Title
Single-channel characteristics of wild-type IKs channels and channels formed with two minK mutants that cause long QT syndrome.

Permalink
https://escholarship.org/uc/item/8nr0s1s3

Journal
The Journal of general physiology, 112(6)

ISSN
0022-1295

Authors
Sesti, F
Goldstein, SA

Publication Date
1998-12-01

DOI
10.1085/jgp.112.6.651

License
https://creativecommons.org/licenses/by/4.0/ 4.0

Peer reviewed
Single-Channel Characteristics of Wild-Type \( \text{I}_\text{Ks} \) Channels and Channels formed with Two MinK Mutants that Cause Long QT Syndrome

Federico Sesti and Steve A.N. Goldstein

From the Section of Developmental Biology and Biophysics, Departments of Pediatrics and Cellular and Molecular Physiology, Boyer Center for Molecular Medicine, Yale University School of Medicine, New Haven, Connecticut 06536-0812

Abstract \( \text{I}_\text{Ks} \) channels are voltage dependent and \( \text{K}^+ \) selective. They influence cardiac action potential duration through their contribution to myocyte repolarization. Assembled from minK and KvLQT1 subunits, \( \text{I}_\text{Ks} \) channels are notable for a heteromeric ion conduction pathway in which both subunit types contribute to pore formation. This study was undertaken to assess the effects of minK on pore function. We first characterized the properties of wild-type human \( \text{I}_\text{Ks} \) channels and channels formed only of KvLQT1 subunits. Channels were expressed in \textit{Xenopus laevis} oocytes or Chinese hamster ovary cells and currents recorded in excised membrane patches or whole-cell mode. Unitary conductance estimates were dependent on bandwidth due to rapid channel “flicker.” At 25 kHz in symmetrical 100-mM KCl, the single-channel conductance of \( \text{I}_\text{Ks} \) channels was \( \sim 16 \) pS (corresponding to \( \sim 0.8 \) pA at 50 mV) as judged by noise-variance analysis; this was fourfold greater than the estimated conductance of homomeric KvLQT1 channels. Mutant \( \text{I}_\text{Ks} \) channels formed with D76N and S74L minK subunits are associated with long QT syndrome. When compared with wild type, mutant channels showed lower unitary currents and diminished open probabilities with only minor changes in ion permeabilities. Apparently, the mutations altered single-channel currents at a site in the pore distinct from the ion selectivity apparatus. Patients carrying these mutant \textit{minK} genes are expected to manifest decreased \( \text{K}^+ \) flux through \( \text{I}_\text{Ks} \) channels due to lowered single-channel conductance and altered gating.

Keywords: KvLQT1 • heart • potassium • long QT syndrome • delayed rectifier

Introduction

Two voltage-dependent \( \text{K}^+ \) currents, \( \text{I}_\text{K} \) and \( \text{I}_\text{Kr} \), bring the heart beat to an end by repolarization of the myocardium (Sanguinetti and Jurkiewicz, 1990, 1991). The \( \text{K}^+ \) channel mediating \( \text{I}_\text{K} \) is formed by coassembly of two gene products, KvLQT1 and minK (Barhanin et al., 1996; Sanguinetti et al., 1996b). KvLQT1 is a 581 residue protein with six transmembrane segments and a classical pore-forming \( P \) domain (Wang et al., 1996b); human minK has just 129 residues and one transmembrane segment (Takumi et al., 1988). While many \( \text{K}^+ \)-selective pores are known to form by symmetrical alignment of four \( \text{P} \) loops around a central pathway (MacKinnon, 1991; Shen et al., 1994; Glowatzki et al., 1995; Doyle et al., 1998), the \( \text{I}_\text{K} \), channel pore incorporates minK residues, a non-\( \text{P} \) loop protein (Goldstein and Miller, 1991; Wang et al., 1996a; Tai and Goldstein, 1998). Six sites in the transmembrane stretch of minK are exposed in the \( \text{I}_\text{K} \) conduction pathway; two adjacent residues flank that portion of the pore that restricts transmembrane movement of \( \text{Na}^+ \), \( \text{Cd}^{2+} \), and \( \text{Zn}^{2+} \) ions; thus, position F55 is in contact with the external solution while G56 is accessible only to the cytosol (Tai and Goldstein, 1998). To delineate the attributes of this unique type of heteromeric ion conduction pathway, we sought to study \( \text{I}_\text{Ks} \) channels formed with wild-type and mutant minK subunits at the single-channel level.

Long QT syndrome (LQTS)\(^1\) is characterized by an abnormally long interval between the QRS complex (reflecting ventricular depolarization) and the T wave (resulting from ventricular repolarization) on the surface electrocardiogram; thus, a long QT interval indicates prolongation of the cardiac action potential. Through an unknown mechanism, this predisposes the myocardium to an arrhythmia called torsade de pointes that can degenerate to ventricular fibrillation and sudden death (Dumaine et al., 1996; Keating and Sanguinetti, 1996a; Roden et al., 1996; Ackerman, 1998). Mutations in cardiac \( \text{K}^+ \) channels that cause LQTS do so by decreasing \( \text{K}^+ \) efflux from affected cells, thereby

---

\(^1\)Abbreviations used in this paper: CHO, Chinese hamster ovary; LQTS, long QT syndrome.
delaying repolarization (Sanguinetti et al., 1995, 1996a; Wang, et al., 1996b; Splawski et al., 1997). LQTS, first associated with mutations in \(KvLQT1\) and \(HERG\) \(K^+\) channel genes has now been linked to mutations in the \(minK\) gene (Wang, et al., 1996b; Schulze-Bahr et al., 1997; Splawski, et al., 1997; Duggal et al., 1998). Inheritance of one mutant \(minK\) allele (encoding D76N or S74L \(minK\)) and one wild-type allele is associated with LQTS and deafness (Schulze-Bahr et al., 1997; Duggal et al., 1998). Splawski et al. (1997) have shown that these mutations decrease \(K^+\) currents by shifting activation of mutant \(I_{Ks}\) channels to more depolarized potentials and by accelerating their deactivation.

