# Review

## Ventricular voltage-gated ion channels: Detection, characteristics, mechanisms, and drug safety evaluation

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### Graphical Abstract

1. The structure, distribution, interaction and function of voltage-gated ion channels (VGICs) maintains a physiological balance of the ionic currents, normal action potential (AP), excitation-contraction coupling, and synchronization in cardiomyocytes.

2. Various genetic mutations and molecular dysregulation of the VGICs could result in ionic imbalance, abnormal AP waveform, and even cardiac arrhythmia.
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Abstract
Cardiac voltage-gated ion channels (VGICs) play critical roles in mediating cardiac electrophysiological signals, such as action potentials, to maintain normal heart excitability and contraction. Inherited or acquired alterations in the structure, expression, or function of VGICs, as well as VGIC-related side effects of pharmaceutical drug delivery can result in abnormal cellular electrophysiological processes that induce life-threatening cardiac arrhythmias or even sudden cardiac death. Hence, to reduce possible heart-related risks, VGICs must be acknowledged as important targets in drug discovery and safety studies related to cardiac disease. In this review, we first summarize the development and application of electrophysiological techniques that are employed in cardiac VGIC studies alone or in combination with other techniques such as cryoelectron microscopy, optical imaging and optogenetics. Subsequently, we describe the characteristics, structure, mechanisms, and functions of various well-studied VGICs in ventricular myocytes and analyze their roles in and contributions to both physiological cardiac excitability and inherited cardiac diseases. Finally, we address the implications of the structure and function of ventricular VGICs for drug safety evaluation. In summary, multidisciplinary studies on VGICs help researchers discover potential targets of VGICs and novel VGICs in heart, enrich their knowledge of the properties and functions, determine the operation mechanisms of pathological VGICs, and introduce groundbreaking trends in drug therapy strategies, and drug safety evaluation.

KEYWORDS
action potentials, cardiac voltage-gated ion channel, cardiovascular safety evaluation, electrophysiological techniques
1 | INTRODUCTION

The cardiac cycle begins when an action potential (APs) is spontaneously generated in the sinoatrial node (SAN), the primary pacemaker in the heart. The coordinated propagation of synchronized electrical impulses relies on effective cooperation among various components in the heart system to maintain cardiac rhythm.1 Specifically, the AP from the SAN passes sequentially through the atria, the atroventricular node (AVN), and His-Purkinje conducting tissue before ultimately reaching the ventricles.1 APs, which are generated and modulated by the opening and closing of ion channels in the plasma membrane, are the fundamental electrical excitation signals responsible for the beating of cardiomyocytes and are distinct among various components in the heart due to various expression of ion channels.1 Among the voltage-gated ion channels (VGICs) involved in ventricular APs, sodium (Na⁺), potassium (K⁺), and calcium (Ca²⁺) channels (Nav, Kv, and Cav channels, respectively) predominate.2 The functions of VGICs, the mechanisms underlying cardiac physiology and pathology, and appropriate diagnostic and treatment strategies have been explored for decades through electrophysiological techniques. Moreover, these techniques have been developed and expanded over time; from laborious, low-throughput methods limited to whole-cell experiments, they have been refined into automated, high-throughput methods. These developments have dramatically augmented the ability of researchers to further explore VGICs. In the real world, aside from genetic mutations affecting VGICs, many drugs can bind to these channels, block ion flow and disrupt the regulation of APs, potentially leading to drug-induced arrhythmia, or “proarrhythmia.”3,4 It is necessary to evaluate the risks of potential drug candidates by using the different techniques mentioned above,5,6 according to the US Food and Drug Administration (FDA) guidelines. In this respect, enhancing the quality of preclinical safety screening is particularly important for validating the safety of therapies to avoid potential adverse effects on ion channels and prevent billions of dollars in losses because of late-stage premarket drug withdrawals in the development process of drug development before marketing.

2 | METHODS FOR DETECTING CARDIAC VGICS

2.1 | Electrophysiological techniques

The manual patch clamp (MPC) technique (Table S1) is the gold standard for analyzing electrophysiological characteristics (APs and specific ion channel currents) in cardiac myocyte research studies under physiological/pathological conditions or in response to drug application. Three main cell models are used: 1) freshly isolated ventricular myocytes from wild-type (WT), diseased or genetically modified animal models; 2) heterologous expression systems specifically expressing the human ion channels of interest; and 3) human induced pluripotent stem cell-derived cardiomyocytes (hiPSC-CMs) from healthy individuals and patients.2,7 In addition, cardiac ion channels can be examined by single-channel MPC recording (Table S1); for example, this technique can be applied to a potential new channel with a putative pore-containing structure,8 or channels that cannot be expressed or trafficked on the cell membrane in heterologous expression systems;9 or channels that are potentially altered in the diseased heart.9 The main limitation of MPC is its low throughput. Therefore, the automated patch clamp (APC) (Table S1) enables much higher-throughput experiments while nevertheless achieving high-quality seals, thereby facilitating the use of the MPC and is now routinely used in cardiac drug discovery and safety testing.10–12

In addition, microelectrode arrays (MEAs) (Table S1) offer an alternative noninvasive that enables noninvasive, high-throughput assays evaluating extracellular field potential (EFP) of excitable cells13; MEAs have also been increasingly used in cardiology to test the safety of drug candidates.13,14 Furthermore, impedance techniques (Table S1) have recently been combined with EFP recording on the same platform to provide a noninvasive, high-throughput and long-term measurement strategy for assessing the synchronous beating of monolayer cardiomyocytes, the duration of EFPs, and the proarrhythmogenic capacity of drug candidates without altering cellular physiology15; this combined approach offers a more comprehensive analysis of excitation-contraction (EC) coupling than either component alone.

Generally, low-throughput MPC is a critical tool for examining the electrophysiological characteristics of cardiac cells and the biophysical properties and functions of ion channels. APC and MEAs have developed into an indispensable platform for pharmaceutical companies and academic laboratories to conduct potential drug target discovery, drug screening, and cardiac safety with high efficiency and accuracy. hiPSC-CM-based APC,16 MEA,17 and impedance15 screening assays are increasingly used to evaluate antiarrhythmic effects, adverse effects or interindividual variations in patients or healthy individuals and to acquire more comprehensive validation data.
2.2 Joint techniques

Cryoelectron microscopy (cryo-EM) (Table S2), which can resolve the structure of macromolecular complexes at the atomic conformation level, has provided researchers with a more in-depth molecular picture of ion selectivity, voltage gating, and intersubunit interactions in channel complexes and thereby provides insights into important biological phenomena, such as electrophysiological feature variations among different VGIC isoforms, feature changes after the application of various compounds, and the mechanisms of mutation-related arrhythmia. Hundreds of disease-associated missense mutations have been mapped onto all major domains in the structure of many VGICs. Cryo-EM structure analysis could provide novel insights into both VGIC-drug interactions and the mechanisms of action of such drugs. Optical imaging techniques can help evaluate whether the functional properties of truncated or mutated VGICs purified for cryo-EM analysis are similar to those of WT full-length VGICs.

Optical imaging methods (Table S2) using voltage- or Ca²⁺-sensitive dyes are less invasive than MPC and are able to measure changes in the MPs, intracellular calcium concentrations, electrical activity and EC coupling of cardiac cells. However, some sensitive dyes are limited by cytotoxicity and short half-lives. Genetically encoded fluorescent Ca²⁺ indicators, such as ArcLight and GCaMP, were developed and applied to cardiac research to monitor functional changes in hiPSC-CMs in a long-term, noninvasive, high-throughput manner. The combination of optical imaging and electrophysiological techniques allows simultaneous recording of optical AP signals and calcium transient signals and permits both high spatial resolution and accurate functional evaluation.

Optogenetics approaches (Table S2), using light to control the perturbation of membrane voltage through the opening of optogenetic channels have been used to modulate cardiomyocyte excitability and heart rate with high precision and to explore the mechanisms of arrhythmia generation. Optogenetic channels can also be used to study the relationship between cardiac myocytes and nonmyocyte cells and provide a feasible way to explore direct evidence of electrical coupling between these cells in normal or injured regions of the heart. Automated frequency-dependent cardiotoxicity screening can be conducted by applying optogenetic stimulation similar to physiological heart rates in hiPSC-derived cardiomyocytes.

3 AP GENERATION AND EC COUPLING OF CARDIOMYOCYTES

3.1 Normal electrophysiology of AP and EC coupling

A typical ventricular AP consists of five phases (P0-P4) that are mediated by different depolarizing and repolarizing ionic currents (Figure 1A). The initial phase (Phase 0) of a cardiac AP occurs after the resting state (Phase IV) of the previous AP and arises from a very large inward \( I_{Na} \) current mediated by Nav channels. Then, \( I_{Kv} \) channels are activated to mediate transient outward potassium currents \( I_o \), leading to partial repolarization in Phase I. During Phase II, L-type Cav channels (LTCCs) are activated, generating an inward \( I_{Cal} \) current. In addition, the \( Na^+ / Ca^{2+} \)-exchanger (NCX) opens in forward mode and mediates an inward \( I_{NCX} \) current by exchanging an influx of 3Na⁺ for an efflux of 1Ca²⁺. On the other hand, the voltage-gated delayed rectifier potassium channels open and mediate outward rectifier currents \( I_{Kr} \) and \( I_{Ks} \). Membrane potential (MP) changes extraordinarily little due to the nearly equal inward and outward currents during this phase, which is also known as the plateau phase. In the late plateau phase, LTCCs are inactivated, and the dominant outward currents, \( I_{Kr} \) and \( I_{Ks} \), result in repolarization in Phase III. Toward the end of Phase III, \( I_{Kr} \) and \( I_{Ks} \) decline, and the inwardly rectifying potassium channels Kir2.1x mediate the \( I_{K1} \) current to drive repolarization and maintain a resting MP (Phase IV).

The beating of the heart relies on EC coupling (Figure 1B). During AP generation, the LTCC-mediated increase in the cytosolic Ca²⁺ concentration instantaneously triggers the opening of the ryanodine receptor 2 (RyR2) channel, a Ca²⁺ channel in the sarcoplasmic reticulum (SR), which causes Ca²⁺ release from the SR, and thereby further increases the cytosolic Ca²⁺ concentration. This Ca²⁺-induced Ca²⁺-release (CICR) prompts Ca²⁺-sensing protein troponin C to initiate contraction (systole). Cytosolic calcium levels are reduced via the Ca²⁺-ATPase type-2a (SERCA2) and the NCX-mediated influx of Ca²⁺ back into the SR and the NCX-mediated influx of Ca²⁺ back to the extracellular space, resulting in the dissociation of calcium and troponin and then muscle relaxation (diastole).

3.2 Abnormal electrophysiology as a trigger of arrhythmias

Disruption of the normal generation and duration of Aps is associated with arrhythmias in the heart. Two types
**FIGURE 1** Normal AP generation and EC coupling of cardiomyocytes; abnormal electrophysiology as a trigger of arrhythmias. (A) A typical ventricular action potential (AP) and the depolarizing and repolarizing ionic currents underlying its different phases (P0-P4). (B) Voltage-gated ion channel (VGIC) distribution and contribution to AP excitation-contraction coupling in cardiomyocytes. Na⁺, Ca²⁺, and K⁺
of afterdepolarizations, early afterdepolarizations (EADs) and delayed afterdepolarizations (DADs), could induce premature APs and contribute to arrhythmias. EADs occur during Phase II or III (Figure 1C). Prolongation of action potential duration (APD) due to the reduction of repolarization currents (e.g., I_{Kr} and I_{Ks}) or the increase in I_{Na,L} current could give rise to abnormal recovery from the inactivation of LTCC channels and further depolarize the membrane due to the reactivation of inward currents I_{Ca,L}. DADs can result from depolarization after the end of AP repolarization (Figure 1C), potentially due to Ca^{2+} overload caused by enhanced SR Ca^{2+} release and the inappropriate activation of the reverse mode of NCX, which mediates outward I_{NCX} current by exchanging an influx of 1Ca^{2+} for an efflux of 3Na^+.1,2

Under physiological conditions, Nav channel activity is regulated by cytosolic Ca^{2+} levels, such as elevation of cytosolic Ca^{2+} levels resulting in destabilization of inactivation and increase of the amount of available channels to open for the next AP.33–35 Na^+ influx, in turn, affects the modulation of cytosolic Ca^{2+} levels.32 However, under pathological conditions, an abnormal increase in Na^+ during diastole can result in inappropriate timing of reverse flow through the NCX channel (3Na^+ influx and 1Ca^{2+} influx), further increasing the cytosolic Ca^{2+} concentration and altering normal EC coupling (Figure 1D).

4 | VENTRICULAR AP-RELATED ION CHANNELS: CLASSIFICATION, STRUCTURE, FUNCTION, REGULATION, AND DISEASE RELEVANCE

4.1 | Nav channels

Cardiac voltage-gated Nav channels initiate AP in electrically excitable cells. The specificities among isoforms (Table 1) are attributed to the distinct α-subunit encoded by the corresponding gene and the different combinations of β subunits.36 β subunits regulate channel surface expression, voltage dependence and gating kinetics.36 SCN5A-encoded Nav1.5 is the most abundantly expressed Nav channel in ventricle and atrium (Table 1) and is responsible for the generation of APs and the conduction of cardiac impulses in cardiomyocytes.18,34,37,38 Additional evidence has shown that other isoforms are also expressed in the ventricular myocytes (Table 1).32,39–48

4.1.1 | Nav1.5

In ventricle cardiomyocytes, Nav1.5 channels are known to be located in lateral membrane, transverse tubules (T-tubules), and intercalated discs, ensuring propagation of
**Table 1** Cardiac voltage-gated Na⁺ channels subtypes

| Subtypes   | Encoding α subunits Gene | Auxiliary subunits | Main location | Subcellular localization in cardiac tissue (V/A/SAN) and region | Cryo-EM structure | TTX sensitivity | Principal physiological functions in human ventricle myocytes |
|------------|--------------------------|--------------------|---------------|---------------------------------------------------------------|-------------------|----------------|-------------------------------------------------------------|
| Nav1.1     | SCN1A                    | β4 encoded by SCN4B | CNS, Heart    | V≈A=SAN; T-tubules                                            | Human Nav1.1-β4 channel | Sensitive       | Cardiac pacemaking and promotes Ca²⁺ dynamics               |
| Nav1.2     | SCN2A                    | β2 encoded by SCN2B | CNS, Heart    | V≈A=SAN T-tubules                                            | Human Nav1.2-β2 subunit | Sensitive       | Contributes small portion to cardiac sodium current          |
| Nav1.3     | SCN3A                    | NR                 | CNS, Heart    | V≈A=SAN T-tubules                                            | NR                 | Sensitive       | Contributes small portion to cardiac sodium current          |
| Nav1.4     | SCN4A                    | β1 encoded by SCN1B | Skeletal muscle, Heart | V≈A≤SAN T-tubules                                      | Electric eel human Nav1.4-β1 subunit | Sensitive       | Contributes small portion to cardiac sodium current          |
| Nav1.5     | SCN5A                    | β1, β2 encoded by SCN1B and SCN2B | Heart | V≈A≤SAN; IDs, lateral membrane, T-tubules                                      | Rabbit Nav1.5-β1, β2 subunits | Resisitant       | Mediates the entry of Na⁺, and triggers overshooting of AP    |
| Nav1.6     | SCN8A                    | β1 encoded by SCN1B | CNS, PNS, Heart | V≈A T-tubules                                               | NR                 | Sensitive       | Contributes small portion to cardiac sodium current; Promote Ca²⁺ dynamics |
| Nav1.8     | SCN10A                   | β2 encoded by SCN2B | PNS, Heart    | V<A T-tubules                                               | NR                 | Resisitant       | Cardiac contraction and conduction                         |

CNS, central nervous system; PNS, peripheral nervous system; IDs, intercalated discs; T-tubules, transverse tubules; Cryo-EM, cryoelectron microscopy; TTX, tetrodotoxin; V, ventricle; A, atrium; SAN, sinoatrial Node; NA: not available; NR, not reported.

electrical impulse in longitudinal, transverse directions of cardiomyocytes, and between adjacent ones, respectively (Figure 1B).⁴⁹ Nav1.5 channels are closed at the resting MP (Phase IV). In response to membrane depolarization, Nav1.5 could be activated. Within 200-300 µs, a large inward peak $I_{Na}$ ($I_{Na,P}$) is formed to trigger overshooting of AP in Phase 0. At the end of this phase, most Nav1.5 channels are rapidly inactivated within 2-5 ms, rendering the channel refractory until repolarization is completed in Phase III. During Phase IV, after recovering from inactivation, the channels are closed and can again be reopened by membrane depolarization (Figure 2A). In Phase II or III, a small population of total Nav1.5 channels could be reactivated before complete inactivation and then generate a relatively small, persistent sodium cardiac inward current called late $I_{Na}$ ($I_{Na,L}$).⁵⁰,⁵¹ Under physiological conditions, $I_{Na,P}$ but not $I_{Na,L}$ plays a central role in ventricular AP generation, while under pathological conditions, $I_{Na,L}$ can play an important role.⁵⁰,⁵¹ Abnormal increases in $I_{Na,L}$ prolong the duration of the AP plateau, triggering EADs or further elevating intracellular Ca²⁺ levels by driving the NCX exchanger to function in reverse mode, thereby inducing DADs and contributing to arrhythmogenesis.⁵¹ To date, some high-resolution structures of Nav channels,²¹,⁵²,⁵³ including rNav1.5C,¹⁸ have been elucidated (Table 1). In general, key structural features of Nav1.5, the structural basis for its physiological function and its dysfunction in cardiac arrhythmias could be learned from the revealed Nav1.5 structures. Unlike other Nav α subunits, in Nav1.5, the regulatory interface with auxiliary β1 and β2 subunits, encoded by SCN1B and SCN2B respectively, is not as strong due to the substitution
CARDIAC VOLTAGE-GATED SODIUM CHANNEL (Nav1.5) STRUCTURE, ACCESSORY PROTEINS AND SIGNALING.

(A) The contribution of \( I_{Na,P} \) (upper) mediated by Nav1.5 to action potential (AP) Phase 0 (lower). (B) The cardiac voltage-gated sodium channel Nav1.5 comprises one \( \alpha \) subunit and one or two auxiliary \( \beta \) subunits. The \( \alpha \) subunit consists of four homologous but nonidentical repeats (DI-DIV) connected by cytoplasmic linkers (ICL I-II, ICL II-III, and ICL III-IV) and is responsible for voltage dependence, pore formation, and surface expression. Each domain contains SI-S6 subunits that are connected by loops located intra- and extracellularly. The carboxyl-terminal domain (CTD) of SI and the amino-terminal domain (NTD) of S6 are located in DI and DIV, respectively. The S4 subunit of each domain contains the voltage sensor. S5 and S6 of each repeat form the pore domain, and the connecting P-loops between the S5 and S6 regions curve back into the pore to form the extracellular selectivity filter (SF), which is responsible for \( Na^+ \) selectivity. ICL III-IV serves as an inactivation gate (IG), which closes the pore within 1-2 ms after opening. The \( \beta \) subunit, consisting of an extracellular domain, an intracellular domain, and a single transmembrane helix, plays an important role in modulating the surface expression, kinetics, and functions of Nav channels. The \( \beta 1 \) and \( \beta 2 \) subunits do not stably associate with the Nav1.5 \( \alpha \) subunit. Y304 in the Nav1.7 \( \alpha \) subunit, which is connected to E48 in the \( \beta \) subunit by a hydrogen bond, is substitute at L316 at the equivalent position in Nav1.5. Similarly, C895, which forms a disulfide bond with the \( \beta 2 \) subunit, is substituted at L869. Several accessory proteins interact with Nav1.5 channels to form macromolecular complexes that regulate ion trafficking, posttranslational modifications, and gating. Nav1.5 activity is driven by Ca\(^{2+}\) levels via its interactions with CaM and CaMKII. Nav1.5 can be activated in response to \( \beta \) adrenergic stimulation via the activation of CaMKII. CaM binds not only to an IQ motif in the CTD of Nav1.5 but also to its IG. The CaMKII inhibitor KN93 was recently reported to alter the kinetics of Nav1.5 inactivation by interrupting the CaM-IG interaction but did not suppress CaMKII function.

