Introduction

G-protein-linked inwardly rectifying potassium channels (GIRKs) mediate important physiological functions in the heart and affect excitability in various neuronal populations (Dascal et al., 1993; Kofuji et al., 1995; Krapivinsky et al., 1995b; Lesage et al., 1995). Thus far, the family consists of four mammalian-cloned members GIRK1–GIRK4 (or Kir3.1–Kir3.4) (Kubo et al., 1993a; Lesage et al., 1994; Krapivinsky et al., 1995b; Dascal, 1997; Jan and Jan, 1997). In cardiac tissue, the GIRK conductance is attributable to activation of hetero-oligomeric channels composed of GIRK1 (G1) and GIRK4 (G4), although homomeric GIRK4 channels may also be functionally significant (Krapivinsky et al., 1995b; Corey and Clapham, 1998). In the CNS, various combinations of subunits are thought to comprise channels found in several brain regions (Kobayashi et al., 1995; Karschin et al., 1996; Bhattacharya et al., 1998; Dascal et al., 2000; Ivanina et al., 2003). In addition to G-protein-coupled receptor (GPCR) systems, including muscarinic, opiate, dopaminergic, adenosine, and serotonin (for review see, Dascal, 1997). After GPCR activation, direct binding of Gβγ subunits to the channel results in activation (Logothetis et al., 1987; Reuveny et al., 1994). Extensive efforts have been made to determine the precise binding site for Gβγ (Logothetis et al., 1987; Huang et al., 1995, 1997; Inanobe et al., 1995, 1997; Inanobe et al., 1995a, 1998; He et al., 1999, 2002; Ivanina et al., 2003; Finley et al., 2004). A consensus identifying the exact Gβγ binding domain has not been determined but is thought to comprise contributions from both the intracellular N termini (NT) and C termini (Huang et al., 1997). Studies of Gβγ binding to GIRK1 and GIRK2 glutathione S-transferase (GST) fusion proteins demonstrated that not all Gβγ binding sites were identical between the two subunits (Ivanina et al., 2003). In addition to Gβγ binding, membrane phospholipids, such as phosphatidylinositol 4,5-bisphosphate, and Na⁺ are thought to participate in GIRK channel activation (Sui et al., 1998; Ho and Murrell-Lagnado, 1999).

Within the GIRK subunit intracellular domains, several motifs have been identified that specify cellular trafficking (Tucker et al., 1996; Woodward et al., 1997; Kennedy et al., 1999; Ma et al., 2002; Mirshahi and Logothetis, 2004).

GIRK channels are coupled to a wide range of G-protein-coupled receptor (GPCR) systems, including muscarinic, opiate, dopaminergic, adenosine, and serotonin (for review see, Dascal, 1997). After GPCR activation, direct binding of Gβγ subunits to the channel results in activation (Logothetis et al., 1987; Reuveny et al., 1994). Extensive efforts have been made to determine the precise binding site for Gβγ (Logothetis et al., 1987; Huang et al., 1995, 1997; Inanobe et al., 1995, 1997; Inanobe et al., 1995a, 1998; He et al., 1999, 2002; Ivanina et al., 2003; Finley et al., 2004). A consensus identifying the exact Gβγ binding domain has not been determined but is thought to comprise contributions from both the intracellular N termini (NT) and C termini (Huang et al., 1997). Studies of Gβγ binding to GIRK1 and GIRK2 glutathione S-transferase (GST) fusion proteins demonstrated that not all Gβγ binding sites were identical between the two subunits (Ivanina et al., 2003). In addition to Gβγ binding, membrane phospholipids, such as phosphatidylinositol 4,5-bisphosphate, and Na⁺ are thought to participate in GIRK channel activation (Sui et al., 1998; Ho and Murrell-Lagnado, 1999).

In this study, we examined specific GIRK1 subunit characteristics. Based on GIRK1 regions identified by previous studies as important sites in Gβγ binding and subunit interaction, we investigated a specific region of the GIRK1 C terminus, which after
deletion or point mutation, prevents functional channel expression when coexpressed with wild-type GIRK4. Recent crystallographic structural analysis of the GIRK1 intracellular domains also indicates that this region of the GIRK1 C terminus may contact adjacent termini and be important in subunit interaction (Nishida and MacKinnon, 2002). The KirBac1.1 potassium channel crystal structure also depicts an interacting region between the intracellular termini (Kuo et al., 2003). In this study, we identified GIRK1 residues that contribute to association and channel function that may be generally important in the translation of the gating mechanism.