Here we compare the properties of single channels formed with \(KvLQT1\) subunits and those of \(I_{Ks}\) channels containing human \(minK\) and \(KvLQT1\) subunits. Thereafter, the effects of \(minK\) mutants on \(I_{Ks}\) channel function are evaluated. Homomorphic \(KvLQT1\) channels are found to have a single-channel conductance fourfold lower than wild-type channels but in conflict with the findings of others (Romey et al., 1995), a gift of Bernd Fakler (University of Tübingen, Tübingen, Germany), and cRNAs transcribed using T3 RNA polymerase and the mMessage mMachine™ kit (Ambion Inc., Austin, TX). Transcripts were quantified by spectroscopy and compared with control samples separated by agarose gel electrophoresis with control samples separated by agarose gel electrophoresis, digested with restriction sites. All sequences were confirmed by automated sequencing and by insertion of mutant gene fragments into translationally silent expression vectors. Transcripts were prepared immediately before injection with a calibrated pipette. Patch currents were recorded 2–3 d after injection using an Axopatch 200A amplifier (Axon Instruments, Foster City, CA), a Quadra 800 computer and ACQUIRE software (Instrutech, Great Neck, NY) and stored unfiltered on VHS tape. The data were filtered through a four-pole Bessel filter for analysis with TAC (Instrutech Corp.) and IGOR (WaveMetrics Inc., Lake Oswego, OR) software packages. Data are displayed without leak subtraction. Unless otherwise stated, the bath solution contained (mM) 100 KCl, 2 EGTA, and 10 HEPES, pH 7.5 with KOH (100 KCl solution). All experiments were performed at 22°C.

Data are presented as mean ± SEM with the number of patches indicated in parentheses. Two-electrode voltage clamp was performed with an Oocyte Clamp™ (Warner Instruments, Hamden, CT), an IBM pentium-based personal computer, and pCLAMP software (Axon Instruments). As reported previously, recorded currents show no evidence for contamination of human \(I_{Ks}\) channels by complexes formed between \(minK\) and the \(KvLQT1\) subunit endogenous to oocytes; the two channel types can be differentiated pharmacologically and no channels containing the endogenous subunit are expressed at times >36 h after cRNA injection (Tai and Goldstein, 1998).

\[ \sigma^2 = \frac{1}{n} \left( \frac{I - V_{rev}}{V_{rev}} \right)^2 + \gamma, \]

where \(\sigma^2\) is variance, \(I\) is mean macroscopic current obtained at different voltages, and \(n\) is number of channels in the patch. Mean macroscopic currents \((I_f)\) and variances at different voltages were determined from all point amplitude histograms constructed from 400 ms of test data fit to the gaussian function:

\[ \exp \left( \frac{- (I - I_f)^2}{2\sigma^2} \right). \]

Progressive reductions (from 400 to 80 ms) in the amount of data subjected to analysis altered measured variance by no more than 5%, indicating that changes in mean current over the sample period were well tolerated. This method differs from that employed by a companion paper by Yang and Sigworth (1998) in that we altered channel open probability \((P_o)\) by varying voltage, whereas they assessed changes in \(P_o\) at a single voltage over time. When the same data sets were evaluated by the two methods, the difference unitary current estimates was <15% \((n = 4\) patches). \(I_{Ks}\) currents are characterized by slow development and failure to reach saturation. Thus, all studies were performed using test depolarizations of many seconds duration from a holding voltage of \(-60\) mV to potentials from 10–80 mV. Currents elicited in the first 120 ms of each test pulse showed no time delay and were ascribed to non-channel dependent; these leak currents and their variances were subtracted. Conductance determinations were based on current and variance data collected near the end of the test depolarizations at times >5 s. This protocol was based on assessments of wild-type \(I_{Ks}\) and \(KvLQT1\) channel conductances at different times during test pulses; in each case, values

**Methods**

**Molecular Biology**

Human \(minK\) and \(KvLQT1\) genes were generous gifts from R. Swanson (Merck, West Point, PA) and M. Keating and M. Sanguinetti (University of Utah, Salt Lake City, UT), respectively (Hice et al., 1994; Sanguinetti et al., 1996a). Mutants of \(minK\) were produced by plaque-forming unit-based mutagenesis (QuikChange™ Site-Directed Mutagenesis Kit; Stratagene Inc., La Jolla, CA), followed by insertion of mutant gene fragments into translationally silent restriction sites. All sequences were confirmed by automated DNA sequencing. Genes were subcloned into pBF2 (Glowatzki et al., 1995), a gift of Bernd Fakler (University of Tübingen, Tübingen, Germany), and cRNAs transcribed using T3 RNA polymerase and the mMessage mMachine™ kit (Ambion Inc., Austin, TX). Transcripts were quantified by spectroscopy and compared with control samples separated by agarose gel electrophoresis and stained with ethidium bromide. Transient transfection of Chinese Hamster Ovary (CHO) cells was achieved by standard DEAE-Dextran/chloroquine/DMSO methodology using either pALTERMax (Promega Corp., Madison, WI) or pcDNA3 (Invitrogen Corp., San Diego, CA) plasmids carrying the genes for \(minK\), \(KvLQT1\), or green fluorescent protein.

