Mutations in Nature Conferred a High Affinity Phosphatidylinositol 4,5-Bisphosphate-binding Site in Vertebrate Inwardly Rectifying Potassium Channels*

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Background: A native Kir channel in a distant relative of vertebrates interacts weakly with PIP₂.

Results: Mutagenesis restores the vertebrate channel sensitivity to PIP₂ in sponge channels.

Conclusion: A basic residue in the tether helix of Kir is required for high affinity PIP₂ regulation.

Significance: Evolution conferred a high affinity interaction of vertebrate Kir channels with PIP₂, which is lacking in a distant relative.

All vertebrate inwardly rectifying potassium (Kir) channels are activated by phosphatidylinositol 4,5-bisphosphate (PIP₂) (Logothetis, D. E., Petrou, V. I., Zhang, M., Mahajan, R., Meng, X. Y., Adney, S. K., Cui, M., and Baki, L. (2015) Annu. Rev. Physiol. 77, 81–104; Fürst, O., Mondou, B., and D’Avanzo, N. (2014) Front. Physiol. 4, 404–404). Structural components of a PIP₂-binding site are conserved in vertebrate Kir channels but not in distantly related animals such as sponges and sea anemones. To expand our understanding of the structure-function relationships of PIP₂ regulation of Kir channels, we studied AqKir, which was cloned from the marine sponge Amphimedon queenslandica, an animal that represents the phylogenetically oldest metazoans. A requirement for PIP₂ in the maintenance of AqKir activity was examined in intact oocytes by activation of a co-expressed voltage-sensing phosphatase, application of wortmannin (at micromolar concentrations), and activation of a co-expressed muscarinic acetylcholine receptor. All three mechanisms to reduce the availability of PIP₂ resulted in inhibition of AqKir current. However, time-dependent rundown of AqKir currents in inside-out patches could not be re-activated by direct application to the inside membrane surface of water-soluble dioctanoyl PIP₂, and the current was incompletely re-activated by the more hydrophobic arachidonoyl stearyl PIP₂. When we introduced mutations to AqKir to restore two positive charges within the vertebrate PIP₂-binding site, both forms of PIP₂ strongly re-activated the mutant sponge channels in inside-out patches. Molecular dynamics simulations validate the additional hydrogen bonding potential of the sponge channel mutants. Thus, nature’s mutations conferred a high affinity activation of vertebrate Kir channels by PIP₂, and this is a more recent evolutionary development than the structures that explain ion channel selectivity and inward rectification.

Inwardly rectifying potassium (Kir)² channels represent a superfamily of one pore domain and two transmembrane domain channels. Kir channels are present in organisms from bacteria to humans and are found throughout the animal kingdom. In animal cells, they play an important role in regulating the resting membrane potential, K⁺ homeostasis, and membrane excitability. An ancestral precursor to all Kir channels is predicted to have originated in prokaryotes followed by changes in channel structure and function at the prokaryotic-eukaryotic transition and, most likely, throughout the time course of animal evolution (3, 4).

Like many ion channels and transporters, the activity of Kir channels from vertebrates, including mammalian and non-mammalian species, is dependent on a membrane lipid, phosphatidylinositol 4,5-bisphosphate (PIP₂) (1, 2, 5–7). PIP₂ is found primarily in the inner leaflet of the plasma membrane (8), and its availability is dynamically controlled by neurotransmitter, hormone, and growth factor receptors that couple to phospholipase C (9, 10). Signaling that activates phospholipase C is expected to lower the availability of PIP₂, which reduces or prevents the activation of vertebrate Kir channels. Understand-

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‡ The abbreviations used are: Kir channel, inwardly rectifying potassium; Aq, Amphimedon queenslandica; PIP₂, phosphatidylinositol 4,5-bisphosphate; TEVC, two-electrode voltage clamp; MD, molecular dynamic; CIVSP, Ciona intestinalis voltage-sensing phosphatase; HK, high K⁺; diC₈, dioctanoyl; PI(4,5)P₂, phosphatidylinositol 4,5-bisphosphate.
**PIP₂ Sensitivity of a Sponge Kir Channel**

... channels are regulated by PIP₂ is of fundamental physiological importance to cellular signaling and K⁺ homeostasis in eukaryotic cells.

The evolutionary origin of PIP₂-mediated regulation of Kir channels is unknown. Prokaryotic Kir channels are inhibited by PIP₂ (11, 12), whereas Kir channels from all eukaryotic cells studied to date are activated by PIP₂ (2, 6, 10, 13). This is an interesting difference, but PIP₂ is not present in the prokaryotic cell membrane (2, 6), and thus its inhibitory effect on bacterial Kir channels lacks physiological relevance. Among eukaryotes, the structural basis for Kir channel activation by PIP₂ has been clarified by many studies (1, 2), culminating in the co-crystallization of PIP₂ with chicken Kir2.2 (7). Among the critical components of the protein-lipid interaction are several basic residues within the channel that provide an electrostatically favorable interaction with the negative charges on the phosphate groups of PIP₂. However, despite the tremendous insight gained by many years of physiology and structural biology experiments, the functional priority of the basic residues in the Kir channel and whether they are sufficient to explain high affinity interactions of the lipid and channel remain untested outside the confines of vertebrate species such as rat, mouse, human, and chicken. Because there is a 400–600 million year evolutionary gap between humans and basal invertebrates such as sponges (14), there is a good possibility that evolutionary changes resulted in structural changes to regulatory sites on ion channels (15).