Recent studies suggest \( \beta 1 \) subunit may differentially control expression and function of \( \alpha \)-subunit via acute and chronic feedback mechanisms. \( \beta 2 \) is pivotal for the correct localization of NaV1.5. Nav1.5 is insensitive to the inhibition of tetrodotoxin (TTX), a selective sodium channel blocker nonprotein toxin, due to the substitution of binding residues at the outer mouth of the Nav1.5 selectivity filter (SF). Nav1.5 is blocked by the class Ic antiarrhythmic drug flecainide with comparatively high affinity and slow binding kinetics compared to class IA and IB antiarrhythmic drugs due to the larger hydrophobic ring structure of flecainide interacting with the central cavity of Nav1.5. In addition, the structural template of Nav1.5 for arrhythmia mutations provides a better understanding of the mechanism of various SCN5A variants in different positions. The traditional view is that a single \( \alpha \)-subunit of Nav1.5 functions as a monomer, while increasing evidence has shown that the \( \alpha \)-subunit of Nav1.5 could be oligomerized into dimers within the first intracellular loop and results in coupled gating properties with the accessory protein14-3-3 interaction. Inhibition of 14-3-3 could abolish the dominant negative (DN) effect and the biophysical coupling between \( \alpha \)-subunits. In addition, several accessory proteins have been demonstrated to interact directly with the \( \alpha \) subunit of Nav1.5 channels (Figure 2B) to form macromolecular complexes with Nav1.5 and modulate the expression, trafficking, and biophysical function of Nav1.5 (Table 2). Calmodulin (CaM), a ubiquitously expressed calcium-binding...
| Accessory proteins | Types | Binding sites | Biophysical function | Techniques |
|--------------------|-------|---------------|---------------------|------------|
| 14-3-3η            | Adaptor protein | ICL_{1,II}     | Regulates Na\(^{+}\) current, negatively shifts the Nav1.5 inactivation curve and postpones recovery from inactivation without affecting the current density or Nav1.5 activation curve\(^{241}\) | MPC        |
| API-γ              | Adaptor protein | Y1810 in the CTD | Forms a recognition site for Golgi to incorporate Nav1.5 into clathrin-coated vesicles and transports to target membrane | MPC, immunostaining, mutagenesis |
| Ankyrin-G          | Anchoring-adaptor protein | ICL_{II,III} | Regulates the accumulation of Nav1.5 on the membrane\(^{63,242}\); E1053K eliminates ankyrin-G binding, displays a faster onset of inactivation and slower recovery from inactivation, and negatively shifts channel activation curve\(^{241}\); 50% peak I\(_{Na}\) is reduced in Ankyrin-G KO myocytes\(^{63}\) | MPC, Co-IP, immunostaining, mutagenesis |
| α-actinin 2        | Cytoskeletal protein | ICL_{III,IV} | Anchors Nav1.5 to the actin cytoskeleton network and increases sodium channel density without affecting gating properties\(^{243}\) | MPC; Co-IP |
| MOG1               | Cofactor | ICL_{II,III} | Regulates the expression of Nav1.5, increases sodium current densities\(^{244}\) | MPC, Co-IP, WB |
| Synapse-associated protein 97 (SAP97) | Anchoring-adaptor protein | PDZ domain in CTD | Forms a complex with ankyrin-G to regulate appropriate Nav1.5 expression, silencing SAP97 significantly reduces Na\(^{+}\) current\(^{245}\) | MPC, WB, immunostaining |
| α1-Syntrophin      | Anchoring scaffold protein | PDZ domain in NTD and CTD | Enhances the expression of Nav1.5 and increases I\(_{Na}\)\(^{246}\) | MPC, Co-IP, WB, immunostaining |
| Calmodulin (CaM), a calcium-binding protein | Regulatory protein | IQ-motif in CTD | Modulates slow inactivation thereby increasing the open probability of Nav1.5\(^{57}\) | MPC, WB, immunostaining |
| CaM-dependent protein kinase II (CaMKII) | Kinase | ICL_{I-II} | Destabilizes inactivation gate and promotes faster recovery from inactivation\(^{57-61}\); a CaMKII inhibitor KN93 interrupting the CaM-IG interaction inhibits Nav1.5 recovery from inactivation without altering the kinetics of inactivation but does not suppress the inhibitory function of CaMKII\(^{64}\) | MPC, WB, immunostaining |
| Protein tyrosine phosphatase (PTPH1) | Enzyme | PDZ domain in CTD | Shifts the Nav1.5 hyperpolarized potentials\(^{251}\) | MPC, WB |
| Caveolin-3         | Scaffolding and regulatory protein | Colocalize in the sarcolemma of human ventricle tissue\(^{252}\); Macromolecular complex including Nav1.5/α1-Syntrophin/ nNOS/Caveolin-3\(^{31}\) | Serves as a negative regulator for cardiac I\(_{Na,L}\) via suppression of nNOS-dependent direct S-nitrosylation of Nav1.5\(^{51}\) | MPC, Co-IP, immunostaining |

CTD, carboxyl terminal domain; NTD, N terminal domain; MPC, manual Qatch clamp; WB: Western blot; co-IP, coimmunoprecipitation.
protein, and CaM-dependent protein kinase II (CaMKII), an adrenergically activated kinase, serve as important components affecting channel function. CaM binds with IQ motif of Nav1.5 carboxyl-terminal domain (CTD) in Ca^{2+}-free forms and Ca^{2+}-bound forms at the basal levels of intracellular Ca^{2+} concentration. While this CaM-Nav1.5 interaction is altered when the elevation of intracellular Ca^{2+} concentration, therefore changing the rate of Nav1.5 inactivation. Increasing evidence has shown that at the high level of intracellular Ca^{2+} concentration, CaM also directly binds to the inactivation gate (IG) of Nav1.5 to destabilize the IG and promote faster recovery from inactivation. CaMKII not only phosphorylates Nav1.5 at key site S571 but also directly interacts with Nav1.5 to regulate the expression and function of Nav1.5. On the other hand, protein phosphatase 2A (PP2A) was recently found to interact with the Nav1.5/ankyrin-G/CaMKII/Biv-spectrin macromolecular complex and balance CaMKII-dependent phosphorylation. The CaMKII inhibitor KN93 but not autacamide-2-related inhibitory peptide (AIP) could interrupt the CaM-IG interaction by forming the ternary complex CaM-IG-KN93 and then inhibit Nav1.5 recovery from inactivation without altering the kinetics of inactivation. Therefore, determining the effects of accessory proteins and signaling pathways on modulating Nav1.5 provides us with a more comprehensive understanding of Nav1.5 roles in cardiac tissues in both health and disease states and is beneficial for the discovery of potential drug targets. Future investigations of the kinetics of CaM-Naıv complexes and the effects of structure-guided mutations on the roles of Nav1.5 in the absence/presence of Ca^{2+} transients will provide us with a more comprehensive understanding of the mechanisms and significance of Ca^{2+}-dependent Naıv roles in cardiac tissues in both healthy and disease states.

Mutations in SCN5A (Table 3) are associated with inherited life-threatening arrhythmias, such as long QT syndrome type 3 (LQT3), and Brugada’s syndrome (BrS). Slower channel inactivation and thus conducting an increase in I_{Na,L} are responsible for the gain-of-function (GOF)-associated LQT3. On the other side, reduction of membrane expression of functional channel due to synthesis deficiency, trafficking defects, impairment of gating (such as slower activation or faster inactivation) or permeation disruption cause SCN5A loss-of-function (LOF)-associated BrS. The α-subunit of Nav1.5 oligomerization also explains the existence of several BrS variants displaying DN effects, providing new therapeutic targets for BrS caused by SCN5A LOF variants. Moreover, LOF mutations in SCN1B and SCN2B are also implicated in BrS. In addition to most isolated GOF or LOF variants of SCN5A which are typically associated with a distinct clinical and electrocardiographic phenotype, variants could also lead to overlapping syndromes77,78 or inherited arrhythmia syndrome different from BrS and LQT3. There is also a category of benign (atypical) SCN5A mutations which shows normal function alone but leads to a reduction in sodium currents when coexpressed with WT in vitro as typical SCN5A BrS mutations do.80 Moreover, missense variants in CAV3-encoded caveolin-3, which forms macromolecules with Nav1.5 and serves as a negative regulator for I_{Na,L}, could result in I_{Na,L} increase and thus cause LQT59, providing new therapeutic strategies to correct I_{Na,L}. Drugs that inhibit I_{Na,L} could shorten the AP duration or QT interval and could therefore be considered a potential treatments for I_{Na,L}-associated diseases.82 Thus, abnormal changes in I_{Na,L} could be considered as a target for drug development and safety evaluation.

High-throughput assays of cardiac Nav1.5 I_{Na,P} have been widely used in cardiac safety screening, but screening studies do not routinely measure I_{Na,L}. However, it is important for potential therapeutic candidates that could minimize I_{Na,L} without affecting I_{Na,P} to be selected. The variety of different protocols and measurement strategies applied in the use of these drugs have contributed to remarkable variations in the reported data on I_{Na,L} and screening results for inhibitory compounds.84,85 I_{Na,L} is small, and studies have had difficulty generating reproducible data; thus, the best choice for an I_{Na,L} enhancer should increase I_{Na,L} with no obvious effect on I_{Na,P}. In addition, it is necessary to double check the median inhibitory concentration (IC_{50}) of potential drugs in the absence of enhancers, eliminating the modification effect of enhancers on the activity of compounds, and to evaluate the IC_{50} of drugs in different stimulation states with regard to variations in the effects on I_{Na,P} and I_{Na,L} in different stimulation states.88

4.1.2 Other Nav channels in the heart

TTX-sensitive Nav channels including neuronal Nav (eg, SCN1A-encoding Nav1.1, SCN2A-encoding Nav1.2, SCN3A-encoding Nav1.3, SCN8A-encoding Nav1.6), which were first identified in neurons, and skeletal muscle Nav (eg, SCN4A-encoding Nav1.4), which was first identified in skeletal muscle, have been unexpectedly found in T-tubules of ventricle myocytes (Table 1), contributing a small portion to the total sodium current under physiological conditions due to their much lower expression level than Nav1.5. While, in inherited forms of cardiac arrhythmia, augmentation of TTX-sensitive Nav channels phosphorylated by β-AR stimulation/CaMKII stimulation, contributes to abnormal increases in I_{Na,L} and arrhythm-
| Subtypes | Encoding subunits gene | Congenital syndrome | Gain or loss of function | Mechanisms underlies the phenotype | Examples of variants |
|----------|------------------------|---------------------|--------------------------|-----------------------------------|---------------------|
| Nav1.1   | SCN1A                  | Cardiac arrhythmia contributed to SIDS with SUDEP | LOF                      | Haploinsufficiency                | R222X<sup>35</sup>: increases transient I<sub>Na</sub> density, incidence of arrhythmogenic AP, EADs, DADs and rates of spontaneous contraction in DS patient iPSC-CMs. |
|          |                        |                     |                          |                                   | G682Y<sup>36</sup>: decreases sodium current in tsA201 cells expressing variant. |
| Nav1.4   | SCN4A                  | Myotonia overlapping with BrS | NR                       | NR                                | V781I<sup>37</sup> |
|          |                        | Myotonia overlapping with prolonged QTc intervals | GOF                      | Gating defects                   | R1448C<sup>46</sup>: shows slower of inactivation and faster recovery time. |
|          |                        |                     |                          |                                   | R1463S<sup>38</sup>: shows slower of inactivation and faster recovery time. |
|          |                        |                     |                           |                                   | V1442M<sup>38</sup>: shows enhancement of fast inactivation. |
|          |                        |                     | LOF                      | Gating defects                   | E1520K<sup>38</sup>: shows reduction of current density. |
| Nav1.5   | SCN5A                  | LQTS3               | GOF                      | Gating defects                   | F1473C<sup>49</sup>: removes complete inactivation and thus conducting increase of I<sub>Na</sub,L. A993T<sup>50</sup>: shows slower inactivation kinetics. |
|          |                        |                     |                          |                                   | W822X<sup>58</sup>: leads to the haploinsufficiency of the Nav1.5 protein and thus resulting in a nearly 50% reduction in Na<sup>+</sup> current amplitude without significant alterations of biophysical properties and any dominant-negative activity on wild type channels. |
| BrS      | LOF                    | Synthesis deficiency |                          |                                   | D1690N<sup>59</sup>: produces a marked DN effect when cotransfected with wild-type channels. |
|          |                        | Trafficking defects |                          |                                   | R878C<sup>50</sup>: nonconductive channel |
|          |                        | Trafficking normal but gating defects |                          |                                   | N1541D<sup>71</sup>: induces an accelerated entry into closed-state inactivation. |
|          |                        | Gating defects      |                          |                                   | R1632C<sup>71,72</sup>: produces an enhanced fast-inactivated state stability because of a pronounced impairment of recovery from fast inactivation. |
|          |                        | Trafficking defects with gating defects |                          |                                   | R1629Q<sup>96</sup>: produces enhanced inactivation properties with a large hyperpolarizing shift in steady-state inactivation, current densities similar to WT. |
|          |                        | Permeation disruption |                          |                                   | A1924T<sup>79</sup>: reduces calmodulin binding and stabilizes Nav1.5 inactivation. |
|          |                        |                          |                          |                                   | G1748D<sup>69</sup>: produces a marked DN effect, positively shifts the activation curve. |
|          |                        |                          |                          |                                   | E90K<sup>77,78</sup> |

(Continues)
| Subtypes                   | Encoding subunits gene | Congenital syndrome | Gain or loss of function | Mechanisms underlies the phenotype | Examples of variants                                                                                                                                 |
|----------------------------|------------------------|---------------------|--------------------------|-----------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------|
| Benign (atypical) BrS      |                         | Normal function as WT | Normal                   |                                   | L567Q<sup>16</sup>: remains relatively unchanged current density, voltage-current relationship, steady-state inactivation, and recovery from inactivation, insufficient to produce a BrS phenotype. While, exerts DN effect on coexpression with WT via deficient trafficking mechanism. |
| Overlap of BrS and LQTS3   |                         | Overlap GOF & LOF |
|                           |                         | Gating defects       |                          | E1784K<sup>77</sup>: exerts LOF effect via hyperpolarizing voltage dependence fast inactivation and accelerating rate of fast inactivation and GOF effect via destabilizing the IFM bound state of the channel to induce noninactivating currents. |
| Different from LQT3 or BrS (In the absence of a distinct ECG phenotype) |                         | Overlap GOF & LOF |
|                           |                         | Gating defects       |                          | C683R<sup>79</sup>: a novel variant, with the GOF effect resulting from a significant increase of the maximal current density and a hyperpolarizing shift of the steady-state activation; without direct effect on \( I_{NaL} \) at baseline or adrenergic stimulation, with the LOF effect resulting from an increased closed-state inactivation. |
| SCN1B                      | BrS                    | LOF                 | Gating defects           |                                   | E87Q<sup>75</sup>: positively shifts the activation curve.                                                                                                                                                       |
| SCN2B                      | BrS                    | LOF                 | Trafficking defects      |                                   | D211G<sup>80</sup>: decreases Nav1.5 cell surface levels.                                                                                                                                                         |
| Nav1.6                     | SCN8A                  | Cardiac arrhythmia contributes to SUDEP in EIEE | GOF                      | No effect on SCN8A transcripts | N1768D<sup>87</sup>: increases calcium transient duration, prolongs the early phase of APD, and increases incidence of DADs but not changes in Nav1.5 expression.                                                                 |
| Nav1.8                     | SCN10A                 | SUD                 | GOF                      | Gating defects                    | P1102S<sup>86</sup>: shows slower inactivation time course, allowing more Na\(^+\) to enter the cell.                                                                                                                                 |
|                           | BrS                    | LOF                 | Gating defects           |                                   | I671I<sup>88</sup>: depolarizes shift of activation and inactivation curve, reduces the sodium current.                                                                                                            |

NR, not reported; DS, Dravet syndrome; SUDEP, sudden unexpected death in epilepsy; SUD, sudden unexplained death; EIEE, early infantile epileptic encephalopathy; LQTS3, long QT syndrome type 3; BrS, Brugada syndrome; GOF, SIDS, sudden infant death syndrome; gain-of-function; LOF, loss-of-function; DN, dominant negative.
mogenic Ca\(^{2+}\) release.\(^{32,41}\) Compared with other TTX-sensitive Nav channels, the location of Nav1.6 (Table 1) is the closest channel to RyR2 (< 100 nm)\(^{41,86}\) indicating that Nav1.6 is capable of impacting Ca\(^{2+}\) cycling proteins and Ca\(^{2+}\) dynamics in both health and disease.\(^{86}\) GOF variants of SCN8A-encoding Nav1.6 (Table 3) potentially leads to sudden unexpected death in epilepsy (SUDEP) due to arrhythmia of the brain and the heart.\(^{87}\) The GOF variant of SCN8A (N1768D) causes hyperexcitability of ventricle myocytes by increasing calcium transient duration, prolonging the early phase of APD, and increasing the incidence of DADs but not by compensatory changes in Nav1.5 expression.\(^{88}\) Selective pharmacological blockade of Nav1.6 and silencing of Nav1.6 indicate that Nav1.6 can potentially contribute to β-AR stimulation-induced \(I_{\text{Na,L}}\) and arrhythmias.\(^{41}\) This explains why catecholaminergic polymorphic ventricular tachycardia (CPVT) models respond to treatment with some Na\(^+\) channel blockers.\(^{41}\) Besides SCN8A, the possible roles of SCN1A, or SCN4A mutations in pathophysiology of cardiac congenital syndrome were also investigated (Table 3).

In addition, TTX-insensitive SCN10A-encoding Nav1.8 (Table 1) channels, which are mainly expressed in the peripheral nervous system, are also found in the heart\(^{42–44,48}\) at a higher level in the atrial myocardium than in the ventricular myocardium,\(^{43}\) exhibiting a more depolarized voltage dependence of inactivation and slower inactivation kinetics than other faster sodium channels like Nav1.5.\(^{44}\) Nav1.8 contributes to abnormal increases in \(I_{\text{Na,L}}\) and consequently prolongs the APD and elevates proarrhythmogenic diastolic SR Ca\(^{2+}\) in cardiac disease.\(^{42}\) Genetic deletion of Nav1.8 produces a smaller \(I_{\text{Na,L}}\) increase than in WT cardiomyocytes during β-AR stimulation.\(^{42}\) LOF and GOF variants in SCN10A (Table 3) are associated with BrS\(^{88}\) and SUDEP,\(^{89}\) respectively. Gating dysfunction with enhanced of inactivation results in LOF of Nav1.8.\(^{88}\) In contrast, dysfunction with slower inactivation could result in GOF of Nav1.8 and then allow more Na current entry.\(^{89}\) Thus, Nav1.8 also plays a significant role in the initiation of proarrhythmic triggers via \(I_{\text{Na,L}}\)-induced SR Ca\(^{2+}\) leakage.

