

Main Manuscript for

A revised mechanism of action of hyperaldosteronism-linked mutations in cytosolic domains of GIRK4 (KCNJ5).

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Classification

Biological Sciences: Physiology.

Keywords

Primary aldosteronism, Kir3, selectivity, K⁺ channel opener, electrophysiology

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Conceptualization, B.S., N.D. and A.K.B; Methodology, B.S., R.H.-J. and N.D.; Investigation and analysis, B.S., R.H.-J, H.P.R. and N.T.; Writing – Original Draft, B.S., H.P.R and N.D.; Writing – Review & Editing, B.S., R.H.-J., N.D.; Funding Acquisition and supervision, N.D. and A.K.B.

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Abstract

G-protein gated, inwardly rectifying potassium channels (GIRK) mediate inhibitory transmission in brain, heart, and adrenal cortex. GIRK4 (KCNJ5) subunits are abundant in the heart and adrenal cortex. Multiple mutations of KCNJ5 cause primary aldosteronism (PA). According to a leading concept, mutations in the pore region of GIRK4 cause loss of K⁺ selectivity; the ensuing Na⁺ influx depolarizes zona glomerulosa cells and activates voltage gated Ca²⁺ channels, inducing hypersecretion of aldosterone. The concept of selectivity loss has been extended to mutations in cytosolic domains of GIRK4 channels, remote from the pore region. We expressed GIRK4_{R52H}, GIRK4_{E246K}, and GIRK4_{G247R} mutants in Xenopus oocytes and human adrenocortical carcinoma cell line (HAC15). Whole-cell currents of heterotetrameric GIRK1/4_{R52H} and GIRK1/4_{E246K} (but not GIRK1/4_{G247R}) channels were greatly reduced compared to GIRK1/4_{WT}. Nevertheless, all heterotetrameric mutants retained full K⁺ selectivity and inward rectification. When expressed as homotetramers, only GIRK4_{WT}, but none of the mutants, produced wholecell currents. Confocal imaging, single channel and Förster Resonance Energy Transfer (FRET) analyses showed: 1) reduction of membrane abundance of all mutated channels, especially as homotetramers, 2) impaired interaction with Gβγ subunits, and 3) reduced open probability of GIRK1/4_{R52H}. VU0529331, a GIRK4 opener, activated homotetrameric GIRK4_{G247R} channels, but not GIRK4_{R52H} and GIRK4_{E246K}. Our results suggest impaired gating (GIRK 4_{R52H}) and expression in plasma membrane (all mutants). We suggest that, contrary to the previously proposed mechanism, R52H and E246K mutants are loss-offunction rather than gain-of-function/selectivity-loss mutants. Hence, GIRK4 openers may be a potential course of treatment for patients with cytosolic N- and C-terminal mutations.

Significance Statement

Mutations in KCNJ5 gene, which encodes for the GIRK4 subunit of G-protein inwardly rectifying K+ channels, are the main cause of primary aldosteronism, a major contributor to secondary hypertension. We report that three mutations in the cytosolic domain of GIRK4 cause loss-of-function, contrary to the prevailing concept that these

mutations cause loss of selectivity and subsequent depolarization, i.e. essentially gain-of-function. Our findings correct the existing misconception regarding the biophysical mechanism that impairs the channel function, and may provide indications for future personalized treatment of the disease.

Main Text

Introduction

G-protein gated, inwardly rectifying potassium channels (GIRK; Kir3) mediate inhibitory signaling via G-protein coupled receptors (GPCR) in the brain, heart, and adrenal cortex. There are four GIRK subunits (GIRK1-4) forming homo- (GIRK2, GIRK4) or heterotetrameric channels (GIRK1/2, GIRK1/3, GIRK1/4, GIRK2/3). G $\beta\gamma$ dimers are the main gating factor of GIRK channels (1-3); upon GPCR activation, G $\beta\gamma$ dissociates from G $\alpha_{i/o}$, binds to the channel and activates it (4). Like in all inward rectifiers, in GIRKs the outward current is smaller than the inward, due to the occlusion of the cytoplasmic pore by Mg²⁺ and polyamines (5). GIRK4 subunits, encoded by *KCNJ5* gene, are mostly abundant in heart and adrenal cortex, forming GIRK4 homo- and/or GIRK1/4 (*KCNJ3/KCNJ5*) heterotetrameric channels (6-9). GIRK channels play critical roles in regulation of heart rate and aldosterone secretion (6, 7). However, subunit composition of GIRK channels in human zona glomerulosa cells of the adrenal cortex is yet to be elucidated. Notably, much lower mRNA levels of GIRK1 compared to GIRK4 have been reported in these cells (6), indicating a possible predominance of GIRK4 homotetramers.

Aldosterone, the steroid hormone secreted by the zona glomerulosa cells, is essential for blood pressure regulation. In physiological conditions, aldosterone secretion is regulated by the interplay between angiotensin II and plasma K⁺ concentration, [K⁺] (10). Angiotensin activates angiotensin II type 1 receptor, leading to depolarization of aldosterone-secreting cells through inhibition of K⁺ channels and Na⁺/K⁺ pump (11, 12). Consequently, L- and T-type voltage-gated Ca²⁺ channels (VGCC) open, Ca²⁺ enters the cell and activates aldosterone secretion. Any change in the level of angiotensin II or [K⁺] (even 1 mM) affects aldosterone secretion (10, 13).

Primary aldosteronism (PA) is a disease characterized by hypersecretion of aldosterone. PA accounts for 90% of secondary hypertension cases, approximately 10% of hypertensive patients worldwide (14). The main causes of PA are somatic and germline mutations in *KCNJ5*, *CACNA1D*, *CACNA1H*, *ATP1A1*, *ATP2B3*, *CTNNB1*, *ARMC5* genes (12). Sporadic and familial mutations in *KCNJ5* account for up to 70% of PA cases, often accompanied by adrenal adenoma (14, 15). *KCNJ5* germline mutations cause familial hyperaldosteronism type III, out of four types (16). In 2011, Choi et al. (6) showed that PA-causing mutations in the GYG motif of the selectivity filter (G151R) or in close proximity to it (T158A and L168R) in GIRK4 cause a loss of selectivity for K⁺, yielding K⁺/Na⁺ non-selective GIRK4 and GIRK1/4 channels (6). It is widely accepted that this results in depolarization caused by the influx of Na⁺ and a consequent influx of Ca²⁺ through L- and T-type VGCCs followed by constitutive aldosterone secretion and hypertension (17, 18). Since 2011, additional PA-linked mutations were discovered in *KCNJ5* gene, in the pore region or in the cytosolic N- and C-terminal domains of GIRK4 (16, 19, 20).

Loss of K⁺ selectivity and inward rectification have been reported for R52H and E246K mutations located in GIRK4 cytosolic N- and C-terminal domains, respectively (20). However, unlike for the well-established effect of mutations in the pore region (21), it is more challenging to comprehend how mutations in cytosolic domains, remote from the pore (Fig. 1), would affect selectivity. Despite that, the same gain-of-function/loss-of-selectivity mechanism suggested for PA induced by pore mutations (6) was extended for these cytosolic domain mutations as well. G247R is another mutation within the cytosolic domain found to be casually related to PA, but the mechanism remains controversial. Both loss-of-function (22) or no alteration in channel activity (20) have been reported for the G247R mutation. Thus, the biophysical and cellular mechanisms of GIRK4 channel malfunction caused by PA-linked mutations in the cytosolic domain remain incompletely understood.