**Electrophysiology**

*Xenopus laevis* oocytes were isolated and injected with 46 nl containing 5 ng \(KvLQT1\) with or without 1 ng \(minK\) cRNA. Mixtures of cRNAs were prepared immediately before injection with a calibrated pipette. Patch currents were recorded 2–3 d after injection using an Axopatch 200A amplifier (Axon Instruments, Foster City, CA), a Quadra 800 computer and ACQUIRE software (Instrutech, Great Neck, NY) and stored unfiltered on VHS tape. The data were filtered through a four-pole Bessel filter for analysis with TAC (Instrutech Corp.) and IGOR (WaveMetrics Inc., Lake Oswego, OR) software packages. Data are displayed without leak subtraction. Unless otherwise stated, the bath solution contained (mM) 100 KCl, 2 EGTA, and 10 HEPES, pH 7.5 with KOH (100 KCl solution). All experiments were performed at 22°C.

Data are presented as mean ± SEM with the number of patches indicated in parentheses. Two-electrode voltage clamp was performed with an Oocyte Clamp™ (Warner Instruments, Hamden, CT), an IBM pentium-based personal computer, and pCLAMP software (Axon Instruments). As reported previously, recorded currents show no evidence for contamination of human \(I_{Ks}\) channels by complexes formed between \(minK\) and the \(KvLQT1\) subunit endogenous to oocytes; the two channel types can be differentiated pharmacologically and no channels containing the endogenous subunit are expressed at times >36 h after cRNA injection (Tai and Goldstein, 1998).

**Noise Variance Analysis**

A variation of the method of stationary variance analysis (Sigworth and Zhou, 1992) was employed assuming linear dependence of single-channel current on membrane voltage; thus,

\[ i_{sc} = \gamma (V - V_{rev}), \]

where \(i_{sc}\) is single-channel current, \(\gamma\) unitary conductance, \(V\) membrane voltage, and \(V_{rev}\) reversal potential. Unless otherwise stated, the data were sampled at 80 kHz and filtered at 25 kHz at voltages of 10–80 mV in 10-mV steps, and unitary conductances determined by fitting to:

\[ \frac{\sigma^2}{I (V - V_{rev})} = \frac{1}{n} \left( \frac{I}{V - V_{rev}} \right)^2 + \gamma, \]

where \(\sigma^2\) is variance, \(I\) is mean macroscopic current obtained at different voltages, and \(n\) is number of channels in the patch. Mean macroscopic currents \((I_f)\) and variances at different voltages were determined from all point amplitude histograms constructed from 400 ms of test data fit to the gaussian function:

\[ \exp \left( \frac{-(I - I_f)^2}{2\sigma^2} \right). \]

Progressive reductions (from 400 to 80 ms) in the amount of data subjected to analysis altered measured variance by no more than 5%, indicating that changes in mean current over the sample period were well tolerated. This method differs from that employed by a companion paper by Yang and Sigworth (1998) in that we altered channel open probability \((P_o)\) by varying voltage, whereas they assessed changes in \(P_o\) at a single voltage over time. When the same data sets were evaluated by the two methods, the difference unitary current estimates was <15% \((n = 4\) patches). \(I_{Ks}\) currents are characterized by slow development and failure to reach saturation. Thus, all studies were performed using test depolarizations of many seconds duration from a holding voltage of \(-60\) mV to potentials from 10–80 mV. Currents elicited in the first 120 ms of each test pulse showed no time delay and were ascribed to non-channel dependent; these leak currents and their variances were subtracted. Conductance determinations were based on current and variance data collected near the end of the test depolarizations at times >5 s. This protocol was based on assessments of wild-type \(I_{Ks}\) and \(KvLQT1\) channel conductances at different times during test pulses; in each case, values

652 \(I_{Ks}\) Channels
Sesti and Goldstein were stable when determined at times longer than 3 s, presumably because large numbers of channels were active at later times and channel currents significantly greater in magnitude than non-channel noise (data not shown).

RESULTS

Unitary Conductance of Wild Type KvLQT1 and I\(_K\) Channels

Cloning the gene for KvLQT1 (Wang et al., 1996b) and its subsequent overexpression (Sanguinetti et al., 1996b) made possible studies of KvLQT1 and I\(_K\) channels in excised membrane patches. Fig. 1 shows a family of macroscopic KvLQT1 currents recorded in a giant inside-out patch excised from an oocyte injected 2 d earlier with KvLQT1 cRNA. In response to 6-s test pulses from \(-60\) mV to a variety of depolarized potentials, KvLQT1 channels showed fast activation of outward currents and fast deactivation of inward currents (Table I). Under the same conditions, I\(_K\) channels formed by coassembly of wild-type human minK and KvLQT1 channels (\(\square\), mean \(\pm\) SEM for three separate patches; see Table I). (D) Variance–current relationship of data in A (\(\square\), KvLQT1 channels) and B (\(\bullet\), I\(_K\) channels); the conductances calculated in these patches were 4.1 pS for A, and 15 pS for B.

![Figure 1](image-url)

**Figure 1.** Currents recorded in excised inside-out patches containing KvLQT1 or I\(_K\) channels in symmetrical 100-mM KCl solution. Patches were clamped to \(-60\) mV, and then depolarized for 6 s. (A) KvLQT1 channels depolarized to test potentials of 0–80 mV in 10-mV steps. (B) I\(_K\) channels depolarized to test potentials of \(-20\) to 80 mV in 10-mV steps. (C) Peak current–voltage relationship for KvLQT1 channels (\(\square\)) and I\(_K\) channels (\(\bullet\)), mean \(\pm\) SEM for three separate patches; see Table I. (D) Variance–current relationship of data in A (\(\square\), KvLQT1 channels) and B (\(\bullet\), I\(_K\) channels); the conductances calculated in these patches were 4.1 pS for A, and 15 pS for B.

