Identification of small-molecule ion channel modulators in C. elegans channelopathy models

Qiang Jiang1,2, Kai Li1,2, Wen-Jing Lu3, Shuang Li2,4, Xin Chen1,2,5, Xi-Juan Liu1, Jie Yuan1,2, Qiuqong Ding2,4, Feng Lan3 & Shi-Qing Cai1

Ion channels are important therapeutic targets, but the discovery of ion channel drugs remains challenging due to a lack of assays that allow high-throughput screening in the physiological context. Here we report C. elegans phenotype-based methods for screening ion channel drugs. Expression of modified human ether-a-go-go-related gene (hERG) potassium channels in C. elegans results in egg-laying and locomotive defects, which offer indicators for screening small-molecule channel modulators. Screening in worms expressing hERG<sup>A561V</sup>, which carries a trafficking-defective mutation A561V known to associate with long-QT syndrome, identifies two functional correctors Prostratin and ingenol-3,20-dibenzoate. These compounds activate PKCe signaling and consequently phosphorylate S606 at the pore region of the channel to promote hERG<sup>A561V</sup> trafficking to the plasma membrane. Importantly, the compounds correct electrophysiological abnormalities in hiPSC-derived cardiomyocytes bearing a heterozygous CRISPR/Cas9-edited hERG<sup>A561V</sup>. Thus, we have developed an in vivo high-throughput method for screening compounds that have therapeutic potential in treating channelopathies.
on channels are the molecular basis for the cell excitability. Malfunction of ion channels causes over 55 different inherited human diseases known as channelopathies1. Although 15% of US Food and Drug Administration (FDA)-approved drugs act through targeting ion channels2–5, there remains an urgent need for developing drugs for many untreated channelopathies. Discovery of new drugs targeting ion channels is known to be notoriously difficult. Assaying ion channel activity with patch-clamp recording of ion channel currents has very low screening throughput, although this problem is now largely alleviated by recent advances in cell line-based fluorescent assays and automated electrophysiology2–5. However, existing in vitro assays using cell lines expressing a particular ion channel could not overcome the problem that the channel in their native tissues exists in a cellular and physiological environment substantially different from that in the cell line, hence exhibits different molecular and functional properties. For example, accessory subunits usually dramatically alter the pharmacological properties of ion channels4,5. An efficient screening tool thus should examine the function of ion channels in the physiological context. Most types of ion channels are found in C. elegans and are known to regulate worm behaviors6–10. Like their human homologs, C. elegans channels usually consist of pore-forming and accessory subunits10,11. Many mechanisms underlying the biogenesis of ion channels are conserved from C. elegans to humans11–13. Based on these findings and cumulative knowledge regarding ion channel mutations associated with human diseases, we hypothesized that expression of disease-relevant human ion channels in C. elegans would disturb normal electrical signaling and hence cause phenotypic defects in the transgenic worms. Compounds that modulate human ion channel function could be identified by examining their effects on the phenotypes of the transgenic worms. Phenotype-based screening in C. elegans has been used in dissecting mechanism of drug action and screening for new drugs14–16. In this study, we specifically targeted ion channel function using phenotype-based screening in C. elegans models of channelopathies, and demonstrated that specific small-molecule modulators could be identified for correcting the defective function of mutant ion channels associated with channelopathies, e.g., Long-QT syndrome (LQTS).

Human ether-a-go-go-related gene17 (hERG) encodes a voltage-gated K+ channel that mediates cardiac IKr currents. Dysfunction of hERG K+ channels due to genetic mutations or drug-induced inhibition results in LQTs18–22, a life-threatening cardiac arrhythmia that affects 1 in 2500 live births and results in about 5000 death each year in US alone23. Due to the fatal side effects of causing LQTS, several drugs have been withdrawn from the market since 1990s and the FDA now requires to test the drug effect on hERG activity during preclinical safety evaluation22,24. Most of hERG mutations underlying type 2 LQTS (LQT2) suppress IKr currents by causing trafficking defects25,26. The mutant hERG K+ channels are usually able to elicit normal or near normal Ikr currents if they are properly transported to the plasma membrane by experimental manipulations, e.g., lowering the incubation temperature27,28. Pharmacological correction of the trafficking defect thus represents a new promising strategy for treating LQTS.

In this study, we generate channelopathy animal models by expressing chimeric hERG or trafficking-defective hERG mutant channels in C. elegans. Phenotype-based screens in the transgenic worms identify aliphatic acid (ALA) as a novel hERG trafficking inhibitor and two protein kinase C (PKC) activators, Prostratin and ingenol-3,20-dibenzoate (IDB), as functional correctors of some LQT2 mutant channels. Thus, we provide an in vivo system for the discovery of ion channel modulators.

