Altered cardiac excitation–contraction coupling in mutant mice with familial hypertrophic cardiomyopathy

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Introduction

Familial hypertrophic cardiomyopathy (FHC) is an inherited autosomal dominant disease characterized by ventricular hypertrophy and myofibrillar disarray. Sudden cardiac death, especially among young athletes, represents the most severe outcome of FHC in affected individuals (1–3). The genetic alterations have been localized to genes that encode various myofibrillar proteins (4, 5). Among these, the α-myosin heavy chain (α-MHC) gene was the first to be implicated and remains the most common cause of FHC. Recently, a transgenic model of FHC in mice with truncated troponin T (TnT) has also been developed (6). These transgenic mice hearts are smaller than wild-type, with fewer myocytes, but they exhibit significant delay in relaxation in response to increased work load with mild systolic dysfunction. Despite these advances, it is not yet clear how the genetic alterations are well defined. We characterized calcium cycling and contractile activation in trabeculae from a mutant mouse model of FHC (Arg403Gln knockin, α-myosin heavy chain). Wild-type mice of the same strain and age (~20 weeks old) served as controls. During twitch contractions, peak intracellular Ca2+ ([Ca2+]i) was higher in mutant muscles than in the wild-type (P < 0.05), but force development was equivalent in the two groups. Ca2+ transient amplitude increased dramatically in both groups as stimulation rate increased from 0.2 to 4 Hz. Nevertheless, developed force fell at the higher stimulation rates in the mutants but not in controls (P < 0.05). The steady-state force–[Ca2+]i relationship was less steep in mutants (Hill coefficient, 2.94 ± 0.27 vs. 5.28 ± 0.64; P > 0.003), with no changes in the [Ca2+]i required for 50% activation or maximal Ca2+-activated force. Thus, calcium cycling and myofilament properties are both altered in FHC mutant mice: more Ca2+ is mobilized to generate force, but this does not suffice to maintain contractility at high stimulation rates.

Excitation–contraction coupling in cardiac muscle of familial hypertrophic cardiomyopathy (FHC) remains poorly understood, despite the fact that the genetic alterations are well defined. We characterized calcium cycling and contractile activation in trabeculae from a mutant mouse model of FHC (Arg403Gln knockin, α-myosin heavy chain). Wild-type mice of the same strain and age (~20 weeks old) served as controls. During twitch contractions, peak intracellular Ca2+ ([Ca2+]i) was higher in mutant muscles than in the wild-type (P < 0.05), but force development was equivalent in the two groups. Ca2+ transient amplitude increased dramatically in both groups as stimulation rate increased from 0.2 to 4 Hz. Nevertheless, developed force fell at the higher stimulation rates in the mutants but not in controls (P < 0.05). The steady-state force–[Ca2+]i relationship was less steep in mutants (Hill coefficient, 2.94 ± 0.27 vs. 5.28 ± 0.64; P > 0.003), with no changes in the [Ca2+]i required for 50% activation or maximal Ca2+-activated force. Thus, calcium cycling and myofilament properties are both altered in FHC mutant mice: more Ca2+ is mobilized to generate force, but this does not suffice to maintain contractility at high stimulation rates.

The recent introduction of an FHC mutation (Arg403Gln) into the mouse α-MHC gene has produced an animal model resembling, in many aspects, human FHC (10). Here, we have studied the physiological properties of muscles from the hearts of these mice. The results indicate that there are alterations in excitation–contraction coupling in mutant mice, with lesions in myofilament contraction and associated changes of intracellular Ca2+ ([Ca2+]i) handling.