### 4.2 Ca channels

In response to membrane depolarization, voltage-gated calcium (Cav) channels activate and mediate extracellular Ca\(^{2+}\) influx into the cytosol, which serves as the second messenger of electrical signaling, initiating many physiological processes, such as excitability, contraction and cell death.\(^{90}\) The Cav1 and Cav3 groups mediate L-type and T-type currents, respectively, and are involved in cardiac function. Cav1 is more highly expressed than Cav3 in ventricular myocytes, while Cav3 is mainly expressed in SAN cells (Table 4).\(^{90}\) Ryanodine receptors (RyRs), intracellular Ca\(^{2+}\) channels in the sarcoplasmic/endoplasmic reticulum (SR/ER), control the rapid release of Ca\(^{2+}\) from SR/ER into the cytoplasm to initiate CICR, a key event that triggers skeletal and cardiac muscle contraction.\(^{91,92}\) Among three mammalian isoforms (RyR1, RyR2, and RyR3), RyR2 is primarily expressed in cardiac muscles,\(^{91–93}\) and higher expressed in the ventricle (Table 4).\(^{37,38}\)

#### 4.2.1 Cav1.2

Cav1.2 channels, located in T-tubules of ventricular myocytes (Figure 1A), are assumed to be the major subtype of Cav1 channels that mediate the entry of Ca\(^{2+}\), which is required for the AP plateau (Figure 3A), and EC coupling, triggering activation of RyR2 and initiating CICR (Figure 1A).\(^{90}\)

The Cav1.2 complex consists of one pore-forming subunit \(\alpha 1c\) and the auxiliary subunits \(\alpha 2\delta\) and intracellular β (Figure 3B).\(^{94,95,96–98}\) The full-length cryo-EM structure of cardiac Cav1.2 has not been revealed, while skeletal Cav1.1 was the first Cav channel to have its full-length cryo-EM structure reported with an overall resolution of 4.2 and 3.6 Å.\(^{99,100}\) The structure of Cav1.1 provides a structural template for the Cav1 family and comparisons for molecular interpretations of the functions and disease mechanisms between eukaryotic Cav and Nav channels.\(^{99,100}\) Because the \(\alpha 1\) subunits from Cav1.1 and Cav1.2 are highly homologous, Cav1.2 modeling could be based on the Cav1.1 structure for analyzing the molecular determinants of opening and closure of Cav1.2.\(^{94}\) Voltage-independent upward movement or voltage-dependent movement of S4 segments maintain the opening or closure of the gating, respectively.\(^{94}\) The voltage sensor S4-S5 are coupled with pore S6 segments by directly interacting with a ring of small residues, which are regarded as interesting sites for studying electromechanical coupling.\(^{94}\) In addition, in complex with channel-selective agonists/antagonists, structural analysis helps to elucidate their specific binding sites and reveal the structural reasons why similar types of molecules (such as nifedipine and Bay K8644) exert opposite antagonist and agonist effects on Cav1.1 channels.\(^{20}\) The auxiliary subunits \(\alpha 2\delta\) and β generally modulate the surface expression and biophysical kinetics of \(\alpha 1c\).\(^{97,98}\) Recent studies have raised a new function for \(\beta\) subunits in hearts: \(\beta\) subunit binding to \(\alpha 1c\) might be dispensable for Cav1.2 trafficking at normal physiological conditions but is essential for the augmentation of Ca\(^{2+}\) current and cardiac contractile response to β-adrenergic stimulation.\(^{96}\)
## Table 4: Cardiac Ca\(^{2+}\) channels subtypes

| Cav types | Subtypes | Encoding \(\alpha\) subunits gene | Auxiliary subunits | Main location | Subcellular localization in cardiac tissue (V/A/SAN)\(^{37,38}\) and region | Cryo-EM Structure | Principal physiological functions in human ventricle myocytes |
|-----------|----------|----------------------------------|-------------------|--------------|----------------------------------------------------------------|-------------------|------------------------------------------------------------|
| L (long lasting and large conductance) Type | Cav1.2 | CACNA1C | \(\alpha 2\)δ and intracellular \(\beta\), KchIP2 | Heart, CNS | V \(\approx\) A > SAN T-tubule\(^{37}\) | NR, refers to Cav1.1 | Mediates the entry of Ca\(^{2+}\), contributes the AP plateau and initiates excitation-contraction coupling in cardiac muscle |
| Cav1.3 | CACNA1D | NR | CNS, Heart | V < A < SAN | NR | Cardiac SAN pacemaker activity |
| T (transient-opening and small conductance) Type | Cav3.1 | CACNA1G | \(\alpha 2\)δ | Heart, CNS | V < A < SAN | Human Cav3.1 complex containing \(\alpha 1\), \(\alpha 2\)δ subunits\(^{253}\); providing the structural reason why less energy is required for Cav3.1 to open the intracellular gate and facilitating the activation at lower voltages\(^{253}\) | Cardiac SAN pacemaker activity |
| Cav3.2 | CACNA1H | NR | Heart, CNS | V \(\approx\) A < SAN | NR | Cardiac SAN pacemaker activity |
| Intracellular calcium channel | RYR2 | RYR2 | NA | Heart | V \(\approx\) A \(\gg\) SAN | Porcine Ryr2\(^{110}\) | Controls rapid release of Ca\(^{2+}\) from SR/ER into cytoplasm to initiate CICR |

CNS, central nervous system; V, ventricle; A, atrium; SAN, sinoatrial node; Cryo-EM, cryoelectron microscopy; NA: not available; NR, not reported.
FIGURE 3  Cardiac voltage-gated L-type calcium channel (Cav1.2) and RyR2 structure, accessory proteins and signaling. (A) The contribution of inward current $I_{Ca,L}$ (upper) mediated by Cav1.2 to action potential (AP) Phase II (lower). (B) The L-type calcium channel Cav1.2 is formed by the interaction of the pore-forming $\alpha_1$ core subunit with auxiliary subunits, including $\alpha_2\delta$ and intracellular $\beta$. The $\alpha_1$ subunit consists of four homologous domains with a voltage sensor S4 and pore-forming S5 and S6 in each domain and connected by cytoplasmic linkers (ICL1, ICL2, and ICL3). The $\beta$ subunit is localized exclusively at the cytosolic face of the channel and its GK domain interacts with the $\alpha$-interaction domain (AID) of the $\alpha_1$ subunit I-II loop to mediate Cav1.2 trafficking by antagonizing ER retention signals. The $\alpha_2\delta$ subunit binds to extracellular regions, including domain III of the $\alpha_1c$ subunit. $\delta$ is linked with a larger $\alpha_2$ polypeptide via a disulfide bridge. The $\alpha_1$ subunit interacts with several proteins, receptors and subunits of other channels. The amino-terminal domain NTD, ICL1 and carboxyl-terminal domain (CTD; LZ motif) of Cav1.2 interact with AKAP, PKA and $\beta_2$-AR CTD also bind to AKAP. PP2A binds to site next to S1928 in the CTD. CaM binds to the IQ motif. KChIP2, an accessory subunit of Kv4.3, directly interacts with the inhibitory module at the NTD of the Cav1.2 $\alpha_1$ subunit. The Cav1.2 $\beta$ subunit interacts with the AID of the $\alpha_1$ subunit I-II loop. The $\beta$-AR/cAMP/PKA, $\beta_1$-AR/CaMKII, TrkB R/CaMKII signaling pathways are involved in modulating the expression and function of Cav1.2 in the heart. PKA-related phosphorylation pathway (red arrows) and CaMKII-related phosphorylation pathway (purple arrows). (C) RyR2 is a large, square, homotetramer in the configuration of a four-leaf clover. Each subunit of the homotetramer consists of a large cytosolic domain (called the foot structure), which is responsible for interaction with protein modulators, and CTTD. Four identical carboxyl-terminal transmembrane domains (CTTDs) are responsible for forming the central ion-conducting pore. RyR2 is located beneath Cav1.2 in T-tubules and is connected with $\alpha_1$ subunit of Cav1.2 by sorcin. In addition, CaM and FKBPI2/12.6 also interact with the cytosolic foot structure. Kinase (PKA, CaMKII) and phosphatase
In addition, several accessory proteins interact with the α1c subunit of Cav1.2 and regulate Cav1.2 expression and function (Table 6). Cav1.2 is involved in the β-AR/cAMP/PKA signaling pathway.95 (Figure 3B). β1-AR/cAMP signaling is diffusive and global, while β2-AR/cAMP is relatively localized.95 The CTD of β2-AR not only binds to A-kinase anchoring protein (AKAP) but also directly binds to Cav1.2 to mediate local signaling via the cAMP-dependent PKA pathway and facilitate localized cAMP signaling.95 The PKA-dependent phosphorylation of amino acids in the CTD of the α1c subunit has been demonstrated to be decisive for the β-AR-mediated upregulation of cardiac I_{Ca,L}.95,101 In addition, some amino acids in the CTD of the α1c subunit are targets of the β1-AR/CaMKII signaling pathway.95 In parallel with the roles of the β-AR system, BDNF-TrkB binding regulates myocardial Ca^{2+} cycling and EC coupling by triggering CaMKII-dependent signaling.102

GOF variants in CACANIC (Table 5) cause Timothy syndrome (TS), which is a multisystemic disorder including LQTS8, autism, and dysmorphic features.103 Complete loss of inactivation kinetics leading to a prolonged calcium influx during action potentials,103 or left shift in the activation curve leading to increase in window currents,104 could result in GOF of the Cav1.2 channel. Variant E1496K slowed inactivation, causing isolated LQTS8 without TS.105 On the other side, LOF variants (Table 5) which disruption of protein trafficking,106,107 gating,108 or Ca^{2+} permeation108,109 account for genotyped BrS cases. These results implicated the importance of the Cav1.2-mediated calcium signaling in human physiology and heart disease.

4.2.2 Ryr2

The near-atomic-resolution cryo-EM structure of RyR2 from porcine hearts has been recently revealed in both the open and closed states,110 or with key modulators,111,112 offering the opportunity to characterize the roles of the structural elements and modulators during gating shifts. Each subunit of the homotetrameric RyR2 consists of a large cytosolic domain, which is responsible for interaction with protein modulators, linking gap between the SR and transverse tubule (T-tubule) membranes, and carboxyl-terminal transmembrane domain (CTTD), four identical of which are responsible for forming the central ion-conducting pore (Figure 3C).110 The cryo-EM structures of the RyR2 complex and the abovementioned Cav1.1 establish a solid foundation for future revealing the Cav1.2 complex, the complex formation between Cav1.2 and RyR2, and excitation-contraction coupling.

Several proteins (Table 6) interact with the cytosolic region of RyR2 to regulate its open probability. For example, CaM inactivates Ryr2 during diastolic cytosolic calcium elevation, thus playing an important role in Ca^{2+} alternans.113 The CaM binding sites on cytosolic sites of Ryr2 will be shifted and dependent on Ca^{2+} concentration binding to CaM.112 Enhancement of CaM function promotes, whereas impairment of CaM function suppresses Ca^{2+} alternans.113 Several enzymes, such as PKA, CaMKII, PP1, and PP2A, interact with Ryr2 and exert phosphorylation/dephosphorylation effects on Ryr2.114 The hyperphosphorylation of RyR2 by PKA and/or by CaMKII causes abnormal Ca^{2+} leakage from the SR. RyR2 is also coupled to proteins at the luminal SR surface, such as type 2 calsequestrin (CASQ2),116,117 which increases the open probability and facilitates high rates of Ca^{2+} efflux during systole.116

Moreover, RyR2 also interact with other channels. RyR2 is located beneath most Cav1.2 (within ~12 nm) in T-tubules and is connected with the α1 subunit of Cav1.2 by sorcin, which is a Ca^{2+}-binding protein reducing the open probability of Ryr2, bridging the gap between SR and the sarcolemma for interchannel cross-talk.118 In addition, trimeric intracellular cation (TRIC) channels represent a recently discovered class of cation channels that were first identified in rabbit skeletal muscle in 2007.119 TRIC-A is a subtype that is abundantly expressed in excitable cells, having slightly higher permeability for K+ than Na+ and mediating counterion movements by releasing Ca^{2+} from the SR.120 The cryo-EM structure of the symmetrical cation channels has been reported.8 Moreover, TRIC-A also directly interacts with the cytosolic region of RyR2 via its carboxyl-terminal tail domains (Figure 3C) to modulate intracellular Ca^{2+} homeostasis and thereby facilitates Ca^{2+} release from the SR.121 The open probability and current amplitude of TRIC-A are increased by a positive shift in the MP but are blocked by exposure to a high-concentration Ca^{2+} bath on the luminal side during the resting state.8,121 TRIC-A gene deletion decreases the sensitivity of individual RyR channels to β-AR/PKA stimulation, eventually resulting in Ca^{2+} release impairment and irregular ECG.121 These studies indicate that TRIC-A promotes the release of Ca^{2+} from the SR via RyR2 and maintains RyR2 function at low Ca^{2+} to neutralize the (PP1 and PP2A) exert phosphorylation/dephosphorylation effects on Ryr2. Type 2 calsequestrin (CASQ2) interacts with the luminal surface of Ryr2 to increase the open probability. In addition, class A of trimeric intracellular cation (TRIC-A) channels on the SR membrane directly interact with the cytosolic region of Ryr2 (Figure 1B) via its carboxyl-terminal tail domains to facilitate Ca^{2+} release from the SR.
### Table 5 Mutations in cardiac Ca\(^{2+}\) channels subtypes associated with congenital syndromes

| Subtypes | Encoding subunits gene | Congenital syndrome | Gain or Loss of function | Mechanisms underlies the phenotype | Examples of variants |
|----------|-------------------------|---------------------|--------------------------|-----------------------------------|---------------------|
| Cav1.2   | CACNA1C                 | TS                  | GOF                      | Gating dysfunction                | G406R\[^1^0^3\]: leads to a prolonged calcium influx during action potentials caused by complete loss of voltage-dependent channel inactivation. G419R\[^1^0^4\]: displays a 4-fold increase in the peak current density and a left shift in the activation curve resulting in increase in window currents. |
|          |                         |                     |                          |                                   | E1496K\[^1^0^5\]: slows inactivation and thus might contribute to prolonged action potential duration. |
|          |                         |                     |                          |                                   | A39V; T320M/Q428E\[^1^0^7\] |
|          |                         |                     |                          |                                   | V2014I\[^1^0^8\]: significantly reduces conductance of the calcium channel at potentials between 0 and +30 mV during activation, shifts half-inactivation voltage to more negative potentials. |
|          |                         |                     |                          |                                   | E1156K\[^1^0^8,1^0^9\]: destroys the calcium selectivity, and instead converts the mutant channel into a channel with a marked increase in sodium-mediated inward currents and potassium-mediated outward currents. |
|          |                         | Isolated LQT8       | GOF                      | Gating dysfunction                | E1496K\[^1^0^5\]: slows inactivation and thus might contribute to prolonged action potential duration. |
|          |                         | without causing TS  |                          |                                   | E1156K\[^1^0^8,1^0^9\]: destroys the calcium selectivity, and instead converts the mutant channel into a channel with a marked increase in sodium-mediated inward currents and potassium-mediated outward currents. |
|          |                         | BrS3                | LOF                      | Trafficking defects               | A39V; T320M/Q428E\[^1^0^7\] |
|          |                         |                     |                          | Gating defects                    | V2014I\[^1^0^8\]: significantly reduces conductance of the calcium channel at potentials between 0 and +30 mV during activation, shifts half-inactivation voltage to more negative potentials. |
|          |                         |                     |                          |                                   | E1156K\[^1^0^8,1^0^9\]: destroys the calcium selectivity, and instead converts the mutant channel into a channel with a marked increase in sodium-mediated inward currents and potassium-mediated outward currents. |
| Cav1.3   | CACNA1D                 | SANDD               | LOF                      | Gating defects                    | 403\_404insGly\[^2^5^4\] |
| Cav3.1   | CACNA1G                 | Bradycardia, atrioventricular conduction block | LOF | NR | NR |
| RYR2     | RYR2                    | CPVT                | GOF                      | Gating defects                    | RI76Q\[^1^0^2^6\]: increases probability of channel opening, increases incidence of spontaneous Ca\(^{2+}\) oscillations thus causing susceptibility to CPVT. S2246L\[^1^0^4\]: disrupts the interdomain interactions after channel activation and increases channel opening. |
|          |                         |                     |                          | Channel instability               | S2246L\[^1^0^4\]: disrupts the interdomain interactions after channel activation and increases channel opening. |
|          |                         | CRDS which could cause SCD without CPVT | LOF | Gating defects | D4646A\[^1^0^8\]: impairs the cytosolic Ca\(^{2+}\) activation and diminishes the luminal Ca\(^{2+}\) activation of single RyR2 channels; suppresses catecholamine-induced SR Ca\(^{2+}\) release and produces systolic arrhythmogenic abnormalities without affecting expression. |

TS, Timothy syndrome; SANDD, sinoatrial node dysfunction and deafness syndrome; LQTS, long QT syndrome; CPVT, catecholamine-induced ventricular arrhythmias; SCD, sudden cardiac death; CICR, Ca\(^{2+}\)-induced Ca\(^{2+}\)-release; CRDS, Ca\(^{2+}\) release deficiency syndrome; GOF, gain-of-function; LOF, loss-of-function.
### Table 6: Accessory Proteins Reported to Interact with and Regulate Cav1.2 and RYR2

| Accessory Proteins | Types                  | Binding Sites                                                                 | Biophysical Function                                                                 | Techniques         |
|--------------------|------------------------|-------------------------------------------------------------------------------|--------------------------------------------------------------------------------------|---------------------|
| **Cav1.2**         |                        |                                |                                                                                     |                     |
| Bridging Integrator 1 (BIN1) | Scaffolding protein | Adjacent to Cav1.2 channels clustered in T-tubules | BIN1 is responsible for Cav1.2 trafficking to T-tubules; knockdown of BIN1 decreases the surface expression of Cav1.2 and calcium transients in mouse cardiomyocytes. BIN1 increases Cav1.2 channel clustering and whole-cell Ca\(^{2+}\) currents in human embryonic stem cell-derived cardiomyocyte (hESC-CM) | MPC, Ca\(^{2+}\) imaging, WB |
| AKAP 79/150        | Anchoring protein      | In the NTD, ICL1-II and LZ motif in the CTD of α1 subunit                  | Forms macromolecular complex with Cav1.2 and takes part in different regulatory pathways by recruiting several signaling molecules, such as PKA to Cav1.2; AKAP interaction is disrupted by the membrane-permeable stearilated peptide Ht31 | WB                  |
| KChIP2             | Accessory subunit of Kv4.3 | In the NTD of the α1 subunit | Modulates the Cav1.2 current without affecting Cav1.2 protein expression or trafficking. | MPC, WB             |
| CaM                | Regulatory protein     | IQ motif in the CTD                                                         | Facilitates the Cav1.2-Cav1.2 channel interactions within a cluster and then work cooperatively | MPC, Ca\(^{2+}\) imaging |
| PKA                | Kinase                 | Recruited via AKAP                                                          | Upregulation of L-type currents by phosphorylates S1700/T1704, S1928 in the CTD | MPC                 |
| PP2A               | Phosphatase            | Between S1928 and LZ motif in the CTD                                       | Antagonizes β-AR/PKA mediates phosphorylation of Cav1.2 and upregulation of L-type currents | MPC                 |
| **RYR2**           |                        |                                |                                                                                     |                     |
| FKP06 binding proteins (FKBP12/12.6) | Regulatory protein | Cytosolic region                                                               | Stabilizes RyR2 in the closed state, reduces its activity, prevent aberrant activation of the channel during the resting phase of the cardiac cycle | Single-channel recordings, MPC |
| Sorcin             | Calcium binding protein | Cytosolic region; CTD of the α1 subunits of Cav1.2                        | Sorcin completely inhibits ryanodine binding to cardiac RyRs, reduces the open probability of RyR2 and Cav1.2, and bridges the gap between SR and the sarcolemma for interchannel cross-talk | WB, Co-IP, single-channel recordings |
| CaM                | Regulatory protein     | Cytosolic region                                                               | Inactivate RyR2 during diastolic cytosolic calcium elevation, thus playing an important role in Ca\(^{2+}\) alternans | Ca\(^{2+}\) imaging, MPC |
| CaMKII             | Kinase                 | Cytosolic region                                                               | Phosphorylates of RyR2, regulates the channel open probability                      | Single-channel recordings, Ca\(^{2+}\) imaging |
| PKA                | Kinase                 | Cytosolic region, S2809                                                       | Phosphorylates of RyR2 and dissociates FKBP12,6, regulates the channel open probability | Single-channel recordings, WB |
| TRIC-A             | Regulatory protein     | Cytosolic region                                                               | Serves as counterion channels that provide the flow of K\(^{+}\) ions into the SR during the acute phase of Ca\(^{2+}\) release and thereby facilitates Ca\(^{2+}\) release from the SR | Single-channel recordings |
| Type 2 calsequestrin (CASQ2) | Regulatory protein | Luminal region                                                                 | Increases the open probability of RyR2                                               | Single-channel recordings, WB |

CTD, carboxyl terminal domain; NTD, N terminal domain; MPC, manual Qatch clamp; WB: Western blot; co-IP, coimmunoprecipitation.
transient luminal negative charge caused by Ca\(^{2+}\) release in cardiomyocytes.

GOF variants in RyR2 (Table 5) are implicated in ventricular tachyarrhythmias, including type 1 of CPVT type (CPVT1), which is characterized by stress-induced ventricular tachycardia in the absence of a structurally abnormal heart.\(^{122}\) GOF variants could induce channel instability by disrupting the interdomain interactions after channel activation,\(^{124}\) or increase the open probability of RyR2 and pathological SR Ca\(^{2+}\) release,\(^{115,125}\) and thus causing susceptibility to CPVT. On the other hand, RyR2 LOF variants have been identified among survivors of cardiac arrest without exhibiting the CPVT phenotype and further regarded as RyR2 Ca\(^{2+}\) release deficiency syndrome (CRDS) via an EAD-mediated mechanism.\(^{126}\)

In CPVT cardiomyocytes with the RyR2 variant R176Q, a viral vector containing a CaMKII inhibitor (autocamtide-2-related inhibitory peptide, AAV9-GFP-AIP) completely suppressed the abnormal increase in spontaneous Ca\(^{2+}\) transients, suggesting that CaMKII suppression represents a potential therapy for CPVT.\(^{127}\) A KN93-mediated increase in RyR2 Ca\(^{2+}\) release in cardiomyocytes was found to be due to disruption of the CaM-RyR2 interaction rather than inhibition of CaMKII.\(^{64}\) Gene transfer of CaM, exhibiting a slower Ca\(^{2+}\) dissociation rate and longer RyR2 refractoriness, alleviated arrhythmias in a CASQ2-associated CPVT mouse model.\(^{128}\) Previous studies have illustrated that flecainide prevents ventricular tachyarrhythmia in patients with CPVT by blocking of the TTX-sensitive Nav channel.\(^{41}\) Recent research has shown that the arrhythmic effect of flecainide mainly relies on blocking RyR2 channels but not TTX-sensitive Nav channels.\(^{129}\) The secondary amine on the piperidine ring in flecainide is necessary for its activity in RyR2 channels.\(^{129}\) In general, the regulation of RyR2 modulators (RyR2-CaM interaction) represents an important therapeutic target for regulating cardiac alternans in cardiac ventricular arrhythmia.

