Octameric Stoichiometry of the $K_{\text{ATP}}$ Channel Complex

S.-L. Shyng and C.G. Nichols

From the Department of Cell Biology and Physiology, Washington University School of Medicine, St. Louis, Missouri 63110

Abstract: ATP-sensitive potassium ($K_{\text{ATP}}$) channels link cellular metabolism to electrical activity in nerve, muscle, and endocrine tissues. They are formed as a functional complex of two unrelated subunits—a member of the Kir inward rectifier potassium channel family, and a sulfonylurea receptor (SUR), a member of the ATP-binding cassette transporter family, which includes cystic fibrosis transmembrane conductance regulators and multidrug resistance protein, regulators of chloride channel activity. This recent discovery has brought together proteins from two very distinct superfamilies in a novel functional complex. The pancreatic $K_{\text{ATP}}$ channel is probably formed specifically of Kir6.2 and SUR1 isoforms. The relationship between SUR1 and Kir6.2 must be determined to understand how SUR1 and Kir6.2 interact to form this unique channel. We have used mutant Kir6.2 subunits and dimeric (SUR1-Kir6.2) constructs to examine the functional stoichiometry of the $K_{\text{ATP}}$ channel. The data indicate that the $K_{\text{ATP}}$ Channel pore is lined by four Kir6.2 subunits, and that each Kir6.2 subunit requires one SUR1 subunit to generate a functional channel in an octameric or tetradimeric structure.

Key words: potassium channel • adenosine triphosphate–binding cassette protein • sulfonylurea receptor • inward rectifier

Introduction

ATP-sensitive potassium ($K_{\text{ATP}}$) channels are unique in requiring two structurally distinct subunits to generate functional channels. One subunit (Kir6.2) is a member of the inward rectifier family of potassium channels (Inagaki et al., 1995; Nichols and Lopatin, 1997), while the other (SUR1, a high affinity sulfonylurea receptor; Aguilar-Bryan et al., 1995) is a member of the ATP-binding cassette (ABC)1 superfamily (Higgins, 1995). Expression of either subunit alone normally fails to generate active channels, while coexpression of Kir6.1 or Kir6.2 subunits with SUR1 or SUR2 subunits forms $K_{\text{ATP}}$ channels with properties identical to those described in pancreatic β cells and cardiac myocytes (Noma, 1983; Ashcroft, 1988; Inagaki et al., 1995a, 1996). The requirement for both subunits raises the possibility that each one contributes to formation of the channel pore in an arrangement other than the homotetrameric array that has been shown for other K+ channels (MacKinnon, 1991; Glowatzki et al., 1995; Yang et al., 1995).

In the present study, we have examined the stoichiometry of channel formation using mutant Kir6.2 subunits and dimeric SUR1-Kir6.2 constructs. Mutation of asparagine to aspartate in the pore of Kir6.2 leads to the generation of strongly inwardly rectifying channels (as previously demonstrated for similar mutations in Kir1.1 channels; Lu and MacKinnon, 1994), as a result of induction of high sensitivity to block by intracellular spermine (Lopatin et al., 1994; Shyng et al., 1997a). By coexpressing wild-type and mutant Kir6.2 subunits, we generated heteromeric channels with intermediate rectification properties. Examination of current–voltage relationships from these heteromeric channels reveals that the $K_{\text{ATP}}$ channel pore is likely to be formed as a tetrameric arrangement of Kir6.2 subunits, as shown previously for voltage-gated K+ channels and homomeric inward rectifier channels (MacKinnon, 1991; Glowatzki et al., 1995; Yang et al., 1995). Expression of dimeric SUR1-Kir6.2 constructs results in functional channels with very similar properties to those formed from coexpression of the parental monomers, consistent with a 1:1 SUR1/Kir6.2 stoichiometry being sufficient to form the $K_{\text{ATP}}$ channel. Additional experiments with coexpression of dimers and monomers indicate that a 1:1 stoichiometry may be preferred, leading to the conclusion that the $K_{\text{ATP}}$ channel is likely to be formed as an octameric complex of four Kir6.2 subunits and four SUR1 subunits.