### Table I

<table>
<thead>
<tr>
<th>Channel (No. of patches)</th>
<th>(\tau), pS</th>
<th>Activation (\tau)</th>
<th>(V_{1/2}), mV</th>
<th>(Z_0)</th>
<th>(I + 60) mV</th>
<th>(-60) mV</th>
<th>Deactivation (\tau)</th>
</tr>
</thead>
<tbody>
<tr>
<td>KvLQT1 (7)</td>
<td>4.0 ± 1.4</td>
<td>0.3 ± 0.1</td>
<td>(-6 ± 2)</td>
<td>2.0 ± 0.1</td>
<td>1.1 ± 0.2</td>
<td>0.4 ± 0.1</td>
<td></td>
</tr>
<tr>
<td>Wild-type I(_K) (15)</td>
<td>15.6 ± 4.3</td>
<td>5.4 ± 1.4</td>
<td>20 ± 3</td>
<td>1.0 ± 0.1</td>
<td>2.2 ± 0.3</td>
<td>1.2 ± 0.1</td>
<td></td>
</tr>
<tr>
<td>G55C I(_K) (3)</td>
<td>11.7 ± 1.9</td>
<td>5.6 ± 0.8</td>
<td>19 ± 1</td>
<td>1.1 ± 0.2</td>
<td>2.1 ± 0.1</td>
<td>1.2 ± 0.1</td>
<td></td>
</tr>
<tr>
<td>D76N + WT I(_K) (5)</td>
<td>8.1 ± 2.5</td>
<td>3.8 ± 1.3</td>
<td>28 ± 2</td>
<td>1.0 ± 0.1</td>
<td>2.5 ± 0.3</td>
<td>0.5 ± 0.1</td>
<td></td>
</tr>
<tr>
<td>D76N I(_K) (8)</td>
<td>4.8 ± 1.4</td>
<td>2.8 ± 0.4</td>
<td>37 ± 6</td>
<td>1.3 ± 0.2</td>
<td>3.2 ± 0.5</td>
<td>0.3 ± 0.1</td>
<td></td>
</tr>
<tr>
<td>S74L + WT I(_K) (5)</td>
<td>13.6 ± 1.5</td>
<td>6.6 ± 1.8</td>
<td>34 ± 4</td>
<td>1.1 ± 0.2</td>
<td>2.1 ± 0.3</td>
<td>1.0 ± 0.2</td>
<td></td>
</tr>
<tr>
<td>S74L I(_K) (7)</td>
<td>8.9 ± 1.5</td>
<td>7.8 ± 1.8</td>
<td>43 ± 2</td>
<td>1.6 ± 0.1</td>
<td>2.1 ± 0.3</td>
<td>0.8 ± 0.1</td>
<td></td>
</tr>
<tr>
<td>D76N + S74L I(_K) (5)</td>
<td>6.6 ± 1.2</td>
<td>4.0 ± 0.3</td>
<td>39 ± 3</td>
<td>1.3 ± 0.2</td>
<td>2.8 ± 0.4</td>
<td>0.4 ± 0.1</td>
<td></td>
</tr>
</tbody>
</table>

Unitary conductance was determined by sampling at 80 kHz and filtering at 25 kHz at test voltages of 10–80 mV in 10-mV steps (METHODS). Activation kinetics were estimated from currents measured in macropatches during a 6-s depolarization to 60 mV from a holding voltage of \(-60\) mV in symmetrical 100-mM KCl solution; currents were fit by a single exponential function: \(I = I_0 e^{-t/\tau}\). Half maximal activation potentials \((V_{1/2})\) and equivalent valences \((Z_0)\) were determined by comparing maximal currents at 60 and \(-60\) mV in symmetrical 100-mM KCl solution. Deactivation traces were fit with the function: \(I = I_0 e^{-t/\tau}\). The values for KvLQT1 and I\(_K\) correspond to unitary currents at 50 mV and 25 kHz of 0.20 and 0.78 pA, respectively.
subunits exhibited slower activation rates, slower deactivation rates (Fig. 1 B and Table I) and increased inward rectification (Fig. 1 C).

To assess the conductive properties of KvLQT1 and IKs channels, a variation of stationary variance analysis (Sigworth and Zhou, 1992) was performed (see methods). Unitary conductances (g) were determined by fitting collected data to Eqs. 1–3 (Fig. 1 D). The average value for KvLQT1 channels was $4.0 \pm 1.4 \ pS$ (mean $\pm$ SEM for seven patches), while that for IKs channels was $15.6 \pm 4.3 \ pS$ (mean $\pm$ SEM for 15 patches, Table I); these correspond to unitary channel currents of 0.20 and 0.78 pA, respectively, at 50 mV. Thus, homomeric KvLQT1 channels passed a unitary current roughly fourfold smaller than IKs channels containing wild-type human minK and KvLQT1 subunits.

These estimates suggested it might be possible to observe IKs channels directly in inside-out patches. Despite many similarities in single-channel activity and macroscopic currents, it was extremely difficult to discern discrete gating transitions of human IKs channels. Thus, single channels were inactive at $-80 \ mV$, activated slowly at 40 mV, and, upon repolarization to $-80 \ mV$, were active briefly before deactivation (Fig. 2 A). While expansion of the time scale made it possible to appreciate channel activity in comparison to baseline, the “flickery” nature of the channel precluded observation of transitions between closed and open states (Fig. 2 B).

Our inability to discern single-channel transitions made it imperative to confirm the activity under scrutiny truly resulted from IKs channel function. Support for this identification came first from similarity of microscopic and macroscopic activation kinetics, as judged from comparison of patches with only a few channels (Fig. 3 A) assembled in a cumulative fashion (Fig. 3 B) and giant patches with many channels (Fig. 3 C). A second line of evidence was the observation that single channels in patches passed K$^+$ but not Na$^+$ ions, as ex-

