N-Terminal Arginines Modulate Plasma-Membrane Localization of Kv7.1/KCNE1 Channel Complexes

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Abstract

Background and Objective: The slow delayed rectifier current (I\textsubscript{K\textsubscript{s}}) is important for cardiac action potential termination. The underlying channel is composed of Kv7.1 \textalpha-subunits and KCNE1 \textbeta-subunits. While most evidence suggests a role of KCNE1 transmembrane domain and C-terminus for the interaction, the N-terminal KCNE1 polymorphism 38G is associated with reduced I\textsubscript{K\textsubscript{s}} and atrial fibrillation (a human arrhythmia). Structure-function relationship of the KCNE1 N-terminus for I\textsubscript{K\textsubscript{s}} modulation is poorly understood and was subject of this study.

Methods: We studied N-terminal KCNE1 constructs disrupting structurally important positively charged amino-acids (arginines) at positions 32, 33, 36 as well as KCNE1 constructs that modify position 38 including an N-terminal truncation mutation. Experimental procedures included molecular cloning, patch-clamp recording, protein biochemistry, real-time-PCR and confocal microscopy.

Results: All KCNE1 constructs physically interacted with Kv7.1. I\textsubscript{K\textsubscript{s}} resulting from co-expression of Kv7.1 with non-atrial fibrillation '38S' was greater than with any other construct. Ionic currents resulting from co-transfection of a KCNE1 mutant with arginine substitutions ('38G-3xA') were comparable to currents evoked from cells transfected with an N-terminally truncated KCNE1-construct ('ΔA-38'). Western-blot analysis of plasma-membrane preparations and confocal images consistently showed a greater amount of Kv7.1 protein at the plasma-membrane in cells co-transfected with the non-atrial fibrillation KCNE1-38S than with any other construct.

Conclusions: The results of our study indicate that N-terminal arginines in positions 32, 33, 36 of KCNE1 are important for reconstitution of I\textsubscript{K\textsubscript{s}}. Furthermore, our results hint towards a role of these N-terminal amino-acids in membrane representation of the delayed rectifier channel complex.

Introduction

The slow delayed rectifier current (I\textsubscript{K\textsubscript{s}}) is important for cardiac repolarization. One of the current’s key functions is to prevent excessive action potential prolongation during adrenergic stimulation. It represents an important constituent of the “repolarization reserve” [1]. The single-transmembrane segment \textbeta-subunit KCNE1 modulates the function of the six-transmembrane segment, pore-forming \textalpha-subunit Kv7.1 [2,3]. Within the heart KCNE1 is the major interacting \textbeta-subunit associating with Kv7.1. The interaction between these proteins determines I\textsubscript{K\textsubscript{s}} properties and modulates current characteristics (eliminating ionic current inactivation, increasing unitary conductance and slowing activation) [4,5]. Most of the interactions underlying this modulation have been localized to the transmembrane domain and the C-terminus for the interaction and ionic current modulation remains largely unexplored.

An N-terminal single nucleotide polymorphism results in an amino-acid change (G38S) in an unconserved KCNE1 position and is highly prevalent (Fig. 1A). It can be found in up to ~50% of individuals in different ethnicities [10,11]. A population study described an association of the common allele KCNE1-38G with atrial fibrillation, a highly prevalent human arrhythmia. Odds ratios for atrial fibrillation occurrence were 2.16 with one ‘38G’ allele and 3.58 with two ‘38G’ alleles [10]. The atrial fibrillation-associated KCNE1-38G allele results in reduced I\textsubscript{K\textsubscript{s}} density possibly due to impaired membrane trafficking of I\textsubscript{K\textsubscript{s}} channels [12]. The underlying structure-function correlation of this N-terminal region has not yet been studied.