Methods

Mouse muscle preparation. Adult male mice (Black Swiss strain, ~20 weeks old, ~35 g; for a detailed description of the mouse model of FHC, see ref. 10) were anesthetized by intraperitoneal injection. The hearts were rapidly excised via midsternal thoracotomy. As described for wild-type FHC, see ref. 10) were anesthetized by intra-abdominal injection of sodium pentobarbital (~10–20 mg), and the hearts were rapid-

room temperature (20–22°C). Force and sarcomere length were measured as described previously (12). Diastolic sarcomere length (SL) was measured by laser diffraction, set to 2.1–2.2 mm, and monitored throughout the experiments. The preparations were field stimulated with 5-ms pulses (Grass S44 Stimulator; Grass Instruments Co., Quincy, Massachusetts, USA). Tetanization was achieved by stimulating the preparations at 10 Hz in the presence of cyclopiazonic acid (50 mM) at varied extracellular Ca^{2+} ([Ca^{2+}]_o) (usually 0.5–15 mM).

**Results**

**Effect of [Ca^{2+}]_o on twitch force**

We first characterized force development and calcium cycling during twitch contractions. Figure 1 compares calcium transients (a and c) and force (b and d) in representative wild-type (a and b) and mutant (c and d) muscles, at various [Ca^{2+}]_o. In both groups, calcium transient amplitude and force increase as [Ca^{2+}]_o is elevated from 2 to 4 to 10 mM. Note, however, that Ca^{2+} transient amplitudes are distinctly greater in the mutant muscle than in wild-type, despite the fact that systolic and diastolic forces were roughly equal at each [Ca^{2+}]_o.

The pooled data in Fig. 2 confirm that the discrepancy between calcium cycling and force was consistent and significant. At any given [Ca^{2+}]_o, more intracellular Ca^{2+} is required in the mutant muscles to generate the same levels of force. As another way of gauging the relationship between [Ca^{2+}]_o and force during twitch contractions, we plotted peak [Ca^{2+}]_i against peak force in each muscle (Fig. 3). Although there is no difference in the slopes (P > 0.2), the relationship from the mutant muscles is shifted to the right relative to the wild-type. Note also that the mutant muscles reach higher values of [Ca^{2+}]_i during the generation of equivalent forces. These data clearly show that excitation-contraction coupling is abnormal in mutant myocardium. Moreover, the changes are distinctive, differing from those described previously in stunned myocardium (15).

**Dynamics of twitches and Ca^{2+} transients in wild-type and mutant muscles at varied [Ca^{2+}]_o**

We next looked for possible alterations in the time course of calcium and force during twitch contractions. Figure 4 plots the times to peak (a and b) and half-relaxation (c and d) for twitch force (a and c) and Ca^{2+} transients (b and d). The times to peak force increased equivalently as [Ca^{2+}]_o was raised in the two groups, whereas the times to peak [Ca^{2+}]_i were not affected by changes in [Ca^{2+}]_o, in either the wild-type or mutant muscles. The only significant kinetic difference between the two groups was a delay of relaxation of force in the mutant muscles, evident at [Ca^{2+}]_o >5 mM. Similar delays in relaxation have recently been reported in intact hearts from the same colony of mutant mice (16).

**Effect of stimulation frequency on [Ca^{2+}]_i and force development**

Mouse hearts normally contract at very high frequencies (>500/min at 37°C), whereas our baseline studies were performed at 0.5 Hz (20–22°C). To look for
but the relative importance and nature of myofilament involvement remain unclear. To evaluate myofilament Ca\(^{2+}\) responsiveness directly, steady-state activation (Fig. 6a) was achieved by bursts of rapid pacing in the presence of cyclopiazonic acid (50 mM), a reversible inhibitor of calcium uptake by the sarcoplasmic reticulum. Figure 6b shows the pooled steady-state force–[Ca\(^{2+}\)]\(_i\) relationships in the two groups. Several features of the activation curves are noteworthy. First, maximal Ca\(^{2+}\)-activated force, or \(F_{\text{max}}\) (48 ± 4.2 mN/mm\(^2\) in wild-type, 45 ± 5.5 mN/mm\(^2\) in mutant) is lower in both groups compared with muscles from younger mice of another strain (11). An overall comparison of the curves from mutant and wild-type mice reveals significant differences (\(P = 0.03\) by multivariate ANOVA). The mid-point of activation is similar in the mutant (Ca\(_{50} = 1.35 ± 0.17\) mM) and wild-type muscles (1.20 ± 0.07 mM; \(P > 0.05\)), but the slopes of the steady-state relationships are shallower in the mutants (5.30 ± 0.64 in wild-type vs. 2.94 ± 0.27 in mutant; \(P < 0.003\)). As a consequence of the decrease in slope without a change in the midpoint, force in the mutants tends to be greater at lower [Ca\(^{2+}\)]\(_i\), but the converse is true at higher [Ca\(^{2+}\)]. In other words, at low [Ca\(^{2+}\)], mutant myofilaments are more sensitive to Ca\(^{2+}\), whereas at higher [Ca\(^{2+}\)], less force is produced until the two curves converge at \(F_{\text{max}}\). Similar behavior has been observed in skinned mutant muscles from this mouse model (17).