To explore the impact of evolutionary pressure on structural changes in ion channel proteins, we studied PIP₂ regulation of AqKir, a K⁺ channel with strong inward rectification, which was previously cloned from the marine sponge *Amphimedon queenslandica* (16). Sponges are an important comparative group for understanding the relationships between extant unicellular protists and the first multicellular organisms from which all animals evolved (14, 17–20). By studying the physiological and structural impact of “nature’s mutations,” we provide a window into the functional consequences of evolution that is represented in distantly related animal species. A Kir channel cloned from a marine sponge (AqKir) lacks two positive charges that are important for high affinity interaction with PIP₂ in vertebrate Kir2 channels (7). We report that, following time-dependent rundown, AqKir channels are insensitive to re-activation by PIP₂ added to the internal surface of an excised membrane patch, despite additional evidence from intact oocytes that suggests that PIP₂ may be required for this channel’s function. In addition, AqKir channels, which we engineered to possess the two positive charges present in the PIP₂-binding site of vertebrate Kir2 channels, showed strong activation by diC₈ PIP₂, with a sensitivity that was even 3-fold greater than that for vertebrate Kir2.1 channels. Our results confirm that certain positively charged residues (corresponding to residues 80 and 189 in cKir2.2) are absolutely necessary to confer high sensitivity to PIP₂ and represent evolutionary changes that are more recent than the evolution of structures that confer K⁺ selectivity and inward rectification, which are conserved from sponge to human (16). Finally, we propose that other structures present in distant relatives of vertebrate Kir channels must facilitate the activation properties conferred by PIP₂ such that restoration of even a single positive charge (N180K in AqKir) recapitulates the high sensitivity to PIP₂ present in vertebrate Kir channels.

**Experimental Procedures**

*Functional Expression in Oocytes—*Wild-type channel constructs included AqKir (16) (GenBank™ accession number FJ375323, previously named AmqKirA), chicken Kir2.2 (cKir2.2, GenBank™ accession number F1NHE9), mKir2.1 (GenBank™ accession number X73052), and mKir2.3 (GenBank™ accession number S71382). A plasmid containing CiVSP (GenBank™ accession number AB183035) was used, and mutation C363S was prepared to destroy the enzymatic activity (21). Mutants were constructed using QuikChange (Agilent Technologies, catalogue no. 200523) and confirmed by DNA sequencing. The preparation of cRNA, isolation of oocytes, and injections were performed using standard methods (22). In brief, plasmids containing sponge channel cDNAs were linearized and capped RNAs synthesized using Ambion (Austin, TX) mMessage Machine RNA polymerase kits. RNA was purified by use of the RNAid kit (Bio 101, Vista, CA) or LiCl precipitation, and concentrations were determined by Nanodrop (ThermoScientific) spectrophotometry. Oocytes were surgically harvested from anesthetized female *Xenopus laevis* (*Xenopus* I, Dexter, MI) frogs. Oocytes were released and defolliculated by gentle agitation in 0.5 mg/ml collagenase A (Sigma) dissolved in a Ca²⁺-free solution containing (in mM) 96 NaCl, 2 KCl, 1 MgCl₂, 5 HEPES, pH 7.4. Oocytes were then washed and selected, and stage V/VI oocytes were microinjected with 36–50 nl of cRNA dissolved in diethyl pyrocarbonate-treated water (0.05–15 ng of cRNA/oocyte), depending on functional activity (21). Mutants were constructed using QuikChange (Agilent Technologies, catalogue no. 200523) and confirmed by DNA sequencing. The preparation of cRNA, isolation of oocytes, and injections were performed using standard methods (22). For two-electrode voltage clamp (TEVC), we used a Geneclamp 500B amplifier (Axon Instruments, Foster City, CA) or an OC-725C amplifier (Warner Instruments, Hamden, CT). Voltage-measuring and current-passing electrodes were backfilled with 3 M KCl and had resistances between 0.3 and 1.0 megohms. Currents were sampled at 5–10 kHz and filtered at 1–2 kHz. All recordings were obtained at room temperature (about 23°C), and the bath was perfused continuously during recordings. The standard external solution contained (in mM) 96 KCl, 4 NaCl, 1 CaCl₂, 2 MgCl₂, 10 HEPES, pH 7.3–7.4.

*Electrophysiology—*Potassium currents were recorded from oocytes using standard electrophysiological methods (22–26). For two-electrode voltage clamp (TEVC), we used a Geneclamp 500B amplifier (Axon Instruments, Foster City, CA) or an OC-725C amplifier (Warner Instruments, Hamden, CT). Voltage-measuring and current-passing electrodes were backfilled with 3 M KCl and had resistances between 0.3 and 1.0 megohms. Currents were sampled at 5–10 kHz and filtered at 1–2 kHz. All recordings were obtained at room temperature (about 23°C), and the bath was perfused continuously during recordings. The standard external solution contained (in mM) 96 KCl, 4 NaCl, 1 CaCl₂, 2 MgCl₂, 10 HEPES, pH 7.3–7.4.

