was used in some experiments to inhibit most of the Kir2.1 currents at -80 mV. To activate Ci-VSP, the following voltage protocol was used in TEVC oocyte recordings: 1-s sweeps composed of a 170-ms ramp from -80 to +80 mV followed by an 830-ms step to +80 mV. To deactivate Ci-VSP, the oocytes were held at -80 mV. Sweeps were applied until the resulting currents reached steady state. ND96 contained 96 mM NaCl, 1 mM KCl, 1.8 mM CaCl<sub>2</sub>, 1 mM MgCl<sub>2</sub>, 5 mM HEPES (pH 7.4 with NaOH). Current amplitudes were measured at +80 mV and -80 mV. Data acquisition and analysis were achieved using pClamp9.2 (Molecular Devices, CA) and Origin 7.5 (Microcal, MA) software.

Macropatch channel activity was recorded from devitellinized oocytes under the inside-out mode of standard patchclamp methods using an Axon 200B patch-clamp amplifier and Clampex 10.0 data acquisition software (Molecular Devices). Electrodes were made from borosilicate glass using a Sutter P-97 microelectrode puller (Sutter Instrument Co., CA) that gave a tip diameter of 5–15  $\mu$ m and had a resistance of 0.5–1 M $\Omega$  when filled with an electrode solution containing 120 mM KCl, 2 mM MgCl<sub>2</sub>, and 10 mM HEPES (pH 7.4 with KOH). Two bath solutions were used: a FVP solution containing 60 mM KCl, 5 mM EDTA-K, 5 mM KF, 0.1 mM Na<sub>3</sub>VO<sub>4</sub>, 10 mM K<sub>4</sub>P<sub>2</sub>O<sub>7</sub>, 10 mM HEPES (pH 7.4 with KOH) and FVP solution with X  $\mu$ M (0–100  $\mu$ M for dose response and 30  $\mu$ M for gating kinetics measurements) diC8 PIP<sub>2</sub>. Data were analyzed using pClamp9.2 (Molecular Devices) and Origin 7.5 (Microcal) software.

*HA Tag Expression Experiments*—We engineered an HA tag to the extracellular loop between the outer transmembrane helix and the P region of Kir2.1 (8). Oocytes were blocked for 60 min in ND96 with 1% BSA at 4 °C, labeled with 0.5  $\mu$ g/ml rat monoclonal anti-HA antibody (3F10, Roche) (in 1% BSA overnight at 4 °C), washed at 4 °C, and incubated with HRPcoupled secondary antibody (goat anti-rat) (in 1% BSA for 60 min). Cells were washed (1% BSA, 4 °C) and transferred to ND96 without BSA. Individual oocytes were placed in 55  $\mu$ l of SuperSignal ELISA (Pierce) and incubated at room temperature for 1 min. Chemiluminescence was quantified in a TD-20E luminometer (Turner Designs, CA). As a control, we used uninjected oocytes.

*Data Analysis*—The *error bars* in the figures represent S.E. Each experiment shown or described was performed on five to seven oocytes of the same batch for TEVC recording and three to five oocytes of the same batch for macropatch recordings. A minimum of three to four batches of oocytes was tested for each experiment shown.

#### RESULTS

The Val-223 and Glu-272 Show High Correlativity with A306 in the G Loop Gate—To explore the mechanism by which the G loop gate relates to the rest of the cytosolic domain during PIP<sub>2</sub>induced Kir channel gating, we performed 10-ns MD simulations of the cytosolic domains of the WT Kir2.1 and analyzed the relative movements between C-terminal residues and the G-loop residue Ala-306. From the free MD simulations, we found that the relative movements between Glu-272 (at the beginning of the GH loop), Ala-306 (at the apex of the G loop), and Val-223 (in the CD loop) were highly correlated (Fig. 1B). Fig. 1C shows the location of residues Val-223, Glu-272, and Glu-306 in the cytosolic portion of the crystal structure. Unlike the CD and G loops that are physically proximal and are thought to interact with each other, the GH loop is located quite far from the other two loops (Fig. 1C). We hypothesized that Val-223 and Glu-272 could play important roles in the PIP<sub>2</sub>-induced Kir channel gating.