Results
Generation of C. elegans animal models of channelopathies. Human ion channels share high similarity with their C. elegans homologs, especially in their transmembrane domains24. To overcome the possible barrier of expressing human ion channels in C. elegans, we first made human-worm chimeric ion channels according to the strategy described in Fig. 1a. Ion channel inhibitors and activators often target at transmembrane domains and/or loops at the membrane periphery of the channels. We thus assumed that a chimeric ion channel consisting of transmembrane domains and their connecting loops of a human ion channel, and N-/C-terminus of its C. elegans homolog, would largely maintain pharmacological properties of the human ion channel and could be functionally expressed in C. elegans. A gain-of-function mutation was introduced into the channel (named as chimeric ion channel 1, Fig. 1a) to induce clear behavioral defects in the transgenic worms expressing the channel, and drugs that alleviate the phenotypic defects could be inhibitors of the channel (Fig. 1b). For screening channel activators or functional correction of a mutant channel, a second mutation, which is a loss-of-function mutation, was introduced into the channel (named as chimeric ion channel 2, Fig. 1a), and transgenic worms expressing this channel should show no or weak behavioral defects due to the reduced channel function. Small molecules that aggravate the worms’ behavioral defects are potential activators of the channel (Fig. 1b).

According to the above strategy, we made C. elegans models of channelopathies for identifying chemical modulators of hERG K+ channels. The hERG K+ channel shows homology with the C. elegans ERG-type K+ channel UNC-103 (Supplementary Fig. 1a), which is widely expressed in C. elegans head neurons and vulva muscles and regulates worms’ locomotion, egg-laying, and male mating behaviors6,7. We first tried to express full-length hERG K+ channels in C. elegans, but failed due to trafficking of the channel. We then constructed a GFP fused chimeric hERG (hERGchimera/AS56W;GFP), which consisted of N-terminus and C-terminus of UNC-103, transmembrane, and cyclic nucleotide-binding domains of hERG, and a gain-of-function mutation AS56W30 in the S4 segment of the hERG K+ channel (Supplementary Fig. 1b). The mutation AS56W, which switches the voltage-dependent activation of hERG K+ channels to a much more negative potential30, was expected to alter cell excitability and consequently induce behavioral defects in worms. Indeed, expression of unc-103 promoter-driven hERGchimera/AS56W::GFP proteins in N2 wild-type worms generated transgenic worms (hERGchimera/AS56W) with clear defects in locomotion and egg-laying (Fig. 1c–e). Fluorescent images suggested that hERGchimera/AS56W::GFP channels could be delivered to the plasma membrane (Fig. 1f). In addition, expression of hERGchimera/AS56W rescued the copulatory spicule protrusion defect caused by a loss-of-function mutation in the unc-103 gene in male worms (Supplementary Fig. 1c). Taken together, these data suggest that hERGchimera/AS56W could affect worms’ behaviors.

Next, we examined whether behavioral defects in worm models of channelopathies could be rescued by known specific blockers of these channels. Cells of C. elegans are relatively inaccessible for most chemical compounds because of their cuticles and xenobiotic efflux pumps in the intestine. To facilitate penetration of compounds into the worm, we generated als-20hERGchimera/AS56W mutant worms by crossing hERGchimera/AS56W with als-20 null mutant worms, whose cuticles have increased permeability for small molecules31. To further enhance the targeting of the compounds, we fed these worms with bacteria expressing double-strand RNAs targeting ifd-2 and c15c7.5, two genes encoding nonessential intestinal intermediate filament proteins that impede the effects of ingested compounds32. We found that the phenotypes in the
hERGchimera/A536W worms were ameliorated by individually adding 10 μM well-known hERG blockers, and these effects were markedly enhanced when the acs-20 null mutation was introduced in the transgenic worms (Supplementary Fig. 2). This result indicates that drugs could be efficiently delivered to their targets, and modified human ion channels expressed in the transgenic worms are functional and the cause of behavioral defects.

A small-molecule screen identifies a new hERG trafficking inhibitor. Next, we examined whether acs-20:hERGchimera/A536W worms could be used for identification of hERG K⁺ channel modulators. To identify unknown hERG inhibitors, we screened about 4000 small molecules for their suppression of behavioral defects in acs-20:hERGchimera/A536W worms (Fig. 2a). Dozens of compounds were found to alleviate the phenotypic defects to some extents. Among them, alphitolic acid (ALA) clearly restored locomotion and egg-laying behaviors of the tested worms (Fig. 2b). Additional test using HEK293T cells expressing wild-type hERG K⁺ channels showed that acute treatment with either 10 or 10 μM ALA did not change the current density (Fig. 2c), suggesting that ALA did not act directly on hERG K⁺ channel permeation. By contrast, long-term (24–32 h) treatment with ALA led to reduced current density of this channel in a dose-dependent manner. The median inhibition concentration (IC₅₀) was 8.2 μM, as determined by analyzing changes in deactivating tail currents at the membrane potential of −140 mV (Fig. 2d–f). Western blotting data showed that hERG proteins appeared as two bands corresponding to the endoplasmic reticulum (ER)-