Materials and Methods

Expression of $K_{\text{ATP}}$ Channels in COSm6 Cells

COSm6 cells were plated at a density of 2.5 x 10^5 cells per well (30-mm six-well dishes) and cultured in Dulbecco’s Modified Eagle Medium plus 10 mM glucose (DMEM-HG), supplemented...
with fetal calf serum (10%). 24 h later, monomeric hamster SUR1 (in pECE vector; Aguilar-Bryan et al., 1995), mouse Kir6.2 (in pCMV vector, Inagaki et al., 1995a), or dimeric SUR1-Kir6.2 (in pECE vector) were cotransfected into the COS7 cells with diethylaminoethyl-dextran (0.5 mg/ml). Cells were incubated for 2 min in HEPES-buffered salt solution containing DMSO (10%), and then for 4 h in DMEM-HG plus 2% FCS and chloroquine (100 μM), and then returned to DMEM-HG plus 10% FCS. For °86Rb+ flux experiments, °86RbCl (1 μCi/ml) was added in fresh DMEM-HG containing FCS (10%) 24 h after transfection. Cells were incubated for 12-24 h before measurement of Rb efflux. For efflux measurements, cells were incubated for 30 min at 25°C in Krebs’ Ringer solution, with or without metabolic inhibitors (2.5 μg/ml oligomycin plus 1 mM 2-deoxy-glucose) or glibenclamide. At selected time points, the solution was aspirated from the cells and replaced with fresh solution. The °86Rb+ in the aspirated solution was counted.

**Patch-Clamp Measurements**

1–3 d after transfection, patch-clamp experiments were made at room temperature in an oil-gate chamber (Lederer and Nichols, 1989). Micropipettes were pulled from thin-walled glass (WPI, New Haven, CT) on a horizontal puller (Sutter Instruments, Co., Novato, CA), fire polished, and the tips were coated with a 1:1 mixture of light mineral oil and Parafilm (American National Can Co., Greenwich, CT), to reduce capacitive currents. Electrode resistance was typically 0.5–1 MΩ when filled with K-INT solution (see below). Microelectrodes were ‘sealed’ onto cells by applying light suction to the rear of the pipette. Inside-out patches were obtained by lifting the electrode, and then passing the electrode tip through the oil gate. Membrane patches were voltage-clamped with an Axopatch 1B patch-clamp (Axon Instruments, Foster City, CA). PClamp software and a Labmaster TL125 D/A converter (Axon Instruments) were used to generate voltage pulses. Data was normally filtered at 0.5–3 kHz, signals were digitized at 22 kHz (Neurocorder; Neurodata, New York) and stored on video tape. Experiments were replayed onto a chart recorder or digitized into a microcomputer using Axotape software (Axon Instruments). The standard bath (intracellular) solution used throughout these experiments (K-INT) had the following composition (mM): 140 KCl, 10 K-HEPES, 1 K-EGTA, with additions as described. The solution pH was 7.3. The pipette solution was either also K-INT, or Na-INT in which 140 mM KCl was replaced through these experiments (K-INT) had the following composition (mM): 140 KCl, 10 K-HEPES, 1 K-EGTA, with additions as described. The solution pH was 7.3. The pipette solution was either also K-INT, or Na-INT in which 140 mM KCl was replaced with fresh solution. The °86Rb+ in the aspirated solution was counted.

**RESULTS**

**K<sub>ATP</sub> Channels Are Assembled From Four Kir6.2 Subunits**

K<sub>ATP</sub> channels in native tissues do not show strong inward rectification (Noma, 1983; Ciani and Ribalet, 1988) and, as demonstrated in Fig. 1a, neither do K<sub>ATP</sub> channels generated by coexpression of Kir6.2 or SUR1 subunits. Work on cloned inward rectifiers has demonstrated that strong inward rectification is controlled by a pore lining residue in the M2 transmembrane domain (Lu and MacKinnon, 1994; Stanfield et al., 1994), and that the presence of a negative charge at this “rectification controller” position confers strong inward rectification by generation of a high affinity site for the voltage-dependent binding of cytoplasmic polyamines or Mg<sup>2+</sup> (Fakler et al., 1994; Ficker et al., 1994; Lopatin et al., 1994). Kir6.2 subunits are homologous to Kir1.1 (ROMK1; Ho et al., 1993) channels in this region of the M2 segment; both contain an asparagine (N160 in Kir6.2) at the rectification controller position. We used a mutant Kir6.2 containing a negatively charged aspartate at this position (Kir6.2[N160D]; Clement et al., 1997; Shyng et al., 1997a). Expression of this mutant with SUR1 subunits results in generation of K<sub>ATP</sub> channels that rectify strongly in the presence of cytoplasmic spermine (Fig. 1b), although single channel conductance is unaltered and channels are still sensitive to inhibition by ATP (K<sub>ATP</sub> = 10.6 μM for wild-type Kir6.2 plus SUR1 channels, and 46.1 μM for Kir6.2[N160D] plus SUR1 channels; Shyng et al., 1997a). Rb<sup>+</sup> flux through both channels is fully inhibited by 1 μM glibenclamide (not shown). The large difference in spermine sensitivity of K<sub>ATP</sub> channels formed from wild-type and Kir6.2[N160D] subunits makes it possible to analyze the hetero-oligomeric assembly of Kir6.2 subunits and directly test the hypothesis that the K<sub>ATP</sub> channel pore involves a tetrameric arrangement of Kir6.2 subunits (MacKinnon, 1991; Glowatzki et al., 1995; Yang et al., 1995).