### 4.3 Kv channels

Cardiac Kv channels play prominent roles in resting potential maintenance, AP repolarization, and the AP plateau phase.\(^{130,131}\) For example, Kv1.4/Kv4.3, Kir2.1, Kv11.1, and Kv7.1 are highly expressed in the ventricular myocytes (Table 7).\(^{37,38}\) Kir2.1 contributes to the maintenance of the resting potential in Phase IV, while Kv4.3 and Kv1.4 contribute to repolarization, specifically the notch (the transient repolarization period) of the AP.\(^{130,131}\) Of particular relevance to the AP plateau is the delayed rectifier current (I\(_{Ks}\)), which includes rapid (I\(_{Kr}\)) and slow (I\(_{Ks}\)) components that are governed by distinct channel subtypes Kv11.1 and Kv7.1, respectively.\(^{130,131}\) Dysfunction of cardiac Kv channels can result in APD changes and the subsequent development of LQTS, SQTS, or other related life-threatening ventricular arrhythmias or sudden cardiac death.\(^{1,132}\)

#### 4.3.1 Kv4.3

The rapidly activated and inactivated transient outward potassium current (I\(_{to}\)) contributes to early ventricular AP repolarization and underlies the initial “notch” before the AP plateau phase in humans and other larger mammals (Figure 4A).\(^{130}\) I\(_{to,fast}\) and I\(_{to,slow}\) are the two distinct components of I\(_{to}\), and are mediated by Kv4.3 and Kv1.4, respectively, in humans and by Kv4.2/Kv4.3 and Kv4.1, respectively, in rodents.\(^{1}\) Unlike in human and mammalian models, ventricular AP in rodent models exhibits fast repolarization without a plateau phase due to I\(_{to}\) rather than I\(_{Kr}\), playing the major role in repolarizing currents.\(^{1,130,133}\) The significant prolongation of repolarization duration, which is affected more by a reduction in I\(_{to}\) than a reduction in I\(_{Kr}\), underlies the mechanism for heart failure with preserved ejection fraction (HFpEF), typical heart failure symptoms with a normal ejection fraction)-related ventricular arrhythmias and sudden cardiac death in rodent models.\(^{134}\) A rabbit ventricular APD could be shortened, and its plateau could be lost when mouse I\(_{to}\) currents were integrated.\(^{1}\) Thus, differences in I\(_{to}\) densities in different species contribute to variations in the waveforms of action potentials.\(^{1,130}\)

Kv4.3 is composed of one pore-forming \(\alpha\) subunit and \(K^+\) channel interacting protein 2 (KChIP2) \(\beta\) subunit and is regulated by several accessory protein interactions (Figure 4B) (Table 8). A reduction in Kv4.3 expression and I\(_{to,fast}\) in heart disease, is associated with \(\beta\)-AR/CaMKII-mediated activation\(^{133}\) and \(\beta\)-AR/NF-kB-mediated activation.\(^{135}\) Moreover, an increasing number of studies have speculated that Kv4.3 and Nav1.5 not only regulate each other’s functions, but also have the ability to interact with each other.\(^{136,137}\) Nav1.5 and Kv4.3 reside is visualized in close proximity (<40 nm) at the membrane.\(^{136}\) Overexpression of Kv4.3 protein significantly decreased AP upstroke and Nav1.5 current density without affecting Nav1.5 total protein expression and its kinetic properties.\(^{137}\) In addition to auxiliary subunit of KChIP2, Navβ1 subunit also associated with Kv4.3\(^{138,139}\) and regulated the I\(_{Na}/I_{to}\) balance by yielding an increase in I\(_{Na}\) and a decrease in I\(_{to}\).\(^{136}\)
| Potassium channel types                                                                 | Subtypes | α Subunits | Auxiliary subunits       | Main location          | Subcellular localization in cardiac tissue (V/A/SAN) | Cryo-EM structure | Principal physiological functions in human ventricle myocytes |
|----------------------------------------------------------------------------------------|----------|------------|--------------------------|------------------------|-----------------------------------------------------|------------------|---------------------------------------------------------------|
| Voltage-dependent K⁺ channels 1-9, Shaker-related channels, containing six transmembrane regions (S1-S6) with a single pore | Kv1.4    | KCNA4      | Kvβ1.2                   | Heart                  | V > A ≈ SAN T-tubules, IDs                           | NR               | Mediates I_{to,slow} and contributes to early AP repolarization |
|                                                                                       | Kv4.3    | KCND3      | KChIP2 encoded by KCNIP2; Navβ1 encoded by SCN1B | CNS, Heart             | V < A > SAN T-tubules                                | NR               | Mediates I_{to,fast} and contributes to early ventricular AP repolarization |
|                                                                                       | Kv7.1    | KCNQ1      | MinK encoded by KCNE1    | Heart                  | V > A > SAN; IDs, lateral membrane, T-tubules        | Frog KCNQ-CaM complex in an uncoupled, PIP2-free state | Mediates Iₙₛ and contributes to Phase II, III AP repolarization and early Phase IV of the AP |
| Voltage-dependent K⁺ channels 10-12, nonshaker-related channels                       | Kv11.1   | KCNH2      | MinK and MiRP encoded by KCNE1 and KCNE2, respectively | Heart, CNS             | V ≈ A > SAN IDs, lateral membrane, T-tubules        | Human Kv11.1     | Mediates Iₙₛ and contributes to Phase II, III AP repolarization and early Phase IV of the AP |
| Inward rectifying K⁺ current, containing only two transmembrane regions and a single pore | Kir2.1   | KCNJ2      | NA                       | CNS, Heart             | V > A > SAN T-tubules                                | NR               | Mediates I_{k,1} and contributes to Phase IV resting MP and the terminal Phase III repolarization |

CNS, central nervous system; IDs, intercalated discs; V, ventricle; A, atrium; SAN, sinoatrial node; Cryo-EM, cryoelectron microscopy; KChIP2, K⁺ channel interacting protein 2; NA, not available; NR, not reported.
FIGURE 4  Cardiac voltage-gated potassium channel structure, accessory proteins and signaling. (A) The contribution of the outward current $I_{o}$ (upper) mediated by Kv4.3 to action potential (AP) Phase I (lower). (B) Kv4.3 is formed by the $\alpha$ subunit and accessory protein $K^+$ channel interacting protein 2 (KChIP2, $\beta$ subunit). A tripartite complex including the anchoring protein SAP97 and kinase CaMKII is formed at the Kv4.3 carboxyl-terminal domain (CTD) via a PDZ domain-binding motif Ser-Ala-Leu (SAL). The channel current is modulated by
## Table 8 Accessory proteins reported to interact with and regulate cardiac Kv

| Accessory proteins | Types | Binding sites | Biophysical function | Techniques |
|--------------------|-------|---------------|----------------------|------------|
| **Kv4.3**          |       |               |                      |            |
| Dipeptidyl peptidase-like protein 6 (DPP6) | Additional β-subunit | Serine proteases but lacks enzymatic activity | Regulates the inactivation and recovery from inactivation properties of Kv4.3 channels. Co-expression of DPP6 with Kv4.3 and KChIP2 produces a similar current kinetics as in human ventricular myocytes. | MPC, WB |
| SAP97              | Anchoring-adaptor protein | PDZ domain-binding motif Ser-Ala-Leu (SAL) in the CTD | Forms a tripartite complex with CaMKII through SAL motif, increase Ito even if in the absence of KChIP and DPP6. | MPC, pull-down assays |
| CaMKII            | Kinase | SAL motif in the CTD | Forms a tripartite complex with SAP through SAL motif. A frequency-dependent reduction in Kv4.3 expression and Ito current is directly linked to increases in CaMKII activation. | MPC, Co-IP |
| **Kv7.1**          |       |               |                      |            |
| Lipid phosphatidylinositol 4,5-bisphosphate (PIP2) | Cofactor | R249 of S4-S5 linker | In the absence of PIP2, voltage-sensing domain activation fails to open the pore. | MPC, WB |
| A-kinase anchoring protein 9 (AKAP9) | Anchoring protein | Binds with a leucine zipper (LZ) motif in the CTD | A substrate for PKA phosphorylation by itself, and also recruits signaling proteins, such as PKA, PP1 to form macro-molecular to mediate Kv7.1 phosphorylation or dephosphorylation regulation upon β1-adrenergic stimulation. | MPC, phosphorylation assays, IP |
| Protein phosphatase 1 (PP1) | Protein phosphatase | Recruited via AKAP9 | Antagonizes PKA-mediated S27 phosphorylation. | MPC, IP |
| Calmodulin (CaM)   | Regulatory protein | IG in ICLIII-IV, IQ-motif in CTD, a coiled coil formed by the proximal A and B helices, competes with and displaces PIP2 binding at K526 and K527 of helix B in the CTD | Regulates channel gating, Negatively shifts the activation curve. | MPC, WB |
### TABLE 8  (Continued)

| Accessory proteins | Types                      | Binding sites                     | Biophysical function                                                                                                                                                                                                 | Techniques |
|--------------------|----------------------------|-----------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------|
| Kv11.1             |                            |                                   | **Accessories**                                                                                                                                                                                                       |            |
| 14-3-3ε            | Adaptor protein            | S283 in NTD and S1137 in CTD      | Accelerates channel activation after phosphorylation by β-AR/PKA, stabilizes and prolongs the phosphorylation state by preventing dephosphorylation[81]                                                                 | MPC, WB    |
| Kir2.1             |                            |                                   | **Accessories**                                                                                                                                                                                                       |            |
| AP1-γ              | Adaptor protein            | CTD                               | Marks Kir2.1 for incorporation into clathrin-coated vesicles at the Golgi[74]                                                                                                                                          | Co-IP, MPC |
| SAP97              | Anchoring-adaptor protein  | NTD, CTD                          | Regulates surface expression of channels and is assembled a macromolecular signaling complex[26]                                                                                                                         | Co-IP, MPC |
| AKAP79             | Anchoring protein          | NTD, CTD                          | Anchors kinase close to channel phosphorylation sites[75]                                                                                                                                                              | Co-IP, MPC |
| Caveolin-3         | Scaffolding and regulatory | NTD                               | Regulates Kir2.1 trafficking and surface expression[283]                                                                                                                                                                | Co-IP, MPC |
| PIP2               | Cofactor                   | CTD, NTD                          | Activates Kir2.1 function[284]                                                                                                                                                                                          | MPC        |

CTD, carboxyl terminal domain; NTD, N terminal domain; MPC, manual Qpatch clamp; WB: Western blot; co-IP, coimmunoprecipitation.
Mutations in KCND3-encoded Kv4.3 or SCN5A-encoded Nav1.5 further showed the functional relationship between Kv4.3 and Nav1.5. GOF and LOF mutations in KCND3 (Table 9) respectively decreased and increased the Nav1.5 current, respectively. On the other hand, SCN5A LOF mutations increased Ito, by facilitating Kv4.3 cell surface expression or by slowing its steady-state inactivation. Thus, during the early phase of ventricular AP repolarization, a fine balance may exist between INa and Ito. GOF mutations in the KCND3 contributes to increase of peak Ito via efficient protein trafficking and gating, resulting in the imbalance of those two currents, the accentuation of the AP notch, and the development of BrS and/or early repolarization syndrome.

4.3.2 | Kv7.1

The slow delayed rectifier current, I\textsubscript{Ks}, is mediated by KCNQ1-encoding Kv7.1 and plays an important role in regulating the repolarization phase that terminates cardiac APs and thereby ends contraction (Figure 4C). In the heart, the KCNE1, encoding the auxiliary \( \beta \)-subunit KchIP2, interacts with Kv7.1 \( \alpha \) chains and affects both voltage-sensing S4 movement and the gate, making the activation of the complex much slower than that of Kv7.1 alone (Figure 4D). Cryo-EM analysis revealed a unique feature of Kv7.1: pore opening requires lipid phosphatidylinositol 4,5-bisphosphate (PIP2) binding during membrane depolarization, thereby increasing current and slowing inactivation. In addition, Kv7.1 is regulated by accessory protein interactions (Table 8), \( \beta \)-AR/PKA-mediated phosphorylation and PP1-mediated dephosphorylation (Figure 4D).

Mutations in KCNQ1 (Table 9) are the leading cause of several congenital cardiac diseases, including LQTS and SQTS. LQT1, the most common genotype-positive LQTS, is associated with LOF mutations in the KCNQ1-encoded Kv7.1 \( \alpha \) subunit and is often triggered by \( \beta \)-AR stimulation. Trafficking defects, \( \beta \)-AR stimulation, and \( \beta \)-AR/PKA-pathway, which phosphorylates Kv7.1 at its amino-terminal domain (NTD; S27). A-kinase anchoring protein 9 (AKAP9) interacts with the LZ motif in the Kv7.1 CTD and is anchored by PKA and PP1. The contribution of the outward current I\textsubscript{Ks} (upper) mediated by the inwardly rectifying potassium channel (Kir2.1) to AP Phase II and III (lower). (D) Kv7.1 is formed from the \( \alpha \)1 subunit, which consists of four homologous domains with voltage sensing S4 segment and pore-forming S5 and S6 segments in each domain. Auxiliary subunit KCNE1 interacts with residue V141 of S1 in KCNQ1 to allow the complex only open from a fully activated S4 conformation by altering the VSD S4-to-gate coupling, and also interacts with F339 in KCNQ1 to reduce the open probability at negative voltages. PIP2 binds to the S4-S5 linker during membrane depolarization. CaM binds to the Kv7.1 C-terminus, where it can compete with and replace PIP2. CaM also interacts with the Kv7.1 voltage sensor domain. Kv7.1 is modulated by the \( \beta \)-AR/PKA pathway, which phosphorylates Kv7.1 at its amino-terminal domain (NTD; S27). A-kinase anchoring protein 9 (AKAP9) interacts with the LZ motif in the Kv7.1 CTD and is anchored by PKA and PP1. The contribution of the outward current I\textsubscript{Ks} (upper) mediated by the inwardly rectifying potassium channel (Kir2.1) to AP Phase II and III (lower). (E) The contribution of the outward current I\textsubscript{Kr} (upper) mediated by the inwardly rectifying potassium channel (Kir2.1) to AP Phase II and III (lower). (F) The \( \alpha \) subunit of Kv11.1 contains six transmembrane helices, with S4 acting as the voltage sensor and S5-S6 forming the pore. The KCNE\( \beta \) subunits KCNE1 and KCNE2 interact with Kv11.1. The adaptor protein 14-3-3 involves in Kv11.1 and CTD, \( \beta \)-AR competes with Kir11.1 for association with 14-3-3 proteins, \( \beta \)-AR/PKA, \( \alpha \)-AR/PKC, and AT1/PKC are involved in Kv11.1 channel modulation. PKA and PKC phosphorylate the \( \alpha \) subunit (red arrows). (G) The contribution of the strong inward rectifier potassium current I\textsubscript{Ki} (upper) mediated by the Kir2.1 channel to AP Phase 0, III, and IV (lower). (H) Kir2.1 channels have two membrane-spanning domains; a-AR which forms the ion selectivity filter and intracellular N- and C-terminal domains. The CTD of Kir2.1 directly associates with API, APB, and SAP97. SAP97 also interacts with \( \beta \)-AR and to modulates the effect of \( \beta \)-AR on Kir2.1. APB and SAP97 also anchor kinases (PKA, PKC), and phosphatase (PP2B) close to Kir2.1 phosphorylation sites. The NTD of Kir2.1 directly interact with, caveolin-3 which regulates Kir2.1 trafficking and surface expression. PIP2 binds to both the CTD and the NTD to modulate channel gating.
| Subtypes | Encoding subunits Gene | Congenital syndrome | Gain or Loss of Function | Mechanisms underlies the phenotype | Examples of variants |
|----------|------------------------|---------------------|--------------------------|-----------------------------------|--------------------|
| Kv4.3    | KCND3                  | BrS                 | GOF                      | Trafficking efficiency             | L450F<sup>13,14</sup>: increases peak $I_{to}$ current density. |
|          |                        |                     | Gating defects           | Trafficking efficiency and gating defects | G600R<sup>14</sup>: increases peak $I_{to}$ current density and slows inactivation. |
| ERS      |                        |                     | GOF                      | Gating defects                     | G306A<sup>12</sup>: significantly increases $I_{to}$ current densities, slows inactivation, and prolongs the recovery from inactivation. |
| SUDS predisposing cardiac arrhythmia syndromes | Overlap GOF & LOF, GOF dominance |                     | Gating defects | V392I<sup>12</sup>: increases peak $I_{to}$ current density and total charge, while slows decay time, indicating a BrS-like $I_{to}$ GOF. While slows the recovery from inactivation. |
| Spinocerebellar ataxia (SCA19/22) | LOF                  |                     | Trafficking defects        |                                   | Δ227F-Kv4.3<sup>11</sup> |

| Kv7.1    | KCNQ1                  | SQTS2               | GOF                      | Gating defects                     | F279I<sup>15</sup>: impairs the association with KCNE1, produces a negative shift in the activation curve and an acceleration of the activation kinetics leading to increase of $I_{ks}$ |
|          |                        |                     | Gating defects           |                                   | R259H<sup>15</sup>: markedly prolongs the time constant of deactivation leading to a GOF in $I_{ks}$ without affects activation and inactivation kinetics. |
| LQTS1    | LOF                    |                     | Trafficking defects      |                                   | R190Q<sup>15</sup>: leads to a 70% reduction in $I_{ks}$. |
|          |                        |                     | Gating defects           |                                   | D242N<sup>15</sup>: removes the inactivation kinetics, slows the activation kinetics by shifting the voltage dependence of activation to more depolarized potentials thus hindering $I_{ks}$ current at physiologically relevant membrane potentials. |
|          |                        |                     |                              |                                   | A371T and S373P<sup>15</sup>: impairs CaM binding and alters channel assembly, thus stabilizing inactivation, and decreasing current density. |
|          |                        |                     |                              |                                   | R555C, R555H, K557E and R562M<sup>15</sup>: markedly reduces the current densities, positively shifts the voltage dependence of activation, slows activation kinetics, increases deactivation rates and reduces interaction with the KCNE1 C-terminus and PIP2 binding. |

| KCNE1    | LQTS5                  | LOF                 | Trafficking defects       |                                  | T322M, T322A, and G325R<sup>15</sup>: cause a complete loss of $I_{ks}$. |

(Continues)
| Subtypes | Encoding subunits Gene | Congenital syndrome | Gain or Loss of Function | Mechanisms underlies the phenotype | Examples of variants |
|----------|------------------------|---------------------|--------------------------|-----------------------------------|---------------------|
| Kv11.1   | KCN\(H2\)             | SQTS\(1\)          | GOF                      | Gating defects                    | N588K\(^{261}\); increases steady-state current and abolishes rectification of the current. |
|          |                        | LQTS\(2\)          | LOF                      | Synthesis deficiency              | Y611H and V822M\(^{266}\) |
|          |                        |                    |                          | Trafficking defects               | 80-90% variants\(^{160-162}\) cause loss of Kv11.1 expression on plasma surface trafficking to the plasma and exert dominant-negative effect. |
|          |                        |                    |                          | Gating defects                    | T421M\(^{267}\); positively shifts the voltage dependence of activation. |
|          |                        |                    |                          | Permeation disruption             | L553N\(^{163}\); produces a dramatically faster deactivation time. |
|          | KCNE\(2\)             | LQTS\(6\)          | LOF                      | Gating defects                    | G628S\(^{161,266,268}\); leads to a reduced selectivity to potassium. |
| Kir2.1   | KCNJ\(2\)             | Borderline for SQTS\(3\) | GOF                      | Gating defects                    | M54T\(^{164}\); increases deactivation rates. |
|          |                        | LQT\(7\) (ATS)     | LOF                      | Trafficking defects               | F58S\(^{187}\); produces an increase of the channel conductance and in its open probability |
|          |                        |                    |                          | Gating defects                    | S114/Y315\(^{181,179}\); produces reduction of membrane expression reduction and has DN effect on WT. |
|          |                        |                    |                          | Permeation disruption             | V77E/M307V\(^{176}\); produces nonconductive Kir2.1 without affecting cell surfaces expression and has DN effect on WT. |

LQTS, long QT syndrome; SQTS, short QT syndrome; BrS, Brugada syndrome; ERS, early repolarization syndrome; SUDS, sudden unexplained death syndrome; ATS, Andersen-Tawil syndrome; GOF, gain-of-function; LOF, loss-of-function.
depolarization in Phase 0-II. Then, it quickly recovers from inactivation and reopens during the initial Phase III repolarization, followed by slow deactivation that permits sustained Phase III and early Phase IV of the AP (Figure 4E).\[158\] Kv11.1 channels exhibit longer-lasting and higher-amplitude tail currents than have been found for other outward current channels that contribute to cardiac AP repolarization and duration.\[159\]