Materials and Methods

Molecular biology. GIRK1 was cloned from a rat insulinoma tumor cell (RIN library and a predicted amino acid sequence identical to the cardiac clone originally described (Kubo et al., 1993b)); GIRK4 and GIRK5 were generous gifts from Dr. D. E. Clapham (Children’s Hospital, Boston, MA), and Kir2.1 was a generous gift from Dr. J. Kyle (The University of Chicago, IL). The m2 muscarinic receptor (m2R) (Clontech, Palo Alto, CA) in the pGEMZ vector was subcloned into the pcDNA3.1Zeo vector for mammalian cell expression, both with and without GFP (green fluorescent protein) fused to the GIRK1 or GIRK4 C terminus. Kir2.1 was subcloned into the pcDNA3.1Zeo + vector (Invitrogen, San Diego, CA). A chimera of GIRK1 was prepared by PCR gene fusion technique in which the 32 amino acid C-peptide region of human proinsulin was transplanted to the 3’ end of the GIRK1 clone (Philipson et al., 1995), to produce an epitope-tagged channel. All GIRK constructs were subcloned into either the pMXT vector, obtained from Dr. P. Kofuji (University of Minnesota, Minneapolis, MN), for oocyte expression or the pEGFPN3 vector (Clontech) for mammalian cell expression. All point mutations for both the oocytes and mammalian constructs were introduced by site-directed mutagenesis (QuikChange; Stratagene, La Jolla, CA). The FLAG epitope was introduced into the GIRK1 subunit between residues 114 and 115 as described previously (Kennedy et al., 1999) using specific primers to introduce the epitope (DYKDDDDK) and also to remove the glycosylation site N119D. The GIRK1Δ330–384 construct was made by introducing silent restriction sites that were subsequently digested, dephosphorylated, and ligated. All point-mutant GIRK1-FLAG constructs were made by site-directed mutagenesis (QuikChange; Stratagene) of the GIRK1-FLAG construct. The N terminus of GIRK1 (residues 1–83) fused with a C-terminal myc tag (EQKLISEEDL), the N terminus of Kir2.1 (residues 1–84) fused with a C-terminal myc tag, the N terminus of GIRK4 (residues 1–89) fused with a C-terminal FLAG tag, and the C terminus of GIRK4 (residues 189–419) fused with a C-terminal hemagglutinin (HA) tag (YPYDVPDYA) were individually subcloned into the pcDNA3.1Zeo + vector. The GIRK1 C terminus [transmembrane 2 (TM2) plus C terminus, residues 158–501], the GIRK1 C terminus Δ330–384, and the Kir2.1 C terminus (TM2 plus C terminus, residues 157–427) were subcloned into a pPEFY (enhanced yellow fluorescent protein) vector (Clontech) generously provided by Dr. G. Thinakaran (The University of Chicago, Chicago, IL) containing a secretion signal, followed by YFP. cDNAs encoding the indicated C termini of Kir2.1 (residues 182–427) and GIRK1 (residues 183–501 plus C-peptide) were generated by PCR and cloned into a carrier plasmid with the GST coding sequence in pGEX-4T (Amersham Biosciences, Arlington Heights, IL). βARK (β-adrenergic receptor kinase) was generously provided by Dr. R. J. Leffkowitz (Duke University, Durham, NC). All point mutants were made by site-directed mutagenesis (QuikChange; Stratagene) of the respective constructs. Integrated DNA Technologies (Corvalle, VA) synthesized all primers, and all constructs were sequenced at the University of Chicago Cancer Research Sequencing facility.

For oocyte experiments, the m2R construct was linearized with HindIII, and cRNA was transcribed using the T7 polymerase mMessage mMachne kit (Ambion, Austin, TX). All GIRK constructs were linearized with SauI, and cRNA was transcribed using T3 polymerase mMessage mMachne kit (Ambion). The cRNA concentration was quantified by UV absorption at 260 nm (A260), and the quality of RNA was examined by agarose gels.

COS-7 cell culture and cDNA transfection. COS-7 cells (American Type Culture Collection, Manassas, VA) were plated in 35 mm dishes, 25 mm coverslips, or 100 mm dishes and grown in DMEM supplemented with 10% fetal bovine serum, 100 U/ml penicillin, and 100 μg/ml streptomycin. One day after plating, cells were transfected using Lipofectamine Plus (Invitrogen) with 0.5 μg of m2R cDNA and 1 μg of cDNA for each GIRK subunit used in electrophysiological and immunostaining experiments, and twice the amount of cDNA for protein expression experiments. In electrophysiological experiments in which EGFP-tagged subunits were not used, pEGFP vector cDNA was cotransfected at a 1:10 ratio for positive identification of transfected cells. For Kir2.1 experiments, 1 μg of wild-type or Kir2.1 L330R cDNA was coexpressed with 100 ng of pEGFP vector cDNA for positive identification of transfected cells. For communoprecipitation experiments, cells were transfected with 2 μg of cDNA of each GIRK construct as indicated.

Electrophysiological recording from oocytes. Oocytes were removed from Xenopus laevis as described previously (Yoshii and Kurihara, 1989) and maintained at 18°C in OR-2 solution that was changed once daily. Oocytes were injected with 50 nl containing equal amounts of each single subunit cRNA (∼5 ng) and m2 receptor (∼2.5 ng). The OR-2 solution contained 90 mM NaCl, 1 mM MgCl2, 1 mM CaCl2, 2.5 mM KCl, and 5 mM HEPES, supplemented with 100 μg/ml gentamycin and 5 mM pyruvate, pH 7.6. Two-microelectrode voltage-clamp recordings were performed 2–3 d after injection at room temperature (RT) using a TURBO TEC-10 amplifier (NI Electronics, Tamm, Germany), an ITC-16 interface (InstruTech, Great Neck, NY), and an IBM-compatible personal computer. Microelectrodes were filled with 3 M KCl and had resistances of 0.5–2 MΩ. Oocytes were continuously superfused with a bath solution of 90 mM NaCl or KCl, 1 mM MgCl2, and 5 mM HEPES, pH 7.6, with NaOH/KOH. G-protein–dependent currents were induced with the addition of 5 μM carbachol (Carb) (Sigma, St. Louis, MO) to the bath solution. In all experiments, the holding potential was −80 mV; test potentials were delivered once every second and stepped between −150 and 60 mV in 30 mV intervals. Data collection and analysis were performed using Pulse/Pulse Fit (Heka, Lambrecht, Germany), and data were plotted using the integrated graphics package IGOR (WaveMetrics, Lake Oswego, OR) or Microsoft (Seattle, WA) Excel. Data are presented as mean ± SEM and are taken from one or several batches of oocytes. Significance of results was performed by the Kruskal–Wallis one-way ANOVA method and Dunn’s test. All experiments were conducted at room temperature.