Kir2.1(V223L) and Kir2.2(V224L) Affect the PIP<sub>2</sub>-induced Gating Kinetics through a Common Mechanism—We first examined the effect of mutations at the Kir2.1 223 position from Val to Leu or Met. Fig. 2A shows a right shift of the dose response curve of V223L responding to the stimulatory effect of PIP<sub>2</sub> (EC<sub>50</sub> for WT, 2.75  $\pm$  0.12  $\mu$ M and for V223L, 8.57  $\pm$  0.76  $\mu$ M). However, unlike other mutants reported to decrease the





FIGURE 1. **MD simulation performed on the C-terminal domain of Kir2.** Shown are highly correlated movements among Val-223, Glu-272, and Ala-306. *A*, two Kir2.2 subunits shown with the SF, M2 gate, and cytosolic G loop gate indicated. Similarly, the CD and GH loops that regulate the stability of the G loop gate have been *highlighted. B*, correlation of movements among C-terminal residues of Kir2.1. Val-223, Glu-272, and Ala-306 are shown as  $\blacklozenge$ ,  $\blacklozenge$ , and  $\clubsuit$ , respectively. *C*, schematic structure of one subunit cytosolic domain where the Kir2.1 residues Val-223, Glu-272, and Ala-306 are shown as *small spheres*. The *pink spheres* in *A* represent the potassium ions in the ion pathway. In *A* and *C*, the CD, GH, and G loops are highlighted in *pink, gray*, and *green*, respectively.

apparent affinity to  $\mathrm{PIP}_{2}$ , the V223L mutant displayed a 20% increase in its peak currents (Fig. 2B). We next measured membrane surface expression of the WT and V223L mutant using HA-tagged channels. Our data indicated a 42% reduction in cell surface expression for V223L (0.58  $\pm$  0.01 when WT Kir2.1 was normalized to 1). These results suggested that the increased whole-cell currents of the V223L mutant were not due to an increase in available channels on the cell surface but likely to increased activity. To further examine this possibility, we measured the gating kinetics of Kir2.1 and its V223L mutant using either TEVC and the lipid phosphatase Ci-VSP (22) in intact cells or inside-out patches of the patch-clamp technique. Use of Ci-VSP allows rapid and reversible control of phosphatase activity through voltage-clamp recordings, which, in turn, has allowed us to discern the macroscopic closing and opening Kir2 gating kinetics induced by  $\mathrm{PIP}_2$  dephosphorylation and rephosphorylation.

Fig. 2, *C* and *D*, shows an acceleration of the gating kinetics of the V223L mutant relative to the WT by inhibition (*left panels*) or activation (*right panels*) of Ci-VSP (kinetics of current inhibition ( $\tau_{off}$ ) of V223L, 4.78  $\pm$  0.51 s *versus* WT, 17.73  $\pm$  0.72 s; kinetics of current recovery ( $\tau_{on}$ ) of V223L, 6.28  $\pm$  0.36 s *versus* WT, 15.72  $\pm$  1.66 s). Ci-VSP reduces membrane PIP<sub>2</sub> levels when the membrane is depolarized (36). Alternatively, we examined the gating kinetics of activation and deactivation of Kir2.1 WT and its V223L mutant by perfusion and washout of exogenous diC8 PIP<sub>2</sub> in inside-out patches. Following patch excision, we perfused the patch with a high-potassium solution (HK) lacking ATP, which results in PIP<sub>2</sub> dephosphorylation and rundown of currents. After complete rundown, diC8 PIP<sub>2</sub> was applied at the intracellular side of patches expressing WT or V223L to reactivate the channels. The kinetics of both the run-

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down ( $\tau_{off}$ ) (*right panels*) and reactivation ( $\tau_{on}$ ) (*left panels*) by diC8 PIP<sub>2</sub> were accelerated by the V223L mutation ( $\tau_{off}$  for WT, 21.97  $\pm$  1.42 s and for V223L, 3.44  $\pm$  2.09 s;  $\tau_{on}$  for WT, 7.25  $\pm$  1.71 s and for V223L, 4.05  $\pm$  0.73 s (supplemental Fig. S2).

We then tested the effects of the V223M mutation on the PIP<sub>2</sub>-dependent gating kinetics. The apparent affinity of the WT and V223M channels to diC8 PIP<sub>2</sub> were indistinguishable (EC<sub>50</sub> for WT, 2.75 ± 0.12  $\mu$ M and for V223M, 2.62 ± 0.46  $\mu$ M) (supplemental Fig. S3A). V223M also showed similar amplitude currents in TEVC as the WT Kir2.1 (supplemental Fig. S3B). Moreover, the kinetics of Ci-VSP-induced changes in currents were also not significantly different for WT *versus* the V223M mutant ( $\tau_{off}$  for WT, 17.73 ± 0.72 s and for V223M, 16.93 ± 0.95 s;  $\tau_{on}$  for WT, 15.72 ± 1.66 s and for V223M, 12.27 ± 0.61 s) (supplemental Fig. S3, *C* and *D*). Thus, V223M, unlike the V223L mutant, displayed similar apparent affinity to PIP<sub>2</sub>, whole-cell current level and gating kinetics as the WT Kir2.1 channel.