Kv11.1 is composed of one pore-forming \( \alpha \) subunit and two \( \beta \) subunits (\( \text{MinK} \) and \( \text{MiRP1} \) encoded by \( \text{KCNE1} \) and \( \text{KCNE2} \)) (Figure 4F). The structure of the \( hERG \) channel with depolarized voltage sensors and open pores was revealed using cryo-EM.\[19\] A small central cavity includes extended pockets, which is specific to Kv11.1, explaining the notable susceptibility of this channel to a wide range of drugs.\[153\] This high-resolution structure of the \( hERG \) channel in the open state also provided the opportunity to investigate the potential mechanisms for the state-dependent blockade of \( hERG \) by drugs.\[22\]

Kv11.1 is regulated by accessory protein interactions and signaling pathways (Figure 4F) (Table 8). Phosphorylation of Kv11.1 could be induced by the stimulation of \( \beta \)-AR/cAMP/PKA or G protein-coupled receptors (such as angiotensin II receptor \( \text{AT1} \) and the \( \alpha \)-adrenoceptors)/PKC signaling pathway, resulting in a decrease in \( I_{Kr} \).\[158\]

LOF mutations in Kv11.1 (Table 9) are characterized by reduced \( I_{Kr} \) and are associated with LQTS2, perhaps due to the disruption of the \( \alpha \) subunits responsible for channel synthesis/translation, a reduction in intracellular transport or the accessory protein interactions required for channel trafficking on the membrane, or the impairment of channel gating structure as well as permeation.\[160-162\] Among those mechanisms, trafficking defects is the dominant one, responsible for approximately 80-90\% of LQT2 by decreasing the folding efficiency of Kv11.1 proteins and increasing their retention in the endoplasmic reticulum (ER).\[160-162\] Comprehensive analysis of hundreds LQT2-linked mutations in four Kv11.1 structural domains and found that deficient protein trafficking is the dominant mechanism for all domains except for the distal C-terminus. Comprehensive and accurate analysis of mutations between normal and abnormal trafficking across multiple structural domains would aid in understanding the deleterious nature of these mutations.\[162,163\] Increasing high-throughput assays are developing and as alternative to traditional western blot assay to collect functional data.\[165\] In addition, LOF mutation in \( \text{KCNE2} \) is associated with LQTS6, a rare type of LQTS.\[155,164\] The allosteric modulation (Table 10) of Kv11.1 was investigated to explore methods of alleviating channel dysfunction and increasing \( I_{Kr} \) current and may represent a useful new approach for treating inherited and drug-induced LQTS2.\[165\]

SQTS1 is caused by GOF mutations (Table 9) in the Kv11.1 channel and is the most prevalent SQTS subtype. Mutations that impair the inactivation of Kv11.1\[166\] might explain the lack of efficacy of many class III antiarrhythmic drugs (such as sotalol and ibutilide\[167\]) in some patients. Interestingly, hydroquinidine, a class I antiarrhythmic drug inhibiting the Nav1.5 channel, could also block Kv11.1, significantly intervene with ventricular tachyarrhythmia induction\[167\] and prolong the QT interval in SQTS patients with Kv11.1 mutations.\[168\] In addition, ivabradine, as a class 0 antiarrhythmic drug inhibiting hyperpolarization-activated cyclic nucleotide-gated (HCN) channels, could also block \( I_{Kr} \) currents by binding in the vicinity of the lipid-facing surface M651 residue, which is directly coupled to the conformational dynamics of residues in the pore helices,\[32,169\] and exert antiarrhythmic effects in SQTS hiPSC-CMs with the N588K mutation.\[170\] This represents one important method by which the efficacy of drugs used for SQTS treatment can be evaluated in hiPSC-CMs with mutant Kv11.1 or multiple ion channels to predict effects in SQTS patients.\[171\] It would be worthwhile to further examine the effects of traditional inhibitors and to develop novel specific inhibitors to expand the clinical options available for these patients.

4.3.4 | Kir2.1

The strong inward rectifier potassium current \( I_{K1} \), primarily mediated by isoforms of the Kir2.x family (\( \text{KCNJ2} \)-encoding Kir2.1/\( \text{KCNJ12} \)-encoding Kir2.2), plays a critical role in stabilizing the resting MP and maintaining the duration of the terminal Phase III repolarization in human ventricle myocytes.\[38\] Kir2.1 is more dominant than Kir2.2 in human ventricle myocytes\[38\] under resting conditions, and Kir2.1 is in an open state and is permeable to K\(^{+}\). Kir2.x is abundantly expressed in ventricle and atrial myocytes, but is absent in SAN cells, allowing a relatively depolarized MP and maintaining pacemaker activity in SAN cells.\[37,172\] In contrast to adult ventricular CMs, a substantial lack of \( I_{K1} \) in hiPSC ventricular CM is regarded as one mechanistic contributor to the immature electrophysiological properties of spontaneous AP. Artificial expression of Kir2.1 might overcome this limitation, render the electrophysiological phenotype to be mature, and ablate proarrhythmic AP traits.\[173\]

The structure of \( K_{ir} \) channels is relatively simple in comparison with Nav, Cav and the members of the voltage-gated \( K \) channels. Each subunit of the Kir2.1 tetramer has only two membrane-spanning helices (S1-S2) but without the four membrane helices that form the voltage sensor in \( K \) channels.\[174\] Kir2.1 is regulated by acces-
| Modulators | Binding site | Major effect | Side effect | Model | Techniques |
|------------|--------------|--------------|-------------|-------|------------|
| VU0405601  | Extracellular domain of the Kv11.1 channel rather than to its central cavity | Reduces the APD-prolonging effect of dofetilide | NR | Primary rabbit hearts, HEK293 Kv11.1 cell lines | MPC, [3H] dofetilide-binding assays; radioligand-binding assays |
| AZSMO-23 (types 2 of Kv11.1 activators) | Removal of inactivation | 1) Shifts the inactivation curve positively; 2) Increases prepulse and tail current | Blocks hKv4.3-hKChIP2.2, hCav3.2 and hKv1.5 and activates hCav1.2/β2/α2δ channels | WT/mutation Kv11.1, and other cardiac ion channels expressed cell lines | APC |
| LUF7244    | Strong affinity, allosteric site topologically distinct from where classic Kv11.1 blockers bind | 1) Slows rate of deactivation; 2) Shifts the inactivation curve positively; 3) Exerts a significant negative allosteric effect on the binding of typical Kv11.1 blockers; 4) A suppressive effect on proarrhythmia in neonatal rat ventricular myocyte monolayers | NR | HEK293 Kv11.1 cell lines | MPC, [3H] dofetilide-binding assays; radioligand-binding assays |
| LUF7346    | Allosteric site topologically distinct from where classic Kv11.1 blockers bind | 1) Slows the rate of deactivation, and 2) shifts activation curve negatively in iPSC-CMs with the N996I mutation, related to trafficking defect in LQT2 | NR | HEK293 Kv11.1 cell lines, iPSC-CMs from LQT2 patient | MPC, MEAs |
| Lumacaftor (LUM, clinical drug) | Allosteric site topologically distinct from where classic Kv11.1 blockers bind | Increases channel trafficking on the cell membrane and to reverse field potential duration prolongation in hiPSC-CMs derived from LQTS2 patients | NR | iPSC-CMs from LQT2 patient | MEAs, calcium imaging |
| SB-335573 (types 4 of Kv11.1 activators, a structural analog of the agonist NS3623) | Binding in either open or closed states of channel | Increases the open channel probability | No blocker effect on Cav1.2 and Nav1.5 | CHO-Kv11.1 stable cell line | APC |
| SKF-32802 (types 3 of Kv11.1 activators) | Binding in either open or closed states of channel | 1) Shifts the inactivation curve positively, and 2) Increases the open channel probability | No blocker effect on Cav1.2 and Nav1.5 | CHO-Kv11.1 stable cell line | APC |

APC, automated patch clamp; MEAs, microelectrode arrays; HEK293, human embryonic kidney 293 cells; CHO, Chinese hamster ovary cells.
sory protein interactions and signaling pathways (Figure 4F) (Table 8). Newly synthesized Kir2.1 could be sent to specific membrane subdomains for functional expression by Golgi according to a recognition site formed by the residues in the CTD and amino-terminal domain (NTD) and interaction with adaptor protein complex 1 (AP1). AKAP79 directly interacts with Kir2.1 through the intracellular N and C domains to promote anchoring other kinases (PKA, PKC) and close to Kir2.1 phosphorylation sites. PIP2 is an essential cofactor for activating Kir2.1 channel function. In addition, Kir2.1 closely interacts with Nav1.5 (Figure 1) and shares a coupled forward trafficking process with Nav1.5. Normal trafficking of Kir2.1 could enhance the functional expression of Nav1.5 compared to Nav1.5 alone, while trafficking-deficient variants disrupt Kir2.1 functional expression at the membrane and also exert a DN effect on Nav1.5 functional membrane expression. Thus, in addition to controlling resting MP, I_K1 could also modify Nav1.5 function and cell excitability. In turn, suppression of Nav1.5 by the CaMKII inhibitor KN93 or by trafficking-defective Nav1.5 variants could trap Kir2.1 channels, thus decreasing I_K1 in addition to I_Na.

Most KCNJ2 LOF mutations (Table 9) are associated with type 1 Andersen-Tawil syndrome (ATS), in which LQTS7 is the primary cardiac manifestation. Inactivation of I_K1 reduction could prolong the terminal phase of the cardiac AP and contribute to the development of DAD and ventricular arrhythmias in ATS. LOF mutations could suppress I_K1 via impairment of PIP2 gating, membrane trafficking, or potassium conduction. On the other hand, GOF mutations in KCNJ2 (Table 9) cause SQT3.

5 | CARDIOVASCULAR SAFETY EVALUATION

5.1 | Drug-induced cardiovascular arrhythmias

In addition to gene mutation-induced congenital arrhythmias, drug therapy could exert side effects on cardiac VGIC and increase the risk of life-threatening arrhythmias, such as drug-induced LQTS (diLQTS) and torsades de pointes (TdPs) that is morphologically distinctive polymorphic ventricular tachycardias with short-long-short cycles pattern. LQTS on the surface electrocardiogram correlates with ventricular AP repolarization prolongation at the cellular level. Drugs can induce AP repolarization prolongation by inhibiting I_Ks, or, more frequently, I_Kr. Due to the robustness of I_Kr, defective I_Ks by blockade of Kv7.1 might produce little AP prolongation in humans and other large mammals but might further prolong AP and induce LQT1 when challenged with β-AR stimulation or reduce repolarizing currents by drugs, especially I_Kr.

Kv11.1 is recognized as a predominant target for diLQTS and TdPs due to its intrinsic arrhythmogenic activity, although it is one of the interests of the development of antitachyarrhythmia drugs. The list of drugs that inhibit Kv11.1 includes not only includes antiarrhythmics (such as dofetilide) but also antipsychotics (such as Pimozide), gastroprokinetic agents (such as cisapride), antihistamines (such as astemizole), and other drug classes. Among all potassium channels, Kv11.1 is unique in having a small central cavity with extended pockets so that it is susceptible to direct blockade by a wide range of drugs. In addition, some drugs could exert inhibitory effects on Kv11.1 trafficking or coexisting effects of channel blocking and trafficking defects, thus causing diLQTS and TdPs.

Due to the increasing attention that diLQTS has attracted from clinics, drug developers, and pharmaceutical regulators, cardiovascular safety concerns are the most common reasons for the withdrawal of approved drugs from the market or the termination of potential drugs during preclinical or clinical trials. For example, the noncardiovascular drug cisapride has been withdrawn from the US market because it produces a modest increase in the QT interval in children, causing TdP; the drug exerts this effect by inhibiting Kv11.1. Since the outbreak of the coronavirus disease 2019 (COVID-19), many repurposed drugs are proposed as potential therapies for this disease; their risks, causing LQTS or TdPs is being evaluated.

5.2 | Development of drug safety evaluation guidelines

Since the guidelines, including the International Council for Harmonisation (ICH) S7B (nonclinical) and E14 (clinical), were announced in 2005, Kv11.1 channel safety screening data of new drug candidates before beginning clinical trials has become a great need in the pharmaceutical industry. However, promising drug candidates might be eliminated by the guidelines because variations in the potency of Kv11.1 blocking could result from varying patch clamp protocols and a poor ability to statistically quantify experimental variability. Moreover, promising drugs might be Kv11.1 blockers but exceptions in terms of causing TdPs or arrhythmia. Some also block other cardiac currents (Table 11) necessary for TdPs development but do not obviously prolong AP repolarization. Thus, in early multichannel studies, a model named multiple ion channel effects (MICE), based on the concentration-dependent responses of Kv11.1, Nav1.5, and Cav1.2 currents to torsadogenic and nontorsadogenic drugs, was
| Drug          | Class | Major effect                          | Effect on other channel | Effect on APD | Techniques | Clinical effect                                                                 | Current clinical trial                                                                 |
|--------------|-------|---------------------------------------|-------------------------|---------------|------------|--------------------------------------------------------------------------------|----------------------------------------------------------------------------------------|
| Quinidine    | Ia    | Nav1.5 open state inhibitor with intermediate dissociation kinetics | Potent Kv11.1 blocker (in S6 segment-, IC50 = 2 μM)289 | APD prolongation | MPC, APC89 | Clinical drug available for patients with SQTS290                              | NCT01873950 Phase I completed: Study of the Electrocardiographic Effects of Ranolazine, Dofetilide, Verapamil, and Quinidine in Healthy Subjects |
| Ranolazine   | New class Id | A potent inhibitor of late INa,L. | Potent Kv11.1 blocker234,258, No effect on Kv11.1 in SQTS1 N588K mutation patients82,291 | Modest APD prolongation | MPC82,291; APC85 | Treatment of angina pectoris82                                                  | 1) NCT01728025 Phase II completed: Long Term Prophylactic Therapy of Congenital Long QT Syndrome Type III (LQT3) With Ranolazine; 2) NCT02133352 Phase IV completed: Treatment of Pulmonary Hypertension Associated with diastolic left ventricular dysfunction; 3) NCT01721967 Phase IV completed: Treatment of Chest Pain in HCM Patients, Hypertrophic Cardiomyopathy; 4) NCT02360397 Phase 2 completed: Ventricular Premature Complexes, Myocardial Ischemia; NCT01349491 Phase III terminated: Prevention of Atrial Fibrillation After Electrical Cardioversion |
| GS-458967    | New class Id | Potent and selective inhibitor of INa,L. No effect on INa,P density82 | Minimal inhibition of IKs, IC50 ratio (IKs / INa,L > 76 folds)84 | No prolongation on APD and QRS interval82; reduction of APD prolongation in SCN5A-1795insD hiPSC-CMs292 | MPC84, 292 | NR, potential antiarrhythmic effects82                                           | No clinical trial yet                                                             |
| Ivabradine   | 0     | HCN channel blocker                   | Kv11.1 blocker166,169,170,293, Nav1.5 inhibition293 | APD prolongation in cardiomyocytes166; reverses APD shortening in N588K SQTS1 hiPSC-CMs292; no prolongation of ventricular-like APs in cardiomyocytes derived from iPSCs293 | MPC293, single-cell contraction measurement1 | Clinical drug available for reduction of heart rate in sinus tachycardia293        | NCT03866395 Phase IV completed: Ivabradine on Residual Myocardial Ischemia After PCI |
| Verapamil    | IV    | L-type Ca2+ channel blocker10         | Potent Kv11.1 blocker294 | No prolongation on APD294; decreases the QT interval293 | MPC and APC294 | Clinical drug available for heart rate control of atrial fibrillation          | NCT01873950 Phase I completed: Study of the Electrocardiographic Effects of Ranolazine, Dofetilide, Verapamil, and Quinidine in Healthy Subjects |

HCN, hyperpolarization-activated cyclic nucleotide.
Clinical trial homepage: https://clinicaltrials.gov.
proposed to be more effective than Kv11.1 assays in predicting TdPs.

Although the current paradigm has largely kept potential torsadogenic drugs off the markets, but a new cardiac safety paradigm with comprehensive model-informed approach rather than exclusively by potency of Kv11.1 block and by QT prolongation is urgent to adopted to improve the deficiencies of current paradigm, more specifically discern a real proarrhythmic risk of promising drugs, and enhance the development of effective and safe products or therapeutics. 

In 2013, several organizations formed a team to develop the Comprehensive In Vitro Proarrhythmia Assay (CiPA) initiative, a new paradigm developed with the goal of presenting a deeper understanding of the mechanism of TdPs and improving the assessment of the proarrhythmic effects of potential drugs. It is driven by mechanistically based in vitro assays of drug effects on multiple cardiac channels coupled in silico reconstruction of cardiac AP, and comparison of predicted and observed responses in human-derived cardiac myocytes. Twenty-eight drugs with well-characterized three torsadogenic risk groups (Table S3) have been selected and considered as test cases to build/calibrate model for testing and validation of in silico and stem cell CIPA models. Several working groups are involved in developing the CiPA:

1. The ion channel group is developing voltage-clamp protocols by MPC or APC for several key cardiac ion channels. It is believed that at least six ion channels are involved in cardiac APs: Nav1.5, Kv4.3, Cav1.2, Kv11.1, Kv7.1, and Kir2.1. A study evaluated the predictive ability of these six ion channels using APC and showed that four ion channels provided good predictions, whereas the analysis of three channels wrongfully predicted one high-risk drug to be safe. Improved systematic approaches for accurately estimating the potency and safety margins are required. Increasing APC-based assays have been explored in Kv11.1, Nav1.5, Cav1.2, Kir2.1, Kv7.1, and Kv4.3 to improve the evaluation strategies.

2. The in silico group is building computer models to reconstruct electrophysiological activities and drug effects on multiple human cardiac currents by integrating experimental data within a heart cell and subsequently outputting the net impact on the cellular APD and QT interval for predicting drug-induced proarrhythmic risks. For example, by using an in silico model, several proposed drugs against COVID-19 are estimated to have a significant risk for LQTS; thus, mandatory monitoring of the QT interval should be performed among patients in use of drugs. In silico models are keeping updated to expand the index for discriminating TdPs compounds and to satisfy a series of general principles for the validation of proarrhythmia risk prediction. Those principles will help shape the future important directions of more accurate prediction models. For example, development of better simulating models to capture the drug response not only in normal humans but also in specific patient populations. With the application of in silico modeling, machine learning could identify cellular electrophysiological phenotypes associated with patients who has certain cardiac diseases and further predict which patients face an elevated risk of ventricular arrhythmias and sudden death. However, information such as comparisons among drugs with similar chemical or affinity profiles is not yet possible incorporated into in silico model. Thus, newer proarrhythmia risk prediction models could be developed to aid in decision making. For example, a computational pipeline was recently developed to predict Kv11.1 blocker proarrhythmic risk from drug chemistry and distinguish drugs that have similar chemistry and effects on the AP and QT interval but different proarrhythmic risk levels.

3. The myocyte group used iPSC-CM assays to evaluate the in vitro and in silico assay results. Native human cardiomyocytes are ideal but with difficulties to obtain, maintain in long-term culture. Native cardiomyocytes from different species have variations in the waveforms of APs and drug responses due to differences of potassium currents densities. Rodent is not an appropriate specie for modelling human repolarization due to dominant I_{Kr}; dogs and rabbits are relative closely to human due to the major role of I_{Ks} in repolarization. Thus, the need for proarrhythmia evaluation in preclinical studies based on human models is emphasized. Currently, hiPSC-CMs have provided a perfect platform for proarrhythmia evaluation and safety evaluation of human cardiomyocytes in preclinical studies, and various AP parameters could be measured using high-throughput systems.

4. The clinical translation group will use clinical Phase I ECGs to evaluate potential unanticipated effects.

In addition, evaluating the effect of compounds on the overall APs rather than a single ion channel current has been proposed to be a more appropriate approach. However, the more depolarized resting MP of hiPSC-CMs than that of primary cardiomyocytes is the limitation and challenge of their use in safety evaluation. This is due to the distinct expression level of ion channels expression compared to primary cardiomyocytes, especially the low expression of the I_{K1} channel protein Kir2.1. Exogenous overexpression of the Kir2.1 or electronic injection of an I_{K1}-like current by dynamic clamp into hiPSC-CMs to compensate and thus achieve more stable AP facilitates
6 | CONCLUSION AND PERSPECTIVE

This review provides detailed descriptions of major ion channels in ventricular myocytes, including their expression, structures, regulators, and contributions to normal excitability and congenital pathology. It has been discussed that the application basic and newly discovered knowledge of cardiac ion channels and the continuous development of techniques employed in studies of cardiac ion channels can lead to more attentions to comprehensive proarrhythmic risk assessment in human cardiomyocytes platform in preclinical studies and promote development of cardiovascular safety evaluation guidelines.