**Fig. 1** C. elegans models of channelopathies show behavioral defects. a Schematic illumination of chimeric ion channels. The chimeric ion channels consist of transmembrane domains and their connecting loops of a human ion channel, and N-/C-terminus of its C. elegans homolog. The gain-of-function mutation was expected to alter cell excitability and consequently induce behavioral defects in transgenic worms. b Schematic illumination of generating animal models of channelopathies and strategy for drug screenings. Transgenic worms expressing chimeric ion channel 1 proteins will exhibit clear behavior defects. Drugs that alleviate the phenotypic defects could be inhibitors of the channel. Transgenic worms expressing chimeric ion channel 2 proteins will show no or weak behavior defects due to the reduced channel function, and drugs aggravate the worms’ behavioral defects are potential activators or functional correctors of the channel. c Microscopy images of wild-type N2 and hERGchimera/A536W transgenic worms. Scale bar, 200 μM. White arrows indicate C. elegans eggs. d, e Locomotive (d) and egg-laying (e) behaviors of of young adult wild-type N2 and hERGchimera/A536W transgenic worms. f Fluorescent images of worms expressing both mCherry and hERGchimera/A536W::GFP proteins that were driven by the unc-103 promoter. Blue arrows indicate head neurons. Scale bar, 10 μm. All data shown are mean ± s.e.m. ***P < 0.001 (Student’s t-tests for d, e).
resident (with core glycosylation, ~135 kD) and post-ER (with fully glycosylation, ~155 kD) forms (Fig. 2g). Long-term treatment with ALA reduced the hERG protein levels of the 155 kD, but not the 135 kD band (Fig. 2g). Furthermore, immunostaining of HEK293T cells expressing hERG K\textsuperscript{+} channels also showed that ALA inhibited protein trafficking of hERG K\textsuperscript{+} channels (Fig. 2h). These results suggest that ALA is a novel hERG protein trafficking inhibitor. Together, screening assays in worms expressing hERG K\textsuperscript{+} channels could identify new hERG inhibitors.

Identification of functional correctors of LQTS2 mutant channels. The majority of LQTS2-related mutations suppress the I\textsubscript{Kr} currents by retaining hERG K\textsuperscript{+} channels in the ER\textsuperscript{25,26}. Pharmacological correction of trafficking defects thus represents a promising therapeutic strategy for LQTS\textsuperscript{27,28,33,34}. In order to generate an animal model of LQTS, a trafficking-defective LQTS2 mutation A561V\textsuperscript{27,35} was introduced to the hERG\textsuperscript{chimera/A536W} protein (Fig. 3a). As expected, the transgenic worms hERG\textsuperscript{chimera/A536W/A561V} (expressing hERG\textsuperscript{chimera/A536W/A561V}:...
**Fig. 3** Small-molecule screens identify functional correctors of hERGA561V. 

**a** Schematic illumination of construction of hERG chimera/A536W/A561V. Mutations A536W and A561V are indicated by a red and a black circle, respectively. aa amino acids.

**b** Localization of hERG chimera/A536W::GFP and hERG chimera/A536W/A561V::GFP expressed in C. elegans neurons. White arrows indicate head neurons. Scale bar, 10 μm.

**c** Schematic illumination of screening strategy for the discovery of functional correctors. Briefly, we first searched for compounds that caused locomotive and egg-laying defects, and then the candidate compounds were further selected according to surface expression of hERG chimera/A536W/A561V::GFP proteins.

**d** Molecular formulas of Prostratin and IDB.

**e**, **f** Representative whole-cell currents (**e**) and I/V curves (**f**) of hERGWT–hERGA561V channels expressed in HEK293T cells treated with DMSO (n = 26), 3 μM Prostratin (n = 21) or 2 μM IDB (n = 25).

**g** Dose-dependent effects of Prostratin and IDB on the current density of hERGWT–hERGA561V channels. About 20 cells per dose were recorded and analyzed. 

**h** Effects of Prostratin and IDB on the protein trafficking of hERGWT–hERGA561V channels. Black, gray, and red arrows indicate 155 kD, 135 kD bands of hERG proteins, and tubulin, respectively.

**i** Effects of Prostratin and IDB on the current densities of LQTS-related hERG mutant channels. The ratio of hERGWT/hERGA561V was 2:1 for **e**, **f**, **g**, and 1:1 for **h**, **i**. The ratio of hERGWT/other hERG mutant was 1:1. All experiments were performed at least three times. Data shown are mean ± s.e.m. *P < 0.05, **P < 0.01, ***P < 0.001 (Student’s t-tests for **f**, one-way ANOVA Dunnett’s test for **i**).
GFP proteins) showed GFP reporters in the neuronal cytoplasm (Fig. 3b), indicating mistrafficking of these channel proteins. Consistently, hERGchimera/A536W/A561V worms exhibited nearly normal behaviors (Supplementary Movies 1–3).

The hERG A561V K+ channels could be functional if they reach to the plasma membrane27. We hypothesized that if a compound promotes the surface expression of hERGchimera/A536W/A561V, the transgenic worms would show locomotion and egg-laying defects. We thus generated acs-20 hERGchimera/A536W/A561V worms by crossing acs-20 null mutant worms with hERGchimera/A536W/A561V, and screened for novel functional correctors of hERGchimera/A536W/A561V according to the strategy depicted in the Fig. 3c. Among 10,600 compounds screened at a final concentration of 20 μM, we identified six hits, among which two compounds Prostratin and IDB (Fig. 3d and Supplementary Fig. 3) were known as PKC activators.