The present study examined the hypothesis that arginines in position 32, 33 and 36 within the KCNE1 N-terminus are linker and subsequently modulates ion channel gating [9]. The role of the KCNE1 N-terminus for delayed rectifier channel interaction and ionic current modulation remains largely unexplored.
Expression of KCNE1 mutants and interaction with Kv7.1

Results

Expression of KCNE1 mutants and interaction with Kv7.1

Analysis of KCNE1 protein expression by Western blotting revealed bands at the apparent molecular weight of ~17 kD for both ‘38S’ and ‘38G’. The molecular weight of ‘Δ1-38’ was slightly smaller due to the truncation and ‘linker’ was heavier due to four additional amino-acids. ‘38S-3xA’ and ‘38G-3xA’ appeared slightly smaller than ‘38S’ and ‘38G’. Fig. 2A shows crude membrane preparations from cells transfected with respective flag-tagged KCNE1 constructs illustrating similar overall protein expression. All constructs effectively co-immunoprecipitated with Kv7.1 indicating physical interaction between Kv7.1 channel complexes and for IKs modulation.

Electrophysiological studies

Ionic currents were recorded from HEK and CHO cells to exclude cell-line specific effects, results were comparable (not shown). Figs. 3A–F illustrate representative currents from HEK and CHO cells transfected with respective flag-tagged KCNE1 constructs. Ionic currents were recorded from HEK and CHO cells. Analysis of KCNE1 protein expression by Western blotting revealed bands at the apparent molecular weight of ~17 kD for both ‘38S’ and ‘38G’. The molecular weight of ‘Δ1-38’ was slightly smaller due to the truncation and ‘linker’ was heavier due to four additional amino-acids. ‘38S-3xA’ and ‘38G-3xA’ appeared slightly smaller than ‘38S’ and ‘38G’. Fig. 2A shows crude membrane preparations from cells transfected with respective flag-tagged KCNE1 constructs illustrating similar overall protein expression. All constructs effectively co-immunoprecipitated with Kv7.1 indicating physical interaction between Kv7.1 channel complexes and for IKs modulation.

Fig. 1. Schematic of KCNE1 and constructs. This figure schematically illustrates KCNE1 structure and mutants used in the present study. A, left, alignment of KCNE1 sequences from various mammalian species. Grey underlines conserved residues. Glycine at position 38 is not strongly conserved among species providing no first-glance evidence for evolutionary importance. A, right, schematic of KCNE1 at the membrane with the N-terminal part oriented towards the cell exterior and C-terminus towards the cytosol. B, schematic of KCNE1 N-terminal constructs and mutations created for the present study. Ten N-terminal amino-acids (AA) illustrate differences between KCNE1 constructs. Position 38 carries a glycine in the prevalent single nucleotide polymorphism. One of the constructs contained an N-terminal truncation (‘Δ1-38’), another one (‘linker’) replaced position 38 by 5 alanines. Additionally, three positively-charged arginines at positions 32, 33 and 36 have been exchanged for alanines in order to probe the role of these AA in KCNE1 function.
Figure 2. Immunodetection of heterologously expressed constructs. Immunodetection of flag-tagged KCNE1 constructs. A, crude membrane preparations from HEK cells transiently transfected with respective KCNE1 constructs. Actin (∼42 kD) is shown as loading control. B, effective co-immunoprecipitation (IP) occurred for Kv7.1 with all flag-tagged KCNE1 constructs. The upper blot shows protein samples from HEK cells precipitated by anti-flag and bands detected by anti-Kv7.1. The lower blot shows samples precipitated by anti-Kv7.1 and bands detected by anti-flag antibody (n = 2 experiments each). C shows respective confocal images of KCNE1 subunits expressed without Kv7.1. Bars represent 5 μm. Images are representative of at least 5 different experiments. NT – non-transfected control, IP – immunoprecipitation.

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Figure 3. Electrophysiological properties. A–F, representative ionic currents elicited by whole-cell patch-clamp with the protocol shown in inset A. G, current-voltage relations of mean ± SEM depolarization induced activating step-current densities from cells transfected with Kv7.1 plus various KCNE1 constructs. H, mean ± SEM current-voltage relationships of repolarization induced tail-current densities. P-values are shown for currents recorded from cells transfected with Kv7.1+’38S’ vs. all other constructs. The remainder of constructs did not lead to significantly different current sizes compared with ‘38G’ (P = n.s.). TP – test potential.