Recent research on potential targets of interest in cardiomyocytes, such as TTXs sensitive or TTX-insensitive Nav and Ryr2 regulatory TRICA channels, has opened new avenues for improving our understanding of the molecular mechanisms of Ca^{2+} homeostasis, EC coupling, and associated cardiac disease pathogenesis. The development of a Nav-selective inhibitor or a heart-specific Nav channel-KO mouse model will be beneficial for further confirming the pathological mechanism of specific Nav channels. The selective inhibition of Nav channels may offer a potential therapeutic target to alleviate arrhythmias during states of Ca^{2+} overload. The development of hiPSC-CM, high-throughput techniques for cellular phenotype detection (such as Aps and contraction), computational simulation models facilitate integration of multiple channels, achieving a comprehensive view of channelopathies as a global phenomenon in human myocytes. Modeling of patient-specific iPSC-CMs provides great benefit for the precision medicine treatment of congenital cardiac arrhythmia and for the screening of promising or already approved drugs to test for mutation-specific antiarrhythmic effects.

Over time, technological developments will certainly further promote the study of an increasing number of scientific questions related to cardiac physiology and pathology and reveal additional ion channels with potential involvement. Based on cryo-EM structures of many VGICs in basic science, a large body of experimental and clinical observations concerning VGICs has been interpreted and summarized by the structural template. The development of clinical and translational medicine could be advanced by the discovery of the potential drug targets within many VGICs, as well as drugs characteristics, targets-drugs interaction, and computational models for integrating and predicting information. Recently, a novel multiscale approach has been developed to predict drug-induced arrhythmia directly based on structural models of drug-channel interactions and kinetics by using integrative experimental and computational modeling and machine-learning from the atom to the rhythm in the heart.

For potential targets, cryo-EM structures map and classify hundreds of clinical arrhythmia variants onto all major domains in the structure of many VGICs reveal the common or distinct clusters of arrhythmia mutations among different types of VGICs or different isoforms of the same VGIC, provide the molecular basis for understanding disease mechanisms, and thus allow the development of structure-based diagnosis and drug discovery for arrhythmias in the future. For clinical or potential drugs, the cryo-EM structure of VGIC-drug interactions provide structural insights into the binding affinity and mechanism of drugs, which is beneficial for modifying the structure of drugs, screening alternatives or synthesizing new compounds. For example, cryo-EM structure of Kv11.1 channel in the open state promotes the investigation of the state-dependent blockade of Kv11.1 by the heart-rate-lowering agent ivabradine, which could also exert antiarrhythmic effects in SQTS1 hiPSC-CMs with the N588K mutation. The development of novel additional pharmacological approaches (eg, activators/allosteric modulators of potassium Kv11.1 and Kv7.1 channels) are needed to counteract both congenital LQTSs, although currently available therapies (implantable cardioverter defibrillators) have yielded good clinical responses. For example, Lumacaftor, a drug already in clinical use for cystic fibrosis, has been demonstrated to interact with a site distinct from where classic Kv11.1 blockers bind, thereby restoring Kv11.1 trafficking defects and alleviating LQTS2. Polyunsaturated fatty acids (PUFAs) and their analogs N-arachidonoyl taurine have been found to speed up Kv7.1

clinical applications, drug discovery, and cardiotoxicity screening. Although it is unclear when the CiPA project will lead to new guidelines (as organizations are generally conservative when considering changes to effective standard protocols), the CiPA initiative and other similar projects worldwide are promoting the development of questions and answers (Q&As) to facilitate the application of the ICH S7B and E14 guidelines. With the development of these techniques, other cardiac safety liabilities, such as dysfunction of EC coupling and contractile and structural cardiotoxicity, may also be added to electrophysiological tests in the same platform to complement CiPA for regulatory use.

In general, with the development of medium- or high-throughput test systems to produce efficient, reliable result output and of basic knowledge of VGICs to update the detection assay designs and analysis methods, drug safety evaluation will receive more attention in preclinical research. Evaluation will be conducted as early as possible to avoid further unnecessary investments in unusable compounds during later stages of drug development.
channel opening and restore channel gating of many different mutant channels. PUFAs and their analogs are effective in shortening the cardiac action potential in pharmacologically prolonged ventricular action potential and QT interval in isolated guinea pig hearts and in hPSC-CM. Therefore, activators of Kv7.1 are also worth developing to treat LQT1 based on structure-function studies on diverse IKs channel mutations. However, PUFAs analogs vary in selectivity and different effects for Kv7.1, Nav1.5, and Cav1.2 through nonidentical mechanisms. It is necessary to determine the specific binding sites of PUFAs analogs among normal VGICs and to further identify the most therapeutically relevant PUFAs and PUFA analogs in the treatment of different LQTS subtypes. Moreover, if negative allosteric modulators are used in combination with patient-specific hPSC-CM, drugs that have been withdrawn from market or excluded from clinical application due to diLQT effects may be reconsidered or even rescued to clinical use after safety validation by electrophysiological approaches.

In addition, site-specific and target-oriented approaches using nanomaterials (NMs) have been increasingly applied but might exert potential toxicity on ion channels and cardiac electrophysiology. Maybe revealing these NM-induced structural changes in ion channels could facilitate the modification of bioactive NPs to optimize NM-based drug delivery and safety. In general, in-depth studies that combine electrophysiological approaches with other technologies are being used to explore the expression, function, mechanism, and structure, and activity modulation of WT VGICs and a broad variety of mutated VGICs, providing critical contributions to our knowledge of the roles of VGICs in both normal and diseased cardiac functions, thus facilitating to the discovery of potential structurally and functionally guided drug targets for the modification of channel function and for the treatment of inherited or drug induced cardiac diseases, providing a basis for structure- and mechanism-based personalized clinical management, prompting safety control committees to establish more integrated strategies for drug screening, and enabling improved prediction of cardiac risks to provide safer and more effective drugs for clinical use.

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AUTHOR CONTRIBUTIONS
Hua Li, Xiangdong Wang, and Junbo Ge proposed the conception, study design, and had the final approval of the manuscript submitted. Lulan Chen and Yue He participated in the data collections and analysis, the drafting of the manuscript, and the submission.

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REFERENCES
1. Bartos DC, Grandi E, Ripplinger, CM. Ion channels in the heart. Comprehen Physiol. 2015;5:1423-1464.
2. Garg P, Garg V, Shrestha R, Kamp TJ, Wu JC. Human Induced pluripotent stem cell-derived cardiomyocytes as models for cardiac channelopathies: a primer for non-electrophysiologists. Circ Res. 2018;123:224-243.
3. Roden DM. Drug-induced prolongation of the QT interval. N Engl J Med. 2004;350:1013-1022.
4. Fermini B, Fossa AA. The impact of drug-induced QT interval prolongation on drug discovery and development. Nat Rev Drug Discov. 2003;2:439-447.
5. Fang L, Sager P, Yang X, et al. Workshop report: FDA workshop on improving cardiotoxicity assessment with human-relevant platforms. Circ Res. 2019;125:885-887.
6. Sager PT, Gintant G, Turner JR, Pettit S, Stockbridge N. Rechanneling the cardiac proarrhythmia safety paradigm: a meeting report from the Cardiac Safety Research Consortium. Am Heart J. 2014;167:292-300.
7. Sallam K, Li Y, Sager PT, Houser SR, Wu JC. Finding the rhythm of sudden cardiac death: new opportunities using induced pluripotent stem cell-derived cardiomyocytes. Circ Res. 2015;116:1989-2004.
8. Wang XH, Su M, Gao F, et al. Structural basis for activity of TRIC counter-ion channels in calcium release. P Natl Acad Sci USA. 2019;116:4238-4243.
9. Voigt N, Li N, Wang Q, et al. Enhanced sarcoplasmic reticulum Ca2+ leak and increased Na+-Ca2+ exchanger function underlie delayed afterdepolarizations in patients with chronic atrial fibrillation. Circulation. 2012;125 2059-2070.
10. Kramer J, Himmel HM, Lindqvist A, et al. Cross-site and cross-platform variability of automated patch clamp assessments of drug effects on human cardiac currents in recombinant cells. Sci Rep. 2020;10:5627.
11. Obergrossberger A, Goetzke TA, Brinkwirth N, et al. An update on the advancing high-throughput screening techniques for patch clamp-based ion channel screens: implications for drug discovery. Expert Opin Drug Discov. 2018;13:269-277.
12. Dunlop J, Bowby M, Peri R, Vasilyev D, Arias R. High-throughput electrophysiology: an emerging paradigm for ion-channel screening and physiology. Nat Rev Drug Discov. 2008;7:358-368.
13. Meyer T, Boven KH, Günther E, Fejtl M. Micro-electrode arrays in cardiac safety pharmacology—a novel tool to study QT interval prolongation. Drug Safety. 2004;27 763-772.
14. Spira ME, Hai A. Multi-electrode array technologies for neuroscience and cardiology. Nat Nanotechnol. 2013;8:83-94.
15. Obergrossbauer A, Juhasz K, Thomas U, et al. Safety pharmacology studies using iPSC-derived cardiomyocytes. J Pharmacol Toxicol Met. 2016;81:223-232.
16. Moller C, Witchel H. Automated electrophysiology makes the pace for cardiac ion channel safety screening. Front Pharmacol. 2011;2:73.

Shinozawa T, Nakamur K, Shoji M, et al. Recapitulation of clinical individual susceptibility to drug-induced QT prolongation in healthy subjects using iPSC-derived cardiomyocytes. Stem Cell Reports. 2017;8:226-234.
18. Jiang D, Shi H, Tonggu Q, et al. Structure of the cardiac sodium channel. Cell. 2020;180(1):122-134.
19. Wang WW, MacKinnon R. Cryo-EM structure of the open human ether-a-go-go-related K+ channel hERG. Cell. 169, 422-(2017).
20. Zhao Y, Huang G, Wu J, et al. Molecular basis for ligand modulation of a mammalian voltage-gated Ca(2+) channel. Cell. 177, 1495-1506 e1412 (2019).
21. Pan X, Li Z, Jin X, et al. Comparative structural analysis of human Nav1.1 and Nav1.5 reveals mutational hotspots for sodium channelopathies. Proc Natl Acad Sci U S A. 118, e210066118 (2021).
22. Perissinotti L, Guo J, Kudaibergenova M, et al. The pore-lipid interface: role of amino-acid determinants of lipophilic access by ivabradine to the hERG pore domain. Mol Pharmacol. 2019;96:259-271.
23. Herron TJ, Lee P, Jalife J. Optical imaging of voltage and calcium in cardiac cells & tissues. Circ Res. 2012;110:609-623.
24. Yang YX, Liu N, He YY, et al. Improved calcium sensor GCaMP-X overcomes the calcium channel perturbations induced by the calmodulin in GCaMP. Nat Commun. 9, 1504 (2018).
25. Shinnawi R, Huber I, Maizels L, et al. Monitoring human-induced pluripotent stem cell-derived cardiomyocytes with genetically encoded calcium and voltage fluorescent reporters. Stem Cell Reports. 2015;5:582-596.
26. Ferenczi EA, Tan XQ, Huang CLH. Principles of optogenetic methods and their application to cardiac experimental systems. Front Physiol. 10, 1096 (2019).
27. Boyle PM, Karathanos TV, Trayanova NA. Cardiac optogenetics: 2018. JACC Clin Electrophysiol. 2018;4:155-167.
28. Bruegmann T, Malan D, Hesse M, et al. Optogenetic control of heart muscle in vitro and in vivo. Nat Methods. 2010;7:897-900.
29. Nussinovitch U, Shinnawi R, Gepstein L. Modulation of cardiac tissue electrophysiological properties with light-sensitive proteins. Cardiovasc Res. 2014;102:176-187.
30. Rehnelt S, Malan D, Juhasz K, et al. Frequency-dependent multi-well cardiotoxicity screening enabled by optogenetic stimulation. Int J Mol Sci. 18, 2634 (2017).
31. Huang CL. Murine Electrophysiological models of cardiac arrhythmogenesis. Physiol Rev.. 2017;97:283-409.
32. Veeraraghavan R, Gyorkes S, Radwanski PB. Neuronal sodium channels: emerging components of the nano-machinery of cardiac calcium cycling. J Physiol. 2017;595:3823-3834.
33. Johnson CN. Calcium modulation of cardiac sodium channels. J Physiol. 598(14), 2835-2846 (2019).
34. Chen-Izu Y, Shaw RM, Pitt GS, et al. Na+ channel function, regulation, structure, trafficking and sequestration. J Physiol. 2015;593:1347-1360.
35. Van Petegem F, Lobo PA, Ahern CA. Seeing the forest through the trees: towards a unified view on physiological calcium regulation of voltage-gated sodium channels. Biophys J. 2012;103:2243-2251.
36. O’Malley HA, Isom LL. Sodium channel beta subunits: emerging targets in channelopathies. Annu Rev Physiol. 2015;77:481-504.
37. Chandler NJ, Greener ID, Tellez JO, et al. Molecular architecture of the human sinus node: insights into the function of the cardiac pacemaker. Circulation. 2009;119:1562-1575.
38. Gaborit N, Le Bouter S, Szuts V, et al. Regional and tissue specific transcript signatures of ion channel genes in the non-diseased human heart. J Physiol. 2007;582:675-693.
39. Maier SK, Westenbroek RE, Schenckman KA, Feigl EO, Scheuer T, Catterall WA. An unexpected role for brain-type sodium channels in coupling of cell surface depolarization to contraction in the heart. Proc Natl Acad Sci U S A. 2002;99:4073-4078.
40. Radwanski PB, Brunello L, Veeraraghavan R, et al. Neuronal Na+ channel blockade suppresses arrhythmogenic diastolic Ca2+ release. Cardiovasc Res. 2015;106:143-152.
41. Radwanski PB, Ho HT, Veeraraghavan R, et al. Neuronal Na+( ) channels are integral components of pro-arrhythmic Na(+)/Ca(2+) signaling nanodomain that promotes cardiac arrhythmias during beta-adrenergic stimulation. JACC Basic Trans Sci. 2016;1:251-266.
42. Dybkova N, Ahmad S, Pabel S, et al. Differential regulation of sodium channels as a novel proarrhythmic mechanism in the human failing heart. Cardiovasc Res. 2018;114:1728-1737.
43. Pabel S, Ahmad S, Tirilomis P, et al. Inhibition of Nav1.8 prevents atrial arrhythmogenesis in human and mice. Basic Res Cardiol. 2020;115:20.
44. Yang T, Atack TC, Stroud DM, Zhang W, Hall L, Roden DM. Blocking Scn10a channels in heart reduces late sodium current and is antiarrhythmic. Circ Res. 2012;111:332-332.
45. Dhar Malhotra J, Chen C, Rivolta I, et al. Characterization of sodium channel alpha- and beta-subunits in rat and mouse cardiac myocytes. Circulation. 2001;103:1303-1310.
46. Pereon Y, Lande G, Demolombe S, et al. Paramyotonia congenita with an SCN4A mutation affecting cardiac repolarization. Neurology. 2003;60:340-342.
47. Westenbroek RE, Lande G, Demolombe S, Maier SK, Catterall WA, Scheuer T. Localization of sodium channel subtypes in mouse ventricular myocytes using quantitative immunocytochemistry. J Mol Cell Cardiol. 2013;64:69-78.
48. Chambers JC, Zhao J, Terracciano CM, et al. Genetic variation in SCN10A influences cardiac conduction. Nat Genet. 2010;42:149-152.
49. Rook MB, Evers MM, Vos MA, Bierhuizen MF. Biology of cardiac sodium channel Nav1.5 expression. Cardiovasc Res. 2012;93:283-409.
50. Antzelevitch C, Nesterenko V, Shroyock JC, Rajamani S, Song Y, Belardinelli L. The role of late I Na in development of cardiac arrhythmias. Handb Exp Pharmacol. 2014;221:137-168.
51. Chadda KR, Jeevaratnam K, Lei M, Huang CL. Sodium channel biophysics, late sodium current and genetic arrhythmic syndromes. Fplgers Arch. 2017;469:629-641.
52. Pan X, Li Z, Huang X, et al. Molecular basis for pore blockade of human Na(+) channel Nav1.2 by the mu-conotoxin KIIIA. *Science*. 2019;363:1309-1313.

53. Pan X, Li Z, Zhou Q, et al. Structure of the human voltage-gated sodium channel Nav1.4 in complex with betal. *Science*. 2018;362(6418):1147-1152.

54. Edokobi N, Isom LL. Voltage-gated sodium channel beta1/beta1B subunits regulate cardiac physiology and pathophysiology. *Front Physiol*. 2018;9:351.

55. Cortada E, Brugada R, Verges M. Trafficking and function of the voltage-gated sodium channel beta2 subunit. *Biomolecules*. 2019;9, 604 (2019).

56. Clatot J, Hoshi M, Wan XP, et al. Voltage-gated sodium channel beta1/beta1B subunits regulate cardiac physiology and pathophysiology. *Front Physiol*. 2018;9:351.

57. Johnson CN, Potet F, Thompson MK, et al. A Mechanism of calmodulin modulation of the human cardiac sodium channel. *Structure*. 2018;26:683-694 e683.

58. Nof E, Vysochek L, Meisel E, et al. Mutations in NaV1.5 reveal calcium-calmodulin regulation of sodium channel. *Front Physiol*. 2019;10:700.

59. Potet F, Chagot B, Anghelescu M, et al. Functional interactions between distinct sodium channel cytoplasmic domains through the action of calmodulin. *J Biol Chem*. 2009;284:8846-8854.

60. Sarhan MF, Van Petegem F, Ahern CA. A double tyrosine motif in the cardiac sodium channel domain III-IV linker couples calcium-dependent calmodulin binding to inactivation gating. *J Biol Chem*. 2009;284:33265-33274.

61. Sarhan MF, Tung CC, Van Petegem F, Ahern CA. Crystallographic basis for calcium regulation of sodium channels. *PNat Acad Sci USA*. 2012;109:3558-3563.

62. El Refaey M, Musa H, Murphy NP, et al. Protein phosphatase 2A regulates cardiac Na(+)-channels. *Circ Res*. 2019;124:737-746.

63. Makara MA, Curran J, Little SC, et al. Ankyrin-G coordinates intercalated disc signaling platform to regulate cardiac excitability in vivo. *Circ Res*. 2014;115:929-938.

64. Johnson CN, Pattanayek R, Potet F, et al. The CaMKII inhibitor KN93-calmodulin interaction and implications for calmodulin tuning of NaV1.5 and RyR2 function. *Cell Calcium*. 2019;62:102063.

65. Wilde AAM, Amin AS. Clinical spectrum of SCN5A mutations: long QT syndrome, Brugada syndrome, and cardiomyopathy. *JACC Clin Electrophysiol*. 2018;4:569-579.

66. Terrenoire C, Wang K, Tung KW, et al. Induced pluripotent stem cells used to reveal drug actions in a long QT syndrome family with complex genetics. *J Gen Physiol*. 2013;141:61-72.

67. Ortiz-Bonnay B, Rinné S, Moss R, et al. Electrophysiological characterization of a large set of novel variants in the SCN5A-gene: identification of novel LQT53 and BrS mutations. *Pflugers Arch*. 2016;468:1375-1387.

68. Keller DI, Barrane FZ, Gouas L, et al. A novel nonsense mutation in the SCN5A gene leads to Brugada syndrome and a silent gene mutation carrier state. *Can J Cardiol*. 2005;21:925-931.

69. Núñez L, Barana A, Amorós I, et al. p.D1690N Nav1.5 rescues p.G1748D mutation gating defects in a compound heterozygous Brugada syndrome patient. *Heart Rhythm*. 2013;10(2):264-272.

70. Clatot J, Ziyadeh-Iseem A, Maugenre S, et al. Dominant-negative effect of SCN5A N-terminal mutations through the interaction of Na(v)1.5 alpha-subunits. *Cardiovasc Res*. 2012;96:53-63.

71. Dharmawan T, Nakajima T, Iizuka T, et al. Enhanced closed-state inactivation of mutant cardiac sodium channels (SCN5A N1541D and R1632C) through different mechanisms. *J Mol Cell Cardiol*. 2019;130:88-95.

72. Nakajima T, Kaneko Y, Saito A, Ota M, Ijiiima T, Kurabayashi M. Enhanced fast-inactivated state stability of cardiac sodium channels by a novel voltage sensor SCN5A mutation, R1632C, as a cause of atypical Brugada syndrome. *Heart Rhythm*. 2015;12:2296-2304.

73. Garcia-Molina E, Lacunza J, Ruiz-Espejo F, et al. A study of the SCN5A gene in a cohort of 76 patients with Brugada syndrome. *Clin Genet*. 2013;83:530-538.

74. Glazer AM, Wada Y, Li B, et al. High-throughput reclassification of SCN5A variants. *Am J Hum Genet*. 2020;107:111-123.

75. Watanabe H, Koopmann TT, Le Scouarnec S, et al. Sodium channel betal subunit mutations associated with Brugada syndrome and cardiac conduction disease in humans. *J Clin Invest*. 2008;118:2260-2268.

76. Riuro H, Beltran-Alvarez P, Tarradas A, et al. A Missense mutation in the sodium channel 2 subunit reveals SCN2B as a new candidate gene for Brugada syndrome. *Hum Mut*. 2013;34:961-966.