Electrophysiological recording from COS-7 cells. Whole-cell recordings were performed at room temperature 48–72 h after transfection in an initial bath solution consisting of the following (in mM): 140 NaCl, 5.4 KCl, 2 CaCl2, 1 MgCl2, and 10 HEPES, pH 7.4. After currents in low-K+ solutions were recorded, cells were superfused with a high-K+ (HK) solution containing the following (in mM): 5.4 NaCl, 140 KCl, 2 CaCl2, 1 MgCl2, and 10 HEPES, pH 7.4. The pipette recording solution contained the following (in mM): 120 KCl, 2 CaCl2, 1 MgCl2, 11 EGTA, 33 KOH, 10 HEPES, 1 NaGTP, and 2 MgATP, pH 7.2. In some experiments, CaCl2 and EGTA were replaced with 10 mM BAPTA-potassium salt in the pipette, and CaCl2 was removed from the extracellular solutions with no difference in function. G-protein–dependent currents were induced with the addition of 10 μM carbachol (Sigma) to the bath solution. K+ currents were blocked with 500 μM BaCl2 added to the high-K+ carbachol solution. In all experiments, the holding potential was −80 mV; test potentials were delivered in steps between −160 and 50 mV in 30 mV intervals. Whole-cell currents were recorded with an EPC-9 (InstruTech) patch-clamp amplifier at 2 kHz and low-pass filtered at 1 kHz. Stimulation and data acquisition were controlled by the PULSE software package, and data analysis was performed with IGOR software (WaveMetrics). Data are presented as means ± SEM. Significance of results was determined using Student’s t test.
Protein expression/Western blotting. COS-7 cells were grown in 100 mm dishes and transfected with 2 μg of each subunit as indicated with Lipofectamine Plus (Invitrogen). Cells were washed and lysed with buffer containing 50 mM Tris, pH 7.4, 150 mM NaCl, 1% NP-40, 1 mM EDTA, 1 mM PMSF, and 1 μg/ml each aprotinin, leupeptin, and pepstatin for 45 min at 4°C; 48 h after transfection. Cells were centrifuged at 14,000 × g for 20 min at 4°C. Supernatant was removed and measured for protein concentration (BCA reagent; Pierce, Rockford, IL), and samples were solubilized with sample buffer containing 10 mM DTT and boiled for 5 min. Samples were resolved by SDS-PAGE, subsequently transferred to polyvinylidene difluoride membrane, and blocked with 5% milk for 30 min at room temperature. Blots were probed with rat α-C-peptide at 1:100 (generously provided by Dr. L. Phillipson, The University of Chicago, Chicago, IL), followed by goat HRP-α-rat (KPL, Gaithersburg, MD) or α-GIRK1 (Alomone Labs, Jerusalem, Israel) at 1:500, followed by goat HRP-α-rabbit (Amersham Biosciences).

GST fusion proteins and βγ binding assay. GST fusion proteins were expressed in BL21, grown to an OD_{600} of 0.6, induced with 1 mM IPTG, and grown for 3 h. Bacteria were pelleted and resuspended in 13 mM of lysis buffer [50 mM Tris-HCl, pH 8.2, 150 mM NaCl, 0.5% Triton X-100, protease inhibitors (1 mM PMSF, 1 mM EDTA) and protease inhibitors (1 mM PMSF, 2 mg/ml chymostatin, 2 mg/ml pepstatin, 2 mg/ml antipain, and 4 mg/ml leupeptin)]. Suspensions were sonicated four times for 15 s. Triton X-100 was added to a final concentration of 1%. Suspension was rotated at RT for 15 min and then centrifuged for 20 min at 20,000 × g at 4°C. Supernatant was aliquoted into 1 ml aliquots and frozen. Quantification of the lysates was performed by adding variable volumes of lysate to 20 μl of 50% slurry of glutathione agarose (Sigma), adjusting the total volume to 1 ml with lysis buffer (plus Triton X-100), and binding by rocking at RT for 15 min. The agarose was washed twice by centrifuging for 10 s, aspirating the supernatant off, and resuspending in 1 ml of GST wash buffer (50 mM Tris-HCl, pH 8.2, 150 mM NaCl, 2 mM EDTA, 1% Triton X-100, and 1 mM PMSF). Fusion protein elution was by adding of 15 μl of protein loading buffer (5% SDS, 5% β-mercaptoethanol, and 5% glycerol) and boiled for 2 min. Samples were resolved by SDS-PAGE and Coomassie blue stained. BSA of known concentration was loaded onto the same gel to determine quantity of fusion protein bound to the agarose. Lysate volume yielding 500 ng of fusion protein was bound to 20 μl of glutathione agarose for each fusion protein as described above. After the wash, the affinity resin was equilibrated in βγ-binding buffer (50 mM Tris-HCl, 150 mM NaCl, and 0.01% polyoxyethylene 9 lauryl ether) (Sigma). After equilibration, the agarose was resuspended in 20 μl of βγ binding buffer [final volume of 40 μl, final concentration of fusion protein of 200 nM (molecular weight of fusion protein 64 kDa), and 1 μl of purified bovine brain βγ (40 μg/ml; Signal Transduction, San Diego, CA)] was added to the affinity agarose to give a final concentration of 40 μM βγ. The sample was mixed every 3 min for 30 min on ice. The fusion protein agarose was then washed four times in 1 ml of βγ binding buffer, resuspended in 15 μl of protein load buffer, boiled for 2 min, and resolved by SDS-PAGE. The primary antibody used for detection was α-Gβ (Santa Cruz Biotechnology, Santa Cruz, CA) at 1:250. After exposure, the blot was stripped and reprobed with α-GST (Santa Cruz Biotechnology) to ensure equal loading of fusion protein transfer.

