Defective cardiac ion channels: from mutations to clinical syndromes

Colleen E. Clancy and Robert S. Kass

Department of Pharmacology, Columbia University, New York, New York, USA


Normal cardiac excitation and relaxation involves a delicate balance of complex dynamic interactions between ionic currents passing through a variety of membrane channels and the cellular environment. Genetic defects, polymorphisms, therapeutic intervention or structural abnormalities can disrupt this balance and underlie severe arrhythmogenic phenotypes that lead to sudden cardiac death. Inheritable gene defects give rise to phenotypic variation and an unpredictable manifestation of syndromes, ranging from silent gene carriers to profoundly symptomatic individuals, even within single families (1–7). As such, realizing the relationship between genetic mutations and clinical syndromes is becoming increasingly complex. In this issue of the JCI, Grant and colleagues (3) investigate the manifestations of phenotypically opposite and overlapping cardiac arrhythmogenic syndromes that surprisingly stem from the same mutation (1–4).

Cardiac excitation reflects membrane depolarization of cardiac myocytes, primarily due to the activation of fast voltage-dependent Na⁺ channels that underlie the action potential upstroke. Activation is followed by a long depolarized plateau phase that permits Ca²⁺-induced Ca²⁺ release from the sarcoplasmic reticulum, binding of Ca²⁺ to contractile proteins on the sarcomeres, and coordinated contraction. Repolarization follows due to the time- and voltage-dependent activation of repolarizing potassium currents. Relaxation of contraction is coupled to the electrical repolarization phase, which allows filling of the ventricles prior to the next excitation. Each of these electrical processes can be detected on the body surface electrocardiogram (ECG) as a signal average of the temporal and spatial gradients generated during each phase (8–11) (Figure 1a). Electrical excitation gradients in the atria (atrial depolarization) manifest on the ECG as P waves, while gradients of ventricular depolarization are seen as the QRS complex. Gradients in ventricular repolarization are reflected in the T wave (Figure 1).

A recently described example of a multi-syndrome genetic defect in the SCN5A gene, encoding the cardiac Na⁺ channel (Figure 2), is the insertion of an aspartic acid, 1795insD, in the C-terminus of the cardiac Na⁺ channel that underlies both Brugada (BrS) and Long-QT (LQTs) cardiac arrhythmic syndromes (1, 2).

Grant and colleagues investigate an even more complex mutation (3). The deletion of lysine, ΔK1500, in the III–IV linker of SCN5A (Figure 2) is associated with BrS, LQTs, and isolated cardiac conduction disease (ICCD). LQTs are typically associated with a gain of Na⁺ channel function that stems from mutation induced destabilization of channel inactivation, leading to a persistent inward Na⁺ current (I_{Na}) during the action potential (AP) plateau and prolonged repolarization (12, 13). Paradoxically, BrS and ICCD are linked to a loss of Na⁺ channel function and a resulting reduction in macroscopic current (3, 6, 7, 14, 15). How can multiple and seemingly contradictory arrhythmnic syndromes arise from a mutation at a single locus?

The heterogeneous myocardial substrate

The solution seems to lie in the intrinsic heterogeneity of the underlying myocardial substrate with which...
mutant Na⁺ channels interact. The ventricular myocardium is comprised of at least three distinct cell types referred to as epicardial, midmyocardial (M), and endocardial cells, which exhibit distinct electrophysiological properties (8).

Epicardial cells display a characteristic spike and dome morphology (Figure 3, left panel, thin line) due to large transient outward K⁺ current (I_{to}), and short action potential duration (APD) resulting from a high density of the slowly activating component of the delayed rectifier K⁺ current (I_{Ks}). Mutations that act to reduce I_{Na} in the presence of these large repolarizing currents (I_{to} and I_{Ks}), result in premature plateau repolarization and APs with distinctive triangular morphology (Figure 3, left panel, thick line) (8–11, 16). Selective loss of the AP plateau in these cells, results in dispersion of plateau potentials across the ventricular wall. This gradient (∇V) generates ST segment elevation on the ECG, which is a diagnostic indicator of BrS (9, 14). Clinically, ST segment elevation is observed in right precordial leads of BrS patients, consistent with the large I_{to} density in right ventricular epicardium (9, 11). In M cells I_{to} and I_{Ks} are smaller, and so unable to overwhelm the mutation induced reduction in I_{Ks} (8, 9, 16). The non-inactivating component of I_{Na}, in the presence of smaller repolarizing currents (I_{to} and I_{Ks}), result in premature plateau repolarization and APs with distinctive triangular morphology (Figure 3, left panel, thick line) (8–11, 16). Selective loss of the AP plateau in these cells, results in dispersion of plateau potentials across the ventricular wall. This gradient (∇V) generates ST segment elevation on the ECG, indicative of LQTs.