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constructs in comparison to ‘38G’, we evaluated the effect of KCNE1 constructs on Kv7.1 and KCNE1 plasma-membrane localization. Fig. 5 shows representative western blots from fractionation experiments of cells transfected with Kv7.1 and respective flag-tagged KCNE1 constructs. Freedom from contamination of plasma-membrane and endoplasmic reticulum fractions is illustrated by absence of spectrin (plasma-membrane marker) in endoplasmic fraction and absence of calnexin (endoplasmic reticulum marker) in membrane fractions. Kv7.1 (Fig. 5A) and KCNE1 (Fig. 5B) were immunodetected in respective fractions and spectrin and calnexin were further used as expression control. Blots illustrate greater Kv7.1 abundance at the plasma-membrane and less in the endoplasmic reticulum when cells were co-transfected with KCNE1-38S than the other constructs (mean ± SEM data: ‘38S’: 1.3 ± 0.2, ‘38G’: 0.9 ± 0.1, ‘Δ1-38’: 0.8 ± 0.1, ‘linker’: 0.7 ± 0.1, ‘38G-3xA’: 0.7 ± 0.1, ‘38G-3xA’): 0.7 ± 0.1 arbitrary units [AU]; P<0.005 vs. ‘38S’ for all, Fig. 5A). Plasma-membrane distribution of respective KCNE1 constructs was modulated in analogy to Kv7.1 distribution (‘38S’: 1.4 ± 0.1, ‘38G’: 0.8 ± 0.1, ‘Δ1-38’: 0.6 ± 0.1, ‘linker’: 0.7 ± 0.1, ‘38G-3xA’: 0.6 ± 0.1, ‘38G-3xA’): 0.8 ± 0.1 AU; P<0.005 vs. ‘38S’ for all; Fig. 5B).

These data are consistent with electrophysiological measurements and suggest that the N-terminus and specifically the three arginines are important for plasma-membrane representation of the KCNE1 β-subunit as results obtained with alanine-scanning constructs were comparable to those obtained with the N-terminally truncated construct.

Immunofluorescent studies

In order to further substantiate these results we applied confocal microscopy (Fig. 6) to double-labelled cells co-transfected with Kv7.1 and respective flag-tagged KCNE1 constructs. Pancadherin staining was performed to mark plasma-membranes. All KCNE1 constructs co-localized with the α-subunit. Upon laser line-scanning with identical gain settings, Kv7.1 and KCNE1 membrane fluorescence appeared consistently greater in the presence of KCNE1-38S compared with the other KCNE1 constructs.

mRNA transcription of Kv7.1 (α-) and KCNE1 (β-subunits)

We excluded variation in levels of transcription in this heterologous system as a confounding factor. All KCNE1 constructs showed similar mRNA levels (‘38S’: 9.7 ± 0.6, ‘38G’: 10.0 ± 1.2, ‘Δ1-38’: 9.8 ± 1.9, ‘linker’: 10.6 ± 2.2, ‘38G-3xA’: 10.0 ± 1.8, ‘38G-3xA’: 9.9 ± 1.4; non-transfected control: 0.2 ± 0.1 AU; P = n.s.; Fig. 7). Co-transfection of KCNE1 constructs did not change Kv7.1 mRNA expression (‘38S’: 8.8 ± 1.0, ‘38G’: 8.6 ± 0.5, ‘Δ1-38’: 8.5 ± 1.9, ‘linker’: 8.7 ± 1.1, ‘38G-3xA’: 8.1 ± 1.0, ‘38G-3xA’: 9.2 ± 1.1; non-transfected control: 0.1 ± 0.0 AU; P = n.s.; Fig. 7).