We next asked whether other Kir2 channels showed similar effects as the Kir2.1(V223L) mutant did. Kir2.2 also has a Val (Val-224) at the analogous position to the Kir2.1(V223) (supplemental Fig. S4A). The Kir2.2(V224L) mutant, like the Kir2.1(V223L) mutant, accelerated both  $\tau_{off}$  (*left panels*) and  $\tau_{on}$  (*right panels*) ( $\tau_{off}$  for WT, 42.87  $\pm$  7.45 s and for V224L, 1.36  $\pm$  0.22 s;  $\tau_{on}$  for WT, 176.88  $\pm$  19.42 s and for V224L, 13.53  $\pm$  0.43 s) (supplemental Fig. S4, *B* and *C*). Unlike Kir2.1, the apparent affinity of the Kir2.2(V224L) mutant for diC8-PIP<sub>2</sub> was left-shifted (EC<sub>50</sub> for WT, 6.13  $\pm$  0.43  $\mu$ M and for V224L, 2.46  $\pm$  0.13  $\mu$ M) (supplemental Fig. S4*D*). The WT and the V224L Kir2.2 mutant showed similar whole-cell currents (supplemental Fig. S4*E*).

V223L and V224L, but not V223M, also Increase the Flexibility of the GH Loop in the Linker between the CD and G Loops— To explore the mechanism by which the V223L and V224L affected the  $PIP_2$ -induced gating kinetics, we performed 10-ns MD simulations of the cytosolic domains of the WT Kir2.1, Kir2.2 channels and the three mutants (V223L, V223M, and V224L) and analyzed the RMSF, which is a measure of the flexibility of each residue for the WT and mutant channels. Fig. 3A shows significant changes in the flexibility of the GH loop (amino acids 270 to 294, which precisely define the boundaries of the GH loop, see supplemental Fig. S1) that were induced specifically by the V223L or V224L mutation compared with the WT or the V223M mutant, respectively.

We asked whether this enhanced flexibility induced by the Val to Leu mutation could be related to its effects on the channel gating kinetics. To address this question, we examined the PIP<sub>2</sub>-induced gating kinetics of the WT Kir2.1 and V223L mutant channels in intracellular solutions with increased viscosity. The dynamics of a protein in solution are intimately coupled to the dynamics of the solvent. The fluctuation amplitudes and relaxation rates, *i.e.* the flexibility of proteins, can be reduced by increases in solution viscosity (37–40). Previous studies have shown that increases in solution viscosity affect the moving parts of voltage-dependent Na<sup>+</sup> channels during gating, causing a decrease in the rate of channel activation (41). We increased the viscosity of the intracellular solution by addition of 2 M sucrose (42) and measured rundown kinetics of currents





FIGURE 2. **The V223L mutation accelerates the gating kinetics of Kir2.** *A*, dose response of diC8 PIP<sub>2</sub> of Kir2.1 and V223L, showing a right shift in the apparent affinity of PIP<sub>2</sub> sensitivity as a result of the mutation. *B*, peak currents of Kir2.1 and V223L do not show a decrease in whole-cell current, as other mutations do, that decrease the apparent channel affinity to PIP<sub>2</sub>. *C*, the representative time course of inhibition (PIP<sub>2</sub> dephosphorylation induced by Ci-VSP, a 3.7-fold effect) and reactivation (PIP<sub>2</sub> rephosphorylation, a 2.5-fold effect) of WT-Kir2.1 and V223L. *D*, bars are the time constants (mean  $\pm$  S.E. of at least six experiments) corresponding to *C*.

from inside-out patches expressing the WT Kir2.1 or the V223L mutant channels. Compared with the normal intracellular solution, the solution with the 2  $_{\rm M}$  sucrose showed a non-significant trend of slowing down the time course of rundown in the WT (-sucrose, 24.41  $\pm$  8.40 s; + sucrose, 27.70  $\pm$  7.90 s) (Fig. 3*B*). In contrast, the rundown kinetics of the V223L mutant were slowed significantly (- sucrose, 3.77  $\pm$  0.53 s; + sucrose, 8.78  $\pm$  0.65 s).