The A561V mutation is known to cause strong dominant-negative suppression of wild-type hERG (hERGWT) protein trafficking, resulting in reduced channel function at the surface35,36. Strikingly, either Prostratin or IDB treatment enhanced the current density of hERG K+ channels in a dose-dependent pattern when we co-expressed hERGWT and hERG A561V at a ratio of 2:1 (Fig. 3e–g). The half maximal effective concentrations (EC50) of Prostratin and IDB were 0.95 μM and 0.042 μM, respectively (Fig. 3g). We then examined whether these compounds affect protein trafficking of hERG mutant channels. In this assay, we co-expressed hERGWT and mutated hERG K+ channels at a ratio of 1:1 because hERG mutations are heterozygous in most LQTS2 patients. Western blotting experiments showed that both Prostratin and IDB promoted trafficking of hERGWT-hERGA561V channels in a dose-dependent pattern, as reflected by the elevated 155 kD hERG proteins (Fig. 3h and Supplementary Fig. 4), which distributed outside the ER. In addition to the A561 site, hERG K+ channels with other LQTS-related mutations such as I31S, G601S, and S818L were found to exhibit enhanced function after treatment with 3 µM Prostratin or IDB and Prostratin on correcting the functional defects in LQTS2 mutant channels (Fig. 5b). To further explore which phosphorylation site is required for the function of IDB and Prostratin, we generated three mutated hERG K+ channels with a lack of PKC phosphorylation sites in the N-terminus, the transmembrane domains, or the C-terminus. We found that only blockage of phosphorylation sites in the transmembrane domains diminished the function of IDB and Prostratin (Fig. 5c–e). Further studies showed that a phosphorylation site of hERG K+ channels at S606 (Fig. 5f), but not S636 and T670 (Supplementary Fig. 7), was required for the function of IDB and Prostratin. We next mutated S606 to E606 to mimic the phosphorylation of hERG K+ channels, and found that the mutation S606E in hERGWT marked increased the current density of hERGWT-hERGA561V (Fig. 5g), suggesting that the phosphorylation of this site promotes the function of the LQTS2 mutant channel. Taken together, our results suggest that IDB and Prostratin activate the PKCε signaling and consequently phosphorylate S606 at the pore region of the channel to correct the defective function of LQTS2 mutant channels.

Identified compounds restore electrophysiology of cardiomyocytes carrying a LQTS2 mutation. The hERG K+ channel plays a major regulatory role in the re-polarization phase of cardiac action potentials. Human-induced pluripotent stem cell-derived cardiomyocytes (hiPSC-CMs) from LQTS2 patients (with a heterozygous A561V mutation in hERG) showed diminished Ik currents and prolonged re-polarization durations27. We then generated a heterozygous A561V mutation in hiPSCs by CRISPR/ Cas9-mediated genome editing (Fig. 6a and Supplementary Fig. 8a), and differentiated these hiPSCs into cardiomyocytes (named hiPSC A561V-CMs). The cardiac ontogeny of both control hiPSC-CMs and hiPSC A561V-CMs was confirmed by analyses of cardiac cell-specific transcriptional, structural, and functional markers (Fig. 6b, c).

We found that endogenous Ik, currents, isolated by the hERG blocker E-4031, in hiPSC A561V-CMs were significantly decreased in comparison with those in control hiPSC-CMs (Fig. 6d, e). In consistence with this result, immunostaining of hiPSC A561V-CMs showed that hERG A561V proteins accumulated in the cytoplasm (Supplementary Fig. 8b). Concurrently, the APD90 (action potential duration at 90% of full re-polarization) of atrial-like and ventricular-like hiPSC A561V-CMs was about two times longer than that of control hiPSC-CMs (Fig. 6f, g). Furthermore, action potentials in 5 out of 22 hiPSC A561V-CMs exhibited early afterdepolarization (EAD), the harbinger of cardiac arrhythmias in LQTS, while none of action potentials in control hiPSC-CMs (n = 25) showed EADs (Supplementary Fig. 8c, d). Thus, the electrophysiological phenotypes in hiPSC A561V-CMs were similar to those in LQTS2 patient-specific hiPSC-CMs27,28.