Here, we investigated GIRK4_{R52H}, GIRK4_{E246K} and GIRK4_{G247R} mutants expressed in Xenopus oocytes and in human adrenocortical carcinoma cell line (HAC15). We demonstrate that all heterotetrameric GIRK1/4 mutants examined remain selective to K⁺ and show an unimpaired inward rectification. In contrast, the activity of heterotetrameric GIRK1/4_{R52H} and GIRK1/4_{E246K} (but not GIRK1/4_{G247R}) channels is severely impaired compared to GIRK1/4_{WT}. Homotetrameric GIRK4_{R52H}, GIRK4_{E246K}, and GIRK4_{G247R} mutants are non-functional. Single channel analysis, confocal imaging and Förster Resonance Energy Transfer (FRET) suggested both impaired gating and Gβγ interaction, and compromised plasma membrane (PM) expression in GIRK4_{R52H} and GIRK4_{E246K}. For GIRK4_{G247R}, the only significant defect was a severe impairment of PM expression for its homotetrameric composition. A homotetrameric GIRK4 channel opener, VU0529331, enhanced GIRK4_{G247R} currents, suggesting a potential strategy for treatment in affected patients. Our results suggest that R52H and E246K mutations are loss of function mutations that impair channel gating and membrane abundance, but not K⁺ selectivity or inward rectification. Hence, we suggest that the mechanism previously proposed for R52H and E246K mutants should be revised.

Results

GIRK1/4WT and GIRK1/4 mutants produce basal and Gβγ-activated currents in Xenopus laevis oocytes

The mutated amino acids (R52, E246, G247) are fully conserved in all GIRK subunits (Fig. 1A), suggesting their functional significance. They are located in the cytosolic domain far from the selectivity filter, as illustrated in Fig. 1B for the homologous amino acids in the crystal structure of GIRK2 (23).

We expressed heterotetrameric GIRK1/4 channels in *Xenopus laevis* oocytes and recorded whole-cell currents using two electrode voltage clamp. For GIRK1/4_{WT} channels, switching from low-K⁺ (2 mM K⁺) ND96 solution to a high-K⁺ (24 mM K⁺) HK24 solution yielded an inward current reflecting the GIRK's basal activity (I_{basal}). GIRK1/4_{WT} was further activated by G $\beta\gamma$, which is evident from the strong increase (6.86±0.42 fold)

in the constitutive K⁺ current ($I_{\beta\gamma}$) in G $\beta\gamma$ -expressing oocytes (Fig. 1C). I_{basal} of GIRK1/ I_{R52H} and GIRK1/ I_{E246K} was significantly smaller than I_{basal} of GIRK1/ I_{WT} , and in some experiments it was indistinguishable from native oocyte's currents, especially for GIRK1/ I_{R52H} (Fig. 1D). To determine GIRK I-V relationships, we used voltage ramp protocols (Fig. 1E,F). The I-V curves of basal and G I_{V} -evoked currents of GIRK1/ I_{WT} showed the distinctive inward rectification pattern (Fig. 1E). I_{basal} of GIRK1/ I_{R52H} and GIRK1/ I_{E246K} often showed seemingly non-rectifying I-V relationships (Fig. 1F, left, and Fig. S1B); however, as we show below, this was an artifact arising from the very small GIRK current, comparable to oocyte's endogenous currents. GIRK1/ I_{G247R} showed distinctive inward rectification, yet I_{basal} was significantly lower compared to GIRK1/ I_{WT} (Fig. 1D, F). On the other hand, I_{I} of GIRK1/ I_{G247R} was similar to I_{I} of GIRK1/ I_{WT} , while I_{I} of GIRK1/ I_{R52H} and GIRK1/ I_{E246K} was significantly lower (Fig. 1D, F).

We also coexpressed GIRK1/4 with the muscarinic acetylcholine receptor M_2 (M_2R). Application of acetylcholine (ACh) evoked additional GIRK current, I_{evoked} (Fig. S1A) (24). Incidentally, however, addition of ACh also activated oocyte's endogenous Ca^{2+} dependent chloride channels, possibly via endogenous G_q -coupled muscarinic receptors occasionally present in the oocytes, or promiscuous coupling of m2R to G_q (25). These CI^- currents were not sensitive to the GIRK channel blocker, Ba^{2+} , decayed slowly after agonist washout, and were blocked by the injection of the Ca^{2+} chelator BAPTA (Fig. S1) (26). Ca^{2+} dependent chloride currents may be a major cause of artifact when estimating the reversal potential (V_{rev}) when the expressed GIRK's currents are small (Fig. S1B, C). To avoid these artifacts, in the following we activated the GIRK channels by co-expressing $G\beta\gamma$.

GIRK4 mutants are selective for K⁺ and retain inward rectification properties

To examine whether GIRK4 mutants lose selectivity or inward-rectification, we coexpressed GIRK1/4 channels with Gβγ and measured whole-cell currents while applying voltage ramps in four concentrations of extracellular K⁺, [K⁺]₀ (Fig. 2A, C). V_{rev} was measured from the intercept of the net GIRK I-V curve with the voltage axis (Fig. 2B, C). Plots of V_{rev} as a function of $[K^+]_0$ (plotted on log_{10} scale, Fig. 2C) gave fairly straight lines. The slopes of the V_{rev} vs. log₁₀[K⁺]₀ plots, in mV per tenfold change in [K₀] (decade), were 53.33 \pm 0.77, 58.49 \pm 1.6, 58.9 \pm 2.96, and 54.36 \pm 0.42 for GIRK1/4_{WT}, GIRK1/4_{R52H}, GIRK1/ 4_{E246K} and GIRK1/ 4_{G247R} , respectively (Fig. 2D) . These values are close to ~58 mV/decade predicted for a selective K⁺ channel (21). In contrast, the slope for GIRK1/4_{Y152C}, a known non-selective pore mutant (27) used as control, was 13.45±4 mV/decade (Fig. 2C,D). Permeability ratios (P_{Na}/P_K) for GIRK1/4_{WT} and the R52H, E246K and G247R were close to zero, as opposed to the pore mutant GIRK4 $_{Y152C}$ (Fig. 2E). Thus, these cytosolic N- and C-terminal mutations do not affect the K⁺ selectivity. We also quantified the extent of inward rectification (Fir) (see Methods and Fig. 2B). Fir was similar in GIRK1/ 4_{WT} and all mutants (Fig. 2F), usually <0.1, except GIRK1/ 4_{Y152C} where F_{ir} was significantly higher, indicating partial loss of rectification in this channel, as reported (27). We conclude that mutated channels are inwardly rectifying to the same extent as WT channels.

R52H, E246K, and G247R mutations in GIRK4 impair channels' surface expression and interaction with GBy

To assess the abundance of GIRK4 mutants in the PM, we expressed GIRK4 channels tagged with YFP at the N-terminus (YFP-GIRK4), with or without Gβy, as homoand heterotetramers with GIRK1 (Fig. 3 and 4, respectively). Since the abundance of GIRK4 vs. GIRK1/4 in aldosterone-secreting cells remains an open question (see Introduction), we first studied the GIRK4 homotetramers (Fig. 3). Each mutant was studied in a separate experiment and compared with YFP-GIRK4_{WT}. Measurement of YFP fluorescence showed a significantly lower PM expression of YFP-GIRK4_{R52H} and YFP-GIRK4_{E246K} compared to YFP-GIRK4_{WT}, with or without coexpression of G β y (Fig. 3A, B). PM expression of YFP-GIRK4_{G247R} without Gβγ was reduced, but with Gβγ it was similar to YFP-GIRK4_{WT} (Fig 3B). Electrophysiological measurements showed that I_{basal} of GIRK4 homotetramers was very small, usually indistinguishable from native oocyte currents. Although I-V relationships occasionally showed inward rectification, the small size of GIRK currents did not allow a rigorous analysis (Fig S2). Conversely, Gβγ-evoked YFP-GIRK4_{WT} currents were quite substantial, ~ 0.5 μA on the average (Fig. 3C) and showed typical GIRK current characteristics (Fig. 3D). In contrast, I_{βν} of all homotetrameric mutants was almost undetectable (Fig. 3C,D). This was true even for the homotetrameric YFP-GIRK4_{G247R}, despite its comparable level of surface expression compared to YFP-GIRK4_{WT}, in presence of G $\beta\gamma$. Thus, homotetramers of the 3 mutants under study show impaired surface expression and appear non-functional.