The E272G Mutant Mimics the Effect of the V223L Mutant— If indeed Val-223 were to be allosterically coupled to the G loop (at Ala-306) via the flexible region (at Glu-272), could an increase in the relative flexibility of Glu-272 mimic the V223L effects on Kir2.1 gating? Gly residues are utilized in protein structures to confer flexibility (*e.g.* the Gly hinge in the inner helix that forms the permeation pathway) (39, 43, 44). We introduced a glycine at Glu-272 and tested whether it increased local flexibility and whether it mimicked the gating effects of

V223L. The RMSF analysis of the E272G mutant indeed showed a significant increase in local flexibility (Fig. 4A). The apparent affinity of the E272G mutant to diC8-PIP<sub>2</sub> was slightly left-shifted (EC<sub>50</sub> values for WT and the E272G mutant were  $2.75 \pm 0.12 \ \mu\text{M}$  and  $2.14 \pm 0.21 \ \mu\text{M}$ , respectively) (Fig. 4B), whereas the mutant displayed similar peak currents (Fig. 4C). Interestingly, and consistent with V223L, the E272G mutant accelerated the gating kinetics of both channel inhibition and reactivation by Ci-VSP ( $au_{
m off}$  for WT, 17.73  $\pm$  0.72 s and for E272G, 9.81  $\pm$  0.68 s;  $\tau_{\rm on}$  for WT, 15.72  $\pm$  1.66 s and for E272G, 7.13  $\pm$  0.61 s) (Fig. 4, *D*–*F*). A Gly scan of this region (from 266 to 272, supplemental Fig. S5A) revealed that Gly mutations at Thr-268 and Ile-269 in the  $\beta$ G strand were not tolerated well (were either functionally dead or displayed very low activity, respectively), whereas mutations (Pro-266 and Ile-267) on the N-terminal side of the GH loop were without effect. Mutants at the beginning of the flexible GH loop (Val-270, His-271, and





FIGURE 3. Induction of flexibility of the linker between the CD and G-loops is coupled with changes in the gating kinetics of Kir channels. *A*, RMSF analysis of the linker between CD and G loops of Kir2.1, Kir2.2, and their mutants. The Val-to-Leu mutations but not the Met mutation cause an increase in the RMSF of Kir2.2 *B*, time constants (mean  $\pm$  S.E. of at least three experiments) of rundown (because of diC8 PIP<sub>2</sub> washout) of WT-Kir2.1 (*left panel*) and V223L (*right panel*) in the presence and absence of sucrose. Only Kir2.1(V223L) shows a significant slowing of the rundown kinetics in the presence of sucrose.

Glu-272) showed significant effects on  $PIP_2$ -induced gating (supplemental Fig. S5, *B* and *C*).

A Conserved Gating Mechanism Linking the CD and G Loops-To gain structural insights of how PIP<sub>2</sub> might induce channel gating via Val-223 of the CD loop in a manner that involves the GH loop, we performed 100 ns free MD simulations on the WT Kir2.2 and V224L, both in the absence (apo) and presence (holo) of PIP<sub>2</sub> using the full-length Kir2.2 crystal structure (13). Supplemental Table S1 shows salt bridge interactions between PIP<sub>2</sub> and Kir2.2 during a 20-ns interval in the second half of the simulations. The simulation showed PIP<sub>2</sub> to interact with residues Gln-51, Arg-78, Arg-80, Lys-183, Lys-188, Lys-189, Arg-219, and Lys-220 (Fig. 5A and supplemental Table S1) in good agreement with a cocrystal structure of Kir2.2 in complex with  $PIP_2$  (17). Comparison of the apo and holo simulations revealed that PIP<sub>2</sub> releases the N terminus from the CD loop by weakening the interactions between Arg-219-Asp-61 and His-222-Gln-51 (Fig. 5, *B* and *C*, Fig. 6, *A* and *B*, and Supplemental Table S2). The N terminus then switches to interact with the  $\alpha$ G helix by strengthening the interactions between Asp-292-Arg-44 and Glu-294-Arg-44 (Fig. 5, B and C, and Fig. 6, A and B). The V224L mutant, with an increased flexibility of the GH loop, reduced the intersubunit interactions of the N terminus with the CD loop residues in the apo state (Gln-51-His-222 and Asp-61-Arg-219) and the interactions of the N terminus with the  $\alpha G$ helix in the holo state (Asp-292-Arg-44 and Glu-294-Arg-44) compared with the WT (Fig. 6, C and D). Thus, the mutant decreased overall interactions of both the CD and GH loops with the N terminus and increased the flexibility of the GH loop, as if interactions with the N terminus normally constrain