Expression of SUR1 cDNA with a 1:1 mixture of wild-type and Kir6.2[N160D] subunit cDNAs gives rise to channels that differ substantially from the parental homo-oligomeric channels in their sensitivity to block by intracellular spermine (Fig. 1c). The relative conductance–voltage (G<sub>REL-V</sub>) curve contains more than two components (Fig. 1c, bottom), and therefore could not arise from a weighted average of parental G<sub>REL-V</sub> curves, indicating that multimerization generates channels with novel phenotypes. We attempted to fit G<sub>REL-V</sub> curves like that in Fig. 1c, bottom with the sum of multiple Boltzmann functions. Acceptable fits could not be obtained with the sum of four or less Boltzmann func-

![Figure 1](image-url)
respond to different combinations of wild-type Kir6.2 and Kir6.2[N160D] in a tetramer, with fitted probability of wild-type incorporation \( P = 0.56 \). For this patch, the unconstrained Boltzmann functions (for \( i = 2–4 \)) components, were fitted with \( V_1 = -48 \) mV, \( z_1 = 2.9; V_2 = +9 \) mV, \( z_2 = 2.9; V_3 = +86 \) mV, \( z_3 = 1.0 \). The individual Boltzmann functions are also plotted, with the contributing number of wild-type and Kir6.2[N160D] subunits indicated beside each curve. (d) Residual errors (calculated as the sum of squares of the difference between the fit and the data, every 1 mV, between -80 to +160 mV, excluding -10 to +10 mV), obtained by fitting with \( n = 2–6 \) subunits \( \Sigma (G_{rel} - G_f)^2 \), where \( G_f \) is the fitted \( G_{rel} \) plotted against the number of assumed subunits \( N \). The dashed line and open symbols correspond to the patch illustrated in c. Solid lines and symbols correspond to averaged data for \( n = 3–5 \) subunit fits for all patches \( n = 9 \) patches, mean \( \pm \) SEM). The trimeric model \( (n = 3) \) is inadequate to fit the data (see d), and better fits are not obtained by increasing \( n \) from 4 to 5. Similar results were obtained when attempting to fit the model to derivative \( \Delta G_{rel}/\Delta V \) data (not shown).
portant. (The ordering of subunits within the tetramer was unim-
mutable subunits should give rise to similar or increas-
ing the reasonable assumption that decreasing numbers of
channels containing 0, 1, 2, 3, or 4 wild-type Kir6.2 sub-
unites, respectively (Fig. 2, a and b), thus allowing
represent channels containing 3, 2, and 1 mutant sub-
units, respectively, and that $V_i < V_5 < V_4 < V_3 < V_2$, and $z_4 > z_5 > z_3 > z_2 > z_1$. These constraints are based on
the reasonable assumption that decreasing numbers of
mutant subunits should give rise to similar or increas-
ingly shallower and weaker spermene block, and that
the ordering of subunits within the tetramer was unim-
portant. (c) We assumed that the fitted amplitude ($A_i$) corre-
sponds to the probability ($P_i$) of formation of each chan-

$$G_{rel} = \sum_{i=1}^{5} A_i \cdot \{1 + \exp[(F/RT) \cdot z_i \cdot (V - V_i)]\}^{-1}$$

(1)

where $A_i$, $V_i$, and $z_i$ are the amplitude, voltage of half-
maximal inhibition, and effective valency, respectively, of the $i^{th}$ component. With the constraints described
below, we did not obtain better fits with the sum of six
or more Boltzmann functions, based on least squares
residuals (Fig. 1 d), leading us to the conclusion that five different species of tetrameric channels are ran-
donally assembled in these mixing experiments, the
channels containing 0, 1, 2, 3, or 4 wild-type Kir6.2 sub-
units are incorporated with equal probability ($p$) af-
ter the binomial distribution:

$$p_x = \binom{n}{x} \cdot p^x \cdot (1 - p)^{n-x}$$

(2)