We next studied heterotetramers of GIRK1 with WT or mutated YFP-GIRK4. The surface expression of GIRK1/YFP-GIRK4_{E246K} was significantly lower than for the wild-type channel, with or without coexpression of G $\beta\gamma$ (Fig. 4A, B). The PM levels of GIRK1/YFP-GIRK4_{R52H} and GIRK1/YFP-GIRK4_{G247R} were not significantly different from GIRK1/YFP-GIRK4_{WT}. However, a significant reduction for both mutants was observed when they were coexpressed with G $\beta\gamma$ (compared to GIRK1/YFP-GIRK4_{E246K} coexpressed with G $\beta\gamma$). In the same experiments, all mutants showed reduced currents compared to WT, with or without G $\beta\gamma$ (Fig. 4C).

To examine whether R52H, E246K, and G247R mutations impair GIRK4 interaction with G $\beta\gamma$, we used spectral FRET (28). We coexpressed CFP-G $\beta\gamma$ (donor; usually at two concentrations) with YFP-GIRK4 (acceptor), and GIRK1 (Fig. 4D,E, S3). YFP-labeled, G protein-insensitive inwardly rectifying K channel IRK1 (Kir2.1; *KCNJ2*) was used as negative control (Fig. S3). To circumvent variability between oocyte batches, we calculated donor/acceptor molar ratio in each oocyte using the double-labeled YFP-GIRK2-CFP (coexpressed with GIRK1) as a molecular caliper (29). In sensitized emission FRET, the FRET efficiency increases with donor/acceptor ratio (30). The FRET titration curve reaches saturation at high donor/acceptor molar ratios (30) (>5 for GIRK-G $\beta\gamma$ FRET

pair (29)). Since in our experiments we have not achieved full saturation (Fig. 4D), we divided the data into bins with comparable donor/acceptor molar ratios, in the range where the representation of each mutant or WT was at least 3 experimental points (Fig. 4E). We found that FRET efficiency (E_{app}), that reports interaction with G $\beta\gamma$, was significantly impaired for GIRK1/YFP-GIRK4_{R52H} and GIRK1/YFP-GIRK4_{E246K} in all donor/acceptor molar ratio ranges. Interaction of G $\beta\gamma$ with GIRK1/YFP-GIRK4_{G247R} was similar or slightly smaller than with GIRK1/YFP-GIRK4_{WT} (Fig. 4E).

Single channel analysis of GIRK4 mutants show reduced P_o of GIRK1/4_{R52H} but not of GIRK1/4_{E246K}

To directly address the possible defects in channel function on molecular level, we measured G $\beta\gamma$ -induced single channel activity using cell-attached patch clamp (Figure 5A, B). We used GIRK1/4 heterotetramers, since mutated GIRK4 homotetramers were found non-functional. GIRK1/4 channels were coexpressed at low surface density, with G $\beta\gamma$ using doses of RNA that usually produce maximal Po in GIRK1/4WT. Figure 5B shows exemplary recordings of GIRK1/4WT, GIRK1/4R52H and GIRK1/4E246K. We found that the open probability (Po) of GIRK1/4R52H was significantly lower than Po of GIRK1/4WT, while Po of GIRK1/4E246K was not significantly different (Fig. 5C). Neither single channel current (isingle) (Figs. 5D and S4B) nor single channel conductance (g) (Figs. 5E and S4C) were affected by the mutations.

Another way to assess mutation-induced changes in P_o is from whole-cell data of Figs. 3 and 4. The whole-cell current, I, is given by the equation $I=i_{single}*P_o*N$ (21), where N is the total number of channels in the PM. N is proportional to fluorescence intensity of the labeled channels (in arbitrary units), E_{AU} . Thus, whole-cell current normalized to PM level, I/E_{AU} , is proportional to I/N, therefore $I/E_{AU} \sim i_{single}*P_o$. Since i_{single} is not affected by mutations, differences in I/E_{AU} reflect differences in P_o . The homotetrameric YFP-GIRK4 mutants exhibited a greatly reduced I/E_{AU} (Fig. S5A), suggesting a great reduction in P_o (impaired gating) in addition to the impaired expression. For heterotetrameric GIRK1/YFP-GIRK4 mutants, the normalization procedure showed an impaired P_o for GIRK1/YFP-GIRK4_{R52H} but not for GIRK1/YFP-GIRK4_{E246K} or GIRK1/YFP-GIRK4_{G247R}, supporting the single channel results and indicating that the E246K and G247R mutations do not impair the gating properties of heterotetrameric GIRK1/4 channels (Fig. S5B).

Expression of GIRK1/4 in HAC15 cells

The type of K⁺ channels in the zona glomerulosa cells varies between species, GIRK4 is not present in the adrenal cortex of rats, and probably also in mice (31). Thus, rodent models are ineffective to study the effect of GIRK4 mutations in vivo (31, 32). Hence, to corroborate that mutation-induced changes in GIRK4 function observed in *Xenopus* oocytes also take place in a more relevant cellular environment, we used the human adrenocortical carcinoma cell line (HAC15). First, we tested the presence of endogenous GIRK1 and GIRK4 using Western blot. Both GIRK1 and GIRK4 proteins were

present in these cells (Fig. 6A). However, whole-cell recordings of untransfected cells, or cells transfected with D₂ dopamine receptor, did not show any basal or dopamine (DA)-evoked GIRK current (Fig. 6B, left). Next, we transfected the cells with GFP-GIRK1 and GIRK4 (WT or mutants), with Gβγ subunits and the D₂ receptor. I_{evoked} was elicited by application of 100 μ M dopamine. The activity of GIRK1/4_{R52H} and GIRK1/4_{E246K} mutants was significantly reduced compared to GIRK1/4_{WT}, while GIRK1/4_{G247R} did not differ from GIRK1/4_{WT} (Fig. 6B).

The GIRK2 and GIRK4 opener VU0529331 activates GIRK4_{G247R} homotetramer

Lastly, we aimed to test whether the activity of mutated channels can be rescued by the recently discovered GIRK2 and GIRK4 opener VU0529331, which activates the channels in a G $\beta\gamma$ -independent manner (33). We used oocytes expressing GIRK4 homotetramers, with or without G $\beta\gamma$. VU0529331 activated GIRK4 $_{WT}$ and GIRK4 $_{G247R}$ in a dose-dependent manner, and the activated channels showed the typical inwardly rectifying I-V relationships (Fig. 7). No response could be seen in oocytes injected with GIRK4 $_{R52H}$ and GIRK4 $_{E246K}$ RNA.

Discussion

Loss of selectivity of GIRK4 (*KCNJ5*) channels is believed to be the leading cause of PA, by producing constitutive depolarization of zona glomerulosa cells (6, 17, 27). This concept was extended from mutations in the GIRK4 pore region (6, 27, 34) to additional mutations in the transmembrane domain of GIRK4 (34, 35) and, eventually, to mutations in the cytosolic N- and C-termini (17, 20), despite the identification of two cytosolic domain mutations (R115W and E246G) that impaired membrane abundance of GIRK4 rather than selectivity (19). We studied PA-linked GIRK4 mutations of three conserved amino acids located in the cytosolic domain (Fig. 1): R52 and E246, that were reported to cause loss of selectivity of GIRK4 channels (20), and G247. Here, we confront the selectivity loss concept for C- and N-terminal mutations; we show that R52H, E246K and (to a milder extent) G247R mutations cause a complete or partial loss of function and/or channel abundance in PM, but do not alter channel selectivity or inward rectification.

Many of the *KCNJ5* mutant studies (including our work) were done in *Xenopus laevis* oocytes used as heterologous expression system (20, 22, 34, 36-38). We first thoroughly investigated possible sources of artifact in this system. Like other cells, *Xenopus* oocytes possess a number of endogenous ion channels that yield small currents, usually in the range of a few tens of nA, but may give rise to artefactual interpretations if the expressed exogenous ion channels show poor expression or function. For instance, activation of M2 receptors by ACh often elicited outwardly-rectifying endogenous Ca²⁺-dependent Cl⁻ currents (25) that were abolished by the Ca²⁺ chelator BAPTA (Fig. S1). Even the remaining leak currents may be mistaken for currents of GIRK mutants if the expressed currents are small. To avoid these artifacts, we

activated GIRK channels by coexpressing G $\beta\gamma$ subunits at doses that maximally activate the WT GIRKs; yet, even under these conditions the homotetrameric GIRK4_{R52H} and GIRK4_{E246K} did not produce detectable currents.