Discussion

Major findings

This study identifies the importance of three arginines in the N-terminus of KCNE1 (positions 32, 33 and 36) for plasma-membrane localization of the IKs channel complex. These residues contribute to membrane representation of the β-subunit together with the α-subunit. Removal of these arginines by alanine-scanning in KCNE1-38S led to a comparable IKs phenotype as observed with the atrial fibrillation-related KCNE1-38G or the construct with N-terminal KCNE1 truncation – suggesting a link between a sterical KCNE1-38G alteration mediated by glycine plasma-membrane localization.
increasing protein mobility by formation of a glycine hinge) and a failure of N-terminal arginines to get in contact with the membrane. Structural disruption at position 38 by introducing a ‘linker’ might have similarly affected the ability of arginines to interact with the plasma-membrane.

Relation to delayed rectifier channel trafficking

Kv7.1 and KCNE1 proteins are highly expressed in human atria and ventricles and the resultant I\textsubscript{Kr} is physiologically relevant for cardiac repolarization [4]. The results of our study illustrate that plasma-membrane localization of the \(\alpha\)-subunit Kv7.1 and the \(\beta\)-subunit KCNE1 was dependent on the presence of positively-charged arginines within the KCNE1 N-terminus. Trafficking of the \(\alpha\)-subunit Kv7.1 can be regulated differently when co-assembled with KCNE1. Krumerman et al. found a mutation (KCNE1-L51H) in the transmembrane domain of the \(\beta\)-subunit that conferred trafficking deficiency. KCNE1-L51H did not leave the endoplasmic reticulum and resultant I\textsubscript{Kr} was diminished [13]. In line with this, subsequent work suggested KCNE1 as a chaperone for Kv7.1 intracellular trafficking by protecting the \(\alpha\)-subunit from proteosomal degradation [14]. Conversely, Chandrasekhar \textit{et al.} documented evidence for the necessity of Kv7.1 for KCNE1 membrane expression arguing that Kv7.1 may in fact be needed to help KCNE1 to the membrane [15].

Our results are consistent with previous work that indicated reduced plasma-membrane localization for the atrial fibrillation-associated ‘38G’ allele reducing resultant I\textsubscript{Kr} [12]. In the present study we identified three arginines at positions 32, 33 and 36 that are important for plasma-membrane representation of the I\textsubscript{Ks} complex. Positively-charged amino-acids may interact with hydrophilic phospholipid components of cellular membranes [16]. Arginine is typically located at membrane-water boundaries [17]. Several studies have identified a respective role for arginine in membrane localization. For instance, Jaskolski \textit{et al.} described splice variants of a glutamate receptor where the variant containing a sequence of two arginines and two lysines showed significantly higher membrane localization [18]. Furthermore, membrane anchoring of saposin-C depends heavily on two lysines of the protein. Substitution of either lysine with neutral or negatively-charged amino-acids led to decreased anchoring while exchange with arginine intensified membrane anchoring [19]. Consistently, substitution of the three N-terminal KCNE1 arginines resulted in reduced current density and plasma-membrane localization in the present work. The reduction of membrane localization was comparable to N-terminal truncation of KCNE1 and glycine substitution suggesting a role of these amino-acids for KCNE1 function.

Furthermore, substitution of serine at position 38 with glycine reduced I\textsubscript{Kr} in a similar fashion as deletion of amino-acids 1-38 –
the greater part of the N-terminus. Results were similar when residue 38 was substituted by a series of five alanines. This construct was synthesized in order to mimic the effect of increased mobility by steric distortion of the protein. Glycine is smaller than serine and does not enable covalent binding [20]. It might therefore act as a hinge rendering the N-terminus more flexible, potentially leading to reduced membrane anchoring due to steric alterations of this KCNE1 region.

Effect of KCNE1 on IKs properties

The KCNE1 transmembrane domain interacts with the pore region [6] and with the transmembrane segment 6 [7,21,22] of the α-subunit Kv7.1.