where $n$ is the total number of subunits in a functional
channel, $x$ is the number, 0 to $n$, of wild-type subunits
in a particular channel, $P$ is the probability of inclusion
of a wild-type subunit, and $(1 - P)$ is the probability of
inclusion of a mutant subunit. Hence, fits were constra-
ined by allowing $P$ to vary. Only patches with cur-
rents $>-200$ pA at $-50$ mV (corresponding to $\sim50$
channels were included in the analysis, to increase the
signal-to-noise ratio to acceptable levels. Leak conduc-
tance in the presence of high [ATP] (5 or 10 mM) was
typically $<10$ pS, and patches were rejected from analy-
sis if the leak conductance was $>100$ pS (i.e., $>5%$ of $K_{ATP}$
conductance). The data were well fitted with $P$ values
close to 0.5 (0.58 $\pm$ 0.04, $n = 4$ for SUR1 plus 1:1 mix-
ture of Kir6.2[N160D] and wild-type Kir6.2 cDNAs).
This agreement between fitted $P$ values and transfected
cDNA ratios is reassuring, since the transfection and expres-
sion efficiency of the wild-type Kir6.2 and Kir6.2[N160D]
cDNAs is not expected to be different.

Each Kir6.2 Subunit Is Normally Associated with One
SUR1 Subunit

We cannot formally discount a role of SUR1 subunits in
generating the channel pore, although none of several
mutations in the cytoplasmic domains of SUR1 have

\begin{figure}
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\caption{Fusion channels show identical voltage dependence to parental monomers. (a) Mean $\pm$ SEM for $z_i$ of single Boltzmann functions fitted to $G_{rel} - V$ relationships obtained from wild-type Kir6.2 or Kir6.2[N160D] subunits coexpressed with SUR1 ( ), or from WFT or NDF fusion proteins ( ), and for $z_i$ of five component Boltzmann functions fitted to $G_{rel} - V$ relationships obtained from mixed wild-type Kir6.2 and Kir6.2[N160D] subunits coexpressed with SUR1 ( ), or from mixed expression of WFT and NDF fusion proteins ( ). For these mixed expressions, only $z_5$.

$z_a$ and $z_b$ were varied, $z_1$ and $z_2$ corresponded to the mean value fitted to homomeric expressions. The dashed line indicates the $z_i$ values used to fit the tetrameric model to data in Fig. 6. (b) Mean $\pm$ SEM for $V_i$ of single Boltzmann functions fitted to $G_{rel} - V$ relationships obtained from wild-type Kir6.2 or Kir6.2[N160D] subunits coexpressed with SUR1 ( ), or from WFT or NDF fusion proteins ( ), and for $V_i$ of five component Boltzmann functions fitted to $G_{rel} - V$ relationships obtained from mixed wild-
type Kir6.2 and Kir6.2[N160D] subunits coexpressed with SUR1 ( ), or from mixed expression of WFT and NDF fusion proteins ( ). For these mixed expressions, only $V_2$, $V_3$, and $V_4$ were varied, $V_5$ and $V_6$ corresponded to the mean value fitted to homomeric expressions. The dashed line indicates the $V_i$ values used to fit the tetrameric model to data in Fig. 6.

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been shown to affect conductance or rectification of expressed channels (Nichols et al., 1996; Gribble et al., 1997; Shyng et al. 1997b). Very recently, evidence has been presented that certain Kir6.2 constructs with deletions of 26 or 36 amino acids from the COOH terminus can actually form functional K\textsubscript{ATP} channels in the absence of the sulfonylurea receptor (Tucker et al., 1997). In that study, however, full length Kir6.2 subunits did not form functional channels without SUR1 coexpression, and even truncated constructs generated more current when coexpressed with SUR1, consistent with a normal requirement for association.

To investigate the stoichiometry of SUR1 and Kir6.2 subunits in functional channels, we used SUR1-Kir6.2 fusion proteins, with a defined 1:1 SUR1/Kir6.2 ratio (Fig. 3), in which the COOH terminus of SUR1 is covalently linked to the NH\textsubscript{2} terminus of Kir6.2 using a hexa-glycine (-Gly\textsubscript{6}-) linker (Clement et al., 1997). Fusion proteins containing wild-type Kir6.2 (wild-type fusion, WTF), or N160D mutated Kir6.2 (N160D fusion, NDF), generate K\textsubscript{ATP} channels that are fully inhibited by 1 \mu M glibenclamide in Rb\textsuperscript{+} efflux assays (data not shown), and by ATP in excised patches. There is no obvious difference in the density of channels formed by expression of the fusion proteins versus coexpression of the parent monomers, based on the density of maximally activated K\textsubscript{ATP} conductances in \textsuperscript{38}Rb\textsuperscript{+} efflux experiments (see Fig. 5). Fig. 4 shows records of currents through WTF and NDF channels to illustrate ATP sensitivity. Although these channels have wild-type single channel conductance in the absence of internal multivalent cations (75 pS in symmetrical 140 mM K\textsuperscript{+}, not shown), they are approximately five- to eightfold less sensitive to ATP than the parental monomers. The estimated K\textsubscript{i} (from \(n = 3\)–5 patches) is 45.3 \mu M for WTF (compare 10.6 \mu M for wild-type Kir6.2 plus SUR1 channels), and 384.6 \mu M for NDF (compare 46.1 \mu M for Kir6.2[N160D] plus SUR1 channels). Because of the reduced ATP sensitivity of fusion protein channels, we routinely used exposure to 10 mM ATP for estimation of leakage current. As shown in Figs. 2 and 3, fusion proteins show identical spermine sensitivity to the chan-