Another source of ambiguity in GIRK4 mutant studies is the subunit composition. In view of the low abundance of GIRK1 RNA in zona glomerulosa (6), it is conceivable that GIRK4 homotetramers are abundant in aldosterone-secreting cells. However, most heterologous expression studies have been done with GIRK1/4 heterotetramers (19, 27, 35, 36, 39). Only two GIRK4 pore mutants have been tested as homotetramers, and reported to be functional and non-selective (6). Homotetrameric GIRK4 $_{G247R}$ mutants was tested on the background of an additional mutation, S143T (22). The homotetrameric GIRK4 $_{S143T}$ is a pore mutant that expresses better than GIRK4 $_{WT}$ and preserves inward rectification (40, 41), but its selectivity has not been carefully characterized. The GIRK4 $_{S143T}$ homotetramers showed large I_{basal} (22), which is different from GIRK4 $_{WT}$ homotetramers that have very small I_{basal} and large $I_{\beta\gamma}$, as we have demonstrated (Fig. S2). To address this conundrum, we examined both homotetrameric and heterotetrameric (with GIRK1) GIRK4, WT and mutants.

Our central finding was that, in GIRK1/4 heterotetrameric context, all mutations strongly reduced the whole-cell basal GIRK current in *Xenopus* oocytes, and R52H and E246K (but not G247R) mutations greatly reduced the G $\beta\gamma$ -induced currents, both in in *Xenopus* oocytes and in HAC15 cells (Figs. 1 and 6). The homotetrameric mutant channels gave practically no detectable currents at all (Fig. 4). These results indicate that all three mutations cause loss of function, which is more severe in homotetrameric than in heterotetrameric channels.

In contrast, we found no changes in K⁺ selectivity or inward rectification (in the heterotetrameric context; Fig. 2). The transmembrane pore of K⁺ channels is a conserved structure that enables K⁺ ions to permeate rapidly through the channel over Na⁺ ions, and mutations in the pore region shift the permeability ratio towards Na⁺ (42, 43). The cytosolic domain of inwardly rectifying K⁺ channels forms an additional, cytoplasmic pore that is a continuation of the transmembrane pore but does not contain a selectivity filter (44-46). Several amino acids of the cytosolic domain lining the pore control channel's inward rectification by altering Mg²⁺ and polyamine block, but not the ion selectivity (5, 46, 47). We measured the reversal potential of Gβy-activated GIRK1/4 current in four external K⁺ concentrations and found a linear dependence between $log[K^+]$ and V_{rev} with a slope close to that predicted by the Nernst equation for a pure K⁺ pore, and identical to that of the wild type channel. Analysis using the Goldman-Hodgkin-Katz equation confirmed that the mutated channels conducted K⁺ but not Na⁺. Evaluation of the degree of inward rectification showed that inwardrectification of mutated channels was similar to WT (48). Our results clearly demonstrate that all mutants studied here are selective to K⁺ over Na⁺, and rectify, to a similar extent.

To understand the mechanisms underlying loss of function of the mutants, we employed several approaches. Confocal imaging in Xenopus oocytes revealed that PM abundance of mutated homo- or heterotetrameric channels was significantly reduced. Analysis of macroscopic whole-cell currents normalized to PM expression indicated impaired gating (in addition to reduced PM abundance) of R52H, and impaired PM abundance, but not gating, of E246K and G247R heterotetramers (Fig. S5). Single channel recordings (Fig. 5) confirmed the reduction in open channel probability (P_0), which indicates impaired gating, of the heterotetrameric GIRK1/4_{R52H}. Finally, FRET experiments suggested an impaired interaction of the R52H and E246K mutant channels with the main gating factor, Gβy (Fig. 4D,E). On the other hand, while the heterotetrameric GIRK1/ 4_{E246K} had an impaired interaction with G $\beta\gamma$ (FRET data), we did not see significant reduction in Po. Hence, we suggest that R52H mutation impairs both gating and PM abundance of homo- and heterotetrameric channels. E246K mutation impairs mainly channel expression, and to a lesser extent channel activity (in the heterotetrameric context), mainly because of the reduced GBy affinity. G247R mutation had the smallest impact on the heterotetrameric channel, and showed mild impairment in activity (I_{basal} , but not $I_{\beta \nu}$) and in PM abundance.

In our study, none of the mutated homotetramers produced detectable whole-cell currents, with or without coexpression of G $\beta\gamma$. Homotetrameric GIRK4 $_{G247R}$ showed PM expression similar to the WT homotetramer, but it was not activated by G $\beta\gamma$. Remarkably, however, it was activated by the GIRK4 opener, VU0529331. Since the heterotetrameric GIRK1/4 $_{G247R}$ does not show a significant loss of function, yet the patients with G247R mutation express a pathologic phenotype, these results indicate that the subunit composition in human aldosterone-secreting cells tends toward homotetrameric GIRK4 channels rather than heterotetrameric GIRK1/4.

On the other hand, GIRK4_{R52H} and GIRK4_{E246K} showed very low expression in the PM compared to GIRK4_{WT}, and this could be the main reason for the absence of current. These two mutants could not be activated by VU0529331 or G $\beta\gamma$. However, since we see some expression, it is possible that other openers will be able to activate these channels.

Conclusions and prospects

Taken together, our results suggest that the mechanism proposed for R52H and E246K mutants (20) should be revised. We propose that, as opposed to pore mutations, N- and C-terminal domain mutations studied here reduce channel activity and PM abundance. This results in a decrease in resting K⁺ conductance, therefore the resting membrane potential becomes more positive, which triggers Ca²⁺ flow into the cell and hypersecretion of aldosterone. These results are supported by the previous study of Cheng et al. (19), which showed reduced activity and PM abundance of two additional N- and C-terminal domain mutants. Knowing the exact biophysical mechanism that impairs the channel is crucial for setting the course of treatment. While patients with

pore mutations, which yield gain-of-function non-selective Na⁺/K⁺ channels, may benefit from treatment with channel blockers, patients with cytosolic domain mutations may potentially benefit from treatment with GIRK4 channel openers to achieve the same effect: prevention of constitutive depolarization of zona glomerulosa secretory cells.

Materials and Methods

DNA constructs and RNA

cDNA constructs of rat GIRK1, N-terminally YFP-tagged GIRK1 (YFP-GIRK1), GIRK4 $_{WT}$, GIRK4 $_{R52H}$, GIRK4 $_{E246K}$, and GIRK4 $_{G247R}$ were inserted into pBS-MXT vector. DNA of human N-terminally YFP-tagged GIRK4 $_{WT}$ (YFP-GIRK4 $_{WT}$) was a gift from Prof. Wolfgang Schreibmayer, and mutations were made to produce YFP-GIRK4 $_{R52H}$, YFP-GIRK4 $_{E246K}$, and YFP-GIRK4 $_{G247R}$. mRNAs were prepared as described previously (24).

The amounts of mRNA injected per oocyte were varied according to the experimental design, in ng/oocyte: 0.5-2 GIRK1, 0.5-2 YFP-GIRK1, 0.25-1 GIRK4, 0.25-5 YFP-GIRK4. For maximal channel activation by G $\beta\gamma$, we injected 5 ng G β and 1-2 ng G γ mRNA. These weight ratios were chosen to keep approximately equal molar amounts of G β and G γ RNA. RNA of the muscarinic 2 receptor (M₂R), if present, was 1 ng. 25 ng of the anti-GIRK5 oligo nucleotide antisense XIR was routinely injected to prevent the formation of GIRK1/5 channels (49).