Brugada syndrome vs. isolated cardiac conduction disease

One of the more compelling aspects in attempting to elucidate the relationship between genotype and the clinical presentation of disease exists in the differentiation of BrS and ICCD (3, 5, 6, 7). Experimental evidence suggests that a reduction of I_{Na} is a common denominator of both syndromes (3, 5–7, 14, 15). Indeed, a number of mechanisms, including reduced rates of recovery from inactivation, faster inactivation subsequent to channel opening, and protein trafficking defects have been shown to underlie the loss of Na⁺ channel function in BrS. This explanation however, fails to reconcile the presence of ST segment elevation (observed in BrS) in the absence of slowed conduction (as seen in ICCD) and vice-versa. A possible explanation for the phenotypic overlap (3) or exclusivity (5) may lie in the notable commonality among the relatively few characterized ICCD mutations; namely, a consistent rightward shift in the voltage dependence of the mutant Na⁺ channel activation curve (5–7) (see Figure 3b in Grant et al.)

Figure 2

The predicted transmembrane topology of domains I–IV of the cardiac Na⁺ channel α subunit encoded by SCN5A showing the location and nature of the mutations inducing LQTs, BrS, and isolated cardiac conduction disease.

Figure 3

Cellular electrical abnormalities and their relation to changes in the ECG. (Left) At fast pacing (heart) rates, mutation-induced changes in epicardial AP morphologies (thick line) cause dispersion of plateau potentials and a voltage gradient (∇V_m, arrow) (WT, thin line). This gradient will manifest on the ECG as ST segment elevation, indicative of BrS. (Middle) Mutations may prolong APD in M cells (thick line) compared to WT (thin line). The delay in repolarization (∆APD = 60ms) is reflected as QT prolongation on the ECG, a hallmark of LQTs. (Right) A rightward shift in the Na⁺ channel activation curve is sufficient to reduce AP upstroke and slow cardiac impulse conduction (wide QRS segment) as observed in ICCD. WT, wild-type.
in this issue). A depolarizing shift of the Na⁺ channel activation curve indicates that mutant channels require a more positive membrane potential to fully activate than wild-type (WT) channels (5). Most likely, this shift results from a reduction in the rate of channel activation or decreased channel sensitivity to the voltage required for activation. In either case, a rightward shift of the Na⁺ channel activation curve may not necessarily lead to decreased current, but rather a different voltage at which the maximum current amplitude occurs. For example, if an IC mutation results in mutant channels that obtain maximum current amplitude at 0 mV, as opposed to –10 mV for WT channels, mutant channels fully activate within the physiological range of the action potential upstroke. However, the mutant channels may require more time to reach the membrane potential at which the maximum current amplitude occurs. The lag in activation of ICCD mutant Na⁺ channels would result in a reduction of the action potential upstroke velocity (Figure 3, right panel; inset: magnification of AP upstroke), a primary determinant of conduction velocity (17, 18). A reduction in the velocity of electrical propagation will manifest on the ECG in a widening of the QRS interval. The defect characterized by Grant and coauthors in this issue of the JCI is unique in that a single amino acid deletion in the cardiac Na⁺ channel III–IV linker gives rise to three distinct cellular phenotypes as well as three separate clinical syndromes (3). The ΔK1500 mutation leads to a significantly increased component of non-inactivating Na⁺ current, which results in prolongation of repolarization and the QT interval on the ECG. The mutation also results in faster inactivation, a property that is consistent with reduced Na⁺ current. In some epicardial cells, where the repolarizing potassium currents are large, the reduced Na⁺ current may be overwhelmed and AP plateau heterogeneously suppressed, resulting in ST segment elevation, a hallmark of BrS. Finally, the ΔK1500 mutation results in a rightward shift of the channel activation curve, which may be responsible for the reduced conduction velocity seen clinically as the broadening of the QRS interval on the ECG of ICCD patients. While the cellular characterization of the ΔK1500 mutation is consistent with the development of LQTs, BrS, and ICCD simultaneously, the clinical observations are less clear-cut. The four generations of mutation carriers evaluated by Grant et al. display a wide variety of clinical presentations and symptoms (3). The explanation for the variability in phenotypic manifestation of AK1500 may well lie in the underlying genetic substrate via mutant channel interactions with modifier genes and environmental factors. **Modifier genes: A role in the reconciliation of multiple phenotypes from a single genotype?** The broad diversity in phenotypic expression of gene defects among affected individuals with identical cardiac ion channel mutations is likely influenced by other genetic traits separate from the primary disease locus. Genotype-phenotype correlation studies show significant variability in phenotypes among affected individuals with identical causal mutations (6, 7, 19, 20). Clinical presentation is determined by complex interactions between causal genes, genetic background (modifier genes), and environmental factors. While individual modifier genes for Na⁺ channel-linked arrhythmias remain largely unknown, potential modifiers of genetically based arrhythmias include, but are not limited to, gender, febrile states, adrenergic stimulation, signaling molecules, channel-associated protein kinases (20–23), channel-associated protein phosphatases (20–22), and individual electrophysiological and morphological substrates. Thus far, studies are limited to polymorphism association studies and electrophysiological characterization in expression systems, where channels are removed from the physiological interactive environment. Identification of modifier genes will complement the current studies that have identified and characterized causative genes, which may improve upon genetically based diagnosis, risk stratification, and implementation of preventive and therapeutic interventions in patients with Na⁺ channel-linked arrhythmias.