![Figure 3](image-url)

**Figure 3.** Fusion channels contain four dimeric SUR1-Kir6.2 dimers. (a-c) (top) Macroscopic inside-out patch currents in response to voltage ramps (300 ms) between +100 and −100 mV, plotted versus membrane potential, for K\textsubscript{ATP} channels formed from SUR1-Kir6.2 fusion proteins, containing either wild-type (a, WTF) or N160D mutant (b, NDF) Kir6.2 sequences, or from equimolar mixed WTF and NDF constructs (c). (top) Currents are shown in the presence and absence of 20 \mu M spermine (spm); in each case, currents are shown after subtraction of leakage currents in the presence of 10 mM ATP (which reduces channel open probability to <1%). (bottom) Relative conductance (\(G_{REL}\))-voltage relationships from the data shown above. The continuous lines in b and c are best fits of single Boltzmann functions. For these patches, the unconstrained Boltzmann functions (for \(i = 5\) and 0, respectively) were fitted with \(V_0 = +200\) mV, \(z_i = 0.4\) (a); \(V_1 = −58\) mV, \(z_1 = 2.9\) (b). In c (bottom), the smooth line that superimposes on the data is the sum of five Boltzmann functions that correspond to different combinations of wild-type Kir6.2 and Kir6.2[N160D] in a tetramer, with fitted probability of wild-type incorporation (\(P\)) = 0.35. For this patch, the unconstrained Boltzmann functions (for \(i = 2–4\)), components were fitted with \(V_2 = −46\) mV, \(z_2 = 2.9\); \(V_3 = +10\) mV, \(z_3 = 2.5\); \(V_4 = +99\) mV, \(z_4 = 1.2\). The individual Boltzmann functions are also plotted, with the contributing number of wild-type and Kir6.2[N160D] subunits indicated beside each curve.
nals formed by expression of SUR1 and the respective Kir6.2 monomer type, and when mixtures of WTF and NDF are coexpressed, the spermine sensitivity of heteromultimers is identical to those formed by mixtures of parental monomers (Figs. 2 and 3). Expression of a 1:1 mixture of WTF and NDF cDNAs gives rise to the same distribution of spermine block phenotypes as a 1:1 mixture of wild-type Kir6.2 and Kir6.2[N160D] monomer cDNAs with SUR1, in accord with the tetrameric arrangement, the data again being well fitted with $P$ values close to 0.5 for both cases ($P = 0.45 \pm 0.05$, $n = 5$ for 1:1 WTF/NDF cDNA mixtures, see Fig. 3; compare $P = 0.58 \pm 0.04$, $n = 4$ for the SUR1 plus nominally 1:1 mixture of Kir6.2[N160D] and wild-type Kir6.2 subunits, above). Averaged data for the fitted $z_i$ and $V_i$ parameters obtained by expression of mixed wild-type Kir6.2 and Kir6.2[N160D] (plus SUR1) subunits, and by expression of WTF and NDF subunits are shown in Fig. 2, a and b. It is clear that there is no consistent difference in the rectification properties of channels formed by expression of monomeric Kir6.2 subunits, or by expression of dimeric SUR1-Kir6.2 fusion proteins.