Discussion

This is the first study of intact cardiac muscle in a mouse model of FHC. Our success in adapting the trabecular force/calcium measurement methodology for use in mouse cardiac muscle (11) enabled us to perform such a study. We have found that there are indeed alterations in the properties of the myofilaments in FHC mutant mice. As will be discussed later in this article, these changes have important implications for the pathophysiology of FHC.

Contraction and Ca\(^{2+}\) transients in mutant muscles. One striking observation from this study was the increased Ca\(^{2+}\) transient amplitude in mutant mice at higher (>3.0...
Figure 4
Effects of $[\text{Ca}^{2+}]_o$ on dynamics of force (a and c) and $\text{Ca}^{2+}$ transients (b and d). Pooled data for time to peak (a and b) and time from peak to 50% relaxation (c and d) of $\text{Ca}^{2+}$ transients and twitch force ($n = 6$ in each group). In both groups, time to peak force increased as $[\text{Ca}^{2+}]_o$ was increased without changes in time to peak $\text{Ca}^{2+}$ transients. In mutant muscles, relaxation of force was significantly prolonged at $[\text{Ca}^{2+}]_o > 6.0$ mM, whereas decay of $\text{Ca}^{2+}$ transients was significantly accelerated. *$P < 0.05$.

Figure 5
Force–frequency relationship of mouse cardiac muscle. Pooled data of $[\text{Ca}^{2+}]_i$ (a and c) and twitch force (b and d) from wild-type (circles and squares; $n = 6$) and mutant (triangles; $n = 7$) muscles are plotted at varied stimulation frequencies at $[\text{Ca}^{2+}]_o = 2.0$ mM. (a and b) Systolic and diastolic $[\text{Ca}^{2+}]_i$ and force. Note the increases in both twitch force and $[\text{Ca}^{2+}]_i$ in wild-type muscles when stimulation frequency increased. In mutant muscles, increases in force was blunted at higher stimulation rate, with elevation of diastolic force. (c and d) Changes of $[\text{Ca}^{2+}]_i$ and developed force as stimulation rate was increased. Note the decrease in developed force in mutant muscles at higher stimulation rates. The two response curves are statistically different ($P < 0.002$) by multivariate ANOVA.
and Hill coefficient in each group. In mutant muscles, there is a less-steep force–[Ca\textsuperscript{2+}]i relationship, which also crosses over that of the wild-type at Steady-state force–[Ca\textsuperscript{2+}]i relationship of mouse cardiac muscle. (Figure 6 a).

\[ \text{Ca}_{50} \]