For inside-out patch clamp recordings, we used an A-M Systems Model 2400 or an Axon Instruments 200B patch clamp amplifier. Macropatches were excised from oocytes following manual removal of the vitelline membrane (24–26). Electrodes were pulled from borosilicate glass using a Sutter P-97 puller and gave a tip diameter of 5–15 μm and a resistance of 0.5–2 megohms when filled with a solution containing (in mM) 96 KMeSO₄, 1 MgCl₂, and 10 HEPES, pH 7.4. The standard bath (internally facing) high K⁺ (“HK”) solution contained 96 KCl, 5 EGTA, and 10 HEPES, pH 7.4, unless otherwise mentioned.
Currents were elicited with a voltage ramp protocol from $-100$ to +100 mV over 1 s. Signals were low pass-filtered at 10 kHz and digitized at 20-$\mu$s intervals.

Electrophysiological data were recorded on computers equipped with Digidata 1320A (Axon Instruments) A/D hardware. Axon’s Clampex acquisition and Clampfit analysis software (versions 7–9) were used.

**Data Analysis and Curve Fitting**—Data were also transferred to Microsoft Excel and Microcal Origin (Northampton, MA) for additional analysis, curve-fitting, and the production of figures. Results were calculated from cells with negligible background currents and for TEVC, only from cells with membrane potentials not more depolarized than $-30$ mV in 5 mM external K$^+$. Results are expressed as mean $\pm$ S.E. with $n =$ number of oocytes tested with a minimum of three batches of oocytes per experiment. The EC$_{50}$ value of PIP$_2$ effects for each channel was obtained by fitting concentration-response data to a Hill Equation:

$$\frac{I/I_{\text{max}}}{[X]} = \frac{[X]_i}{EC_{50}^n + [X]_i}$$

where $I/I_{\text{max}}$ is the normalized current; $[X]_i$ is the PIP$_2$ concentration applied from the intracellular side; $n$ is the Hill coefficient, and EC$_{50}$ is the concentration required to reach half of the maximum activation.

In cases with multiple comparisons to a single set of control values, a single factor ANOVA with a post-hoc Dunnett’s test was used with $p < 0.05$ considered to be significant. Other statistical evaluations included paired or unpaired Student’s two-tailed $t$-tests, as appropriate, with $p < 0.05$ considered to be significant.

**Chemicals**—AASt (“long chain”) PIP$_2$ was purchased from Avanti Polar Lipids (Alabaster, AL). diC$_8$ PIP$_2$ was purchased from Avanti Polar Lipids or Echelon Biosciences (Salt Lake City, UT). The lipids were prepared as described previously (23, 25, 27). All other chemicals were purchased from Sigma or VWR.

**Modeling of AqKir/PIP$_2$ Channel Complexes**—The AqKir channel model was constructed based on the chicken Kir2.2 x-ray crystal structure template (PDB code 3SPI (7)). The sequence alignment between Kir2.2 and AqKir channels was generated by ClustalW2. The sequence identity between Kir2.2 and AqKir is $\sim$40%. We used the program MODELLER 9 version 12 (28) to generate 10 initial homology models. The model assessment and selection were based on the internal objective function, 1,2-dioleoyl-sn-glycero-3-phosphoethanolamine, and GA341 assessment scores in the MODELLER program to select models with the lowest free energy potentials. The full-length PIP$_2$ molecules were added into the AqKir model by superimposition with the short-chain derivative of PIP$_2$ in the Kir2.2 crystal structure (7). Models for AqKir channel mutants were constructed by a mutation function using Discovery Studio version 3.5 molecular modeling program (Accelrys Software Inc., San Diego).

**Molecular Dynamics Simulations**—The PIP$_2$-AqKir channel complexes were immersed in an explicit palmitoyl oleoyl phosphatidylcholine bilayer generated from the VMD membrane package (29). After being solvated with SPc water molecules, neutralized by KCl (150 mM) as the counter ions and including K$^+$ located in the selectivity filter as obtained from the crystal structures, each system involved $\sim$140,000 atoms in the MD simulations. GROMACS version 4.5.4 (30) was used to conduct the simulation with the GROMOS96 53a6 force field (31). The parameters for PIP$_2$ and lipid were generated as described previously (32). Long range electrostatics were calculated using the particle mesh Ewald (PME) method (33) with a 12 Å cutoff. van der Waal interactions were modeled using Lennard-Jones 6–12 potentials with 14-Å cutoff. All simulations 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 method (34). All bonds were constrained with the LINCS algorithm (35). 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 the steepest descent were carried out on each system followed by a 0.5-ns two-step equilibration process. In the first 0.2 ns, channels, K$^+$ ions, and PIP$_2$ were position-restrained using a constant force of 1000 kJ/mol/nm$^2$, allowing lipid and water molecules to move freely. The restraint was weakened to 10 kJ/mol/nm$^2$ in the following 0.3-ns equilibration. An electrical field of 0.02857 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. A 40-ns production run was conducted on each system, and coordinates were saved every 10 ps for analysis.