The formation of functional $K_{ATP}$ channels by SUR1-Kir6.2 fusion proteins is consistent with a 1:1 SUR1/Kir6.2 stoichiometry being sufficient for channel formation. To examine whether such a stoichiometry is required, we coexpressed fusion proteins with Kir6.2 monomers to allow generation of channels with stoichiometries where Kir6.2 > SUR1. In $^{86}$Rb$^+$ efflux experiments (Fig. 5), transfection of SUR1 and Kir6.2 in approximately equal cDNA ratios produced similar efflux to that resulting from expression of the WTF fusion protein cDNA (at equivalent molar ratio). Coexpression of monomeric Kir6.2 with the fusion protein, at a ratio of Kir6.2/fusion protein cDNA of ~1:1 (keeping the total amount of Kir6.2 cDNA constant, whether in monomeric or fusion protein construct), suppressed the $K_{ATP}$ conductance. This apparent dominant-negative effect may result from incorporation of Kir6.2 monomers and fusion proteins into nonfunctional complexes without a requisite 1:1 (Kir6.2/SUR1) stoichiometry (see Fig. 7). If such an interpretation is correct, then further addition of monomeric SUR1 subunits might rescue the nonfunctional complexes by permitting the requisite stoichiometry to be achieved. Strikingly, this is the case, as shown by $^{86}$Rb$^+$ efflux measurements (Fig. 5). Even though coexpression of pore-forming Kir6.2 subunits with fusion proteins reduces the $K_{ATP}$ conductance, the additional expression of regulatory SUR1 subunits restores the conductance (cDNA ratio 1:1:1). The fusion/Kir6.2 cDNA ratio was the same in both cases, so, although additional expression of SUR1 subunits might conceivably have reduced the total amount of expressed fusion protein and Kir6.2 subunits (due to overload of the biosynthetic machinery), it could not have altered the expressed fusion protein/ Kir6.2 subunit ratio, and could not have increased the total number of expressed Kir6.2 subunits (i.e., the sum of Kir6.2 monomers and Kir6.2 in fusion proteins). The only reasonable conclusion then is that the increased number of functional channels when SUR1 is coexpressed results from rescue of otherwise nonfunctional Kir6.2 subunits as a result of their association with SUR1 subunits.

**Figure 4.** Fusion channels show reduced sensitivity to ATP. Representative currents recorded from inside-out membrane patches containing WTF or NDF $K_{ATP}$ channels (as indicated) at $-50$ mV. Patches were exposed to differing [ATP] as indicated (mM). Inward currents are shown as upward deflections.

**Figure 5.** Monomeric Kir6.2 inhibits fusion protein conductance. $^{86}$Rb$^+$ efflux from untransfected COSm6 cells, and cells expressing SUR1-Kir6.2 fusion proteins with or without additional Kir6.2 and SUR1 subunits. Graphs show percent Rb$^+$ released into the medium as a function of time in the presence of metabolic inhibitors (see methods) for a typical experiment ($n = 3$). The approximate cDNA molar ratio was 1:1:1 for SUR1-Kir6.2 fusion protein/Kir6.2/SUR1.
Insight into the nature of the currents formed by mixed expression of the fusion protein with monomers was obtained from inside-out membrane patches. Channels were formed by coexpression of WTF or NDF fusions with Kir6.2[N160D] or wild-type Kir6.2 monomers, respectively, at equimolar cDNA ratios. As expected from the $^{86}$Rb$^+$ efflux experiments, the observed current density in membrane patches was lower than that seen with expression of fusion proteins alone. Although $I_{\text{ATP}}$ currents were present in 15/15 patches from cells expressing only the fusion proteins, measurable $I_{\text{ATP}}$ currents were present in only 2/12 patches when Kir6.2[N160D] was coexpressed with WTF (1:1 molar ratio of cDNAs), and in only 3/13 patches when wild-type Kir6.2 was coexpressed with NDF. When the molar ratio of fusion protein/Kir6.2 cDNAs was increased to 8:1, measurable currents were present in 13/21 patches (combined data for both WTF plus Kir6.2[N160D], and for NDF plus wild-type Kir6.2). The phenotype of active channels was predominantly that expected from the fusion protein (Fig. 6, a and b). Using estimates of $z_i$ and $V_i$ (spermine sensitivity of heteromeric channels containing 0–4 Kir6.2 subunits) obtained above (Fig. 2, a and b, points connected by dashed lines), we fitted the tetrameric model to data from heteromeric channels composed of Kir6.2 and fusion protein, varying only the apparent ratio of wild-type to mutant Kir6.2 subunits (Fig. 6 and Table I). At all fusion/Kir6.2 cDNA