the flexibility of the GH loop. To examine the prediction from the simulations of the importance of the Asp-292 interactions, we tested the effects of the D292G mutation on the Kir2.2 gating kinetics. The D292G mutation accelerated the gating kinetics of both channel inhibition and reactivation by Ci-VSP ( $\tau_{\text{off}}$  for WT, 42.87  $\pm$  7.45 s and for D292G, 20.74  $\pm$  2.93s;  $\tau_{\text{on}}$  for WT, 176.88  $\pm$  19.42 s and for D292G, 104.82  $\pm$  7.66 s) (Fig. 5, D-G). We also made the E294G mutation, but that caused a significant reduction in whole-cell currents (from 6.56  $\pm$  0.73  $\mu$ A for the WT to 0.68  $\pm$  0.03  $\mu$ A for E294G), making it difficult to characterize it further.

#### DISCUSSION

 $PIP_2$ -induced channel gating is an intrinsically dynamic process that is difficult to understand by the few available static crystal structures that are missing key transition steps connecting them. Here we apply a combination of computational and electrophysiological approaches to discern the role of important structural elements in the  $PIP_2$ -induced gating of Kir2 channels, focusing on the cytosolic G-loop gate. We started with MD simulations performed on the crystal structure of the WT Kir2.1 cytosolic domains. Following the MD simulation, we identified that the movements of Ala-306 (of the G loop) correlated highly with Val-223 (of the CD loop) and Glu-272 (of the GH loop).

Mutagenesis of Val-223 and kinetic electrophysiological measurements showed that V223L, but not V223M, right-shifted the diC8-PIP<sub>2</sub> dose response, increased currents and caused faster gating kinetics by accelerating both the inhibition and recovery rates. The corresponding mutant in Kir2.2,





FIGURE 4. The Kir2.1 (E272G) also accelerates the gating kinetics albeit to a smaller extent than V223L. *A*, The RMSF of Kir2.1 and E272G in the GH loop, the linker between the CD and G-loops. The mutant causes an increase in RMSF relative to the control. *B*, dose-responses of diC8 PIP<sub>2</sub> on Kir2.1 and the E272G mutant reveal a small leftward shift in apparent affinity caused by the mutation. *C*, peak currents of Kir2.1 and the E272G mutant appear comparable. *D*, the representative time course of current inhibition (PIP<sub>2</sub> dephosphorylation induced by Ci-VSP, a 1.8-fold effect) of WT-Kir2.1 and E272G. *E*, the representative time course of reactivation (PIP<sub>2</sub> rephosphorylation, a 2.2-fold effect comparable with the inhibitory effect shown in *D*) of WT-Kir2.1 and E272G. *F*, summary bars (*left* and *right*) represent the time constants (mean  $\pm$  S.E. of at least six experiments) corresponding to *D* and *E*, respectively.

V224L, showed similar kinetic effects as V223L. Additional MD simulations performed on Kir2.1 (V223L) and Kir2.2 (V224L) showed similar increases in the flexibility of residues corresponding to the GH loop of Kir2 channels (*i.e.* the 270–294 residues of the Kir2.1 or the 265–294 residues of the Kir2.2).

A number of crystal structures of KirBac3.1 (21) revealed interactions between the N terminus and the CD loop, forming the "latched" or closed state. In a recent computational study with the Kir3.1 chimera, we revealed that  $PIP_2$  weakens these interactions, forcing the latched N terminus to transition to an "unlatched" or open state (21, 45). The fact that the N-terminal interactions underlie the transition of the G-loop gate from the closed to the open state in such different channels as the Kir3.1 chimera and KirBac3.1 suggest that the critical role of the N

terminus in Kir channel gating has been conserved from prokaryotes to eukaryotes.