**Interaction Energy Calculations**—The Discovery Studio 3.5 molecular modeling program (Accelrys Software Inc.) was used to conduct energy minimizations and calculate the interaction energy between the PIP$_2$ and the AqKir channels. All structures were subjected to energy minimization using Smart Minimizer algorithm (2000 steps) and Generalized Born implicit solvent model using the CHARMM force field. Interaction energies between the PIP$_2$ and the channel were calculated using a distance-dependent dielectric constant ($\varepsilon = 5\varepsilon_0$) implicit solvent model. The decomposed interaction energy contribution of residues to the total interaction energy was analyzed to identify the critical interacting residues in the protein complexes. A residue was considered to be critical if it bound the ligand with an interaction energy lower than $\sim 1$ kcal/mol.

**Results**

**Sequence Comparisons**—An amino acid sequence comparison between sponge and vertebrate Kir channels shows that AqKir is missing two canonical positively charged residues at positions 78 and 180, within the outer and tether helices, respectively (Fig. 1A), of the PI(4,5)P$_2$-binding site in the crystal structure of chicken Kir2.2 (7) and is represented in a homology model of AqKir (Fig. 1B). Both positive charges are present in members of all but one of the seven vertebrate Kir channel subfamilies (Fig. 1A). Thus, AqKir is the only known functional Kir channel that lacks multiple residues previously shown to be important for regulation by PIP$_2$ (27, 36). A BLAST search for additional distant relatives of the vertebrate Kir channels found three putative Kir channels in the genome of the sea anemone Nematostella vectensis (Nv). All three lack a positive charge
Within the tether helix region of the PIP₂-binding site in cKir2.2, although the positive charge in the outer helix region, which is missing in the sponge channel, is preserved in the three sequences from sea anemone (Nv1–3) (Fig. 1A). As both sponges and sea anemones are modern day representatives of more primitive metazoans, these structural similarities provide evidence for the impact of evolution on the function of vertebrate Kir channels.

Experiments on Intact Cells—Based on the sequence differences between AqKir and vertebrate Kir channels, we used TEVC on whole oocytes to probe the requirement for PIP₂ in the maintenance of the AqKir current. We reduced the PIP₂ concentrations in whole oocytes in three ways (37). First, activation of a co-expressed voltage-sensing phosphatase (CiVSP) reversibly reduced the inward AqKir current in a time-dependent fashion (Fig. 2, A and B).

Within the tether helix region of the PIP₂-binding site in cKir2.2, although the positive charge in the outer helix region, which is missing in the sponge channel, is preserved in the three sequences from sea anemone (Nv1, His; Nv2–3, Arg; Fig. 1A). As both sponges and sea anemones are modern day representatives of more primitive metazoans, these structural similarities provide evidence for the impact of evolution on the function of vertebrate Kir channels.
phorylate PI(4,5)P₂ upon membrane depolarization (38, 39).

The time to reach 50% inhibition by CiVSP (Fig. 2A) was 2.66 ±

1.03 s for AqKir (n = 4), which was 10-fold faster than that

measured for Kir2.1 (25.3 ± 6.10 s; n = 6). Likewise, the t₀.₅ for

Kir current recovery (Fig. 2B), using hyperpolarizing pulses to
de-activate the enzyme, was about 3-fold slower for AqKir

(57.2 ± 13.2 s; n = 4) than for Kir2.1 (22.0 ± 1.39 s; n = 2). In

several experiments, we tested a catalytically inactive mutant of

giVSP (C363S (21) and observed no change in Kir current

amplitudes upon membrane depolarization (data not shown).

As a second test for Pip₃ dependence, the amplitude of resting

Kir currents was measured after preincubation with 20 μM

dworntmannin, a concentration that inhibits phosphatidylinositol

4-kinase activity and thus lowers endogenous levels of PI(4,5)P₂

(40). Compared with time-matched, vehicle-treated (DMSO)

tolos using oocytes from the same frogs, wortmannin pre-
treatment resulted in a significant reduction in the resting

AqKir and Kir2.3 currents but not Kir2.1 currents (Fig. 2C).

Notably, Kir2.1 has a higher affinity for Pip₃ interaction than

dostrKir2.3 (41). Third, when muscarinic (M₁) acetylcholine

receprors were co-expressed with AqKir and activated by ace-
tylcholine, the inward AqKir K⁺ currents were inhibited (Fig. 2,

D and E), whereas Kir2.1 currents were not inhibited (Fig. 2E).

The muscarinic inhibition of AqKir mimics the established
effect on Kir2.3 currents by M₁ receptor activation of a G protein-
coupled phospholipase C (41). Together, these three results

from intact oocytes suggest that, despite the lack of posi-
tively charged residues in key locations of the vertebrate Kir2
channel Pip₃-binding site (Fig. 1), AqKir demonstrates known
characteristics of Pip₃-dependent channel regulation. How-

ever, the effects of CiVSP and wortmannin suggest that the

AqKir-Pip₃ interaction is weaker than that for the Kir2.1-Pip₃
interaction.