Coimmunoprecipitation experiments. COS-7 cells were grown in 100 mm dishes and transfected with 2 μg of each subunit indicated with Lipofectamine Plus (Invitrogen). Cells were washed three times with PBS (Sigma) and solubilized with 500 μl of lysis buffer (100 mM NaCl, 0.5% (w/v) Igepal CA-350, 1 mM EDTA, 1 mM PMSF, and 1 μg/ml each aprotinin, leupeptin, and pepstatin) 48 h after transfection for 50 min rotating at 4°C. Samples were centrifuged at 14,000 × g for 20 min at 4°C. The supernatant was removed and measured for protein concentration, and an aliquot was removed (lysate). The supernatant was precleared with 40 μl of Protein G-Plus Agarose (Santa Cruz Biotechnology) or 40 μl of Protein A-agarose (Sigma) for 20 min rotating at 4°C, and subsequently the agarose was removed by centrifugation. The supernatant was incubated with antibody [rabbit α-Kir2.1 NT (generously provided by Dr. C. A. Vandenbark, University of California, Santa Barbara, CA)], mouse c-myc (9E10) antibody (Santa Cruz Biotechnology), mouse FLAG (M2) antibody (Sigma), or rabbit HA antibody (Clontech) for 2 h at 4°C. Protein G-Plus Agarose or Protein A-Agarose was added, and samples were rotated overnight at 4°C. Samples were centrifuged for 5 min at 4°C to isolate agarose/antibody/protein complex. Immunoprecipitated samples were washed with PBS (containing 1 mM EDTA, 1 mM PMSF, and 1 μg/ml each aprotinin, leupeptin, and pepstatin), boiled in sample buffer (100 mM Tris-HCl, pH 6.8, 4% SDS, 0.2% bromophenol blue, and 20% glycerol), and resolved by SDS-PAGE. Mouse α-GFP (Living Colors Av monoclonal antibody, JL-8; Clontech) was used to detect the tagged GIRK1 C-terminal constructs or the tagged Kir2.1 C-terminal constructs. Rabbit HA antibody (Clontech) was used to detect the tagged GIRK4 C-terminal constructs.

Immunostaining. Cells were grown on 25 mm coverslips and transiently transfected with 1 μg of each subunit indicated with Lipofectamine Plus (Invitrogen). Cells were washed with PBS and fixed with 4% paraformaldehyde for 10 min at room temperature 48 h after transfection. When needed, cells were permeabilized with 0.1% Triton X-100 (in PBS) for 3 min at room temperature. Cells were blocked with 0.5% bovine serum albumin (Sigma) and 2% goat serum (Sigma) in 1× PBS for 30 min at room temperature. Cells were incubated with either rat α-C-peptide at a 1:100 dilution or mouse α-FLAG (Sigma) at a 1:1000 dilution for 1 h at room temperature, washed with 1× PBS for 30 min, and incubated with FITC-conjugated goat α-rat IgG (gift from Dr. J. Berkhart, The University of Chicago), Alexa Fluor 488-conjugated goat α-mouse IgG (Molecular Probes, Eugene, OR), or rhodamine-conjugated goat α-mouse IgG (Molecular Probes) for 1 h at room temperature. Cells without exposure to the primary antibody or nontransfected cells were used as control. The coverslips were mounted on a slide and observed using an Olympus Optical (Melville, NY) Fluoview 200 laser-scanning confocal microscope. Images were analyzed with NIH Image J software and MetaMorph (Universal Imaging Corporation, Downingtown, PA). Compiled confocal slices of the entire cell surface intensity was averaged per cell area (square micrometers) and used for quantification of surface protein expression.