Ethical approval of Xenopus laevis, oocyte preparation, and electrophysiology

Experiments have been approved by Tel Aviv University Institutional Animal Care and Use Committee (permit #01-16-104). Maintenance and surgery of female frogs were as described previously (50). All materials were from Sigma unless indicated otherwise. Oocytes were defolliculated by collagenase, injected with RNA and incubated for 3 days at 20-22°C in ND96 solution (low K⁺) (in mM: 96 NaCl, 2 KCl, 1 MgCl₂, 1 CaCl₂, 5 HEPES, pH 7.5), supplemented with 2.5 mM sodium pyruvate, 100 µgml⁻¹ streptomycin and 62.75 µgml⁻¹ penicillin (or 50 µgml⁻¹ gentamycin) (24). Whole-cell GIRK currents were measured using standard two-electrode voltage clamp method at $20-22^{\circ}$ C, in different K⁺ concentrations (8, 24, 48, 72 or 96 mM K⁺ (HK)) (51). Different concentrations of K⁺ solutions were obtained by mixing ND96 with a 96 mM K⁺ solution containing, in mM: 96 KCl, 2 NaCl, 1 CaCl₂, 1 MgCl₂, 5 HEPES, pH adjusted to 7.5 with KOH. Acetylcholine (ACh) was added to HK solution as agonist at 10 μ M, Ba²⁺ (2.5 mM) was used to block the channel. Current-voltage (I-V) relations were obtained using 2 s voltage ramps from -120 mV to +50 mV. Net GIRK I-V relationships were obtained by subtracting the current remaining after blocking all GIRK activity with 2.5 mM Ba²⁺. In one series of experiments (Fig. 2), we obtained the net GIRK I-V relationship by subtracting averaged I-V relationship from native oocytes injected with only GBy. For small GIRK currents (in low [K⁺]_{out}) this procedure is superior over subtraction of Ba²⁺blocked currents, avoiding possible inaccuracies caused by different extent of Ba²⁺ block of inward and outward GIRK currents (51).

VU0529331 (Alomone Labs; V-155), an opener of homotetrameric GIRK2 and GIRK4 channels, was dissolved in 100% DMSO to a final concentration of 25 mM. To measure GIRK4 response to VU0529331, the drug was diluted into HK24 solution to get HK24 + 60 μ M VU0529331 solution. Consecutively, this solution was diluted again to get 20 μ M, and so on for all concentrations.

HAC15 cells culture, transfection, biochemistry and electrophysiology

Cells were acquired from ATCC (ATCC® CRL3301 ™) and cultured as described (52). Cells were grown in Dulbecco's Modified Eagle's/Ham's F-12 medium (DMEM/F-12) (Gibco #11330-05) containing 10% Cosmic calf serum (Hyclone #SH30), 1% L-Glutamine (Sigma-Aldrich; A7506), 1% ITS (Becton Dickenson - FAL354352) and penicillin—streptomycin (Sigma-Aldrich; P4333).

For Western blot, cultured HAC15 cells were treated as described previously (53). GIRK1/4 RNA-injected oocytes were used as positive control. Oocytes were homogenized on ice in homogenization buffer (20 mM Tris, pH 7.4, 5 mM EGTA, 5 mM EDTA, and 100 mM NaCl) containing protease inhibitor mixture (Roche Applied Science). The homogenates were centrifuged at $3000 \times g$ for 5 min at 4 °C and the pellet was discarded. Protein samples (35µg) were electrophoresed on 12% polyacrylamide-SDS gel and transferred to nitrocellulose membranes for Western blotting with antibodies; anti-GIRK1 at 1:300 dilution (Alomone Labs, APC-005) or anti-GIRK4 at 1:100 dilution (Alomone Labs, APC-027), and GAPDH in 1:1000 or 1:2000 dilution (Cell Signaling Technology, P04406). Goat Anti-Rabbit IgG Antibody, (H+L) HRP conjugated secondary antibody at 1:40,000 dilution was applied (Jackson ImmunoResearch Labs 111-035-144, RRID:AB_2307391).

For electrophysiological experiments, cells were plated in a 6-well plate and were transfected with combinations of the following DNAs as designed for specific experiments: 0.5 ng GFP-GIRK1, 0.5 ng GIRK4_{WT}, 0.5 ng GIRK4_{R52H}, 0.5 ng GIRK4_{E246K}, 0.5 ng GIRK4_{G247R}, 0.5-1 ng G β_1 , 0.5-1 ng G γ_2 (all in pcDNA3.1); 0.5 ng D γ_2 receptor in pXOOM (the latter was kindly provided by Kristoffer Sahlholm). 24-36 hours after transfection cells were moved onto 13 mm cover slips, coated with fibronectin and electrophysiological measurements were performed in the next 24-48 hours.

Whole-cell currents were measured using Axopatch 200B (Molecular Devices, Sunnyvale, CA). Holding potential was -80 mV, and voltage ramps were from -120 to +20 mV, no correction for junction potential (~13 mV) was made. Currents were recorded in low K⁺ solution (in mM): NaCl 136, KCl 4, CaCl₂ 2, MgCl₂ 2, HEPES 10, NaH₂PO₄ 0.33, Glucose 10, pH 7.4, or in high K⁺ (HK48) solution (in mM): NaCl 92, KCl 48, CaCl₂ 2, MgCl₂ 2, HEPES 10, NaH₂PO₄ 0.33, Glucose 10, pH 7.4. 100 μ M dopamine was used to measure GPCR response. Electrode solution contained (in mM): NaCl 6, KCl 22, K-Gluconate 110, MgCl₂ 2, HEPES 10, EGTA-HOH 1, ATP-K₂ 2, GTP-Tris 0.5, pH 7.2.

Cell-attached single channel recordings in *Xenopus* oocytes

Patch clamp experiments were done using Axopatch 200B as described (54). Currents were recorded at -80 mV, filtered at 2 kHz and sampled at 20 kHz. Patch pipettes had resistances of 1.4–3.5 MΩ. Pipette solution contained, in mM: 146 KCl, 2 NaCl, 1 MgCl₂, 1 CaCl₂, 1 GdCl₃, 10 HEPES/KOH (pH 7.5). GdCl₃ completely inhibited the stretch-activated channels. The bath solution contained, in mM: 146 KCl, 2 MgCl₂, 1 EGTA, 10 HEPES/KOH (pH 7.5). To obtain single channel recordings, oocytes were injected with low doses of RNA of GIRK1 (10-100 pg/oocyte), and RNA of GIRK4 was half of that (5-50 pg); the amount of GIRK1/ 4_{WT} RNA was 10-20/5-10 pg, respectively. The amounts of the GIRK1/4_{R52H} and GIRK1/4_{E246K} channels' RNA were 20-40/10-20 pg and 40-100/20-50 pg, respectively. In addition, 25 ng of the antisense DNA oligonucleotide against the endogenous GIRK5 channel was injected together with the RNAs (49). Single channel current (isinale) was calculated from all-point histograms of the original records (55), and open probability (Po) was obtained from event lists generated using idealization procedure based on 50% crossing criterion (56). Number of channels was estimated from overlaps of openings during the whole time of recording (at least 5 min). Po was calculated only from records that contained up to 3 channels. Thus, the probability of missing a channel was negligible. For channels activated by co-expressed Gβγ there was no decrease in Po over >4 minutes, and the Po was averaged from the first 4 minutes of the record. Single-channel conductance (g) was measured from I-V relationships, constructed from values of isingle measured at potentials ranging from -120 mV to -40 mV at 20 mV increments.

Confocal imaging

Confocal imaging of the oocytes and analysis were performed as described (29), with a Zeiss 510 META confocal microscope, using a 20× objective. In whole oocytes, the image was focused on an oocyte's animal hemisphere, at the equator. Images were acquired using spectral (λ)-mode. YFP was excited with the 514 nm line of the argon laser and sampled at 534–546 nm. Fluorescence signals were averaged from three regions of interest using Zeiss LSM Image Browser, background and the average signal from uninjected oocytes were subtracted.

Förster resonance energy transfer (FRET).