We then explored whether Prostratin and IDB could correct the electrophysiological defects of hiPSC A561V-CMs. We found that long-term (48 h) administration of Prostratin (3 μM) or IDB (2 μM) markedly enhanced the Ik, currents in hiPSC A561V-CMs (Fig. 6d, e). Immunostaining results suggested that Prostratin and IDB ameliorated the trafficking defects of hERG A561V proteins in hiPSC A561V-CMs (Supplementary Fig. 8b). More importantly, these two compounds shortend the action potential duration of hiPSC A561V-CMs to a level comparable with that of control hiPSC-CMs (Fig. 6f, g) and prevented the occurrence of EADs (Supplementary Fig. 8c, d). By contrast, these two compounds did not affect wild-type Ik, currents and the duration of action potentials in control hiPSC-CMs (Supplementary Fig. 9a–d). We further examined the effect of Prostratin and IDB on the transcription of hERG splice variants and other channels expressed in hiPSC A561V-CMs, and found that administration

Prostratin and IDB restore the function of LQTS2 mutant channels through PKCε signaling. Both Prostratin and IDB are PKC activators. Prostratin is non-selective, and IDB is specific for the PKCe. We then examined whether the rescue of trafficking defects by Prostratin and IDB results from activation of PKCe. We found that downregulation of PKCe, but not PKCβ, abolished the effect of IDB and Prostratin on the current density (Fig. 4a–c and Supplementary Fig. 6) and protein trafficking of hERGWT-hERGA561V channels (Fig. 4d). Furthermore, overexpression of PKCe enhanced current densities and promoted protein trafficking of hERGWT-hERGA561V channels (Fig. 4e, f). Together, these results demonstrate that Prostratin and IDB recover trafficking of hERG mutant channels through the PKCe signaling.

Previous studies27 have shown that hERG contains 18 putative PKC-dependent phosphorylation sites (Fig. 5a). To explore whether the action of Prostratin and IDB depends on phosphorylation of hERG mutant K+ channels, 17 out of 18 PKC phosphorylation sites (except the T74 site, which prevents functional expression of hERG K+ channel57) of the channel were mutated to alanine to block the phosphorylation of these sites. We found that these mutations largely abolished the effects of IDB and Prostratin on correcting the functional defects in LQTS2 mutant channels (Fig. 5b). To further explore which phosphorylation site is required for the function of IDB and Prostratin, we generated three mutated hERG K+ channels with a lack of PKC phosphorylation sites in the N-terminus, the transmembrane domains, or the C-terminus. We found that only blockage of phosphorylation sites in the transmembrane domains diminished the function of IDB and Prostratin (Fig. 5c–e). Further studies showed that a phosphorylation site of hERG K+ channels at S606 (Fig. 5f), but not S636 and T670 (Supplementary Fig. 7), was required for the function of IDB and Prostratin. We next mutated S606 to E606 to mimic the phosphorylation of hERG K+ channels, and found that the mutation S606E in hERGWT markedly increased the current density of hERGWT-hERGA561V (Fig. 5g), suggesting that the phosphorylation of this site promotes the function of the LQTS2 mutant channel. Taken together, our results suggest that IDB and Prostratin activate the PKCε signaling and consequently phosphorylate S606 at the pore region of the channel to correct the defective function of LQTS2 mutant channels.
HERG channels expressed in HEK293T cells. Black and gray arrows indicate 155 kD and 135 kD bands of HERG proteins, respectively. The ratio of HERGWT/HERGAS61V was 2:1 for HEK293T cells treated with control or negative control (NC).

**Discussion**

The discovery of new ion channel drugs could be greatly facilitated by the availability of an economic, reliable, robust, and high-throughput in vivo assay. In this study, we generated *C. elegans* models of channelopathies by expressing disease-related HERG channels in the worms, and developed phenotype-based screening assays by which we successfully identified some novel channel modulators, illustrating the usefulness of this in vivo high-throughput screening method.

Theoretically, inherited mutations could result in dysfunctions of ion channels by the following mechanisms: abnormal transcription/translation, deficient assembly or protein trafficking, defective channel gating/kinetics, and altered channel permeability. Different disease-related mutant channels may need different strategies to rescue their function. For instance, the function of an ion channel mutant with abnormal transcription, translation, or defective trafficking may not be fully corrected by activators of ion channels. Current drug screenings, however, mainly focus on identification of compounds that modulate the gating and kinetics of ion channels. In this study, we have demonstrated that different screening methods could be designed by introducing specific disease-related channel mutations into *C. elegans*.

**Fig. 4** The action of Prostratin and IDB requires activation of PKCe signaling. a The efficiency of siRNAs targeting PKCe. A scrambled siRNA was used as a negative control (NC). b, c Effects of Prostratin and IDB on whole-cell currents (b) and current densities (c) of HERGWT-HERGAS61V channels expressed in HEK293T cells treated with control or PKCe siRNAs. d Effects of Prostratin and IDB on protein trafficking of HERGWT-HERGAS61V channels expressed in HEK293T cells treated with control or PKCe siRNAs. e, f Effects of PKCe over-expression on current densities (e) and protein trafficking (f) of HERGWT-HERGAS61V channels expressed in HEK293T cells. Black and gray arrows indicate 155 kD and 135 kD bands of HERG proteins, respectively. The ratio of HERGWT/HERGAS61V was 2:1 for b, c, e, and 1:1 for d, f. All experiments were performed at least three times. Data shown are mean ± s.e.m. **P < 0.01, ***P < 0.001 (one-way ANOVA Dunnett’s test for c, Student’s t-tests for e, f).
channels with a lack of the S606 PKC phosphorylation site. 