Results

C-Terminal deletion and mutant expression in oocytes

To determine the regions critical for channel function, we examined the GIRK1 C terminus to identify possible domains involved in protein–protein interactions. The GIRK1 subunit consists of 501 amino acids and contains intracellular N and C termini (Dascal, 1997). Sequence alignment of all mammalian GIRK subunits (GIRK1–GIRK4) and the Kir2.1 subunit indicates a high degree of homology throughout the transmembrane regions and proximal C termini. Hydrophobic heptad repeats have been shown to form amphipathic structures that may interact with other proteins (Lupas et al., 1991; Lupas, 1996). The GIRK1 C terminus was found to contain several regions of hydrophobic residue repeats potentially forming interaction domains. Initial experiments were performed using two-electrode voltage clamp of oocytes injected with cRNA of the corresponding subunits indicated and a G-protein-coupled receptor, the muscarinic m2 receptor, which has been shown to couple to GIRK activation in the heart as well as heterologous expression in cell culture (Dascal et al., 1993; Kofuji et al., 1995). Expression of the GIRK1 subunit in oocytes results in the formation of a functional heterotetrameric channel consisting of the GIRK1 subunit and the endogenous GIRK5 oocyte subunit (Hedin et al., 1996). Using this strategy, we found that deletion of the C-terminal region (residues 330–384) of GIRK1 resulted in nonfunctional channel expression (data not shown) (Fig. 1A, sequence alignment). GIRK1 subunits expressed alone do not form functional channels in mammalian expression systems; consequently, GIRK1 subunits were tested by coexpression of the GIRK4 subunit. Coexpression
of GIRK1Δ330–384 with wild-type GIRK4 resulted in a complete loss of channel function compared with GIRK1 plus GIRK4, as summarized in Figure 1B (peak HK current and HK plus Carb-induced current at $-150\,\text{mV} \pm 0.72\,\text{SEM}$ for G1 + G4, $-0.44 \pm 0.07\,\mu\text{A}$, respectively, and for G1Δ330–384 + G4, $-0.03 \pm 0.16\,\mu\text{A}$, respectively). To determine whether the proposed GIRK1 region represents an important functional domain (Fig. 1A), amino acids within the first seven residues were mutated to disrupt the hydrophobic nature of the proposed domain. Point mutations were introduced and assayed by two-microelectrode voltage clamp after coinjection into oocytes with cRNA of the wild-type GIRK4 subunit and m2R. Mutations of GIRK1 V330 did not alter GIRK channel function (data not shown). However, mutations of isoleucine 331 to a charged residue led to a similar loss of function (Fig. 1B; peak HK and HK + Carb current at $-150\,\text{mV} \pm 0.72\,\mu\text{A}$, respectively, for G1Δ330–384 + G4, $-0.12 \pm 0.02\,\mu\text{A}$, and for G1 I331R + G4, $-0.12 \pm 0.05\,\mu\text{A}$, respectively). Mutation of the G1 I331 to a hydrophobic residue such as methionine or leucine did not significantly disrupt GIRK channel function (Fig. 1B; peak HK and HK + Carb current at $-150\,\text{mV} \pm 0.72\,\mu\text{A}$, respectively, for G1 I331M + G4, $-0.97 \pm 0.35\,\mu\text{A}$, and for G1 I331M + G4, $-5.07 \pm 1.42\,\mu\text{A}$).

### GIRK expression in COS-7 cells

To avoid the GIRK5 subunit and to investigate GIRK channel expression in a mammalian system, GIRK subunits and m2R were expressed by transient expression in COS-7 cells. Coexpression of G1Δ, G4, and m2R resulted in a loss of channel function, consistent with results obtained in oocytes (Fig. 2A, B). Peak HK and HK + Carb current density for G1 + G4 was $-20.81 \pm 6.70\,\text{pA} / \text{pF}$ and $-42.47 \pm 8.66\,\text{pA} / \text{pF}$ compared with $-4.07 \pm 2.30$ and $-5.25 \pm 2.27\,\text{pA} / \text{pF}$ for G1Δ + G4 (Fig. 2B). Mutational analysis in oocyte experiments indicated that the isoleucine point mutation to a positively charged residue (G1 I331R) lead to a significant decrease in channel function similar to the deletion of the entire 330–384 region. Therefore, we focused on the effects of this particular mutation when expressed in COS-7 cells. Coexpression of G1 I331R and G4 also resulted in a decrease in functional channel expression (Fig. 2A, B; peak HK and HK + Carb current density for G1 I331R + G4, $-10.59 \pm 3.39$ and $-15.41 \pm 4.26\,\text{pA} / \text{pF}$). To determine whether channel function was altered because of a global decrease in expression of the protein, Western blots of COS-7 whole-cell lysates transfected with subunits were performed to determine subunit expression. The G1Δ and G1 I331R expressed to a similar degree as the wild-type G1 (Fig. 2C). Subunit expression and localization was also investigated by immunostaining of subunits in permeabilized COS-7 cells. Consistent with biochemical experiments, all subunits (including G4) were expressed (Figs. 2D, 3); however, the distribution of G1Δ and G1 I331R appeared qualitatively slightly more perinuclear, indicating a possible alteration in subunit trafficking.

### Trafficking of GIRK subunits

GIRK subunits have been shown to assemble as tetramers and localize to the membrane as a functional oligomer (for review, see Ma and Jan, 2002). Interaction between subunits also appears critical for proper expression of channels at the cell surface, as well as specific trafficking dependent on the subunits expressed (Kennedy et al., 1999; Ma et al., 2002; Mirshahi and Logothetis, 2004). To determine whether the mutation of the G1 isoleucine resulted in a possible disruption of subunit localization to the membrane, a FLAG epitope was inserted into the extracellular region of the GIRK1 subunit P-loop. G1-FLAG coexpressed with the G4 subunit and m2 receptor in COS-7 cells did not alter the level of protein expression or channel function (data not shown). Surface expression of the G1-FLAG subunit and point-mutant subunit (G1 I331R-FLAG) when expressed alone and when coexpression with G4 was analyzed using COS-7 cells. Permeabilized COS-7 cells exhibited a diffuse distribution of G1-FLAG when expressed alone and colocalization when expressed with G4-EGFP. Similar distribution was seen with expression of G1 I331R-FLAG alone and when coexpressed with G4-EGFP (Fig. 3, top row). We monitored surface expression of the epitope-tagged subunits in nonpermeabilized COS-7 cells transfected with the indicated subunits. When expressed alone, neither the G1-FLAG nor the G1 I331R-FLAG were positive for surface expression, consistent with results obtained by Kennedy et al. (1999) that demonstrated a lack of GIRK1 subunit surface expression when expressed alone. However, as a result of coexpression of the wild-type G4, both the G1-FLAG and G1 I331R-FLAG exhibited a punctate surface staining (Fig. 3, bottom row). Quantification of total surface intensity for the G1-FLAG was not different from the
G1 I331R-FLAG (data not shown). Trafficking of the point mutation (G1 I331R-FLAG) to the plasma membrane was unaltered.