Amino-acids 57–59 of KCNE1 importantly modulate IKs activation kinetics [23]. In subsequent work these investigators found that a hydroxylated central amino acid (threonine 58) was necessary for the slow sigmoidal activation produced by KCNE1 together with Kv7.1. An aliphatic amino acid substituted at position 58 of KCNE1 led to a pronounced change in activation properties [7]. The more rapid activation of IKs resulting from co-transfection of Kv7.1 and 38G in this paper cannot be explained by these sterical alterations.

Recent work indicated an interaction of KCNE1 with the Kv7.1 voltage sensor [24,25]. Furthermore, IKs is very sensitive to mutations in the KCNE1 C-terminus [26,27] and several of these mutations have been linked to the long QT syndrome [28]. The C-terminus of KCNE1 interacts directly with the pore region of Kv7.1 and modulates IKs kinetics and current amplitude [29]. Studies using disulfide bonding also indicated potential interaction of Kv7.1 transmembrane segment 1 with the KCNE1 N-terminus (α-amino-acids 40, 41) [30].

Two of the KCNE1 constructs studied in the present paper (‘linker’ and ‘38S-3xA’), slowed channel deactivation. A study by...
Melman et al. located a sequence in the KCNE1 transmembrane domain (amino-acids 46–56) that was important for the deactivation process of IK\(_{\text{a}}\) [23]. Changes in the KCNE1 N-terminus on the other hand reduced IK\(_{\text{a}}\) but had no effects on kinetics. Kinetic changes observed in our study were only evoked by two of the constructs and not by the N-terminally truncated protein. Slowed deactivation in these cases might have been due to indirect conformational changes potentially influencing the aforementioned transmembrane sequence but further studies are needed to understand this phenomenon.

**Relevance for atrial fibrillation**

Chen et al. reported a gain-of-function mutation in Kv7.1 (S140G) that was associated with increased atrial fibrillation prevalence among members of a large Chinese family [31]. In fact, this was the first description of a monogenetically inherited form of atrial fibrillation. While IK\(_{\text{a}}\) gain-of-function may abbreviate action potential duration and refractoriness and therefore represent a pro-arrhythmic substrate, loss-of-function is less well intuitively correlated with the occurrence of atrial fibrillation. In clinical studies, the KCNE1 single nucleotide polymorphism G330S has been associated with atrial fibrillation [10]. Previous modelling studies, the KCNE1 single nucleotide polymorphism G330S has been associated with atrial fibrillation [10].

Clinical observations provide consistent evidence for a physiological consequence of the single nucleotide polymorphism as fibrillatory rates in atrial fibrillation-patients were slower for those with the ‘38G’ allele in comparison to patients with the ‘38S’ allele [32]. Slower fibrillatory rates relate to prolonged refractory periods may well be associated with reduced repolarizing currents.

**Limitations**

The conclusion of the present study relies on indirect evidence that positively charged amino-acids may need to be in contact with the membrane for increasing membrane representation of the IK\(_{\text{a}}\) delayed rectifier channel complex. Direct evidence from crystallography would be highly desirable but is currently unavailable.

Further, with the present study we cannot exclude slower degradation of ion channel complexes with the KCNE1-30S isoform underlying increases in IK\(_{\text{a}}\) and enhanced channel membrane representation.

**Conclusion**

We have performed a detailed electrophysiological and biochemical analysis of structural determinants underlying reduced IK\(_{\text{a}}\) with the atrial fibrillation-related single nucleotide polymorphism KCNE1-30G. The results of this study suggest that the KCNE1 N-terminus is engaged in IK\(_{\text{a}}\) ion channel complex plasma-membrane representation. N-terminal positively-charged arginine residues 32, 33 and 36 play a specific and important role in the reconstitution of IK\(_{\text{a}}\). Presence of glycine at position 38 and substitution of arginines mimic properties (reduced current density and plasma-membrane localization) as seen with N-terminal truncation of KCNE1 arguing for the specific importance of these arginines for physiological IK\(_{\text{a}}\) function.