Fig. 5 The action of Prostratin and IDB depends on the phosphorylation of the S606 at the pore region of the hERG channel. a Schematic illumination of putative PKC phosphorylation sites and point mutations in hERG proteins. b Effects of Prostratin and IDB on the current densities of hERGWT-hERGΔS606 channels with a lack of 17 PKC phosphorylation sites (indicated as 17(A)), PKC phosphorylation sites (T526A, S606A, S636A, and T670A) in the transmembrane domain (indicated as T(A)), or PKC phosphorylation sites (S26A, T162A, T174A, S179A, S250A, S278A, S354A, and T371A) in the N-terminus (indicated as N(A)). c Effects of Prostratin and IDB on the current densities of hERGWT-hERGΔS606 channels with a lack of the S606 PKC phosphorylation site. d Effects of the mutation S606E on the current densities of hERGWT-hERGΔS606V channels. The ratio of hERGWT/hERGΔS606V was 2:1. All experiments were performed at least three times. Data shown are mean ± s.e.m. **P < 0.001, n.s., not significant (one-way ANOVA Dunnett’s test for b, c, d, e, and f; Student’s t tests for g).

elegans disease models. Phenotype-base screens in these animal models could discover compounds that correct abnormal channel phenotypes caused by specific mutations.

The cumulative evidence suggests that defective protein biogenesis represents a rather common mechanism underlying many channopathies, including LQTS, cystic fibrosis, episodic ataxia type 1, epilepsy, and hearing loss. Despite extensive interests in pharmacological correction of defective protein biogenesis of ion channels, few such compounds have been found. A successful example is that a combination administration of CFTR-AF508 protein trafficking corrector VX-809 and CFTR chloride channel potentiator VX-770 was approved for the
treatment of cystic fibrosis\textsuperscript{42}. Some hERG blockers have been reported to promote protein trafficking of LQTS mutant channels\textsuperscript{26,43}, however, the blockage action of drugs may diminish the current conducted by rescued hERG K\textsuperscript{+} channels, consequently nullify their therapeutic potential. In this study, our phenotype-based screening assays in \textit{C. elegans} model of LQTS led to identification of compounds that not only promote protein trafficking of LQTS mutant channels, but also correct the electrophysiological abnormality of cardiomyocytes carrying a LQTS2 mutation. Thus, the \textit{C. elegans} models of LQTS provide useful...
in vivo systems for reliable identification of functional correctors of LQTS2 mutations.

Our screening has identified two hERG modulators Prostratin and IDB that are PKC activators. The PKC signaling is known to act on cardiac sodium channels, K_A,TP channel, KCNQ1, and hERG in regulating the re-polarization of cardiac action potentials.44–46 We found that these two compounds restored the normal function of LQTS-related mutant hERG K+ channels by correcting the bioenergetics defect, without affecting wild-type hERG and cardiomycyte physiology. The PKCs may be a specific isoform of PKC that targets biogenic processing of hERG through phosphorylating the pore region of the channels. The selective and effective action of Prostratin and IDB is particularly important for their potential clinical applications. Prostratin is a natural product found in the bark of the mamala tree, and it has a potential in treatment of AIDS by specifically targeting the latent HIV reservoirs.47–48 Whether Prostratin and IDB are effective for treating LQTS patients remains to be further studied.

In summary, we demonstrate that C. elegans models of channelopathies are excellent systems for the discovery of ion channel drugs. The robustness of this method is evidenced by the fact that about ten hERG activators have been reported previously.49 Since mutations for channelopathies are often found, we proposed that C. elegans models of channelopathies could be generated for other channels besides the hERG K+ channel. Phenotypes of C. elegans channelopathy models are easy to be quantified, and hence a reasonable high-throughput screen could be done in a regular ion channel lab. Small-molecule screens in this study were done manually with about 250 drugs per person per day. Several high-throughput behavioral analysis platforms for C. elegans have been developed for different purposes.50–52 With proper automation in behavioral screening, we expect that a high-throughput screening system based on our method will become an efficient tool for drug discovery for channelopathies.

**Methods**

**Worm strains.** C. elegans strains were cultured on Nematode Growth Medium (NGM) at 20 °C. The wild-type N2 Bristol strain and the acs-20(nm3223) strain were obtained from Caenorhabditis Genetics Center and National BioResource Project, respectively. hERG*chimera/A536W/A561V transgenic worms were generated by expressing Punc-107::hERG*chimera/A536W::gfp and Punc-107::hERG*chimera/A536W/A561V::gfp plasmids in N2 worms, respectively. The Punc-107::mcherry plasmid was co-injected with the above plasmids to express mCherry as a cytoplasmic protein marker in these transgenic worms. The transgenic worms were integrated by the UV-TMP method53 and outcrossed for three times.

**Molecular biology.** The hERG*chimera was constructed by fusing UNC-103 (isoform a) N-terminus (1–799 amino acids) with hERG transmembrane/cNBD domains (369–889 amino acids) and UNC-103 C-terminus (599–829 amino acids). PCR product of HA-hERG cDNA was subcloned into the pCI-neo vector to construct plasmids that could be expressed in mammalian cell lines. Mutations were generated by site-directed mutagenesis. All constructs were confirmed by DNA sequencing.