Loss of Activity in Excised Patches—To further test the pre-
diction of Pip₃ dependence of AqKir channels, we examined

the impact of membrane patch excision on current activity of

AqKir channels. In oocytes expressing AqKir currents, patch

excision first resulted in a transient increase in current ampli-
tude, which was quickly followed by decay of the current such
that channel activity was lost within 2 min (Fig. 3A). In 90% of

the patches, the “rundown” of AqKir current demonstrated two
phases, as observed in Fig. 3A (black circles). Overall, the loss of

AqKir current was faster than the well known, time-dependent
decay of vertebrate Kir channels, cKir2.2, and particularly

Kir2.1, measured under the same ionic conditions. We found

that the time to decay to 50% of the initial on-cell AqKir current

was 12-fold faster than for Kir2.1 and 4-fold faster than for

cKir2.2 currents (Fig. 3B). As rundown of Kir channels is com-
monly explained by the loss of Pip₃ from the excised patch due
to phosphatase-mediated dephosphorylation of Pip₃ (42), the
faster rate of AqKir current decay suggests that a higher con-
centration of Pip₃ is needed to sustain the sponge Kir channels
than the vertebrate Kir channels.
PIP₂ Sensitivity of a Sponge Kir Channel

A. On-cell Current Magnitude—On-cell patch recordings obtained before patch excision showed that the average magnitude of the AqKir channel currents was 24-fold lower than that found for Kir2.1 (Figs. 3C and 4C). Likewise, the whole-cell current amplitudes measured in TEVC showed a markedly lower current level for AqKir (−1.9 ± 0.39 μA) when compared with cKir2.2 (−4.6 ± 1.0 μA) or Kir2.1 (−60 ± 4.8 μA; recordings in HK solution). However, a double mutant I78R/N180K restored large AqKir current amplitudes in TEVC (−65 ± 10 μA; n = 7–17) to values comparable with the Kir2.1 channels. The reduced magnitude of the wild-type AqKir channel activity in the on-cell patch and in whole-cell recordings is consistent with the hypothesis that wild-type AqKir channel activity is limited by weaker interactions of PIP₂ with the wild-type AqKir channel than with mutant AqKir channels.

Application of PIP₂ to Inside-out Patches—To further test this hypothesis, we used direct application of two forms of PIP₂ to the inside-facing surface of inside-out membrane patches. The lipids were prepared fresh from frozen stocks and applied in the HK bath solution using a gravity-driven perfusion system. Long chain (AASt) PIP₂ contains the 20-carbon polyunsaturated fatty acid, arachidonic acid (AA) and the 18-carbon saturated stearic acid (St) in its two acyl tails. The hydrophobic nature of AASt-PIP₂ limits its use to single dose applications to membranes or bilayers. In contrast, water-soluble diC₈ PIP₂ contains two 8-carbon unsaturated fatty acid tails and can be applied reversibly, multiple times; it is most useful in quantifying the concentration dependence of PIP₂ and comparing the relative sensitivities for the lipids in different channel constructs.

The experiment in Fig. 4 tests the ability of direct application of AASt PIP₂ to re-activate Kir currents following rundown due to patch excision (i/o arrow). At a concentration of 10 μM, AASt PIP₂ completely restored the Kir2.1 current (Fig. 4A) but even 25 μM incompletely re-activated AqKir (Fig. 4B). In these recordings, the on-cell current magnitude was 24-fold higher for Kir2.1 when compared with AqKir (Fig. 4C), similar to previous results shown in Fig. 3C. To negate the impact of the initial (on-cell) current value on the strength of the re-activation by PIP₂ in each construct, we compared the magnitude of the recovered current after AASt PIP₂ (Fig. 4D, black bars) with the on-cell current amplitude before rundown (Fig. 4D, gray bars). AASt-PIP₂ (10 μM) reactivated all of the Kir2.1 current but only about 25% of the AqKir WT current (Fig. 4D). This incomplete reactivation of the sponge Kir channel activity is consistent with results from our TEVC experiments suggesting that PIP₂ interactions with AqKir are weaker than for Kir2 channels.

Direct application of diC₈ PIP₂ to the intracellular face of the membrane patches expressing AqKir also did not restore channel activity (Fig. 5). The lowest tested concentrations of diC₈ PIP₂ (0.1 and 1 μM) showed only very small increases in AqKir currents after rundown (Fig. 5A), and higher concentrations showed no activity. Clearly, diC₈ PIP₂ could not restore the sponge channel activity. On average, the current activity with diC₈ PIP₂ was so low that AqKir WT current amplitudes reached, at most, 10% of the on-cell patch current. Thus, a concentration-response relationship for channel activation could not be determined for the wild-type sponge Kir channel (see Fig. 6C, black circles), although Kir2.1 showed characteristic re-activation by diC₈ PIP₂ using the same recording conditions and same preparations of the lipid (Fig. 5, B and D). Current-voltage relationships for the two constructs showed that the strongly rectifying inward AqKir currents were not recovered by even 100 μM diC₈ PIP₂ (Fig. 5C), whereas Kir2.1 was maximally activated by 10 μM diC₈ PIP₂ (Fig. 5D).

