Mechanistic complexity of contractile dysfunction in hypertrophic cardiomyopathy

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Hypertrophic cardiomyopathy (HCM) is a relatively common disease in which the muscular wall of the heart becomes thickened. It is estimated to affect 1 in every 500 people—men and women equally—and can cause cardiac arrest in young people, including athletes. In 1990, a genetic linkage between a mutated protein found within cardiac muscle sarcomeres and the HCM phenotype was described (Geisterfer-Lowrance et al., 1990). Since then, the familial form of hypertrophic cardiomyopathy (fHCM) has been ascribed to mutations found in at least ten different sarcomere proteins within cardiac muscle (Seidman and Seidman, 2011; Haas et al., 2015). Although ~80% of fHCM mutations identified as causal are found in thick filament proteins (myosin heavy chain, myosin regulatory light chain, and myosin binding protein C), a significant number have also been identified in thin filament proteins that regulate cardiac muscle contraction (tropomycin and tropomyosin; Tardiff, 2011). Of the mutations identified thus far in the cardiac troponin complex, ~60% occur in the troponin T (cTnT) subunit, with the remainder primarily in the troponin I (cTnI) subunit and a few putative mutations in tropomin C (cTnC; Cheng et al., 2015). In a recent issue of the Journal of General Physiology, Reda and Chandra investigated the effect of a recently identified fHCM mutation in cTnT on the contractile function of cardiac muscle fibers.

Biophysical characterization can confirm when an identified fHCM mutation may be pathogenic and provide mechanistic insight into the disease by determining how it alters function at scales that range from individual cardiac proteins to the contractile apparatus, whole cells, and tissues. Characterization of contractile dysfunction for disease-related mutations has been performed in animal, tissue, and cell models. Until recently, most emphasis has been on the later, or end stages, of heart failure. Perhaps the most common measurement in characterizing how the “Ca$^{2+}$ sensitivity” of the contractile apparatus is usually defined as the pCa$_{50}$ (negative log concentration of Ca$^{2+}$ level that produces half-maximal force) of this force–pCa relationship. For many, but certainly not all fHCM mutations studied, there is an increase in the Ca$^{2+}$ sensitivity of force development (Harris et al., 2011; Tardiff, 2011; Cheng et al., 2015, 2016) These measurements, although relatively easy to make, provide a very limited picture of contractile performance and do not present a clear picture of how ventricular function is compromised. Several studies have also examined the effect of mutations on the kinetics of contractile activation and relaxation, as well as other indices of myosin–actin cross-bridge cycling and cellular contractile behavior. For several fHCM-associated mutations, there are reports that relaxation kinetics are altered and that the effects of PKA-mediated phosphorylation of myofilament proteins (β-adrenergic mechanisms) are reduced or blunted (Belus et al., 2008; Witjas-Paalberends et al., 2014a; Cheng et al., 2015, 2016).

An important feature of cardiac function is the Frank-Starling mechanism, which allows for changes in venous return to be matched by changes in stroke volume. This mechanism is often disrupted in cardiomyopathies (Schwinger et al., 1994; Vahl et al., 1998). The cellular basis for the Frank-Starling mechanism results from the length dependence of cardiac myofilament activation, which is often studied by measuring the dependence of the Ca$^{2+}$ sensitivity of force development on sarcomere length (SL). In the Reda and Chandra (2018) study, the effect of an fHCM-associated mutation in the central region of cTnT (F88L) on the SL dependence of cardiac myofilament activation was investigated. Reconstituted recombinant troponin, containing cTnT with or without F88L, was exchanged into skinned guinea pig cardiac muscle preparations (papillaries). They then measured the Ca$^{2+}$ dependence of force, ATPase production, and instantaneous force increase with rapid stretch at short (1.9 µm) and long (2.3 µm) SL. An important aspect of the study is that guinea pigs express predominantly the β-myosin heavy chain (β-MHC) isoform in ventricular myocytes (similar to humans) as opposed to...
of increased wall stiffness during diastole that resists ventricular filling at the end of systole, thus increasing a cross-bridge component that merit further thought and investigation. The first is that some mutations may not allow complete deactivation of thin filaments that result in enhanced Ca\(^{2+}\) binding. For example, an engineered mutation in cardiac troponin C (L48Q) also causes a greater increase in the Ca\(^{2+}\) sensitivity of force generation at short SL than at long SL and therefore reduces the SL dependence of contractile activation (Korte et al., 2012). Interpretation of the results from rapid stretch measurements led Reda and Chandra (2018) to conclude that at least part of the greater increase in Ca\(^{2+}\) sensitivity at short SL resulted from a reduced impact of negatively strained cross-bridges on positive force-generating cross-bridges, as well as an enhanced cross-bridge to thin filament-cooperative activation mechanism mediated by the F88L mutation. Further analysis suggested that cross-bridge detachment rate and tension cost were not affected by the mutation at either SL. This would seem to indicate that, for this cTnT mutation, there is no uncoupling between work and energy utilization. Other mutations in cTnT have been reported to increase tension-dependent ATP consumption in transgenic mice, as have thick filament protein mutations in animal models and human fHCM (Chandra et al., 2005; He et al., 2007; Ferrantini et al., 2009; Witjas-Paalberends et al., 2014a,b; Wilder et al., 2015; Birch et al., 2016). The results of the Reda and Chandra (2018) study suggest that uncoupling between contractile function and energetics may not be a universal feature of fHCM, though this needs to be further investigated.

Reda and Chandra (2018) conclude that the F88L cTnT mutation might have a negative impact on the Frank-Starling mechanism. This may be true, but may translate to a decrease in cardiac performance only during exercise, when greater venous return results in longer SL at the end of diastole. Indeed, this mutation may result in increased basal myocardial performance, when cardiomyocytes are operating at shorter SLs, and thus could be a compensatory mechanism. Although cross-bridge recruitment was measured only during maximal Ca\(^{2+}\) activation, the authors also concluded that, at physiological levels of Ca\(^{2+}\), an increase in cross-bridge recruitment at longer SL may result in greater systolic muscle force development. This would prolong systolic ejection time, leading to a delay in relaxation and greater resistance to ventricular filling during diastole. There are two concepts here that merit further thought and investigation. The first is that some mutations may not allow complete deactivation of thin filaments at the end of systole, thus increasing a cross-bridge component of increased wall stiffness during diastole that resists ventricular filling. This has been reported by others for mutations in cardiac and skeletal muscle (Greenberg et al., 2009; Racca et al., 2015). The second is that fHCM mutations in troponin proteins may alter the kinetics of relaxation during basal function and/or during stress (exercise), as mentioned above (Cheng et al., 2015, 2016).

In conclusion, the work of Reda and Chandra (2018) agrees well with other recent publications demonstrating that the mechanism underlying contractile dysfunction resulting from fHCM-associated mutations is more complex than simple changes in the Ca\(^{2+}\) sensitivity of contractile activation. Mutation-related alterations in cross-bridge recruitment and cycling kinetics, impaired relaxation, and the SL dependence of these properties likely all play a role. There is also increasing evidence that unique differences in dysfunctional sarcomere behavior may depend on whether mutations are located in thick versus thin filament proteins, or even when mutations are located in different regions of the same protein (Montgomery et al., 2001; He et al., 2007; Tardiff, 2011). Better understanding of the complexity and diversity of mutation-mediated changes in sarcomere structure and function are needed to develop new and more specific targeting strategies for treating fHCM patients.

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