**Immunoblot analysis.** Western blot analyses were performed as described below. Briefly, 16 h after transfection, HEK293T cells were treated with drugs and on the next day cells were lysed in the NP-40 lysis buffer (1% NP-40, 150 mM NaCl, 1 mM NaF, 50 mM Tris, pH 7.6) supplemented with protease inhibitor mixture (05892791001, Roche). After centrifugation at 12,000 × g at 4 °C for 10 min, the supernatants were collected for Western blot analyses (full blots found in Supplementary Fig. 10). The primary antibodies used in this study were rat anti-HA (118672301, Roche) with a dilution 1:3000, rabbit anti-PKCα (sc-214, Santa Cruz) with a dilution 1:1000, mouse anti-β-Tubulin (M30109, Abmart) with a dilution 1:3000.

**Immunocytochemistry.** The HEK293T cells or hiPSC-CMs were fixed with 4% paraformaldehyde for 20 min. After washed for three times by PBS, the fixed cells were permeabilized with 0.1% Triton X-100 for 10 min and were blocked with 10% bovine serum albumin for 1 h. For immunocytochemistry of HEK293T cells, mouse anti-HA (M20003, Abmart; 1:400), and rabbit anti-Calnexin were used as primary antibodies (ab22595, Abcam; 1:400); for immunocytochemistry of hiPSC-CMs, rabbit anti-tropinin 1 (sc-1368a, Santa Cruz; 1:500) or anti-hERG (ab196301, Abcam; 1:200) antibodies and mouse anti-phospho ERK (sc-7383, Sigma) were used as primary antibodies. Anti-mouse Alexa Fluor 594 and anti-rabbit Alexa Fluor 488-conjugated secondary antibodies were used for immunofluorescence detection.

**Behavioral assays.** Synchronized young adult worms were used for behavioral assays. For egg-laying assays, individual late-L4 hermaphrodite was picked on a NGM plate with E. coli strain OP50 and was picked away 24 h later. Total eggs and juveniles on the plate were counted. The locomotion speed was measured on the NGM plate without OP50. About 25 young adult worms were picked on the plate and allowed to recover for 1 min, and then their spontaneous movement was recorded for 1 min. The locomotion speed was calculated by using wrMTrck software. For male’s copulatory spicule protration assays, virgin L4 males were isolated on NGM plates seeded with OP50. After 24 h, male spicule protration was scored if at least one spicule partially extended from the cloaca. At least 40 worms per strain were tested in each experiment. The investors were blinded to the genotypes during C. elegans behavioral assays.

**Small-molecule screen.** Chemical libraries used in this study include the Prestwick Chemical Library, the Spectrum Collection, the Natural Products Library, the ICBN Known Bioactive Library, the IB screening Library, the NIH Clinical Collection, the FDA Approved Drug Library, the SIGMA LOPAC, and the Analyticon Discovery. 48-well screening plates were prepared by adding 500 μL NGM containing 1 mM IPTG and 25 μg ml−1 carbicin in each well. A mixture of HT115 E. coli expressing double-strand RNAs targeting tfl-2 and cIs5.7 was then seeded on NGM plates and allowed to grow for 2 days. Small molecules dissolved in DMSO were then added to NGM plates with a final concentration of 20 μM. After 4 h, synchronized L4 acs-20(hERG*)/A536W or acs-20(hERG*)/A536W/A561V worms were rinsed from their cultivating plates and deposited into each well of these screening plates. Phenotypes of tested worms were examined 36 h later.

**Electrophysiology.** Whole-cell patch-clamp was performed by using Axon 200 B. For recording currents in HEK293T cells expressing hERG channels, pipette solution consisted of (in mM) 135 KCl, 10 EGTA, 5 Mg-ATP and 10 Hepes (pH 7.2 with KOH), and bath solution consisted of (in mM) 130 NaCl, 5 KCl, 10 Hepes, 1 MgCl2, 10 glucose, and 2 CaCl2 (pH 7.4 with NaOH). Recording was performed at room temperature by using a standard protocol that membrane potential was first pre-depolarized to +40 mV to inactivate the channel, and then repolarized voltage from −140 mV to +40 mV (in 20 mV increments) to obtain tail current.

To measure I_Ca currents and cardiac action potentials, hiPSC-CMs were cultured on 0.1% gelatin coated coverslips in a 24-well plate. 3–8 days after seeding, hiPSC-CMs were treated with 2 μM IDB (sc-202663, Santa Cruz) or 3 μM Prostratin (P0077, Sigma) for 48 h. The I_Ca currents of hiPSC-CMs were isolated by applying a holding potential of hERG blocker E-4031 (MS060, Sigma), I_Ca was measured using a protocol that initially depolarizing the membrane potential from −60 mV to +60 mV in 30 mV increments, then repolarizing to −30 mV to elicit tail current. The composition of the pipette solution was (in mM): 120 KCl, 1 MgCl2, 3 Mg-ATP, 10 Hepes (pH 7.2 with KOH), and bath solution consisted of (in mM): 126 choline chloride, 5.4 KCl, 1.8 CaCl2, 1.0 MgCl2, 10 Hepes, and 10 glucose (pH 7.4). Nikalide (5 μM; N7634, Sigma) and chromanol (10 μM; C2613, Sigma) were added in the bath solution to suppress potential interference of I_Ca and I_Ks, respectively.