**Homologous point mutation in GIRK4 and non-G-protein-dependent inward rectifier**

In an effort to demonstrate whether the hydrophobic residue (I331) was of unique importance for a functional G1 subunit or whether it had universal functional significance among GIRK subunits, a homologous point mutation in the G4 subunit, L337R, was made (Fig. 1). The homologous point mutation in the GIRK4 subunit (L337R) also resulted in a decrease in GIRK channel function when coexpressed with the wild-type G1 subunit and m2R (Fig. 4A) (peak HK and HK + Carb current density for G1 + G4, −20.81 ± 6.70 and −42.47 ± 8.66 pA/pF, and for G1 + G4 L337R, −4.98 ± 1.29 and −5.93 ± 1.53 pA/pF). To determine whether the decrease in channel function was dependent on Gβγ binding, the homologous hydrophobic residue in the non-G-protein-dependent inward rectifier Kir2.1 was mutated to a positively charged arginine (Fig. 1). Kir2.1 subunits form homomeric tetrayers and are constitutively active when expressed in heterologous expression systems. Expression of the Kir2.1 L337R resulted in a significant decrease in channel function, as shown in Figure 4B (current density) (peak current at −160 mV, −0.03 ± 0.01 nA/pF) compared with wild-type Kir2.1 channel function (−0.55 ± 0.17 nA/pF). To test for direct binding, GST fusion proteins of the G1 C terminus and point mutations were made and tested for βγ binding. The amount of interacting β subunit was similar for wild-type GST-G1 CT and GST-G1 CT point mutations (Fig. 5).

**Arginine mutation disrupts intracellular domain association**

Because mutated subunits affected channel function in multiple GIRK subunits and different inward rectifier families, we set out to determine whether the arginine mutation in the C terminus of GIRK1 would disrupt association between the intracellular N- and C-terminal domains, indicating a possible conformational importance of the C-terminal residue in context of channel function. Localization of the I331 residue in the GIRK1 subunit structure indicates that the residue lies between adjacent C termini and the N terminus within a subunit (Nishida and MacKinnon, 2002) (Fig. 6A, B). Although the I331 homologous residue of the KirBac1.1 subunit is a methionine (by sequence alignment), the residue lies adjacent to an unresolved region and the interacting region between the N terminus of one subunit and the C terminus of the adjacent subunit (Kuo et al., 2003) (Fig. 6C). We first examined the association of the full-length subunits by coimmunoprecipitation. G1Δ and G1 I331R associated with the G4-GFP to a similar extent as the wild-type G1 (Fig. 7A). Although the subunits still appear to associate, we reasoned that the molecular interactions between the intracellular regions of the subunits may be affected. To test for intracellular domain association, tagged-constructs of the N- and C-terminal domain were made and expressed in COS-7 cells. G1 NT-myc (residues 1–83) and YFP-G1 CT (including second transmembrane domain, residues 158–501) were coexpressed in COS-7 cells. G1 NT-myc was coimmunoprecipitated with the YFP-G1 CT (Fig. 7B). Coexpression of the G1 NT-myc with either the YFP-G1 CTΔ or YFP-G1 CT I331R failed to coimmunoprecipitate and resulted in a loss of association (Fig. 7B, lanes 3–6). The association between the G1 NT-myc and YFP-G1 CT mutations I331M or I331L was unaffected (data not shown).

Next, we set out to determine whether the associations between subunits were affected as a result of residue mutation. We have shown that the N- and C-terminal association of the GIRK1 subunit is disrupted after mutation of the I331 to an arginine. The mutation of the homologous GIRK4 residue L337R also resulted in decreased coimmunoprecipitation with the G4 NT-FLAG, although the wild-type association was relatively weak (Fig. 8A, 8B, 8C).
the association of the N- and C-terminal domains of Kir2.1 (Fig. 8C).

**Mutation of N-terminal residue decreases channel function**

Structural data pinpoint possible residues that may be important in interactions between N- and C-terminal domains. The GIRK1 I331 appears to lie in a hydrophobic pocket in proximity to several C-terminal residues of the adjacent subunit, as well as the N terminus of the subunit (Fig. 6B). We speculated that the GIRK1 N-terminal phenylalanine residue (F46) was a possible candidate for hydrophobic interaction with the I331. Other possible interacting residues lie in the adjacent C-terminal domain, as indicated in the GIRK1 subunit structure. However, because N- and C-terminal associations appeared to be more critical than the C-termini associations in our studies, we focused on mutations within the GIRK1 N terminus. Mutation of the F46 to an arginine or an aspartate resulted in the mixed ability to coimmunoprecipitate with the wild-type GIRK1 CT (Fig. 9A). The G1 NT F46R appeared to weakly interact with the G1 CT, whereas the G1 NT F46D mutation did not associate with the G1 CT domain (Fig. 9A). Like the G1 I331R subunit, the G1 F46 mutations did not inhibit the expression of the subunit at the cell surface (Fig. 9B). Because the GIRK1 F46 mutation did result in a decrease in the interaction between the GIRK1 N and C termini, we also examined the mutational effects on channel function. Mutation of the full-length subunit F46 to either an aspartate or arginine coexpressed with GIRK 4 resulted in disruption of channel function (Fig. 9C,D; peak HK and HK + Carb current density for G1 + G4, −20.81 ± 6.70 and −42.47 ± 8.66 pA/pF, for G1 F46R + G4, −6.33 ± 2.48 and −15.58 ± 4.62 pA/pF, and for G1 F46D + G4, −2.98 ± 1.71 and −2.84 ± 1.56 pA/pF). Mutation of another hydrophobic residue, F46, within this interaction region of the termini resulted in a similar decrease in functional channel expression, as well as decreased association between intracellular domains.