FRET experiments were performed using sensitized emission spectral method (28) as described previously (29) (see also Supplemental Fig. S3). Two spectra were collected from the animal hemisphere of each oocyte, with 405nm (CFP excitation) and 514nm (YFP excitation) laser lines. Oocytes with low signal of one of the fluorophores (usually <100 AU), or with coefficient of variation of A parameter at 524, 535,545,566 nm of more than 0.25, were excluded from analysis. Net FRET signal was calculated in the YFP emission range (with the 405 nm excitation) by consecutive subtraction of a scaled CFP-only spectrum (giving the A ratio parameter) and then of the ratio A0, which reports the direct excitation of YFP by the 405 nm laser, as in Equations 1 and 2:

(Eq. 1) Ratio
$$A = \frac{F_{405}}{F_{514}} = \frac{F_{direct}}{F_{514}} + \frac{F_{FRET}}{F_{514}}$$

$$(Eq. 2) \quad (A - A_0) = \frac{F_{FRET}}{F_{514}}$$

Because of the use of different experimental settings in independent experiments, the fluorescence intensities or A ratios cannot be used to construct FRET titration curves. We imaged a double-labeled protein, expressing both CFP and YFP at a 1:1 stoichiometry (YFP-GIRK2-CFP, DL-GIRK2) in each experiment to convert the fluorescence of CFP and YFP into their molar ratio (the average donor/acceptor ratio, in arbitrary units (AU) corresponds to a molar ratio of one) (29). The molar ratio collected and from many experiments can then be implemented into one titration curve (29).

The apparent FRET efficiency in an individual cell, E_{app}, was calculated as in Equation 3:

(3)
$$E_{app} = \left(\frac{RatioA}{RatioA_0} - 1\right) \frac{\epsilon_A}{\epsilon_D}$$

where ϵ_D and ϵ_A are molar extinction coefficients for the donor and acceptor, respectively, at the donor excitation wavelength (29, 30).

Data analysis

I-V curves were analyzed using scripts written in-house and from Mathworks forum for analyzing .abf files produced by Clampex (Molecular Devices), MATLAB R2015a (MathWorks, USA), and with Clampfit 10.7 software (Molecular Devices). The reversal potential (V_{rev}) was determined from the intercept of the net I-V curve with voltage axis. The extent of inward rectification (F_{ir}) was determined by dividing the current at 50 mV positive to V_{rev} by the current at 50 mV negative to V_{rev} (see Fig. 2B) (48):

$$F_{ir} = \frac{I_{V_{rev} + 50}}{I_{V_{rev} - 50}}$$

Estimated reversal potential (V_{rev}) was calculated using the Nernst equation:

$$V_{rev} = \frac{RT}{zF} \ln(\frac{[X]_{out}}{[X]_{in}})$$

Permeability ratio of sodium and potassium (pNa⁺/pK⁺) was determined from Goldman–Hodgkin–Katz equation:

$$E_{rev} = \frac{RT}{zF} \log(\frac{p_{K^+}[K^+]_o + p_{Na^+}[Na^+]_o}{p_{K^+}[K^+]_i + p_{Na^+}[Na^+]_i})$$

Statistical analysis

Protein expression imaging data have been normalized as described previously (57). For analysis across several experiments, fluorescence intensity in each oocyte was normalized to the average signal in the oocytes of the control group of the same experiment. This procedure yields average normalized intensity as well statistical variability (e.g. SEM) in all treatment groups as well as in the control group. Statistical analysis was performed using SigmaPlot 11 (Systat Software, Inc.) and GraphPad Prism version 8 for Windows (GraphPad Software, La Jolla California USA). If the data passed the Shapiro–Wilk normality test and the equal variance test, two-group comparisons were performed using t-test, while multiple group comparison was performed using one-way ANOVA. If not, the Mann–Whitney rank sum test, or Kruskal-Wallis test were performed, respectively. The data in the bar graphs are presented as mean ± SEM. In box plots, boxes show 25th and 75th percentiles and whiskers show minimal and maximal values, black horizontal line – median, green line -mean.

Acknowledgments

We are grateful to Alomone Labs for the generous gift of VU0529331 and to Mariam Ashkar for assistance in some of the experiments. This work was supported by the joint Israel-India grant ISF # 2255/15 and the Israel Science Foundation grant # 1282/18.

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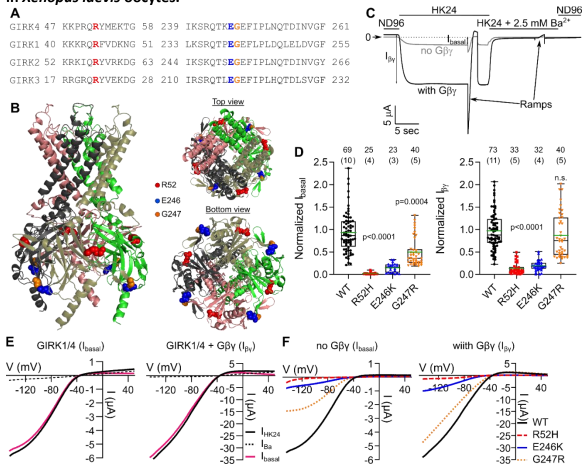
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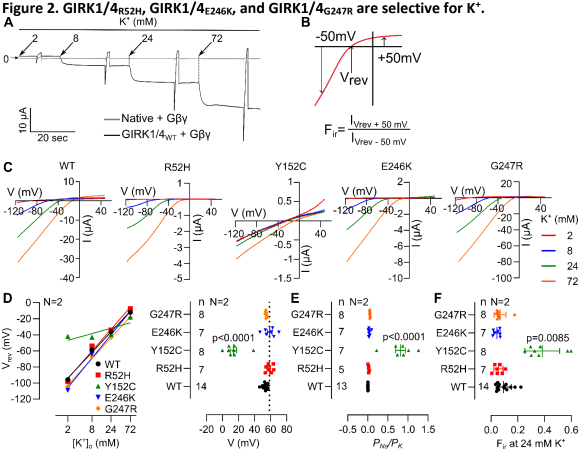
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Figures

Figure 1. Basal and G $\beta\gamma$ -induced activity of heterotetrameric GIRK1/4 $_{WT}$ and mutants in *Xenopus laevis* oocytes.



(A) Alignment of GIRK4 with GIRK1, GIRK2, and GIRK3 shows that R52, E246, and G247R are conserved amino acids. (B) Crystal structure of GIRK2 (23) marked with highlighted positions of the mutations; R52 – red, E246 – blue, and G247 – orange (numbering corresponds to GIRK4). (C) Example of the recording protocol in GIRK1/4_{WT} injected oocytes without (gray) or with (black) coinjected G $\beta\gamma$. Horizontal bars show the corresponding external solutions. Voltage ramps were applied in HK24 and HK24 + 2.5 mM Ba²⁺ solutions. (D) Normalized I_{basal} (left) and I $\beta\gamma$ (right). Currents were measured at V_m=-80 mV. (E) Representative I-V relationships of currents of GIRK1/4 (I_{basal}; left) and GIRK1/4 + G $\beta\gamma$ (I $\beta\gamma$; right). Net GIRK currents were obtained by subtraction of I-V curve in Ba²⁺ ("Ba²⁺-subtraction" procedure). (F) Representative net I-V relationships of GIRK1/4 and mutants; I_{basal} (left) and I $\beta\gamma$ (right).



(A) Examples of current records from oocyte expressing GIRK1/4_{WT} with G $\beta\gamma$ (black) and oocyte expressing only G $\beta\gamma$ (gray). (B) Example of F_{ir} calculation. (C) Exemplary I-V relationships of GIRK1/4_{WT} and mutants in 4 K⁺ concentrations. (D) Left panel, V_{rev} vs. $log_{10}[K^+]_o$ graph of GIRK1/4_{WT} and mutants in representative oocytes. Right panel, average of slopes from the V_{rev} vs. $log_{10}[K^+]_o$ graph, in mV/decade (tenfold change in $[K^+]_o$), dotted line corresponds to a slope of 58.5 mV. (E) P_{Na}/P_K of GIRK1/4_{WT} and mutants. (F) Factor of inward rectification (F_{ir}) of GIRK1/4_{WT} and mutants.