![Figure 2](image-url)
pected of $I_{Ks}$ channels (data not shown). A third experimental support came from studies of a site-directed minK mutant that forms $I_{Ks}$ channels sensitive to extracellular Cd$^{2+}$ (Tai and Goldstein, 1998). Previous whole cell studies showed that wild-type $I_{Ks}$ channels were insensitive to Cd$^{2+}$, while channels formed with G55C human minK (altering glycine 55 to cysteine) were blocked by the metal through a pore-occlusion mechanism (Tai and Goldstein, 1998). Here, single $I_{Ks}$ channels containing G55C minK were studied in excised patches. The G55C $I_{Ks}$ channels appear essentially unaltered from wild type in their gating and unitary conductance (Fig. 4, Table I). However, while $I_{Ks}$ channels carrying wild-type minK were Cd$^{2+}$ insensitive (Fig. 4 A), single channels containing G55C minK were reversibly inhibited by 1 mM Cd$^{2+}$ (Fig. 4, B–D). $I_{Ks}$ channels containing G55C minK, but not wild-type minK, were also sensitive to blockade by external Zn$^{2+}$ (data not shown). The similarities of microscopic and macroscopic currents in their gating kinetics, ion selectivity, current rectification, and pharmacology (Figs. 1–4) argued strongly that the rapidly flickering channels under study in excised patches were $I_{Ks}$ channels.

**Gating and Current Rectification of Wild-Type KvLQT1 and $I_{Ks}$ Channels**

Whole-cell studies previously showed that KvLQT1 currents activated rapidly, reached saturation, and deactivated rapidly, whereas $I_{Ks}$ currents activated slowly, did not saturate, and deactivated slowly (Wang et al., 1996a; Splawski et al., 1997; Tai et al., 1997; Tzounopoulos et al., 1998). Gating behavior of single channels in excised patches reflected these whole-cell findings. Thus, homomeric KvLQT1 channels activated ~15- and deactivated ~4-fold faster than $I_{Ks}$ channels (Table I). These differences were apparent to the eye (Fig. 1, A and B) and were quantified by comparison of time constants determined from exponential fits to the data (Fig. 5, A and B, Table I). In agreement with earlier re-

---

**Figure 3.** Activation of wild-type $I_{Ks}$ channels in inside-out patches in symmetrical 100-mM KCl solution. (A) Four traces from a patch showing channels opening after depolarization to 40 mV (arrow) from a holding voltage of −80 mV; 600 ms of the 6-s pulse is shown; data sampled at 4 kHz and filtered at 1 kHz. (B) Cumulative current of 30 sweeps of the patch shown in A. (C) Single trace of another patch with many channels by the protocol in B.
ports of whole-cell current activation (Sanguinetti et al., 1996b), I\textsubscript{k}s channels in excised patches were less sensitive to voltage than KvLQT1 channels, showing a half-maximal activation potential 25 mV more positive and a normalized open probability–voltage relationship that was more shallow (Fig. 5 A, Table I). Normalized tail current plots make apparent that KvLQT1 channels deactivated more rapidly than wild-type I\textsubscript{k}s channels (Fig. 5 B, Table I).

Unitary Conductance of I\textsubscript{k}s Channels Containing MinK Mutants that Cause LQTS

Recently, mutant genes encoding D76N minK and S74L minK were identified in patients with LQTS (Splawski et al., 1997; Duggal et al., 1998). In some patients, LQTS was inherited in an autosomal dominant fashion and affected individuals possessed one normal and mutant allele. Thus, we first studied the influence of minK mutants on I\textsubscript{k}s function using oocytes injected with cRNAs for KvLQT1 subunits and a 1:1 mixture of wild-type and mutant minK subunits. Thereafter, to study a single population of channels, oocytes were injected with cRNAs for KvLQT1 subunits and a minK mutant in the absence of wild-type minK. The current in Fig. 6 A was recorded in an excised patch from a cell injected with cRNAs for KvLQT1, D76N minK, and wild-type minK (to approximate the subunit composition of myocytes of individuals with D76N minK-associated LQTS). In this case, the conductance was a weighted average from a mixture of I\textsubscript{k}s channels (some with only wild-type minK, some only mutant minK, others with both). The apparent conductance was 8.1 ± 2.5 pS (Table I, Fig. 6 D). Clearly, this experiment did not offer insight into the properties of any individual channel type in the patch.

Fig. 6 B shows the current from oocytes expressing I\textsubscript{k}s channels formed with KvLQT1 and D76N minK subunits. Current–variance analyses estimated the unitary conductance of these channels to be 4.8 ± 1.4 pS (Table I, Fig. 6 C). D76N I\textsubscript{k}s channels also showed greater inward rectification than wild-type channels (Table I, Fig. 6 D).

Fig. 6 E shows the effect of progressively increasing the relative amount of D76N minK cRNA with respect to wild type. Increasing the proportion of D76N minK subunits caused a decreased average unitary conductance. This indicated that D76N I\textsubscript{k}s channels and mixed D76N/wild-type I\textsubscript{k}s channels were both functional and that more than one minK subunit was present in each channel complex. A quantitative consideration of this data in relation to subunit stoichiometry is below (DISCUSSION).
A second group of patients with LQTS carries one mutant gene encoding S74L minK subunits and one wild-type gene. Currents resulting from coexpression of \( \text{KvLQT1} \) cRNA and a 1:1 mixture of S74L and wild-type \( \text{minK} \) cRNAs gave an average conductance of 13.6 ± 1.5 pS (Table I, Fig. 7, A and D). As before, this was an average conductance for a number of channel types. Analyses carried out on oocytes injected with just \( \text{KvLQT1} \) and S74L \( \text{minK} \) cRNAs showed S74L \( I_{\text{K_s}} \) channels to have a unitary conductance of 8.9 ± 1.5 pS (Table I, Fig. 7, B and D). As before, this was an average conductance for a number of channel types. Analyses carried out on oocytes injected with just \( \text{KvLQT1} \) and S74L \( \text{minK} \) cRNAs showed S74L \( I_{\text{K_s}} \) channels to have a unitary conductance of 8.9 ± 1.5 pS (Table I, Fig. 7, B and D). Unlike D76N \( I_{\text{K_s}} \) channels, S74L channels showed current rectification similar to wild type (Table I, Fig. 7 C).