**Methods**

**Site-directed mutagenesis**

KCNE1-38G (‘38G’, GenBank NM_000219) was flanked by restriction sites HindIII and EcoRI by polymerase chain reaction (PCR) using primers shown in Table 1. These primers were used to introduce restriction sites into all constructs (Fig. 1B). The KCNE1 polymorphism ‘38S’ was already subcloned into pcDNA3.1 [12]. Standard cloning technique (Table 1) was used. Additionally, each construct was flag-tagged C-terminally. All constructs were checked by sequencing.

Both naturally occurring alleles of the polymorphism (‘38S’ and ‘38G’) had previously been cloned [12]. Alanine-substitution of the three N-terminal KCNE1 arginines was performed to disrupt these positively charged amino-acids that might play a role in membrane anchoring (‘38S-3xA’, ‘38G-3xA’). N-terminal truncation of KCNE1 was performed to study the role of the greater part of the N-terminus (‘Δ1-38’). Glycine substitution at position 38 by four alanines (‘linker’) was performed in order to mimic the effect of increased mobility by steric distortion of the β-subunit. Glycine might act as a hinge to allow flexibility of the N-terminus,.

**Table 1. Primers for KCNE1 mutagenesis.**

| Primer | Constructs | Comments | Sequence |
|--------|------------|----------|----------|
| Fw1 | all constructs | introduction of restriction site | 5’-CCGAAGCTTATGATCCGTGTC-3’ |
| Fw2 | Δ1-38 | truncation of AA 1-38 | 5’-CCGAAGCTTATGACGGCAAGCTGGAG-3’ |
| Fw3 | linker | overlap extension | 5’-CAAGCAGCAGGACGAGCGGCAAGCTGGAG-3’ |
| Fw4 | Intermediate construct 38S-R36A | overlap extension | 5’-CACCGTCGGCGAGCGGCAAGCTGGAG-3’ |
| Fw5 | Intermediate construct 38G-R36A | overlap extension | 5’-CACCGTCGGCGAGCGGCAAGCTGGAG-3’ |
| Fw6 | 38S-3xA | overlap extension | 5’-CACCGTCGGCGAGCGGCAAGCTGGAG-3’ |
| Fw7 | 38G-3xA | overlap extension | 5’-CACCGTCGGCGAGCGGCAAGCTGGAG-3’ |
| Rev1 | all constructs | introduction of restriction site | 5’-CTCAGAAATTCTCAATGGAGAAGGC-3’ |
| Rev2 | linker | overlap extension | 5’-CTCAGAAATTCTCAATGGAGAAGGC-3’ |
| Rev3 | Intermediate construct 38S-R36A | overlap extension | 5’-CTCAGAAATTCTCAATGGAGAAGGC-3’ |
| Rev4 | Intermediate construct 38G-R36A | overlap extension | 5’-CTCAGAAATTCTCAATGGAGAAGGC-3’ |
| Rev5 | 38S-3xA, 38G-3xA | overlap extension | 5’-CTCAGAAATTCTCAATGGAGAAGGC-3’ |
| Rev6 | flag-tagged constructs | C-terminal flag-tag | 5’-CTCAGAAATTCTCAATGGAGAAGGC-3’ |

Mutation containing sequences in italic, underlined. 

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potentially leading to reduced membrane anchoring due to steric alterations of this KCNE1 region.

Cell culture

Human embryonic kidney (HEK) and Chinese hamster ovary (CHO) cells (ATCC, Manassas, VA, USA) were cultured at 37°C with 5% CO₂ in MEM or F12-medium (Invitrogen, Carlsbad, CA, USA) supplemented with 10% heat-inactivated fetal bovine serum. Transient transfections were obtained with polyethylenimine (Polysciences Inc., Warrington, PA, USA) and a total of 1 μg cDNA for electrophysiological measurements, 0.5 μg for protein-biochemistry and 0.25 μg cDNA for confocal microscopy. Co-transfected green fluorescent protein (0.2 μg cDNA) served as a transfection marker for electrophysiological recordings. All experiments were performed 48 h after transfection.