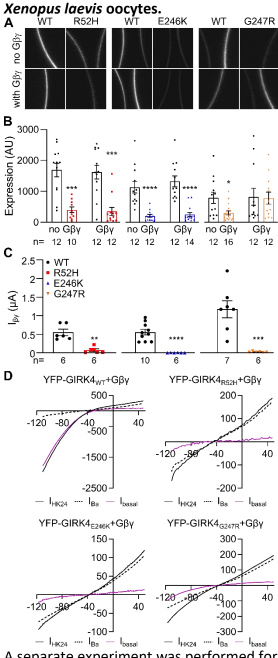


Figure 3. Membrane abundance of homotetrameric YFP-GIRK4 and mutants in

A separate experiment was performed for each mutant. (A) Representative confocal images of YFP-GIRK4 expressing oocytes, with or without G $\beta\gamma$. (B) YFP-GIRK4 expression in the PM; except for YFP-GIRK4 $_{G247R}$ +G $\beta\gamma$, all mutants express much less than YFP-GIRK4 $_{WT}$. (C) I $_{\beta\gamma}$ of YFP-GIRK4 homotetramers. All mutants were not active as homotetramers. (D) Representative I-V relationships of YFP-GIRK4 (WT and mutants) coexpressed with G $\beta\gamma$.

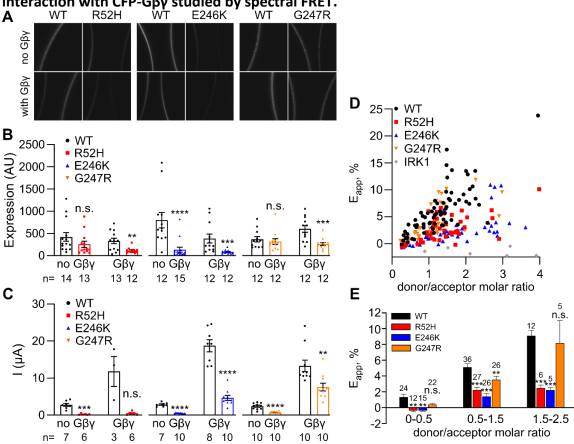


Figure 4. Membrane abundance of heterotetrameric GIRK1/YFP-GIRK4 and their interaction with CFP-Gβγ studied by spectral FRET.

(A) Representative confocal images of GIRK1/YFP-GIRK4 expressing oocytes, with or without G $\beta\gamma$. (B) GIRK1/YFP-GIRK4 expression in PM; mutated channels express less well than GIRK1/YFP-GIRK4_{WT}. A separate experiment was performed for each mutant. (C) Currents of GIRK1/YFP-GIRK4_{WT} and mutated channels. (D) Raw data from FRET experiments, showing E_{app} as a function of donor/acceptor molar ratio. Each symbol corresponds to E_{app} from one oocyte, from 4 experiments. (E) Binned analysis of E_{app} shows that, at all donor/acceptor molar ratios, the interaction of CFP-G $\beta\gamma$ with GIRK1/YFP-GIRK4_{E246K} was impaired, whereas the interaction of CFP-G $\beta\gamma$ with GIRK1/YFP-GIRK4_{G247R} was more similar to GIRK1/YFP-GIRK4_{WT}. Molar ratios >2.5 were not included in the analysis because of insufficient data points for GIRK1/YFP-GIRK4_{WT}.

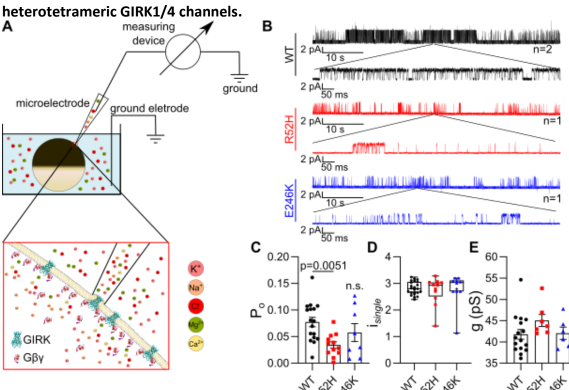


Figure 5. Cell attached patch clamp recording and single channel analysis of

(A) A simplified scheme of the cell attached patch clamp recording. (B) Representative recordings of GIRK1/4_{WT}, GIRK1/4_{R52H}, and GIRK1/4_{E246K} with a high dose of G $\beta\gamma$ (5 ng G β and 1 ng G γ RNA). (C) P_o of GIRK1/4_{R52H} is significantly reduced compared to GIRK1/4_{WT}. (D) i_{single} of mutants and (E) single channel conductance of mutants are similar to GIRK1/4_{WT}.

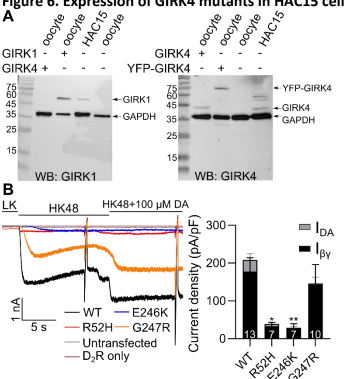
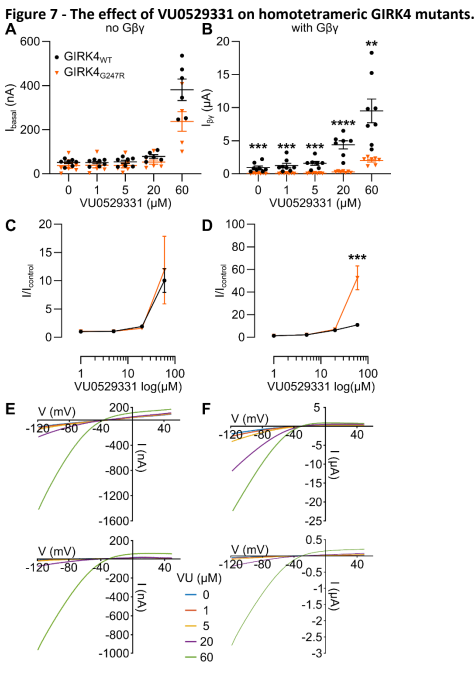


Figure 6. Expression of GIRK4 mutants in HAC15 cells.

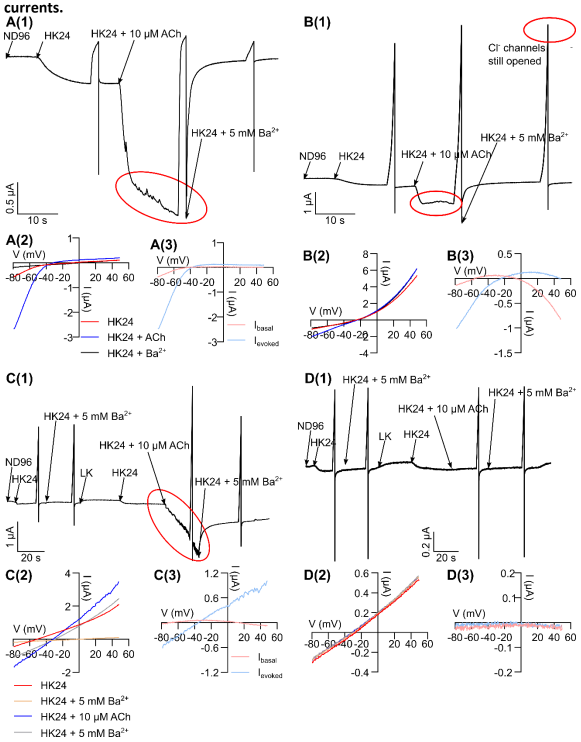
(A) Western blots of HAC15 cells with GAPDH antibody (both panels) and GIRK1 antibody (left) and GIRK4 antibody (right) show that HAC15 cells express GIRK1 and GIRK4 channels. Oocytes expressing GIRK1, GIRK4, and YFP-GIRK4 were used as positive controls, naïve oocytes as negative control, and GAPDH as loading control. (B) Examples (left) and summary (right) of currents in HAC15 cells expressing GFP-GIRK1/GIRK4_{WT}, GFP-GIRK1/GIRK4_{R52H}, GFP-GIRK1/GIRK4_{E246K}, and GFP-GIRK1/GIRK4_{G247R}. Channels were expressed with D_2 dopamine receptor and $G\beta\gamma$ ($I_{\beta\gamma}$), thus the small I_{DA} .