A third group of patients (with Jervell and Lange-Nielsen Syndrome or Romano-Ward Syndrome) manifest both LQTS and hearing defects (Schulze-Bahr et al., 1997; Duggal et al., 1998); these individuals carry two mutant alleles of \( \text{minK} \). When oocytes were coinjected with cRNAs for \( \text{KvLQT1} \) and a 1:1 mixture of D76N and S74L \( \text{minK} \) cRNAs (as seen in some of these individuals), the resulting channels had an average unitary conductance of 6.6 ± 1.2 pS (Table I, Fig. 8, A and C). These channel mixtures showed average conductance and rectification properties intermediate in phenotype to pure D76N and S74L \( I_{\text{K_s}} \) channels (Table I, Fig. 8, B and C).

Gating of \( I_{\text{K_s}} \) Channels Containing \( \text{minK} \) Mutants

While activation of wild-type \( I_{\text{K_s}} \) channels proceeded about half as fast as D76N channels and twice as fast as S74L channels, all \( I_{\text{K_s}} \) channels activated more slowly than homomeric \( \text{KvLQT1} \) channels (Table I). Compared with wild-type \( I_{\text{K_s}} \) channels, the half-maximal activation potential for D76N and S74L channels was shifted to more depolarized voltages (Fig. 9 A, Table I). Deactivation of D76N \( I_{\text{K_s}} \) channels was significantly faster and S74L channels moderately faster than wild-type channels (Fig. 9 B, Table I), in good agreement with an earlier report (Splawski et al., 1997). Patches from cells expressing both D76N and S74L \( \text{minK} \) showed activation kinetics that were intermediate to those of pure D76N or S74L \( I_{\text{K_s}} \) channels (Table I), but speeded deactivation kinetics similar to D76N channels (Fig. 9 B, Table I). These results were consistent with the idea that \( \text{KvLQT1} \) could assemble with either D76N or S74L \( \text{minK} \) subunits to form functional mutant \( I_{\text{K_s}} \) channels characterized by small unitary currents, altered gating kinetics and marked inward rectification (Table I).

Selectivity of \( I_{\text{K_s}} \) Channels Containing \( \text{minK} \) Mutants that Cause LQTS

As coassembly altered unitary conductances, we sought evidence for changes in ion selectivity, another pore-determined channel attribute. The relative permeabilities of monovalent cations similar to K⁺ ion were assessed using biionic reversal potential measurements in whole-cell configuration. Wild-type \( I_{\text{K_s}} \) channels showed the same permeability series as \( \text{KvLQT1} \) channels (K⁺ > Rb⁺ > NH₄⁺ > Cs⁺ > Li⁺, Na⁺), but a slightly greater selectivity against NH₄⁺ and Cs⁺ (Table II), as found by others (Wollnik et al., 1997). While mutation

---

**Figure 5.** Gating attributes of \( \text{KvLQT1} \) and \( I_{\text{K_s}} \) channels in excised patches. (A) Normalized open probabilities \( (I/IMax) \) for \( \text{KvLQT1} \) channels (○) and \( I_{\text{K_s}} \) channels (■). Peak tail current at −60 mV was measured after a 6-s prepulse to voltages of −20 to 80 mV. The curves are normalized to 80-mV prepulse value. Theoretical lines are constructed with the Boltzmann function: \( I/IMax = 1/(1 + \exp[z(V_1/2 - V)/kT]) \), where \( z \) is the equivalent valence, \( e \) the elementary charge, \( V_1/2 \) the voltage at which the curve is half maximal (see Table I), \( k \) the Boltzmann constant, and \( T \) the absolute temperature; for \( \text{KvLQT1} \) channels, \( z = 2.0 ± 0.1 \); for \( I_{\text{K_s}} \) channels, \( z = 1.0 ± 0.1 \). (B) Normalized tail currents in a patch at −60 mV. Currents were elicited by a 6-s pulse to 80-mV voltage (inset). Theoretical lines are single exponential fits to the deactivation time course; results from many patches are in Table I.
of human minK position G56 increased permeation of \( I_{Ks} \) channels by \( Na^+ \) ion fivefold (Tai and Goldstein, 1998), and mutation of rat minK position F55 increased permeation by \( NH_4^+ \) and \( Cs^+ \) threefold (Goldstein and Miller, 1991), neither D76N nor S74L channels showed changes in ion discrimination (Table II). Thus, these LQTS-associated mutations at positions S74 and D76 altered single-channel conductance (and gating) without effecting function of the ion selectivity filter.

**Unitary Conductance of Wild-Type KvLQT1 and \( I_{Ks} \) Channels in CHO Cells**

To ascertain whether unitary current magnitudes varied when channels were in mammalian cells rather than oocytes, we reassessed the current variance of KvLQT1 and \( I_{Ks} \) channels expressed in CHO cells (Fig. 10). Unitary conductances in CHO cells were 2.3 ± 0.6 pS (\( n = 4 \) cells) and 7.9 ± 1.3 pS (\( n = 3 \) cells) for KvLQT1 and \( I_{Ks} \) channels, respectively; this corresponds to unitary currents of 0.20 and 0.79 pA with a 50-mV driving force, the same as we found for the channels in oocytes at 25 kHz (Table I, Fig. 1).