Action potentials were elicited by stimulating cardiomyocytes with depolarizing 10 ms-impulses from 0 pA to 40 pA in 4 pA increments. The frequency of stimulus for the protocol was about 0.5 Hz. The K+ solution contained (in mM) 140 NaCl, 5.4 KCl, 1.8 CaCl2, 1.0 MgCl2, 10 Hepes, and 10 glucose (pH 7.4). The action potential solution contained (in mM) 120 KCl, 1 MgCl2, 3 Mg-ATP, 10 Hepes, and 10 EGTA (pH 7.2).
repleted on 10 cm tissue-culture plates at around 20,000 cells per plate to allow for recovery in growth medium. Post-FACS, the cells were allowed to recover for a week, and then single colonies were manually picked and replated individually to wells of 96-well plates. Colonies were allowed to grow to near full confluence over the next 7 days, at which point they were split and replica-plated. Genomic DNA of colonies was extracted and genotyping at the CRISPR target site was identified by PCR amplification and DNA sequencing. Clones with confirmed heterozygous mutation allele or wild-type allele were expanded for further experiments.

Generation of cardiomyocytes from hiPSC. Generation of cardiomyocytes from hiPSC was performed by following the previously reported method with modifications. Cardiomyocyte chimically defined cardiac differentiation kit (CA2004500, CellSync) was used for differentiation of hiPSC into cardiomyocytes. Briefly, human hiPSC were split into 0.5 mM EDTA PBS solution at 1:10 ratio and cultured with PSeasy chemically defined medium for 3 days. When reached ~80% confluence, hiPSC was cultured in Induction Medium II for 2 days then in Induction Medium III. Induction Medium III was changed every other day afterwards. Spontaneously contracting cells were observed from the 8 day after induction. The derived cardiomyocytes were purified with Cardiomyocyte purification medium at the day 15 after induction.

Statistical analysis. Statistical analysis was done using GraphPad Prism 6.0. Data shown are mean ± s.e.m. For the C. elegans behavior data, unpaired Student’s t-test was used to determine significant differences between samples. For electrophysiological data, one-way ANOVA Dunnett’s test was used when comparing multiple groups. Significance levels: *P < 0.05, **P < 0.01, ***P < 0.001. The ALA inhibition curve was fitted with equation: \( y(x) = \frac{y_{max} + y_{min}}{1 + e^{(x-x_0)/V}} \).

Data availability All data supporting the findings of this study are included in this article (and its Supplementary Information files). Further data are available from the corresponding author upon reasonable request.

Received: 20 March 2018 Accepted: 31 August 2018

Published online: 26 September 2018

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Author contributions
Q.J., K.L. and S.-Q.C. designed all the experiments. Q.J. and K.L. performed most of the experiments. Q.J. generated the C. elegans models of channelopathies, performed small-molecule screening assays, and the electrophysiological experiments. K.L. performed the biochemical assays. W.-J.L. conducted the cardiomyocyte differentiation of the hiPSCs. S.L. and Q.J. generated the CRISPR/Cas9-edited hiPSC clones. X.C. performed the electrophysiological recordings of some of the hERG mutant channels expressed in HEK293T cells. J.Y. and X.-J.L. assisted in carrying out the small-molecule screenings. Q.J. and K.L. performed the behavioral assays. F.L. supervised the cardiomyocyte differentiation experiments and analyzed the data. Q.D. supervised the CRISPR/Cas9 editing experiments and analyzed the data. Q.J., K.L., F.L., Q.D. and S.-Q.C. prepared the figures and wrote the manuscript.

Additional information
Supplementary Information accompanies this paper at https://doi.org/10.1038/s41467-018-06514-5.

Competing interests: W.-J.L. is a co-founder of Beijing Cellapy Biological Technology Co., Ltd. The remaining authors declare no competing interests.

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Acknowledgements
We thank Moming Poo and Ying Liu for critical reading of the manuscript, Mei-Yu Ruan for technical assistance, the C. elegans Genetics Center (funded by NIH Office of Research Infrastructure Programs P40 OD010440) and S. Mitani (Tokyo Women’s Medical University School of Medicine) for providing us strains and National Compound Resource Center for providing us small-molecule libraries. This work was supported by grants to S.-Q.C. (the Strategic Priority Research Program of Chinese Academy of Science, XDBS1020100 and the National Natural Science Foundation of China, 31471149 and 81527901), to F.L. (the National Natural Science Foundation of China, 81422003), and to Q.D. (the National Key R&D Program of China, 2017YFA0102800 and 2017YFA0103700).

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