Electrophysiological recordings

Currents were recorded with whole-cell patch-clamp at physiological temperature (36.5 ± 0.5°C) with an Axopatch 200B amplifier (Molecular Devices, Sunnyvale, CA, USA). Borosilicate glass electrodes had 1.5 to 3.0 MΩ tip resistances when filled with internal solution. Compensated cell capacitances were 27.0 ± 2.1 pF with no statistical differences between cells transfected with respective KCNE1 constructs. Liquid junction potentials (7.5 ± 0.5 mV) were not corrected. The extracellular solution was composed of (mM) NaCl 136, KCl 5.4, MgCl₂ 1, CaCl₂ 1, NaH₂PO₄ 0.33, HEPES 5 and dextrose 10 (pH 7.35 with NaOH). Internal solution contained (mM) K- aspartate 110, KCl 20, MgCl₂ 1, MgATP 5, GTP (lihium salt) 0.1, HEPES 10, Na-phosphocreatine 5 and EGTA 5.0 (pH 7.3 with KOH). Currents were elicited with 1 s depolarizing pulses from a holding potential of −80 mV and an inter-pulse interval of 12 s. Tail currents were observed during 2 s repolarizations to −50 mV from various depolarizing potentials.

Clampfit (Molecular Devices, Sunnyvale, CA, USA) and GraphPad Prism (GraphPad Software, San Diego, CA, USA) were used for data analysis. Data are presented as mean ± SEM. P < 0.05 by two-way ANOVA (for comparison among groups) or Student’s t-test (for comparison of individual voltage steps) was considered to indicate statistical significance. We assumed Student's t-test to be sufficiently precise even in the case of unequal variance as sample sizes were similar among different groups.

Protein biochemistry

HEK cells were washed with ice-cold phosphate-buffered saline (pH 8.0) to remove media and scraped into lysis-buffer composed of (mM) Tris-HCl (pH 7.4) 50, NaCl 150, EDTA 1, EGTA 1, PMSF 1 and 1% NP-40, 0.5% sodium deoxycholate, 0.1% SDS as well as protease inhibitors as above. Again all steps were at 4°C. After 1 h incubation under rotation the cells were centrifuged for 10 min at 2,800 g. Samples were precleared with 25 μl protein-G-agarose beads (Pierce Biotechnologies, Rockford, IL, USA) for 30 min under rotation. Beads were removed by 1 min centrifugation at 1,000 g. 1 μg of antibody was added to 2 mg of supernatant and incubated overnight. Another 30 μl protein-G-agarose beads were added and incubated for 5 h. The beads were sedimented by 1 min centrifugation at 1,000 g, washed and suspended in Laemmli buffer before loading onto SDS gels.

Confocal microscopy

Transfected HEK cells were grown on plastic coverslips (ibidi, Martinsried, Germany) and fixed (20 min) with 2% paraformaldehyde (Sigma-Aldrich, St. Louis, MO, USA). After 3 washes (5 min) with phosphate-buffered saline cells were blocked with 5% normal donkey serum (Jackson Immunoresearch Laboratories, West Grove, PA, USA) and 5% bovine serum albumin (BSA, Sigma-Aldrich, St. Louis, MO, USA). Then cells were permeabilized with 0.2% Triton X-100 (Sigma-Aldrich, St. Louis, MO, USA) for 1 h. Primary antibodies were incubated overnight at 4°C: rabbit anti-flag (Sigma Aldrich, St. Louis, MO, USA), goat anti-Kv7.1 (Santa Cruz Biotechnology, Santa Cruz, CA, USA) and mouse anti-pan-cadherin (Abcam, Cambridge, UK) at 1:200 dilution, followed by 3×5 min washes and a 1 h incubation with secondary antibodies (fluorescence-labelled ‘Alexa-Fluor-555’ [anti-rabbit] for KCNE1 and ‘Alexa-Fluor-488’ [anti-goat] for Kv7.1). Confocal microscopy was performed with a Zeiss LSM-510 system. KCNE1 (green) and Kv7.1 (red) were excited at 555 and 495 nm with Helium/Neon- and Argon-Lasers, respectively, emitting fluorescence at 565 and 519 nm. Confocal microscopy experiments of different groups were performed on the same experimental day with identical parameters for all groups. Control experiments omitting primary antibodies and with non-transfected cells revealed absent or very low-level background staining.