![Figure 6](image-url)
KATP channels are unique amongst known potassium channels in requiring an unrelated ABC protein subunit (SUR1) in addition to an inward rectifier K channel (Kir6.2) subunit (Inagaki et al., 1995a). In other cloned inward rectifiers, strong inward rectification is controlled by a pore-lining residue in the M2 transmembrane segment (Fakler et al., 1994; Ficker et al., 1994; Lopatin et al., 1994; Lu and MacKinnon, 1994; Stanfield et al., 1994). Mutation of the corresponding residue in Kir6.2 from asparagine to aspartate results in channels in requiring an unrelated ABC protein subunit (Kir6.2) in addition to an inward rectifier K channel (Kir6.2) subunit (Inagaki et al., 1995a). In other cloned inward rectifiers, strong inward rectification is controlled by a pore-lining residue in the M2 transmembrane segment (Fakler et al., 1994; Ficker et al., 1994; Lopatin et al., 1994; Lu and MacKinnon, 1994; Stanfield et al., 1994). Mutation of the corresponding residue in Kir6.2 from asparagine to aspartate results in generation of KATP channels that rectify strongly in the presence of cytoplasmic spermine (Fig. 1b; Clement et al., 1997; Shyng et al., 1997), single channel conductance being unaltered and channels remaining sensitive to inhibition by ATP (Shyng et al., 1997). The requirement for SUR1 to form active channels still raises the possibility that the receptor might also contribute to the pore, and perhaps reduce or otherwise alter the number of Kir6.2 subunits involved. The large difference in spermine sensitivity of KATP channels formed from wild-type and Kir6.2[N160D] subunits made it possible for us to directly analyze the hetero-oligomeric assembly of Kir6.2 subunits. As shown in Fig. 1d, the data could not be fit by assuming a trimer or less, and the fits of pentameric or hexameric models were not better than the tetrameric model. The analysis that we used (Fig. 1) was previously used by Glowatzki et al. (1995), after the original approach of MacKinnon

**TABLE I**

| Subunits in Coexpression with Fusion Proteins | Nominal molar ratio of cDNAs (fusion protein/Kir6.2 monomer) | 1:1 | 8:1 | 18:1 |
|---------------------------------------------|-------------------------------------------------------------|-----|-----|-----|
| (f6.2 = monomer/total)                      |                                                             |     |     |     |
| −SUR1 subunits (f6.21)                      |                                                             | 0.26(w)* | 0.15(m) | 0.06(m) |
| (f6.22)                                     |                                                             | 0.27(m) | 0.16(m) | 0.06(w) |
| mean                                        |                                                             | 0.26 | 0.15 ± 0.02 | 0.06 |
| +SUR subunits† (f6.21)                      |                                                             | 0.93(m) | 0.84(m) | 0.60(m) |
| (f6.22)                                     |                                                             | 0.93(w) | 0.83(m) | 0.69(m) |
| mean                                        |                                                             | 0.93 ± 0.01 | 0.84 ± 0.01 | 0.58 ± 0.03 |
| Fold increase in monomer incorporation with SUR1 = f6.22/(1−f6.22)/(f6.21/(1−f6.21)) |                                                             | 37.8 | 29.8 | 21.6 |

*(w) and (m) refer to wild-type fusion (plus Kir6.2[N160D] monomer), and N160D fusion (plus wild-type Kir6.2 monomer), respectively. †SUR1/Kir6.2 at nominal 1:1 molar cDNA ratio.

**DISCUSSION**

KATP channels are unique amongst known potassium channels in requiring an unrelated ABC protein subunit (SUR1) in addition to an inward rectifier K channel (Kir6.2) subunit (Inagaki et al., 1995a). In other cloned inward rectifiers, strong inward rectification is controlled by a pore-lining residue in the M2 transmembrane segment (Fakler et al., 1994; Ficker et al., 1994; Lopatin et al., 1994; Lu and MacKinnon, 1994; Stanfield et al., 1994). Mutation of the corresponding residue in Kir6.2 from asparagine to aspartate results in generation of KATP channels that rectify strongly in the presence of cytoplasmic spermine (Fig. 1b; Clement et al., 1997; Shyng et al., 1997), single channel conductance being unaltered and channels remaining sensitive to inhibition by ATP (Shyng et al., 1997). The requirement for SUR1 to form active channels still raises the possibility that the receptor might also contribute to the pore, and perhaps reduce or otherwise alter the number of Kir6.2 subunits involved. The large difference in spermine sensitivity of KATP channels formed from wild-type and Kir6.2[N160D] subunits made it possible for us to directly analyze the hetero-oligomeric assembly of Kir6.2 subunits. As shown in Fig. 1d, the data could not be fit by assuming a trimer or less, and the fits of pentameric or hexameric models were not better than the tetrameric model. The analysis that we used (Fig. 1) was previously used by Glowatzki et al. (1995), after the original approach of MacKinnon
(1991) to demonstrate that homomeric Kir4.1 (BIR10) channels are formed as tetramers. In both cases, the strong rectifying phenotype of the N160D (or equivalent) mutation is dominant, channels containing three wild-type subunits are approximately intermediate between wild-type and homotetrameric mutant channels in spermine sensitivity (Figs. 2 and 3). This suggests that Kir6.2 subunits form tetrameric $K_{\text{ATP}}$ channel pores in a very similar arrangement to those of homomeric Kir channel pores (Glowatzki et al., 1995; Yang et al., 1995).