Recapitulating Mutations in Nature in AqKir—Channel proteins that associate with PIP₂ are known to engage in electrostatic interactions with the negatively charged headgroup of PIP₂ and positively charged residues in the protein (2,7). To test the critical nature of the two basic residues in vertebrate Kir channel-PIP₂ interactions (Fig. 1), we used site-directed mutagenesis to add the positive charges to AqKir, individually and together. Although wild-type whole cell and on-cell AqKir currents were small and failed to be restored by diC₈ applied to excised patches (Fig. 5A, 6A), even a single mutation, N180K, restored the capability of re-activation by diC₈ PIP₂ (Fig. 6, A and C). The effects of diC₈ on N180K were dose-dependent, reversible, and saturated by 30 μM (Fig. 6, A and C); this mim-
The double mutant AqKir N180K has an apparent affinity similar to the vertebrate Kir2.1 channel, whereas the double mutant I78R/N180K has a higher sensitivity to diC₈ than Kir2.1, i.e. EC₅₀ of 0.04 μM for the double mutant versus 2 μM for Kir2.1. Prior to the restoration of I78R/N180K currents by diC₈ as shown in Fig. 6, we also observed a slow time course of rundown in the double mutant (t₅₀ of 250 ± 38.5 ms, n = 22; compared with 17.5 ± 3.86 ms for AqKir WT from Fig. 3B); AAST-PIP₂ quickly reactivated the I78R/N180K current (data not shown).

An additive effect on diC₈ sensitivity of I78R in the double mutant (Fig. 6C, red circles) points to an important role of this arginine residue although, in macropatches, we could not reliably measure currents from the single mutant, I78R, following diC₈ application. However, the mutant I78R produced measurable currents for TEVC recordings (Figs. 6B and 7) and was capable of re-activation by AAST-PIP₂ in inside-out patch recordings (Fig. 8, C and D).

We also examined the AqKir mutant channels for changes in CiVSP-mediated inhibition of whole oocyte current. The previously documented fast inhibition of whole-cell wild-type AqKir currents by depolarization-mediated CiVSP activation (Fig. 2A) was slowed by single or double mutations at residues Ile-78 and Asn-180 (Fig. 7, A and C). The time course of recovery with hyperpolarization to de-activate CiVSP showed the inverse effect; recovery was slow in WT AqKir but faster with mutation of Ile-78 or Asn-180 or both (Fig. 7, B and D). Indeed, quantitative comparisons of the time to 50% inhibition (t₅₀) and the t₅₀ for recovery following inhibition point to an important role for each positive charge alone and an additive effect of the two mutations (Fig. 7, C and D). Overall, the data suggest that the loss of PIP₂ due to phosphatase activity has a stronger impact on WT than on mutant AqKir channels, which further demonstrates that these critical positive charges support the high sensitivity of PIP₂ interaction with the channel. Furthermore, our results suggest that PIP₂ concentrations in the membrane of intact cells (frog oocytes) falls to very low levels upon CiVSP activation, less than that required to sustain the activity of the double mutant AqKir channels for which 1 μM diC₈ PIP₂ maximally activated them.

Neomycin Inhibition—Neomycin, an amino-glycosidic antibiotic, is a positively charged molecule that binds with high affinity to PIP₂; its application to the inside surface of the membrane is expected to inhibit PIP₂-dependent currents by binding to the anionic headgroup of the lipid (43, 44). Thus, we used neomycin to further examine the relative sensitivities to PIP₂ of AqKir mutant channels and the vertebrate Kir2.1 channel. Kir channel currents were allowed to rundown in inside-out patch recordings, and long chain AAST-PIP₂ was applied to restore the Kir currents to sustained levels. We then measured the inhibition of Kir2.1 (Fig. 8A) and AqKir mutant channels (Fig. 8B) by increasing concentrations of neomycin in a stepwise fashion. The fractional current inhibition for five constructs was plotted as a function of neomycin concentration, and each data set fitted to a standard Hill equation (Fig. 8C). The fitted Kᵣ values for neomycin inhibition quantify the strength of PIP₂ binding (Fig. 8D). The strong neomycin inhibition measured for
FIGURE 5. Water-soluble diC₈ PIP₂ fails to re-activate AqKir currents in inside-out patches. A, inward AqKir WT currents measured at −100 mV in on-cell patch mode and after patch excision at the time indicated (arrow i/o). HK + 2 mM MgCl₂ (open bars) was interspersed with applications of increasing concentrations of diC₈ PIP₂ (0.1 – 100 μM in HK without Mg²⁺, gray bars) to test for recovery of current. B, under the same conditions, the dose-dependent reactivation of Kir2.1 by diC₈ PIP₂ is shown. A and B, the dotted lines indicate the zero current level. C, current-voltage relationships elicited by voltage ramps from −100 to +100 mV for a representative AqKir WT on-cell recording (black trace), immediately following patch excision (red trace), and following application of diC₈ PIP₂ as follows: 1 μM (blue), 10 μM (pink), and 100 μM (teal). D, similar recordings for Kir2.1 show the currents in the on-cell patch (black trace) and after re-activation by diC₈ PIP₂ as follows: 0.3 μM (teal), 1 μM (pink), 3 μM (blue), and 10 μM (red) following washout/rundown.