Oocytes expressed GIRK4 mutants and GIRK4_{WT}, with or without Gβγ. Cells were exposed to 4 concentrations of VU0529331 (1, 5, 20, and 60 μM). Only GIRK4_{WT} and GIRK4_{G247R} mutant responded to VU0529331. (A) I_{basal} of GIRK4_{WT} and GIRK4_{G247R}. GIRK4_{WT} and GIRK4_{G247R} react similarly to VU0529331. (B) $I_{βγ}$ of GIRK4_{WT} and GIRK4_{G247R}. GIRK4_{WT} had larger current at 0 μM VU0529931, while GIRK4_{G247R} did not show any response to concentrations lower than 20 μM VU0529331. (C) Normalized I_{basal} (I/ $I_{control}$). GIRK4_{WT} and GIRK4_{G247R} were activated similarly by VU0529331. (D) I/ $I_{control}$ of $I_{βγ}$ show that GIRK4_{G247R} was activated by VU0529331 stronger than GIRK4_{WT}, in relative terms. (E) I-V relationship of GIRK4_{WT} (top) and GIRK4_{G247R} (bottom) with Gβγ. (F) I-V relationship of GIRK4_{WT} (top) and GIRK4_{G247R} (bottom) with Gβγ.

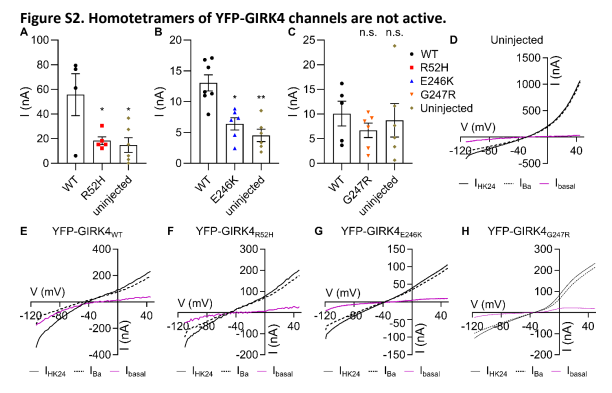
Supplementary figures

Figure S1. In some oocyte batches, application of ACh induces Ca²⁺-dependent Cl



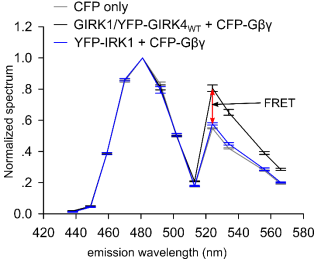
(A) ACh evokes Ca^{2+} -dependent Cl^- currents in oocytes expressing GIRK1/4 and M2R. (A1) Example of a recording from an oocyte coinjected with RNAs of GIRK1/4_{WT} and M₂R. Application of ACh evoked GIRK's I_{evoked} but also a Ca^{2+} -dependent Cl^- current

(marked with red circle). (A2) I-V relationships for I_{basal} and I_{evoked} before Ba²⁺ subtraction. (A3) Net I-V relationships of I_{basal} and I_{evoked} after Ba²⁺ subtraction. Cl⁻ currents were masked by the large currents of GIRK1/4_{WT}. (B) ACh-induced Cl⁻ currents significantly contribute to the overall small currents in an oocyte expressing GIRK1/4_{R52H} and m2R. (B1) The full current record. Application of ACh evoked a Ca²⁺ dependent Cl⁻ current (marked with red circle), which was not blocked by Ba²⁺ and decayed slowly, over minutes. Since the currents of GIRK1/4_{R52H} were small, the Cl⁻ current contributed a large part of the total current. (B2) I-V relationship before Ba²⁺ subtraction. (B3) I-V relationship of I_{basal} and I_{evoked} after Ba²⁺ subtraction. Because Cl⁻ currents (which are outwardly rectifying and thus larger at positive voltages) are still present in Ba2+ after ACh washout, the subtraction of I-V curve recorded in the presence of Ba²⁺ results in an I-V curve showing inward currents at positive voltages, which is an artifact. (C) ACh may evoke Cl⁻ currents in naïve (uninjected with RNA) oocytes. (C1) Example of a recording from an uninjected oocyte; here, Ba²⁺ was applied after HK24, then washed with ND96, then HK24 was applied again followed by ACh, and ACh was washed out with ND96 + 5 mM Ba^{2.}. This protocol was developed so we could get net I_{basal} and I_{evoked}, and yet, the artifacts remained. (C2) I-V relationship before Ba2+ subtraction. (C3) I-V relationship of Ba²⁺ subtracted I_{basal} and I_{evoked}. (D) BAPTA eliminated the Ca²⁺ dependent Cl⁻ current. (D1) Example of a recording from an uninjected oocyte ~1 hour after the injection of BAPTA. (D2) I-V relationship before subtraction; all I-V relationships were linear and almost identical in all solutions. (D3) I-V relationship of Ibasal and Ievoked after Ba²⁺ subtraction. There were no clearly detectable currents, demonstrating the absence of any GIRK-like currents.



(A-C) Currents in oocytes injected with RNAs of YFP-GIRK4 mutants, measured at -80 mV, were indistinguishable from currents recorded in uninjected oocytes. (D-H) Representative I-V relationships of uninjected oocytes (D) and homotetrameric YFP-GIRK4 $_{WT}$ (E), YFP-GIRK4 $_{R52H}$ (F), YFP-GIRK4 $_{E246K}$ (G), YFP-GIRK4 $_{G247R}$ (H).

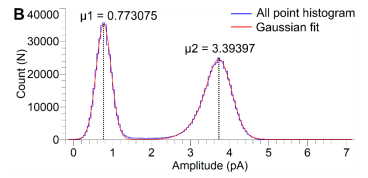
Figure S3. Emission spectra used to measure FRET.



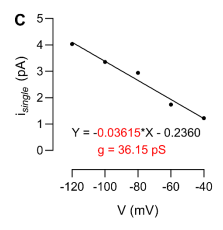
The graph shows emission spectra of GIRK1/YFP-GIRK4 $_{WT}$ + CFP-G $\beta\gamma$ (black line), IRK1-YFP + CFP-G $\beta\gamma$ (light gray line) used as negative control, and CFP-G $\beta\gamma$ only (dark gray line), when the donor CFP was excited with the 405 nm laser. Values on X axis are midpoints of 11 nm-wide sampling windows in the spectral mode of Zeiss 501 Meta confocal microscope. GIRK1/YFP-GIRK4 $_{WT}$ + CFP-G $\beta\gamma$ show large emission increase in YFP emission range (520 –575 nm) compared to the negative control and CFP only groups, demonstrating the energy transfer between the donor and acceptor molecules. Experimental points shown are mean±SEM from a representative experiment.

Figure S4. Estimation of i_{single} and single channel conductance from GIRK1/4 single channel recordings.



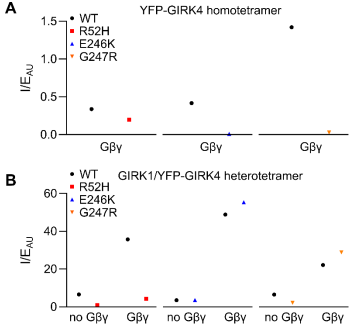


 $i_{single} = 3.39397 - 0.773075 = 2.620895$



(A) Example of a segment of the record that was used for the all point histogram in B. (B) All point histogram fitted with two Gaussians. The left peak is the background noise where the channel was closed, the right peak corresponds to the open state of the channel. Subtraction of the baseline (μ 1) from the open state (μ 2) gives i_{single}. (C) Single channel conductance was derived from the slope of I-V relationship graph; i_{single} was calculated per each voltage in increments of 20 mV.

Figure S5. Analysis of whole-cell currents normalized to expression, for YFP-GIRK4 homotetramers and GIRK1/YFP-GIRK4 heterotetramers.



(A) Normalization of current to expression of YFP-GIRK4 homotetramers indicates an impaired gating of the mutated channels. The normalization procedure included the division of mean value of current on mean value of channel expression in each experiment, giving one value for each experiment. Therefore, no statistical analysis could be performed. (B) Normalization of current to expression of GIRK1/YFP-GIRK4 heterotetramers indicates an impaired expression and gating of GIRK1/YFP-GIRK4_{R52H} heterotetramers; on the other hand, E246K and G247R mutations appear to have the same open probability suggesting unimpaired gating.