There are reports of Kir6.1 and Kir6.2 (COOH-terminal truncated) expressing $K_{\text{ATP}}$ channels in the absence of SUR1 (Inagaki et al., 1995b; Tucker et al., 1997), and there are also reports of “promiscuous” coupling of SUR1 to homomeric Kir1.1 channels that form active K currents in the absence of SUR1 (Ammala et al., 1996). While conflicting results have been reported by others (e.g., Inagaki et al., 1995a), these results suggest a non-obligatory and perhaps nonstoichiometric relationship of SUR1 to Kir6.2. By analyzing channels formed by deliberately generated 1:1 stoichiometry (in SUR1-Kir6.2 fusion proteins), and by altered stoichiometry (mixing fusion proteins with monomeric subunits), we can show (Figs. 2–6, Table I) that a 1:1 stoichiometry is possible and, moreover, may be strongly preferred, indicating a normally octameric assembly. Nevertheless, the results do indicate that a low frequency incorporation of monomeric Kir6.2 subunits in functional channels can still occur when coexpressed with fusion proteins (Fig. 6, a and c). This may be explained by assuming that the necessary SUR1 subunit is provided by a fusion protein, with its Kir6.2 portion “hanging out” of the complex (Fig. 7) as has been suggested to explain formation of Kv and Kir channels by expression of pentameric and trimeric concatamers (Liman et al., 1992; Yang et al., 1995). However, since monomeric SUR1 is 20- to 40-fold more efficient at generating channels with Kir6.2 monomers than is the SUR1-Kir6.2 dimer (Table I), this is clearly an unfavorable arrangement, and not expected to be a significant component of channels formed with equal ratios of Kir6.2 and SUR1 subunits. Using a biochemical approach, Clement et al. (1997) have shown that SUR1 and Kir6.2 are physically associated in COSm6 cells and form a complex with an estimated molecular mass of $\sim$950,000 D, which is reasonably consistent with four SUR1 subunits ($\sim$170,000 D) and four Kir6.2 subunits ($\sim$45,000 D). Estimates for the size of the complex formed by expression of the fusion protein are similar, consistent with the octameric, or tetrameric [$\text{SUR1}-\text{Kir6.2}$] structure that we predict. Using essentially the same biophysical approach used here, Clement et al. (1997) also show that a tetrameric model can fit $G_{\text{REL}}-V$ curves from mixed expression of monomeric SUR1 and triple fusion (SUR1-Kir6.2-Kir6.2) constructs (see also below).

The presence of nucleotide binding folds in the sulfonylurea receptor (Aguilar-Bryan et al., 1995) has led to a widespread expectation that ATP inhibition will be shown to occur through binding to these sites. However, while studies with mutations in the nucleotide binding folds of SUR1 have demonstrated that these binding folds are involved in MgADP and diazoxide stimulation of channel activity (Nichols et al., 1996; Gribble et al., 1997; Shyng et al., 1997b), they provide no evidence for any role in intrinsic ATP inhibition. The recent evidence that the Kir6.2 subunit can control ATP sensitivity, and that a COOH-terminally truncated Kir6.2 (Kir6.2[ΔC26]) forms functional $K_{\text{ATP}}$ channels in the absence of SUR1 (Shyng et al., 1997a; Shyng et al., 1997b; Tucker et al., 1997) raises the possibility that ATP inhibition occurs through binding to Kir6.2 subunits. Examination of the kinetics of ATP sensitivity of $K_{\text{ATP}}$ channels in cardiac myocytes suggested that four ATP-binding sites were involved in the hallmark inhibition of the channel by ATP (Nichols et al., 1991). These results, together with the present demonstration of a tetrameric arrangement of Kir6.2 subunits, suggests a model in which ATP inhibition results from one ATP molecule binding to each of the four Kir6.2 subunits within the channel complex. The generation of K$^+$ channels by the complexing of Kir channel subunits and an ABC protein is unprecedented, but other ABC proteins, including human $\alpha$-glycoprotein and cystic fibrosis transmembrane conductance regulators are also known to be associated with separate ion channels (Higgins, 1995), and the bacterial Sap/Trk systems are ABC proteins associated with K$^+$ transport in Salmonella typhimurium and Escherichia coli (Parra-Lopez et al., 1994). It is an intriguing possibility that the stoichiometric association between SUR1 and Kir6.2 to generate the $K_{\text{ATP}}$ channel may give insight to the architecture of other ion channels generated by complexing ABC proteins with pore-forming subunits. Inagaki et al. (1997) have used a similar strategy with SUR1-Kir6.2 and SUR1-Kir6.2-Kir6.2 fusion constructs to indicate a 1:1 stoichiometry between SUR1 and Kir6.2.

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