FIGURE 6. Effects of AqKir mutations on re-activation by diC₈ PIP₂. A, application of diC₈ PIP₂ (0.01–100 μM) to inside-out patches from oocytes expressing AqKir WT (top) or the mutants N180K (middle) or I78R/N180K (bottom). Lipids were applied (horizontal bars) after rundown in a Mg²⁺-free HK solution. Currents were measured at −80 mV; the dotted line shows the zero current level. B, comparison of whole-cell current amplitudes (in A) for AqKir WT and three mutants measured in TEVC at −80 mV in a bath solution of 96 mM K⁺. The RNA for AqKir WT was injected at 2–4 times the concentration for the mutants. Bars represent mean ± S.E. for n = 7–16 cells for each construct. C, fitted concentration-response data for diC₈ PIP₂ reactivation of Kir channels as follows: AqKir WT (solid black circles); N180K (open blue circles); I78R/N180K (open red circles); and Kir2.1 (solid gray squares). Currents were normalized to the maximum on-cell current amplitude in each patch. D, fitted EC₅₀ values for diC₈ PIP₂ reactivation of AqKir WT, two AqKir mutants, and Kir2.1. C and D, values are mean ± S.E. for n = 3–7 patches for each construct.
AqKirWT was made weaker with each of the single mutants, and the $K_d$ value for neomycin inhibition of the lipid-reactivated double mutant channels was even higher than that measured for Kir2.1. These data are consistent with the strengthened diC$_8$ PIP$_2$ interactions measured for N180K and the double mutation in AqKir (Fig. 7).
**Docking and Energy Analysis**—To understand how the mutations affect the AqKir channel and PIP₂ interactions, we developed homology models of AqKir, both wild-type and single and double mutants, in complex with PIP₂. Fig. 9 depicts a snapshot of AASt-PIP₂ bound to AqKir WT (Fig. 9A) and AqKir I78R/N180K (Fig. 9B). In the double mutant, the positive charges on Lys-180 and Arg-78 orient more closely to the PIP₂ molecule.

We conducted MD simulations on each model system in a membrane environment with explicit solvent and calculated the energy of the PIP₂-channel interactions. Our results show that PIP₂ docked in AqKir WT, at a site that maps to the Kir2.2 PIP₂-binding site, has a total interaction energy of −90.6 kcal/mol. This interaction became more favorable with the I78R/N180K mutant, which had a total interaction energy of −127.4 kcal/mol. Each single mutant also improved the overall favorability of interaction with PIP₂; the total interaction energies for I78R and N180K were −113.4 and −121.0 kcal/mol, respectively.

To assess quantitatively the importance of specific chemical bonds in the proposed interaction of PIP₂ within the sponge Kir channel, we used energy decomposition analysis (Table 1). Addition of one basic residue to AqKir increases the total interaction energy by 2-fold for I78R and 50-fold for N180K. Overall, each AqKir mutant has stronger hydrogen bonding potential than the WT AqKir channel, which is consistent with an increase in the apparent affinity for PIP₂ in the mutant AqKir channels.

**Discussion**

To better understand the impact of evolutionary changes in ion channel structure and function, we studied the PIP₂ regulation of a sponge Kir channel using mutagenesis, electrophysiology, and MD simulations using a homology model. PIP₂ is a known regulator of many ion channel and transport molecules (1, 2). The solution of two crystal structures of vertebrate Kir channels complexed with PIP₂ highlights the interaction of positively charged residues between the transmembrane domain and the C terminus with the negatively charged headgroup phosphates (7, 45). The sponge Kir channel, AqKir, a distant relative of vertebrate Kir channels, lacks two basic residues in the PIP₂-binding site. This structural difference predicts low sensitivity to PIP₂ regulation in the sponge Kir channel.

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**Fig. 9. Homology models showing one AASt-PIP₂ molecule bound to tetrameric Kir channels.** Schematic representation is shown of AqKir WT (wild type) (A) or AqKir I78R/N180K channel (tetramers, two subunits are cyan and two subunits are pale cyan) (B) binding to AASt-PIP₂ (spheres, green carbon backbone) illustrating the position of residues 78 and 180 adjacent to the phospholipid headgroup. The basic (blue) side chains of the mutant channel form additional, favorable intermolecular interactions to stabilize the channel-ligand interaction. A, the amine group of Ile-78 is part of the secondary structure of the outer helix and is not shown in this representation. The images are zoomed in for clarity and thus lack the regions that include the most extracellular portions of the channel and most of the cytoplasmic portions.

**Table 1**

| Construct and residue | Total | van der Waals | Electrostatic |
|-----------------------|-------|---------------|---------------|
| WT                    |       |               |               |
| Residue Ile-78        | −6.77938 | −5.48137   | −1.29801     |
| Residue Asn-180       | −0.51548 | −0.46793    | −0.04755     |
| Mutant I78R           |       |               |               |
| Residue Arg-78        | −11.5338 | −2.60115    | −8.93265     |
| Residue Asn-180       | −0.33572 | −0.53888    | 0.203162     |
| Mutant N180K          |       |               |               |
| Residue Ile-78        | −2.45379 | −1.78845    | −0.66534     |
| Residue Lys-180       | −25.1472 | 0.879979    | −26.0272     |
| Double mutant I78R/N180K | |             |               |
| Residue Arg-78        | −7.89737 | −2.88633    | −5.01104     |
| Residue Lys-180       | −26.2704 | −3.94693    | −22.3235     |