Table 2. Primers for quantitation of Kv7.1 and KCNE1 mRNA.

| Primer | GenBank accession no. | Sequence |
|--------|-----------------------|----------|
| Kv7.1  | NM_000218             | F: 5’-AGACGTGGGTCGGGAAGAC-3’<br>R: 5’-CAAGAAGGAGGTGCAAAAGCA-3’ |
| KCNE1  | NM_000219             | F: 5’-CGGCCCTCTAGCTCTGCTG-3’<br>R: 5’-GTAGCCCGAGCTTGAAAG-3’ |
| β-actin | NM_001101             | F: 5’-GGATCAGCAAGGAGTATG-3’<br>R: 5’-GCATTTCGCGTCCAGGATG-3’ |

F – forward primer, R – reverse primer. doi:10.1371/journal.pone.0026967.t002

Immunoprecipitation

Transfected HEK cells were washed with phosphate-buffered saline and scraped into 1 ml buffer containing (mM) Tris-HCl (pH 7.4) 50, NaCl 150, EDTA 1, EGTA 1, PMSF 1 and 1% NP-40, 0.5% sodium deoxycholate, 0.1% SDS as well as protease inhibitors as above. Again all steps were at 4°C. After 1 h incubation under rotation the cells were centrifuged for 10 min at 2,800 g. Samples were precleared with 25 μl protein-G-agarose beads (Pierce Biotechnologies, Rockford, IL, USA) for 30 min under rotation. Beads were removed by 1 min centrifugation at 1,000 g. 1 μg of antibody was added to 2 mg of supernatant and incubated overnight. Another 30 μl protein-G-agarose beads were added and incubated for 5 h. The beads were sedimented by 1 min centrifugation at 1,000 g, washed and suspended in Laemmli buffer before loading onto SDS gels.

Cells were treated with the ‘Membrane Protein Extraction Kit’ (Promokine, Heidelberg, Germany) to obtain plasma-membrane and endoplasmic reticulum fractions. Equal loading and accuracy of fractionation was probed by immunodetection of calnexin as endoplasmic reticulum marker and spectrin for plasma-membranes. Protein expression was normalized to calnexin or spectrin.
RNA extraction, reverse transcription and RNA quantitation

RNA was extracted from transfected HEK cells using the RNeasy Mini Kit (Qiagen, Valencia, CA, USA). cDNA was synthesized from 1 μg of total RNA. Reactions (20 μl) contained 1.25 μM random hexamers, 1.25 μM oligo-dTs, 0.25 mM each of dNTPs, 200 U reverse transcriptase (Invitrogen, Carlsbad, CA, USA), reverse transcriptase buffer, 10 mM dithiothreitol (DTT) and 40 U RNase inhibitor. RNA, random hexamers, oligo-dTs, dNTPs, and RNase inhibitor were incubated (65°C, 5 min), then put on ice (5 min). Reverse transcriptase buffer and DTT were added and incubated (37°C, 2 min). This was followed by 30 min incubation at 37°C and then 15 min incubation at 70°C. Real-time PCR was performed using the 7300 Fast Real-Time PCR System and the Fast SybGreen Master Kit (Applied Biosystems, Foster City, CA, USA). Reactions (20 μl) contained cDNA (1 μl) and 0.3 μM of each forward and reverse primer (Table 2). A “20 s denaturation” at 95°C was followed by 40 cycles of denaturation (1 s) at 95°C and annealing and elongation (20 s) at 60°C.

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Author Contributions

Conceived and designed the experiments: ZG PB IT SHH JRE. Performed the experiments: ZG PB IT CS. Analyzed the data: ZG PB JRE. Contributed reagents/materials/analysis tools: SHH JRE. Wrote the paper: ZG PB IT SHH JRE.

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