Our 100-ns-long MD simulations were performed on the Kir2.2 WT or the V224L channel in the presence or absence of PIP<sub>2</sub>. There was excellent agreement between the predicted Kir2.2 interacting residues with PIP<sub>2</sub> (Gln-51, Arg-78, Arg-80, Lys-183, Lys-188, Lys-189, Arg-219, and Lys-220) and those reported for Kir2.1 from mutagenesis and electrophysiological measurements (8) (Kir2.2. corresponding residues Gln-51, Arg-65, Lys-183, Arg-186, Lys-188, Lys-189, Arg-190, Arg-219, Lys-220, Arg-229, and Arg-313). A similar agreement exists with Kir2.2 residues shown in a recent crystal structure (17) (Arg-65, Arg-78, Trp-79, Arg-80, Lys-183, Arg-186, Lys-188, Lys-189, Arg-190, and Arg-219). Thus, six of the eight residues predicted





FIGURE 5. **Partial crystal structure model of Kir2.** The WT shows detailed interactions between the CD-loop, N terminus, and the  $\alpha$ G helix. *A*, the residues that interact with PIP<sub>2</sub> from our 100-ns-long MD simulations. PIP<sub>2</sub> is shown in a *ball-and-stick representation*. *B* and *C* show the detailed structures of WT Kir2.1 with (holo) and without (apo) PIP<sub>2</sub>, respectively, to underscore changes in the relationship of the N terminus with the CD and GH loops. *D* and *F* are representative trace currents of Kir2.2 WT and D292G, one of the GH loop residues predicted by our simulations to interact with PIP<sub>2</sub>. PIP<sub>2</sub> dephosphorylation of the D292G mutant induced by Ci-VSP (a 2.1-fold effect) and PIP<sub>2</sub> rephosphorylation (a 1.7-fold effect) relative to the control, respectively. *E* and *G* are summary data (mean  $\pm$  S.E. of at least six experiments).



FIGURE 6. **PIP<sub>2</sub> modulates gating of Kir channels by altering interactions between the CD loop, N terminus, and the \alphaG helix of the GH loop. Shown is the effect of PIP<sub>2</sub> on averaged hydrophobic (***Hp***) interactions between Arg-219-Asp-61 (***A***) and salt bridge (***SB***) (***B***) interactions between His-221-Gln-51, Glu-292-Arg-44, and Glu-294-Arg-44 in the Kir2.2 WT.** *C***, effect of PIP<sub>2</sub> on averaged hydrophobic interactions between Arg-219-Asp-61 (***C***) and salt bridge (***D***) interactions between His-221-Gln-51, Glu-292-Arg-44, and Glu-292-Arg-44, and Glu-292-Arg-44, and Glu-292-Arg-44, and Glu-292-Arg-44, and Glu-292-Arg-44, and Glu-294-Arg-44 in Kir2.2 V224L.** 

to interact with  $PIP_2$  were shown to affect  $PIP_2$  sensitivity experimentally (supplemental Table S1). Other residues shown experimentally to affect  $PIP_2$  sensitivity could be doing so in an allosteric manner. The interaction network between the CD loop, the N terminus, and the loop segment around the  $\alpha$ G helix of the GH loop revealed that there were appreciable interactions between the N terminus and the CD loop (His-222-Gln-51 and Arg-219-Asp-61) in the absence of  $PIP_2$  stabilizing the closed state.  $PIP_2$  weakened these two pairs of interactions, interacting with Arg-219 directly and stabilizing interactions between the N terminus and the GH loop: Asp-292-Arg-44 and Glu-294-Arg-44. Interestingly, Asp-292 and Glu-294 are



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exactly at the other end of the GH loop from Glu-272 that affected PIP<sub>2</sub>-induced gating kinetics. Thus, the N terminus appears to have switched from stabilizing the CD loop in the closed state to stabilizing the  $\alpha$ G helical region within the GH loop in the open state.

The V224L mutation not only stabilized the CD-loop and N-terminal interactions with PIP<sub>2</sub> but also weakened all four pairs of interactions between the N terminus and the CD loop or the GH loop. The weakened His-221-Gln-51 and Arg-219-Asp-61 interactions would make it easier for the N terminus to switch from stabilizing the closed to stabilizing the open state, accelerating the activation kinetics of Kir2 channels. The weakened interactions between the N terminus and  $\alpha$ G helical region of the GH loop allowed for the increased flexibility of the GH loop, ensuring that the channel can easily transition from the open to the closed state, thus accelerating the deactivation kinetics as well. The flexibility of the GH loop (residues 270 to 294), reflecting the weakened interactions between the N terminus and the GH loop, showed greater effects on the deactivation kinetics ( $\tau_{off}$ ).

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