**Discussion**

To gain insight into the mechanisms underlying their function, we have analyzed the currents recorded from channels formed only with KvLQT1 subunits and those coassembled from human minK and KvLQT1 subunits to form \( I_{Ks} \) channels. Noise-variance analyses showed the mixed channel complex to have a unitary conductance fourfold greater than that of homomeric KvLQT1 channels (Table I). As expected, \( I_{Ks} \) channels were found to activate and deactivate more slowly, to be less sensitive to voltage, and to select slightly more effectively against \( Cs^+ \) and \( NH_4^+ \) ions than KvLQT1 channels (Tables I and II). When mutant \( I_{Ks} \) channels were formed with two minK subunits associated with LQTS
Figure 7. Current–variance relationships of I\textsubscript{Ks} channels formed with wild-type and S74L minK subunits. (A) Family of currents recorded in a patch excised from an oocyte coinjected with KvLQT1 cRNA and a 1:1 mixture of wild-type and S74L minK cRNAs. The patch was clamped at −60 mV and depolarized from −20 to 100 mV in steps of 10 mV. (B) Family of currents recorded from oocytes coinjected only with KvLQT1 and S74L minK cRNAs. The protocol was as in A with steps from −20 to 90 mV. (C) Current–voltage relationship of peak tail currents measured for S74L I\textsubscript{Ks} channels, mean ± SEM for three patches. (D) Current–variance relationships computed from the patches in A (▽) and B (▼); the conductances calculated in these patches were 14 and 8.7 pS, respectively.

Figure 8. Current–variance relationships of I\textsubscript{Ks} channels formed with S74L and D76N minK subunits. (A) Family of currents recorded from a patch excised from an oocyte coinjected with KvLQT1 cRNA and a 1:1 mixture of D76N and S74L minK cRNAs. The patch was clamped at −60 mV and depolarized from −20 to 100 mV in 10-mV steps. (B) Current–voltage relationship of peak tail currents measured for channels, mean ± SEM for three separate patches. (C) Current–variance relationship computed from the patch in A; the conductance calculated in this patch was 6.0 pS.
Single IK<sub>s</sub> Channels and compared with wild-type IK<sub>s</sub> channels, they showed lower unitary conductances, a requirement for larger depolarizations to activate, and more rapid deactivation but no significant change in the relative permeability of monovalent cations (Tables I and II).

Our results generally agree with those of Yang and Sigworth (1998), who found coassembly of minK and KvLQT1 to increase single-channel conductance approximately fivefold. While our conductance values are higher than theirs (~16 vs. 5 pS for human IK<sub>s</sub> channels in oocytes at 25 kHz), this can largely be understood by their use of 7 mM external KCl where we employ symmetrical 100 mM KCl solutions. Indeed, unitary currents should be similar with strong depolarization despite external KCl differences and this was found; Yang and Sigworth (1998) estimated a unitary current for human IK<sub>s</sub> channels at 50 mV and 25 kHz of 0.6 ± 0.2 pA where we found 0.8 ± 0.2 pA (Table I). These results stand in stark contrast to a report that KvLQT1 channels in mammalian cells have a unitary conductance 13-fold larger than IK<sub>s</sub> channels (which were attributed a value of just 0.076 pA) (Romey et al., 1997). To ascertain whether this discrepancy was due to expression of the channels in mammalian cells, we performed current–variance analysis using CHO cells. Consistent with our findings in oocytes (Table I), unitary IK<sub>s</sub> currents in CHO cells were ~0.8 pA at 25 kHz, fourfold larger than KvLQT1 channels (Fig. 10). There was also reasonable agreement, considering species and methodologic differences, between the conductance of native IK<sub>s</sub> channels studied in guinea pig cardiac myocytes (Walsh et al., 1991) or stria vascularis (Shen and Marcus, 1998) and our estimate for human channels, as they were ~50% smaller and ~20% larger, respectively.

A potential source of error in our measurements was contamination by endogenous currents. This was especially important to consider, as human IK<sub>s</sub> channels were flickery, making it difficult to discern single-channel transitions. However, the channels under study were firmly identified based on their presence in oocytes only after co-injection of minK and KvLQT1 cRNAs, similarity of their gating kinetics to macroscopic IK<sub>s</sub> currents, their selectivity for K<sup>+</sup> ions, and their sensitivity to block by transition metals when formed with G55C minK, in keeping with earlier macroscopic studies (Tai and Goldstein, 1998).

Another source of concern, given the volatile nature of the channel (and our inability to compute amplitude histograms), was the validity of current–variance analyses. In particular, could noise analysis differentiate changes in channel gating from changes in unitary conductance? Could KvLQT1 channels have a conductance as large as IK<sub>s</sub> channels, but flicker more rapidly? To address this issue, we studied the dependence of the variance on experimental bandwidth. According to Eq. 2, the variance and unitary conductance depend on filter frequency in the same fashion. Thus, increasing the bandwidth should cause similar changes in the variance and conductance until the bandwidth increases above a level sufficient to record complete open–closed transitions; at this point, the conductance should “level-off” and approach its maximal “true” value. Fig. 11 shows such an experiment in which data was sampled at 80 kHz, noise-variance was measured, and unitary con-
ductance calculated as a function of bandwidth from 0.25 to 25 kHz. Above 10 kHz, the calculated conductances for KvLQT1 channels and wild-type and mutant IKs channels appeared to approach maximal values. Moreover, the conductances appeared to rise in parallel, suggesting that flicker kinetics were the same in the four channel types studied (homomeric KvLQT1, wild-type IKs, D76N IKs, and S74L IKs channels). This conclusion is also supported by observation that single IKs channels (Figs. 2–4) exhibit the same apparent single-channel conductance as was determined by noise analysis at the analogous bandwidth. Thus, IKs channel activity at 1 kHz (Figs. 2–4) showed a unitary current of ~0.3 pA, a bandwidth where the current–variance