**Enhanced total and electrostatic interaction energy of this mutant.** The WT channel may form only one weak (3.2–4.0 Å) hydrogen bond with PIP₂, whereas the mutants form a mixture of multiple weak and moderate (2.5–3.2 Å) hydrogen bonds with PIP₂. Overall, each AqKir mutant has stronger hydrogen bonding potential than the WT AqKir channel, which is consistent with an increase in the apparent affinity for PIP₂ in the mutant AqKir channels.
In whole-cell (TEVC) recordings, blocking PIP2 synthesis or stimulating PIP2 hydrolysis or dephosphorylation reduced AqKir currents, effects typically associated with a requirement for regulation by PIP2. Direct application of PIP2 to inside-out membrane patches failed to restore AqKirWT channel activity after rundown, consistent with an even lower apparent affinity for PIP2 following excision, as could occur due to loss of a post-translational modification that is not maintained under excised patch conditions (46). Consistent with this interpretation of our results, in TEVC and excised patches, AqKirWT currents were typically smaller than the AqKir mutant channels that displayed much higher affinity interactions with PIP2.

The partial reactivation of wild-type AqKir by long chain AASt PIP2, but not diC8 PIP2, suggests that hydrophobic interactions with the acyl chain of the lipid could be important in the mechanism of PIP2 activation of AqKir current, but alone are insufficient. It is also possible that other phospholipids maintain the WT current due to species-dependent differences in lipid specificity (47–49), which has not yet been examined in this channel.

Multiple positively charged residues located in the N and C termini of vertebrate Kir channels are important in coordinating with the negatively charged headgroup phosphates of PIP2 (1, 2, 7, 45). Do the various basic residues play equivalent roles? We have clarified the molecular significance by studying a channel that natively lacks these positively charged residues. Vertebrate levels of PIP2 sensitivity could be restored by mutational insertion of just one of the basic residues in AqKir, N180K (represented in 4-fold symmetry by the tetrameric channel). These results signify a particularly important role for this residue. MD simulations provided an explanation (see Fig. 10) in that N180K had the greatest impact on strengthening electrostatic interactions. Interestingly, a search for homologs of the sponge Kir channel in other distantly related metazoan species discovered putative Kir channels in sea anemones, which also lack the critical positive charge in the tether helix region. We suggest that mutation at this residue (Lys-K189 in cKir2.2) over the course of vertebrate evolution was perhaps the most critical final change that brought about the PIP2 dependence of vertebrate Kir channels.

Of course, measurements of channel activity in the excised patches following PIP2 application cannot determine directly whether the mutations affect PIP2 binding or channel gating. To this end, the MD simulations provide an explanation that highlights the more energetically favorable molecular interactions following recapitulation of one or both of the positive charges (Fig. 10 and Table 1).

Whether the sponge Kir channel is physiologically regulated by changes in PIP2 concentrations in its native environment is unknown, but the possibility surely exists. We ran a KEGG analysis (50) of the sponge genome and confirmed that all of the required components for inositol phosphate metabolism are present, and molecular cloning has previously reported that Gαq, phospholipase C, and protein kinase C exist in sponges (51). Thus, the capability for PIP2 regulation of Kir channels or other proteins via phospholipase C signaling pathways is evolutionarily old, with origins at the earliest branch point of extant animal phyla.

Conclusion

Nature’s mutations over the course of animal evolution have resulted in a high affinity PIP2 regulatory site within vertebrate Kir channels. However, critical features that confer high sensitivity to PIP2 in the vertebrate channels are missing in Kir channels from basal metazoans such as the channel from A. queenslandica studied here. The greatly enhanced PIP2 sensitivity of the AqKirWT channels would correlate with channel-specific properties of PIP2, which are likely to be distinct from those of mammalian PIP2.

FIGURE 10. Hydrogen bonding pattern for AqKir channels and the PIP2 headgroup. Stick representation of the hydrogen bonding pattern for AqKir residues 78 and 180 in WT (A), I78R (B), N180K (C), and I78R/N180K (D) channels to the PIP2 headgroup, based on MD simulations with homology models (see “Experimental Procedures” details). Distances between potential hydrogen bonding partners, demarcated by dotted lines, are given in green text. Atoms are colored by standard convention. The WT channel may form only one weak (3.2–4.0 Å) hydrogen bond with PIP2, whereas the mutants form a mixture of multiple weak and moderate (2.5–3.2 Å) hydrogen bonds with PIP2.
PIP₂ Sensitivity of a Sponge Kir Channel

AqKir mutants containing two positive charges confirms the absolute necessity of these basic residues for PIP₂ regulation, which is consistent with our homology model and MD simulations of AqKir complexed with PIP₂. Furthermore, our results support a functional priority for a single positive charge (residue 180 in AqKir) in recapitulating the high sensitivity to PIP₂.

Changes in the binding site over time, in response to forces that drive evolution, could include changes in the bioavailability of PIP₂ (15) or molecules involved in cell-cell signaling. Conservation of Kir channel selectivity and inward rectification mechanisms (16) have been retained over millions of years of evolution, but acquisition of a new function, PIP₂ regulation by high affinity interaction, is more recent, occurring after the prokaryotic-eukaryotic split.

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