77. Peters CH, Watkins AR, Poirier OL, Ruben PC. E1784K, the most common Brugada syndrome and long-QT syndrome type 3 mutant, disrupts sodium channel inactivation through two separate mechanisms. *J Gen Physiol*. 152, e202012595 (2020).

78. Makita N, Behr E, Shimizu W, et al. The E1784K mutation in SCN5A is associated with mixed clinical phenotype of type 3 long QT syndrome. *J Clin Invest*. 2008;118:2219-2229.

79. Steinberg C, Pilote S, Philippon F, et al. SCN5A-C683R exhibits combined gain-of-function and loss-of-function properties related to adrenaline-triggered ventricular arrhythmia. *Exp Physiol*. 2021;106:683-699.

80. Hoshi M, Du XX, Shinnalapattayatorn K, et al. Brugada syndrome disease phenotype explained in apparently benign sodium channel mutations. *Circ Cardiovasc Genet*. 2014;7:123-131.

81. Cheng J, Valdivia CR, Vaidyanathan R, Balijepalli RC, Ackerman MJ, Makielki JC. Caveolin-3 suppresses late sodium current by inhibiting nNOS-dependent S-nitrosylation of SCN5A. *J Mol Cell Cardiol*. 2013;61:102-110.

82. Chorin E, Hu D, Antzelevitch C, et al. Ranolazine for congenital long-QT syndrome type III: experimental and long-term clinical data. *Circ Arrhythm Electrophysiol*. 2016;9:e004370.

83. Erdemli G, Kim AM, Ju HS, Springer C, Penland RC, Hoffman PK. Cardiac safety implications of hNav1.5 blockade and a framework for pre-clinical evaluation. *Front Pharmacol*. 2012;3:63.

84. Belardinelli L, Liu GX, Smith-Maxwell C, et al. A novel, potent, and selective inhibitor of cardiac late sodium current suppresses experimental arrhythmias. *J Pharmacol Exp Ther*. 2013;344:26-32.

85. Guo D, Jenkinson S. Simultaneous assessment of compound hNav1.5 channels by an novel voltagesensor SCN5A mutation, R1632C, as a cause of atypical Brugada syndrome. *Heart Rhythm*. 2015;12:2296-2304.

86. Riuro H, Beltran-Alvarez P, Tarradas A, et al. A Missense mutation in the sodium channel 2 subunit reveals SCN2B as a new candidate gene for Brugada syndrome. *Hum Mut*. 2013;34:961-966.
86. Struckman HL, Baine S, Thomas J, et al. Super-resolution imaging using a novel high-fidelity antibody reveals close association of the neuronal sodium channel NaV1.6 with ryanodine receptors in cardiac muscle. *Microsc Microanal.* 2020;26:157-165.

87. Frasier CR, Wagnon JL, Bao YO, et al. Cardiac arrhythmia in a mouse model of sodium channel SCN8A epileptic encephalopathy. *Proc Natl Acad Sci USA.* 2016;113:12838-12843.

88. Behr ER, Savio-Galimberti E, Barc J, et al. Role of common and rare variants in SCN10A: results from the Brugada syndrome QRS locus gene discovery collaborative study. *Cardiovasc Res.* 2015;106:520-529.

89. Gando I, Williams N, Fishman GI, Sampson BA, Tang Y, Coetzee WA. Functional characterization of SCN10A variants in several cases of sudden unexplained death. *Forensic Sci Int.* 2019;301:289-298.

90. Catterall WA. Voltage-gated calcium channels. *Cold Spring Harb Perspect Biol.* 2011;3:a003947.

91. Sorrentino V, Volpe P. Ryanodine receptors-how many, where and why. *Trends Pharmacol Sci.* 1993;14(3):98-103.

92. Lanner JT, Georgiou DK, Joshi AD, Hamilton SL. Ryanodine receptors: structure, expression, molecular details, and function in calcium release. *Cold Spring Harb Perspect Biol.* 2010;2:a003996.

93. Otsu K, Willard HF, Khanna VK, Zorzato F, Green NM, MacLennan DH. Molecular cloning of cDNA encoding the Ca2+ release channel (ryanodine receptor) of rabbit cardiac muscle sarcoplasmic reticulum. *J Biol Chem.* 1990;265:13472-13483.

94. Hering S, Zangerl-Plessl EM, Beyl S, Hohaus A, Andraou S. Regulation of the heart expressing the Ser1700A/Thr1704A channel Cav1.1 complex. *Science* 2015;348:13472-13483.

95. Dai S, Hall DD, Hell JW. Supramolecular assemblies and localized regulation of voltage-gated ion channels. *Physiol Rev.* 2009;89:411-452.

96. Davies A, Hendrich J, Van Minh AT, Wratten J, Douglas L, Dolphin AC. Functional biology of the alpha(2)delta subunit of voltage-gated Ca2+ channels. *Nature* 2009;572:347-354.

97. Wei J, Yao J, Belke D, et al. Ca(2+)-CaM dependent inactivation of RyR2 underlies Ca(2+) alternans in intact heart. *Circ Res.* 2013;112:163-163.

98. Poomvanicha M, Matthes J, Domes K, et al. Beta-adrenergic regulation of voltage-gated ion channels. *Pflugers Arch.* 2015;470:1291-1309.

99. Meyers MB, Puri TS, Chien AJ, et al. Sorcin associates with the pore-forming subunit of voltage-dependent L-type Ca2+ channels. *J Biol Chem.* 1998;273:18930-18935.

100. Meyers MB, Puri TS, Chien AJ, et al. Sorcin associates with the pore-forming subunit of voltage-dependent L-type Ca2+ channels. *J Biol Chem.* 1998;273:18930-18935.

101. Zhang L, Kelley J, Schmeisser G, Kobayashi YM, Jones LR. Complex formation between junctin, triadin, calsequestrin, and the ryanodine receptor. Proteins of the cardiac junctional sarcoplasmic reticulum membrane. *J Biol Chem.* 1997;272:23389-23397.

102. Meyers MB, Puri TS, Chien AJ, et al. Sorcin associates with the pore-forming subunit of voltage-dependent L-type Ca2+ channels. *J Biol Chem.* 1998;273:18930-18935.
120. Zhou XY, Lin PH, Yamazaki D, et al. Trimeric intracellular cation channels and sarcoplasmic/endoplasmic reticulum calcium homeostasis. Circ Res. 2014;114:706-716.

121. Zhou X, Park KH, Yamazaki D, et al. TRIC-A channel maintains store calcium handling by interacting with type 2 ryanodine receptor in cardiac muscle. Circ Res. 2020;126:417-435.

122. El-Ajouz S, Venturi E, Witschas K, et al. Dampered activity of ryanodine receptor channels in mutant skeletal muscle lacking TRIC-A. J Physiol. 2017;595:4769-4784.

123. Priori SG, Napolitano C, Memmi M, et al. Clinical and molecular characterization of patients with catecholaminergic polymorphic ventricular tachycardia. Circulation. 2002;106:69-74.

124. George CH, Jundi H, Walters N, Thomas NL, West RR, Lai FA. Arrhythmogenic mutation-linked defects in ryanodine receptor autoregulation reveal a novel mechanism of Ca2+ release channel dysfunction. Circ Res. 2006;98:88-97.

125. Kannankeril PJ, Mitchell BM, Goonasekera SA, et al. Mice with the R176Q cardiac ryanodine receptor mutation exhibit catecholamine-induced ventricular tachycardia and cardiomyopathy. Proc Natl Acad Sci U S A. 2006;103:12179-12184.

126. Sun B, Yao J, Ni M, et al. Cardiac ryanodine receptor calcium release deficiency syndrome. Sci Transl Med. 13, eaba7287 (2021).

127. Bezzarides VJ, Caballero A, Wang S, et al. Gene therapy for catecholaminergic polymorphic ventricular tachycardia by inhibition of Ca(2+)-calmodulin-dependent kinase II. Circulation. 2019;140:405-419.

128. Liu B, Walton SD, Ho HT, et al. Gene Transfer of engineered calmodulin alleviates ventricular arrhythmias in a calsequestrin-associated mouse model of catecholaminergic polymorphic ventricular tachycardia. J Am Heart Assoc. 7, e008155 (2018).

129. Kryshtal DO, Blackwell DJ, Egly CL, et al. RYR2 channel inhibition is the principal mechanism of flecainide action in CPVT. Circ Res. 2021;128:321-331.

130. Neronne JM, Kass RS. Molecular physiology of cardiac repolarization. Physiol Rev. 2005;85:1205-1253.

131. Grandi E, Sanguinetti MC, Bartos DC, et al. Potassium channels in the heart: structure, function and regulation. J Physiol. 2017;595:2209-2228.

132. Chiamvimonvat N, Chen-Izu Y, Clancy CE, et al. Potassium currents in the heart: functional roles in repolarization, arrhythmia and therapeutics. J Physiol. 2017;595:2229-2252.

133. Niwa N, Neronne JM. Molecular determinants of cardiac transient outward potassium current (I-to) expression and regulation. J Mol Cellular Cardiol. 2010;48:12-25.

134. Cho JH, Zhang R, Kilfoil PJ, et al. Delayed repolarization underlies ventricular arrhythmias in rats with heart failure and preserved ejection fraction. Circulation. 2017;136:2037-2050.

135. Panamà BK, Korogji AS, Aschar-Sobbi R, et al. Reductions in the cardiac transient outward K+ current I-to caused by chronic beta-adrenergic receptor stimulation are partly rescued by inhibition of nuclear factor kappa B. J Biol Chem. 2016;291:4156-4165.

136. Clatot J, Neyroud N, Cox R, et al. Inter-regulation of Kv4.3 and voltage-gated sodium channels underlies predisposition to cardiac and neuronal channelopathies. Int J Mol Sci. 21, 5057 (2020).

137. Porto V, Wilders R, Casini S, Charpentier F, Verkerk AO, Remme CA. Kv4.3 expression modulates NaV1.5 sodium current. Front Physiol. 2018;9:178.

138. Deschenes I, Armoundas AA, Jones SP, Tomaselli GF. Post-transcriptional gene silencing of KCNIP2 and Navbeta1 in neonatal rat cardiac myocytes reveals a functional association between Na and Ito currents. J Mol Cell Cardiol. 2008;45:336-346.

139. Nguyen HM, Miyazaki H, Hoshi N, et al. Modulation of voltage-gated K+ channels by the sodium channel betal subunit. Proc Natl Acad Sci U S A. 2012;109(45):18577-18582.

140. Giudicessi JR, Ye D, Tester DJ, et al. Transient outward current (I(to)) gain-of-function mutations in the KCND3-encoded Kv4.3 potassium channel and Brugada syndrome. Heart Rhythm. 2011;8:1024-1032.

141. Takayama K, Ohno S, Ding WG, et al. A de novo gain-of-function KCND3 mutation in early repolarization syndrome. Heart Rhythm. 2019;16:1698-1706.

142. Giudicessi JR, Ye D, Kritzberger CJ, et al. Novel mutations in the KCND3-encoded Kv4.3 K+ channel associated with autopsy-negative sudden unexplained death. Hum Mutat. 2012;33:989-997.

143. Barro-Soria R, Ramentol R, Llini SI, Perez ME, Kass RS, Larsen HP. KCNE1 and KCNE3 modulate KCNQ1 channels by affecting different gating transitions. Proc Natl Acad Sci U S A. 114:E7367-E7376 (2017).

144. Sun J, MacKinnon R. Cryo-EM structure of a KCNQ1/CaM complex reveals insights into congenital long QT syndrome. Cell. 2017;169:1042-1054.

145. Bartos DC, Giudicessi JR, Tester DJ, et al. A KCNQ1 mutation contributes to the concealed type 1 long QT phenotype by limiting the Kv7.1 channel conformational changes associated with protein kinase A phosphorylation. Heart Rhythm. 2014;11:459-468.

146. Marx SO, Kurokawa J, Reiken S, et al. Requirement of a macromolecular signaling complex for beta adrenergic receptor modulation of the KCNQ1-KCNE1 potassium channel. Science. 2002;295:496-499.

147. Hedley PL, Jorgensen P, Schlamowitz S, et al. The genetic basis of long QT and short QT syndromes: a mutation update. Hum Mutat. 2009;30:1486-1511.

148. Wu J, Ding WG, Horie M. Molecular pathogenesis of long QT syndrome type 1. J Arrhythm. 2016;32:381-388.

149. Moretti A, Bellin M, Welling A, et al. Patient-specific induced pluripotent stem-cell models for long-QT syndrome. New Engl J Med. 2010;363:1397-1409.

150. Moreno C, Oliveras A, Bartolucci C, et al. D242N, a Kv7.1 LQTS mutation uncovers a key residue for IKs voltage dependence. J Mol Cell Cardiol. 2017;110:61-69.

151. Dvir M, Strulovich R, Sachyani D, et al. Long QT mutations at the interface between KCNQ1 helix C and KCNE1 disrupt (IKs) regulation by PKA and PIP2. J Cell Sci. 2014;127:3943-3955.

152. Shamgar L, Ma LJ, Schmitt N, et al. Calmodulin is essential for cardiac I-KS channel gating and assembly—impaired function in long-QT mutations. Circ Res. 2006;98:1055-1063.

153. Burgess DE, Bartos DC, Reloj AR, et al. High-risk long QT syndrome mutations in the Kv7.1 (KCNQ1) pore disrupt the molecular basis for rapid K(+)-permeation. Biochemistry. 2012;51(45):9076-9085.
154. Harmer SC, Tinker A. The role of abnormal trafficking of KCNE1 in long QT syndrome.  *Biochem Soc Trans*. 2007;35:1074-1076.

155. Splawski I, Shen J, Timothy KW, et al. Spectrum of mutations in long-QT syndrome genes. KVLQT1, HERG, SCN5A, KCNE1, and KCNE2.  *Circulation*. 2000;102:1178-1185.

156. Moreno C, Oliveras A, de la Cruz A, et al. A new KCNQ1 mutation at the S5 segment that impairs its association with KCNE1 is responsible for short QT syndrome.  *Cardiovasc Res*. 2015;107:613-623.

157. Wu ZJ, Huang Y, Fu YC, et al. Characterization of a Chinese KCNQ1 mutation (R259H) that shortens repolarization and causes short QT syndrome 2.  *J Geriatr Cardiol*. 2015;12:394-401.

158. Vandenberg JI, Perry MD, Perrin MJ, Mann SA, Ke Y, Hill AP. hERG K+ channels: structure, function, and clinical significance.  *Physiol Rev*. 2012;92:1393-1478.

159. Kodirov SA. Tale of tail current.  *Prog Biophys Mol Biol*. 2020;150:78-97.

160. Smith JL, Anderson CL, Burgess DE, Elayi CS, January CT, Abbott GW, Sesti F, Splawski I, et al. MiRP1 forms IKr potas-

161. Zhao Z, Li X, El-Battrawy I, et al. Modeling short QT syndrome using human-induced pluripotent stem cell-derived cardiomyocytes.  *J Heart Assoc*. 7, e007394 (2018).

162. Dhamoon AS, Jalife J. The inward rectifier current (IK1) controls cardiac excitability and is involved in arrhythmogenesis.  *Heart Rhythm*. 2005;2(3):316-324.

163. Liew HK, Fu JD, Chiamvimonvat N, et al. Mechanism-based facilitated maturation of human pluripotent stem cell-derived cardiomyocytes.  *Circ Arrhythm Electrophysiol*. 2013;6:191-201.

164. Ma D, Taneja TK, Hagen BM, et al. Goli gp export of the Kir2.1 channel is driven by a trafficking signal located within its terti-

165. Dart C, Leyland ML. Targeting of an A kinase-anchoring protein, AKAP79, to an inwardly rectifying potassium channel, Kir2.1.  *J Biol Chem*. 2001;276:20499-20505.

166. Giustetto C, Schimpf R, Mazzanti A, et al. Long-term follow-

167. Perez-Hernandez M, Matamoros M, Alfayate S, et al. Cardiac Kir2.1 and Na(v)1.5 channels traffic together to the sar-

168. Priori SG, Pandit SV, Rivolta I, et al. Novel form of short QT syndrome (SQT3) is caused by a mutation in the KCNJ2 gene.  *Circ Res*. 2005;112:519-528.

169. Ponce-Balbuena D, Guerrero-Serna G, Valdivia CR, et al. Cardiac Kir2.1 and Na(v)1.5 channels traffic together to the sarcolemma to control excitability.  *Circ Res*. 2018;122:1501-1516.

170. Perez-Hernandez M, Matamoros M, Alfayate S, et al. Brugada syndrome trafficking-defective Nav1.5 channels can trap cardiac Kir2.1/2.2 channels.  *JCI Insight*. 3(2018).

171. Plaster NM, Tawil R, Tristani-Firouzi M, et al. Mutations in Kir2.1 cause the developmental and episodic electrical pheno-

172. Nagase S, Sugasawa Y, Yoshihara M, Ohe T. Electrophysiologic characteristics of an Andersen syndrome patient with KCNJ2 mutation.  *Heart Rhythm*. 2007;4:512-515.

173. Pegan S, Arrabit C, Zhou W, et al. Cytoplasmic domain structures of Kir2.1 and Kir3.1 allow gating and rectification.  *Nat Neurosci*. 2005;8:279-287.

174. Ma D, Tang XD, Rogers TB, Welling PA. An Andersen-Tawil syndrome mutation in Kir2.1 (V302M) alters the G-loop cyto-

175. Lees-Miller JP, Guo J, Wang Y, Perissiniotti LL, Noskov SY, Duff HJ. Ivasradine prolongs phase 3 of cardiac repolarization and blocks the hERG (KCNH2) current over a concentration-range overlapping with that required to block HCN4.  *J Mol Cell Cardiol*. 2015;85:71-78.

176. Zhao Z, Li X, El-Battrawy I, et al. Drug testing in human-induced pluripotent stem cell-derived cardiomyocytes from a patient with short QT syndrome type 1.  *Clin Pharmacol Ther*. 2019;106:642-651.

177. El-Battrawy I, Lan H, Cyganek L, et al. Modeling short QT syn-

178. Jost N, Virag L, Bitay M, et al. Restricting excessive cardiac action potential and QT prolongation: a vital role for IKs in human ventricular muscle.  *Circulation*. 2005;112:1392-1399.
191. Roden DM. Long QT syndrome: reduced repolarization reserve and the genetic link. *J Intern Med*. 2006;259:59-69.

192. So PP, Backx PH, Dorian P. Slow delayed rectifier K+ current block by HMR 1556 increases dispersion of repolarization and promotes Torsades de Pointes in rabbit ventricles. *Br J Pharmacol*. 2008;155:1185-1194.

193. Kuryshhev YA, Ficker E, Wang L, et al. Pentamidine-induced long QT syndrome and block of hERG trafficking. *J Pharmacol Exp Ther*. 2005;312:316-323.

194. Rajamani S, Eckhardt LL, Valdivia CR, et al. Drug-induced long QT syndrome: hERG K+ channel block and disruption of protein trafficking by fluoxetine and norfluoxetine. *Br J Pharmacol*. 2006;149:481-489.

195. Khonphiathanayothin A, Lane J J, Thomas D, Yen L, Chang D, Bubolz B. Effects of cisapride on QT interval in children. *J Pediatr*. 1998;133:51-56.

196. Mohammad S, Zhou Z, Gong Q, January CT. Blockage of the HERG human cardiac K+ channel by the gastrointestinal prokinetic agent cisapride. *Am J Physiol*. 273:H2534-2538 (1997).

197. Michaud V, Dow P, Al Rihani SB, et al. Risk assessment of drug-induced long QT syndrome for some COVID-19 repurposed drugs. *Clin Transl Sci*. 2021;14:20-28.

198. Food & Drug Administration, H.H.S. International Conference on Harmonisation; guidance on E14 Clinical Evaluation of QT/QTc Interval Prolongation and Proarrhythmic Potential for Non-Antiarrhythmic Drugs; availability. Notice. *Fed Regist*. 2005;70:61134-61135.

199. Food & Drug Administration, H.H.S. International Conference on Harmonisation; guidance on S7B Nonclinical Evaluation of the Potential for Delayed Ventricular Repolarization (QT Interval Prolongation) by Human Pharmaceuticals; availability. Notice. *Fed Regist*. 2005;70:61133-61134.

200. Ridder BJ, Leishman DJ, Bridgland-Taylor M, et al. A systematic strategy for estimating hERG block potency and its implications in a new cardiac safety paradigm. *Toxicol Appl Pharmacol*. 2020;394:114961.

201. Lei M, Wu L, Terrar DA, Huang CL. Modernized classification of cardiac antiarrhythmic drugs. *Circulation*. 2018;138(17):1879-1896.

202. Redfern WS, Carlson L, Davis AS, et al. Relationships between preclinical cardiac electrophysiology, clinical QT interval prolongation and torsade de pointes for a broad range of drugs: evidence for a provisional safety margin in drug development. *Cardiovasc Res*. 2003;58:32-45.