<table>
<thead>
<tr>
<th>Channel</th>
<th>Na⁺</th>
<th>Li⁺</th>
<th>Ca⁺</th>
<th>NH₄⁺</th>
<th>Rb⁺</th>
<th>K⁺</th>
</tr>
</thead>
<tbody>
<tr>
<td>KvLQT1</td>
<td>−105 ± 9</td>
<td>−97 ± 7</td>
<td>−52 ± 6</td>
<td>−38 ± 5</td>
<td>−14 ± 5</td>
<td>−1 ± 1</td>
</tr>
<tr>
<td>Wild-type IKs</td>
<td>−97 ± 9</td>
<td>−92 ± 9</td>
<td>−67 ± 5</td>
<td>−51 ± 5</td>
<td>−17 ± 6</td>
<td>2 ± 3</td>
</tr>
<tr>
<td>D76N IKs</td>
<td>−95 ± 8</td>
<td>−95 ± 8</td>
<td>−48 ± 4</td>
<td>−47 ± 6</td>
<td>−7 ± 6</td>
<td>−1 ± 4</td>
</tr>
<tr>
<td>S74L IKs</td>
<td>−100 ± 7</td>
<td>−101 ± 8</td>
<td>−54 ± 6</td>
<td>−42 ± 4</td>
<td>−7 ± 4</td>
<td>4 ± 4</td>
</tr>
</tbody>
</table>

Reversal potentials were measured with 100 mM of the indicated cation in the external solution; whole cell currents were elicited by a 6s command pulse to 30 mV from a holding voltage of −80 mV, followed by repolarization to various test potentials and the slopes of tail currents calculated. Permeability ratios (in parenthesis) are calculated according to $P_K/P_X = \exp(-\Delta V_{rev}/RT)$, where $R$, $T$, and $F$ have their usual meanings and $P_K$ and $P_X$ are the permeability of $K^+$ and the test ion, respectively; this assumes that $K^+$ is the only permeant ion inside the oocyte. Values are mean ± SEM for four to nine oocytes.

**Table II**

| Biionic Reversal Potentials and Relative Permeabilities of Wild-Type and Mutant IKs Channels |
|---------------------------------|-----------------|-----------------|-----------------|-----------------|--------------------|
| Channel                        | Na⁺             | Li⁺             | Ca⁺             | NH₄⁺            | Rb⁺               |
| KvLQT1                         | −105 ± 9        | −97 ± 7         | −52 ± 6         | −38 ± 5         | −14 ± 5           |
| Wild-type IKs                  | −97 ± 9         | −92 ± 9         | −67 ± 5         | −51 ± 5         | −17 ± 6           |
| D76N IKs                       | −95 ± 8         | −95 ± 8         | −48 ± 4         | −47 ± 6         | −7 ± 6            |
| S74L IKs                       | −100 ± 7        | −101 ± 8        | −54 ± 6         | −42 ± 4         | −7 ± 4            |

**Figure 10.** Currents recorded in CHO cells expressing KvLQT1 or IKs channels by whole-cell configuration. Cells were clamped to −80 mV, and then depolarized for 6 s to test potentials from −60 to 10 mV (KvLQT1) or −60 to 50 mV (IKs) in 10-mV steps. The solutions were (mM): bath: 20 KCl, 100 NaCl, 1 CaCl₂, 2 MgCl₂, 10 HEPES, pH 7.5; pipette: 120 KCl, 2 MgCl₂, 2 EGTA, 10 HEPES, pH 7.5. (A) KvLQT1 channels. (B) IKs channels. (C) Variance–current relationships of data in A (□, KvLQT1 channels) and B (■, IKs channels). The calculated conductances in these patches were 2.2 and 8.2 pS, respectively.
and that the distribution, half-life, and assembly of IKs probability with different minK subunit compositions, and variance as:

\[ \minK \text{ subunits in a complex and express the current can estimate an upper limit for the number} \]

\[ (\text{where } \minK \text{ is the cRNA fraction that is mutant). After substituting Eqs. 4 and 5 into Eq. 2, the variance–mean fit yields an apparent single-channel conductance:} \]

\[ I = I_{\minK} \left[ \frac{f_{\text{wt}} \gamma_{\text{wt}} - f_{D76N} \gamma_{D76N}}{\gamma_{\text{wt}}} \right] \]

\[ \text{and} \]

\[ \sigma^2 = \sigma_{\minK}^2 \left[ \frac{f_{\text{wt}} \gamma_{\text{wt}}^2 - f_{D76N} \gamma_{D76N}^2}{\gamma_{\text{wt}}^2} \right] \]

where \( f_{\text{wt}} = [\text{wt}]^n \) is the fraction of injected cRNA that is wild type and \( f_{D76N} = 1 - [\text{wt}]^n \) is the cRNA fraction that is mutant. While our estimates argue against a very large number of minK subunits in each complex (Tzounopoulos et al., 1995), the assumptions underlying the models are unproven. It seems reasonable to conclude that \( \minK \) channels contain at least two but not more than four minK subunits.

There were three effects of forming IKs channels with mutant minK subunits: single-channel conductance was decreased, \( V_{1/2} \) for activation was shifted to more depolarized voltages, and deactivation kinetics were faster. Each of these effects decreased \( K^+ \) flux through \( \minK \) channels, the same mechanism by which mutations in other \( K^+ \) channels have been demonstrated to induce LQTS (Sanguinetti et al., 1995, 1996a,b; Keating and Sanguinetti, 1996a,b). The result of decreased \( K^+ \) flux is to slow phase 3 repolarization, prolong the cardiac action potential, and increase the QT interval measured on a surface electrocardiogram (Rodén et al., 1996).

The data also provide some indirect information about the structure of the \( \minK \) channel pore. In the past, we found that rat minK residues 55, 56, 57, and 59 (in the midst of single transmembrane segment) were exposed in the ion conduction pathway of \( \minK \) channels (Tai and Goldstein, 1998); these sites appeared to reside in close proximity to the ion selectivity filter as 55 and 56 were accessible only from the external solution, 57 and 59 were accessible only from the cytosol, and changes at 56 altered \( Na^+ \) permeability. Here we find that the mutation of residue 74 or 76 changes single-channel conductance without altering ion selectivity. This suggests these two sites are in the pore but reside at a distance from the selectivity-determining apparatus.
REFERENCES