**Figure 3.** Trafficking of GIRK1 subunits is unaffected by mutation. COS-7 cells were either permeabilized or nonpermeabilized and probed with α-FLAG antibody to detect FLAG-tagged GIRK1 wild-type or mutant subunit expression. Top row, Permeabilized cells were transfected with only the GIRK1 subunit indicated or in addition to the GIRK4-GFP to monitor protein localization. Cotransfected cells were probed with the α-FLAG antibody and Alexa Fluor 488-conjugated secondary antibody to monitor surface expression of GIRK1-FLAG (surface protein expression is represented in red for consistency). Images are 0.5 μm slices. Bottom row, Nonpermeabilized cells were transfected with the GIRK1-FLAG subunit indicated and an untagged GIRK4 subunit and subsequently probed with the α-FLAG antibody and Alexa Fluor 488-conjugated secondary antibody to monitor surface expression of GIRK1-FLAG (surface protein expression is represented in red for consistency). Images are 0.5 μm slices.

**Figure 4.** Homologous GIRK4 and Kir2.1 mutations disrupt channel function. A, Average HK and HK + Carb peak currents at −160 mV were normalized per cell (p < 0.05). B, Basal currents were recorded from COS-7 cells transfected with the indicated subunit and pEGFP. Summary of peak basal currents (at −160 mV) recorded from Kir2.1 and Kir2.1 L330R (p < 0.05) is shown.

**Figure 5.** G-protein binding to Kir cytoplasmic domains. GST fusion proteins of the C-terminal domains of G1 wild type, mutants, Kir2.1, or βARK were tested for Gαi association. The blot was probed with α-Gαi and secondary antibody and visualized with chemiluminescence.
Discussion
Mutations do not alter surface subunit expression
In this study, we examined the functional importance of a hydrophobic region in the cytoplasmic termini, particularly of GIRK subunits. Channel subunit mutations may affect channel function by altering protein expression or subunit localization. The mutation GIRK1 I331R resulted in comparable subunit expression and did not alter cell surface expression. However, we cannot exclude the possibility of a modification in channel subunit composition at the plasma membrane. In the heart, GIRK1 and GIRK4 subunits are thought to comprise functional GIRK channels in a 1:1 stoichiometry (Corey et al., 1998). Several subunit compositions form functional channels in the CNS both as homomultimers and heteromultimers of GIRK subunits (Krapivinsky et al., 1995b; Inanobe et al., 1999; Jelacic et al., 2000; Leaney, 2003). We have shown that the arrangement of the GIRK1 subunit relative to the GIRK2 weaver subunit within a given channel dictates functional channel properties (Hou et al., 1999). It has been demonstrated that the GIRK4 subunit contains forward trafficking motifs that result in surface localization of other GIRK subunits (Ma et al., 2002). Our results suggest that the GIRK1 I331R mutant subunit does not disrupt overall recognition between the GIRK1 and GIRK4 subunits and furthermore does not appear to inhibit channel assembly and membrane localization.

C-terminal domain mutations alter channel function
Mutation of GIRK1 I331 or the homologous GIRK4 and Kir2.1 residues resulted in a decrease in channel function. It has been well established that GIRK channels are activated during binding of G-protein βγ subunits (Logothetis et al., 1987; Reuveny et al., 1994; Huang et al., 1995). The G-protein complex appears to bind several GIRK subunit regions of the intracellular N and C termini, including a portion of the hydrophobic region identified in this study (Huang et al., 1995, 1997; He et al., 1999, 2002; Ivanina et al., 2003). Several studies have shown that the bending of the TM2 likely results in the opening of the pore, much like the gating mechanism deduced from structural studies of the bacterial Ca2+-activated K+ channel MhK (Methanobacterium thermoautotrophicum) K+ channel (Sadja et al., 2001; Jiang et al., 2002; Jin et al., 2002). Fluorescence resonance energy transfer (FRET) analysis demonstrated that the intracellular domains of the GIRK1 and GIRK4 subunits are rearranged during activation (Riven et al., 2003). In the KirBac1.1 crystal structure, cytoplasmic domains are linked to TM domains via linkers and an amphipathic helix, which the authors predict are involved in the coupling between movement of intracellular and TM domains (Kuo et al., 2003). Together, these results support the hypothesis that rearrangements of intracellular domains translate a signal that results in channel opening.

Because the C-terminal domain is well conserved among inward rectifiers, we reasoned that the mutation might result in a defect in the general gating transduction mechanism independent of Gβγ binding. Our data, as well as the data of Chen et al. (2002), which demonstrated that the mutation E304A resulted in a disruption of channel function in both GIRK and Kir2.1 channels containing the homologous mutation, support the latter hypothesis. Other residues within the C terminus of GIRK subunits have also been shown to mediate function. The mutation GIRK4 L339E (and GIRK1 L333E) resulted in a reduction in agonist G-protein-dependent current expression and a significant reduction in direct Gβγ binding. The authors concluded that this residue (the homologous Kir2.1 residue is a glutamate) is responsible for receptor stimulated Gβγ activation (He et al., 1999). Functionally, GIRK1 L333E/GIRK2 L344E channels show a decrease in basal current and a significant loss in agonist-induced GIRK current with a smaller decrease in Gβγ binding (Ivanina et al., 2003). Recently, Finley et al. (2004) demonstrated a similar loss in GIRK2L344E/G347H channel function and little change in Gβγ binding to the double-mutated channel. In this study, we did not detect a difference in Gβγ binding to the mutant domains. However, we found a decrease in both agonist-independent and carbachol-dependent GIRK channel activation and disruption of the non–G-protein-dependent activation of Kir2.1. We have not further characterized differences between agonist-independent and –dependent decreases in channel function because the mutation alters function in both channel types. Furthermore, our results indicate that a disruption in Gβγ binding does not underlie
the functional effects, unlike previous studies examining the role of various residues in this region. Although we cannot rule out the possibility that Gβδ binding is reduced for the GIRK1 I331R subunit in an assembled, intact channel, a similar effect in Kir2.1 argues that it is highly likely that the structural integrity of this region is involved in the transduction event resulting in channel gating. The change in structural interactions involving I331 (or L337 in GIRK4) may indirectly alter Gβδ binding to residues identified in previous studies. Our results implicate an additional residue in this region that is important for channel activation.