203. Aiba T, Shimizu W, Inagaki M, et al. Cellular and ionic mechanism for drug-induced long QT syndrome and effectiveness of verapamil. *J Am Coll Cardiol*. 2005;45:300-307.

204. Antzelevitch C, Belardinelli L, Zygmunt AC, et al. Electrophysiological effects of ranolazine, a novel antiarrhythmic agent with antiarrhythmic properties. *Circulation*. 2004;110:904-910.

205. Kramer J, Obejero-Paz CA, Myatt G, et al. MICE models: superior to the HERG model in predicting Torsade de Pointes. *Sci Rep*. 2013;3:2100.

206. Mirams GR, Cui Y, Sher A, et al. Simulation of multiple ion channel block provides improved early prediction of compounds’ clinical torsadogenic risk. *Cardiovasc Res*. 2011;91:53-61.

207. Colatsky T, Ferrini B, Gintant G, et al. The comprehensive in vitro proarrhythmia assay (CiPA) initiative—update on progress. *J Pharmacol Toxicol Methods*. 2016;81:15-20.

208. Crumb WJ, Jr., Vicente J, Johannessen L, Strauss DG. An evaluation of 30 clinical drugs against the comprehensive in vitro proarrhythmia assay (CiPA) proposed ion channel panel. *J Pharmacol Toxicol Methods*. 2016;81:251-262.

209. Okada J, Yoshinaga T, Kurokawa J, et al. Screening system for drug-induced arrhythmogenic risk combining a patch clamp and heart simulator. *Sci Adv*. 2015;1:e1400142.

210. Kuryshhev YA, Brown AM, Duzic E, Kirsch GE. Evaluating state dependence and subtype selectivity of calcium channel modulators in automated electrophysiology assays. *Assay Drug Dev Technol*. 2014;12:110-119.

211. Sanson C, Schombert B, Filoche-Romme B, Partisetti M, Bohme GA. Electrophysiological and pharmacological characterization of human inwardly rectifying Kir2.1 channels on an automated patch-clamp platform. *Assay Drug Dev Technol*. 2019;17:89-99.

212. Elkins RC, Davies MR, Brough SJ, et al. Variability in high-throughput ion-channel screening data and consequences for cardiac safety assessment. *J Pharmacol Toxicol Methods*. 2013;68:112-122.

213. Romero L, Cano J, Gomis-Tena J, et al. In silico QT and APD prolongation assay for early screening of drug-induced proarrhythmic risk. *J Chem Inf Model*. 2018;58:867-878.

214. Li Z, Mirams GR, Yoshinaga T, et al. General principles for the validation of proarrhythmia risk prediction models: an extension of the CiPA in silico strategy. *Clin Pharmacol Ther*. 2020;107:102-111.

215. Rogers AJ, Selvalingam A, Alhusseini MI, et al. Machine learned cellular phenotypes in cardiomyopathy predict sudden death. *Circ Res*. 2021;128:172-184.

216. Yang PC, DeMarco KR, Aghasafari P, et al. A computational pipeline to predict cardioxicity: from the atom to the rhythm. *Circ Res*. 2020;126:947-964.

217. Mann SA, Heide J, Knott T, et al. Recording of multiple ion current components and action potentials in human induced pluripotent stem cell-derived cardiomyocytes via automated patch-clamp. *J Pharmacol Toxicol Methods*. 2019;100:106599.

218. Gintant G, Sager PT, Stockbridge N. Evolution of strategies to improve preclinical cardiac safety testing. *Nat Rev Drug Discov*. 2016;15:457-471.

219. Lopez-Izquierdo A, Warren M, Riedel M, et al. A near-infrared fluorescent voltage-sensitive dye allows for moderate-throughput electrophysiological analyses of human induced pluripotent stem cell-derived cardiomyocytes. *Am J Physiol Heart Circ Physiol*. 307:H1370-1377 (2014).

220. Vaidyanathan R, Markandeya YS, Kamp TJ, Makiecki JC, January CT, Eckhardt LL. IK1-enhanced human-induced pluripotent stem cell-derived cardiomyocytes: an improved cardiomyocyte model to investigate inherited arrhythmia syndromes. *Am J Physiol Heart Circ Physiol*. 2016;310:H1611-1621.

221. Goverseen B, Becker N, Stoelzl-Feix S, et al. A hybrid model for safety pharmacology on an automated patch clamp platform: using dynamic clamp to join iPSC-derived cardiomyocytes and simulations of I-K1 ion channels in real-time. *Front Physiol*. 8, 1094 (2018).
222. Yang X, Papoian T. Moving beyond the comprehensive in vitro proarrhythmia assay: use of human-induced pluripotent stem cell-derived cardiomyocytes to assess contractile effects associated with drug-induced structural cardiotoxicity. *J Appl Toxicol*. 2018;38:1166-1176.

223. Takasuna K, Asakura K, Araki S, et al. Comprehensive in vitro cardiac safety assessment using human stem cell technology: Overview of CSAHI HEART initiative. *J Pharmacol Toxicol Methods*. 2017;83:106595.42-54.

224. Itzhaki I, Maizels L, Huber I, et al. Modelling the long QT syndrome with induced pluripotent stem cells. *Nature*. 2011;471:225-U113.

225. Sala L, Yu ZY, Ward-van Oostwaard D, et al. A new hERG allosteric modulator rescues genetic and drug-induced long-QT syndrome phenotypes in cardiomyocytes from isogenic pairs of patient induced pluripotent stem cells. *Embo Mol Med*. 2016;8:1065-1081.

226. Leifert A, Pan Y, Kinkeldey A, et al. Differential hERG ion channel function of IKs channels with diverse long QT mutations. *Elife*. 2016;5:e20272.

227. Liin SI, Silvera Ejneby M, Barro-Soria R, et al. Polyunsaturated fatty acid analogues differentially affect cardiac NaV, CaV, and KV channels through unique mechanisms. *Elife*. 2020;9:e51453.

228. Bohannon BM, de la Cruz A, Wu X, et al. Polyunsaturated fatty acid analogues differentially affect cardiac NaV, CaV, and KV channels through unique mechanisms. *Elife*. 2020;9:e51453.

229. Wu L, Yong SL, Fan C, et al. Identification of a new co-factor, MOG1, required for the full function of cardiac sodium channel Nav1.5. *J Biol Chem*. 2008;283:6968-6978.

230. Petitprez S, Zmoos AF, Ogrodnik J, et al. SAP97 and dystrophin macromolecular complexes determine two pools of cardiac sodium channels Nav1.5 in cardiomyocytes. *Circ Res*. 2011;108:294-304.

231. Matamoros M, Perez-Hernandez M, Guerrero-Serna G, et al. Nav1.5 N-terminal domain binding to alpha-syntrophin increases membrane density of human Kir2.1, Kir2.2 and Nav1.5 channels. *Cardiovasc Res*. 2016;110:279-290.

232. Tan HL, Kupershmidt S, Zhang R, et al. A calcium sensor in the sodium channel modulates cardiac excitability. *Nature*. 2002;415:442-447.

233. Hund TJ, Koval OM, Li JD, et al. A beta(IV)-spectrin/CaMKII signaling complex is essential for membrane excitability in mice. *J Clin Invest*. 2010;120:3508-3519.

234. Glynn P, Musa H, Wu XQ, et al. Voltage-gated sodium channel phosphorylation at Ser571 regulates late current, arrhythmia, and cardiac function in vivo. *Circulation*. 2015;132:567-577.

235. Wagner N, Dybkova N, Rasenack ECL, et al. Ca2+/calmodulin-independent protein kinase II regulates cardiac Na+ channels. *J Clin Invest*. 2006;116:327-3138.

236. Jespersen T, Gavillet B, van Bemmelen MX, et al. Cardiac sodium channel Na(v)1.5 interacts with and is regulated by the protein tyrosine phosphatase PTPH1. *Biochem Bioph Res Co*. 2006;348:1455-1462.

237. Vatta M, Ackerman MJ, Ye B, et al. Mutant caveolin-3 induces persistent late sodium current and is associated with long-QT syndrome. *Circulation*. 2006;114:2104-2112.

238. Zhao Y, Huang G, Wu Q, et al. Cryo-EM structures of apo and antagonist-bound human Cav3.1. *Nature*. 2019;576:492-497.

239. Bissay V, van Malderen SCH, Keymolen K, et al. SCN4A variants and Brugada syndrome: phenotypic and genotypic overlap between cardiac and skeletal muscle sodium channelopathies. *Eur J Hum Genet*. 2016;24:400-407.

240. Mannikko R, Wong L, Tester DJ, et al. Dysfunction of Nav1.4, a skeletal muscle voltage-gated sodium channel, in susceptible infant death syndrome: a case-control study. *Lancet*. 2018;391:1483-1492.

241. Wilde AA, Brugada R. Phenotypical manifestations of mutations in the genes encoding subunits of the cardiac sodium channel. *Circ Res*. 2011;108:884-897.

242. Mohler PI, Rivolta I, Napolitano C, et al. Nav1.5 EI053K mutation causing Brugada syndrome blocks binding to ankyrin-G and expression of Nav1.5 on the surface of cardiomyocytes. *Proc Natl Acad Sci U S A*. 2004;101:17533-17538.

243. Ziane R, Huang H, Moghadasazadeh B, Beggs AH, Levesque G, Chahine M. Cell membrane expression of cardiac sodium channel Na(v)1.5 is modulated by alpha-actinin-2 interaction. *Biochemistry*. 2010;49:166-178.

244. Wu L, Yong SL, Fan C, et al. Identification of a new co-factor, MOG1, required for the full function of cardiac sodium channel Nav1.5. *J Biol Chem*. 2008;283:6968-6978.

245. Petitprez S, Zmoos AF, Ogrodnik J, et al. SAP97 and dystrophin macromolecular complexes determine two pools of cardiac sodium channels Nav1.5 in cardiomyocytes. *Circ Res*. 2011;108:294-304.

246. Matamoros M, Perez-Hernandez M, Guerrero-Serna G, et al. Nav1.5 N-terminal domain binding to alpha-syntrophin increases membrane density of human Kir2.1, Kir2.2 and Nav1.5 channels. *Cardiovasc Res*. 2016;110:279-290.

247. Tan HL, Kupershmidt S, Zhang R, et al. A calcium sensor in the sodium channel modulates cardiac excitability. *Nature*. 2002;415:442-447.

248. Hund TJ, Koval OM, Li JD, et al. A beta(IV)-spectrin/CaMKII signaling complex is essential for membrane excitability in mice. *J Clin Invest*. 2010;120:3508-3519.

249. Glynn P, Musa H, Wu XQ, et al. Voltage-gated sodium channel phosphorylation at Ser571 regulates late current, arrhythmia, and cardiac function in vivo. *Circulation*. 2015;132:567-577.

250. Wagner N, Dybkova N, Rasenack ECL, et al. Ca2+/calmodulin-independent protein kinase II regulates cardiac Na+ channels. *J Clin Invest*. 2006;116:327-3138.

251. Jespersen T, Gavillet B, van Bemmelen MX, et al. Cardiac sodium channel Na(v)1.5 interacts with and is regulated by the protein tyrosine phosphatase PTPH1. *Biochem Bioph Res Co*. 2006;348:1455-1462.

252. Vatta M, Ackerman MJ, Ye B, et al. Mutant caveolin-3 induces persistent late sodium current and is associated with long-QT syndrome. *Circulation*. 2006;114:2104-2112.

253. Zhao Y, Huang G, Wu Q, et al. Cryo-EM structures of apo and antagonist-bound human Cav3.1. *Nature*. 2019;576:492-497.

254. Baig SM, Koschak A, Lieb A, et al. Loss of Ca(v)1.3 (CACNA1D) function in a human channelopathy with bradycardia and congenital deafness. *Nat Neurosci*. 2011;14:77-84.

255. Hong TT, Smyth JW, Chu KY, et al. BIN1 is reduced and Cav1.2 trafficking is impaired in human failing cardiomyocytes. *Heart Rhythm*. 2012;9:812-820.

256. Hong TT, Smyth JW, Gao DC, et al. BIN1 localizes the L-type calcium channel to cardiac T-tubules. *Plos Biol*. 2010;8.
257. De La Mata A, Tajada S, O’Dwyer S, et al. BIN1 induces the formation of T-tubules and adult-like Ca(2+)/release units in developing cardiomyocytes.  *Stem Cells*. 2019;37:54-64.

258. Thomsen MB, Wang C, Ozgen N, Wang HG, Rosen MR, Pitt GS. Accessory subunit KChIP2 modulates the cardiac L-type calcium current.  *Circ Res*. 2009;104:1382-1389.

259. Dixon RE, Moreno CM, Yuan C, et al. Graded current encoded by Kv4.3 as a putative beta-subunit of human cardiact transient outward sion and function of dipeptidyl-aminopeptidase-like protein 6.  *J Physiol*. 2006;544-5449.

260. Xu H, Ginsburg KS, Hall DD, et al. Targeting of protein phosphatases PP2A and PP2B to the C-terminus of the L-type calcium channel Ca v1.2.  *Biochemistry*. 2010;49:10298-10307.

261. Hall DD, Feekes JA, Arachchige Don AS, et al. Binding of protein phosphatase 2A to the L-type calcium channel Cav1.2 next to Ser1928, its main PKA site, is critical for Ser1928 dephosphorylation.  *Biochemistry*. 2006;45:3448-3459.

262. Wehrens XH, Lehnart SE, Huang F, et al. FKBP12.6 deficiency form a tripartite complex with the anchoring protein Yotiao modulates the cardiac L-type calcium channel Cav1.2.  *J Biol Chem*. 2011;286:31347-31352.

263. Lokuta AJ, Meyers MB, Sander PR, Fishman GI, Valdivia HH. Binding of protein phosphatase receptors by sorcin.  *J Biol Chem*. 2003;278:829-840.

264. Lokuta AJ, Meyers MB, Sander PR, Fishman GI, Valdivia HH. Modulation of cardiac ryanodine receptors by sorcin.  *J Biol Chem*. 1997;272:25333-25338.

265. Balijepalli SY, Lim E, Concannon SP, et al. Mechanism of loss of KV1.1 K+ current in mutant T421M-Kv11.1-expressing rat ventricular myocytes: interaction of trafficking and gating.  *Circulation*. 2012;126:2809-2818.

266. Zhou Z, Gong Q, Epstein ML, January CT. HERG channel dysfunction in human long QT syndrome. Intracellular transport and functional defects.  *J Biol Chem*. 1998;273:21061-21066.

267. Balijepalli SY, Lim E, Concannon SP, et al. Mechanism of loss of Kv11.1 K+ current in mutant T421M-Kv11.1-expressing rat ventricular myocytes: interaction of trafficking and gating.  *Circulation*. 2012;126:2809-2818.

268. Es-Salah-Lamoureux Z, Xiong PY, Goodchild SJ, Ahern CA, Fedida D. Blockade of permeation by potassium but normal gating of the G628S nonconducting hERG channel mutant.  *Biophys J*. 2011;101:662-670.

269. Radicke S, Cotella D, Graf EM, Ravens U, Wettwer E. Expression and function of dipeptidyl-aminopeptidase-like protein 6 as a putative beta-subunit of human cardiac transient outward current encoded by Kv4.3.  *J Physiol*. 2005;565:751-756.

270. El-Haou S, Balse E, Neyroud N, et al. Kv4 potassium channels form a tripartite complex with the anchoring protein SAP97 and CaMKII in cardiac myocytes.  *Circ Res*. 2009;104:758-769.

271. Xiao L, Coutu P, Villeneuve LR, et al. Mechanisms underlying rate-dependent remodeling of transient outward potassium current in canine ventricular myocytes.  *Circ Res*. 2008;103:733-742.

272. Zaydman MA, Silva JR, Delaloye K, et al. Kv7.1 ion channels require a lipid to couple voltage sensing to pore opening.  *P Natl Acad Sci USA*. 2013;110:13180-13185.

273. Zaydman MA, Cui J. PI2P regulation of KCNQ channels: biochemical and molecular mechanisms for lipid modulation of voltage-dependent gating.  *Front Physiol*. 2014;5:195.

274. Tobelaim WS, Dvir M, Lebel G, et al. Competition of calcified calmodulin N lobe and PI2P to an LQT mutation site in Kv7.1 channel.  *Proc Natl Acad Sci U S A*. 2017;114:E869-E878.

275. Chen L, Kurokawa J, Kass RS. Phosphorylation of the A-kinase-anchoring protein Yotiao contributes to protein kinase A regulation of a heart potassium channel.  *J Biol Chem*. 2005;280:31347-31352.

276. Kurokawa J, Motoki HK, Rao J, Kass RS. Regulatory actions of the A-kinase anchoring protein Yotiao on a heart potassium channel downstream of PKA phosphorylation.  *Proc Natl Acad Sci U S A*. 2004;101:16374-16378.

277. Heijman J, Spatjens RL, Seyen SR, et al. Dominant-negative control of cAMP-dependent IKs upregulation in human long-QT syndrome type 1.  *Circ Res*. 2012;110:211-219.

278. Ghosh S, Nunziato DA, Pitt GS. KCNQ1 assembly and function is blocked by long-QT syndrome mutations that disrupt interaction with calmodulin.  *Circ Res*. 2006;98:1048-1054.

279. Sachyani D, Dvir M, Strulovich R, et al. Structural basis of a Kv7.1 potassium channel gating module: studies of the intra-cellular c-terminal domain in complex with calmodulin.  *Structure*. 2014;22:1582-1594.

280. Howard RJ, Clark KA, Holton JM, Minor DL. Structural insight into KCNQ (Kv7) channel assembly and channelopathy.  *Neuron*. 2007;53:663-675.

281. Kagan A, Melman YF, Krumerman A, McDonald TV. 14-3-3 amplifies and prolongs adrenergic stimulation of HERG K+ channel activity.  *EMBO J*. 2002;21(8):1899-1898.

282. Vaidyanathan R, Taffet SM, Vikstrom KL, Amunowo JO. Regulation of cardiac inward rectifier potassium current (I(K1)) by synapse-associated protein-97.  *J Biol Chem*. 2010;285:28000-28009.

283. Vaidyanathan R, Reilly L, Eckhardt LL. Caveolin-3 microdomain: arrhythmia implications for potassium inward rectifier and cardiac sodium channel.  *Front Physiol*. 2018;9:1548.

284. Lopes CM, Zhang H, Rohacs T, et al. Alterations in conserved Kir channel-Pi2P interactions underlie channelopathies.  *Neuron*. 2002;34:933-944.

285. Potet F, Lorinc AN, Chaigne S, et al. Identification and characterization of a compound that protects cardiac tissue from human ether-a-go-go-related gene (hERG)-related drug-induced arrhythmias.  *J Biol Chem*. 2012;287:39613-39625.

286. Mannikko R, Bridgland-Taylor MH, Pye H, et al. Pharmacological and electrophysiological characterization of AZSMO-23, an activator of the hERG K(+) channel.  *Br J Pharmacol*. 2015;172:3112-3125.

287. Qile M, Ji Y, Golden TD, et al. LUF7244 plus dofetilide rescues aberrant Kv11.1 trafficking and produces functional IKv11.1.  *Mol Pharmacol*. 2020;97:355-364.

288. Donovan BT, Bandyopadhyay D, Duraiswami C, Nixon CJ, Townsend CY, Martens SF. Discovery and electrophysiological characterization of SKF-32802: a novel hERG agonist found through a large-scale structural similarity search.  *Eur J Pharmacol*. 2018;818:306-327.

289. Gillie DJ, Novick SJ, Donovan BT, Payne LA, Townsend C. Development of a high-throughput electrophysiological assay for the human ether-a-go-go related potassium channel hERG.  *J Pharmacol Toxicol Methods*. 2013;67:33-44.
290. Mazzanti A, Maragna R, Vacanti G, et al. Hydroquinidine prevents life-threatening arrhythmic events in patients with short QT syndrome. *J Am Coll Cardiol*. 2017;70:3010-3015.

291. Du C, Zhang Y, El Harchi A, Dempsey CE, Hancox JC. Ranolazine inhibition of hERG potassium channels: drug-pore interactions and reduced potency against inactivation mutants. *J Mol Cell Cardiol*. 2014;74:220-230.

292. Portero V, Casini S, Hoekstra M, et al. Anti-arrhythmic potential of the late sodium current inhibitor GS-458967 in murine Scn5a-1798insD+/− and human SCN5A-1795insD+/− iPSC-derived cardiomyocytes. *Cardiovasc Res*. 2017;113:829-838.

293. Haechl N, Ebner J, Hilber K, Todt H, Koenig X. Pharmacological profile of the bradycardic agent ivabradine on human cardiac ion channels. *Cell Physiol Biochem*. 2019;53:36-48.

294. Orvos P, Kohajda Z, Szlovak J, et al. Evaluation of possible proarrhythmic potency: comparison of the effect of dofetilide, cisapride, sotalol, terfenadine, and verapamil on hERG and native IKr currents and on cardiac action potential. *Toxicol Sci*. 2019;168:365-380.

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