### Structural implications support a role in termini interactions

The recent crystal structure of linked GIRK1 subunit intracellular domains suggests that the GIRK1 C-terminal region examined in this study is involved in N- and C-termini interactions. GIRK1 I331 lies in close proximity to the β-sheet formed between the N and C termini of one subunit and other residues of an adjacent GIRK1 subunit (Nishida and MacKinnon, 2002). GIRK subunits are highly homologous throughout this region, suggesting a similar structure in the assembled heteromultimeric channel. The KirBac1.1 crystal structure includes intracellular and transmembrane domains in which the N terminus of one subunit forms a β-sheet with the C terminus of the adjacent subunit. Regardless of which structure more accurately depicts the true termini interactions, it is evident in both structures that the interaction between adjacent N and C termini is likely to play a role in channel gating (Kuo et al., 2003). Finley et al. (2004) also postulated that interactions between GIRK subunit C-terminal regions (GIRK2 residues 344–347) and the adjacent subunit N-terminal domain are involved in activation. It has been shown that the GIRK1 TM domains and C terminus are important for interaction (Woodward et al., 1997). Protein–protein interactions between purified components of the intracellular GIRK1 and GIRK4 termini have also been demonstrated (Huang et al., 1997). Because the loss of channel function appears to be dependent on a hydrophobic residue that maps to an interacting region between cytoplasmic domains in the GIRK1 subunit structure, we examined possible associations between intracellular domains of the subunits. Although our studies do not measure direct interactions, they clearly showed that the GIRK1 N and C termini coimmunoprecipitated and that the mutation significantly decreased this association. In addition, N- and C-terminal association within the GIRK4 subunit or between subunits is decreased when mutated, but the C termini associations are not altered. Although these results are a product of determining domain associations in an expression system, the results mirror the loss of channel function.

Several possibilities may account for both the decrease in association of cytoplasmic domains and a disruption in channel function. In an assembled channel, the I331 may directly interact with...
other adjacent N-terminal residues (such as F46) or the adjacent C-terminal subunit. Mutations may disrupt direct interactions resulting in channel dysfunction. Alternatively, the mutation may disrupt the overall conformation of the region hindering the interaction of other proximal residues that then directly interact with the N terminus. Interactions between N- and C-terminal domains of the Kir6.2 subunit have been localized to a region in the N terminus that, when mutated, resulted in a decrease in channel function (Tucker and Ashcroft, 1999). The N-terminal residue identified is a conserved glycine and maps to a similar region of interest. Also, the Kir6.2 N terminus binds to several C-terminal regions, including a domain that contains the homologous GIRK1 I331 residue (Jones et al., 2001). Another possibility is that the arginine mutation establishes an alternative interaction that stabilizes the channel in a closed conformation. We speculate that the intracellular interface between termini of individual subunits and among adjacent subunits comprises a portion of the pathway of conformational repositioning that eventually results in opening of the pore.

**Hydrophobic nature is critical for channel function**

Our results argue strongly that the conformational arrangement of the isoleucine is highly important and suggests a general mechanism for the dysfunction of channel activation. Moreover, because Kir2.1 channels are constitutively open, mutation of this conserved glycine and maps to a similar region of interest. Also, the Kir6.2 N terminus binds to several C-terminal regions, including a domain that contains the homologous GIRK1 I331 residue (Jones et al., 2001). Another possibility is that the arginine mutation establishes an alternative interaction that stabilizes the channel in a closed conformation. We speculate that the intracellular interface between termini of individual subunits and among adjacent subunits comprises a portion of the pathway of conformational repositioning that eventually results in opening of the pore.

Of the Kir2.1 channel, we would expect a decrease in the association of the N- and C-terminal intracellular domains of the Kir2.1, which we did not observe. Therefore, although similar mutations affect the function of both channels, the conformational changes in the cytoplasmic domains are most likely different.

Together, our results suggest that the structural integrity of this specific region is important for channel function and that the introduction of a charged residue in this hydrophobic region destabilizes the structure. Mutational studies that compare the effects of hydrophobic substitutions will provide additional information on the nature of the interacting region. Additional studies analyzing the interaction between the intracellular components of the GIRK channel and other inward rectifiers will shed light on the importance of these intracellular domain interactions.

**References**


Sui JL, Petit-Jacques J, Logothetis DE (1998) Activation of the atrial K\textsubscript{ATP} channel by the βγ subunits of G proteins or intracellular Na\textsuperscript{+} ions depends on the presence of phosphatidylinositol phosphates. Proc Natl Acad Sci USA 95:1307–1312.