[Ca\textsuperscript{2+}]i. All other force levels were normalized with respect to their own maximal values. The dashed lines are the Hill fits, based on the means of \text{Ca}_{50} (\text{Ca}\textsubscript{2+}o to obtain different levels of tetanization. (b) One muscle in each group. Tetanizations were achieved by stimulating the muscle at 10 Hz in the presence of cyclopiazonic acid (50 mM) at varied closed circles mM [Ca\textsuperscript{2+}]o, despite the fact that twitch force remained the same as in wild-type (Fig. 2). This may intuitively lead one to believe that Ca\textsuperscript{2+} responsiveness of mutant muscle is decreased, because more [Ca\textsuperscript{2+}]i was required to produce equivalent levels of force, as in stunned myocardium (15). Nevertheless, several features of mutant muscles distinguish them phenotypically from stunned myocardium. First, FHC muscle exhibits the adaptive response of an increase in [Ca\textsuperscript{2+}], to maintain force; i.e., more Ca\textsuperscript{2+} is more mobilized. This may be due either to increased entry of Ca\textsuperscript{2+} or increased release of Ca\textsuperscript{2+} from the sarcoplasmic reticulum, or both. Second, the specific features of the decrease in Ca\textsuperscript{2+} responsiveness in FHC mice are unique: when peak force and peak [Ca\textsuperscript{2+}] were plotted against each other (Fig. 3), there was no significant difference between the slopes of the relationships, but rather a parallel rightward shift of the relationship in mutant muscles. Obviously, a different mechanism than the one responsible for stunning is responsible for the decreased Ca\textsuperscript{2+} responsiveness observed here. Primary alterations of cross-bridge cycling kinetics can be manifested as changes in steady-state myofilament Ca\textsuperscript{2+} responsiveness (18); further studies will be required to ascertain the mechanism of these changes and their relationship to specific steps in cross-bridge cycling.

Changes in relaxation of mutant muscles. In both groups of muscles, relaxation became slower as [Ca\textsuperscript{2+}]i was raised (Fig. 4). This slowing of relaxation was apparently unrelated to [Ca\textsuperscript{2+}]i, because the decay of Ca\textsuperscript{2+} transients was, paradoxically, accelerated. As discussed previously (11), the delay of relaxation of force in the face of accelerated relaxation of Ca\textsuperscript{2+} transients is likely the result of force-dependent changes in the affinity of the myofilaments to Ca\textsuperscript{2+} at higher levels of force (19). However, if an increase in the myofilament affinity for Ca\textsuperscript{2+} were the sole mechanism, one would also expect higher peak force in mutant muscle, which was not observed. Increases in [Ca\textsuperscript{2+}]i have multiple effects that feed back upon relaxation. Phosphorylation of troponin I by Ca/calmodulin–dependent protein kinase II (CaMKII) decreases force development and accelerates relaxation (20); the kinase also increases Ca\textsuperscript{2+} reuptake by the sarcoplasmic reticulum (21). Although the idea is reasonable and merits further study, the possible involvement of protein kinase systems in the abnormal relaxation of mutant myocardium remains conjectural.

Effect of stimulation frequency in mutant muscles. Normal mouse cardiac muscle was found to have a positive force–frequency relationship (11), and this feature was also observed in the wild-type muscles studied here (Fig. 5b). The frequency-dependent potentiation of force was less pronounced in the present study, possibly owing to the use of older animals and/or to genuine differences between inbred mouse strains. In mutant muscles, diastolic force increased with stimulation frequency without concomitant changes of [Ca\textsuperscript{2+}]i. The increased diastolic force may simply reflect the previously discussed impairment of relaxation when Ca\textsuperscript{2+} is high (Fig. 4). On the other hand, the failure to produce higher systolic force presumably reflects a defect of force-generating ability of the mutant myofilaments, because Ca\textsuperscript{2+} transients did not change. The negative force–frequency relationship provides a specific rationale for heart failure, especially during exercise. The delayed relaxation may underlie the diastolic dysfunction of FHC as well.

Steady-state force–[Ca\textsuperscript{2+}], relationship in mutant muscles. The steady-state force–[Ca\textsuperscript{2+}], relationship in FHC myocardium is clearly different from that of wild-type (Fig. 6b). This difference is mainly caused by a decrease in the cooperativity of the myofilaments. Because nor-
mal mouse muscle has high myofilament cooperativity (Fig. 6b and ref. 11), the decrease is expected to have a retarding effect on force development, especially when \([\text{Ca}^2+]\) is submaximal. Another alteration in the relationship worth noting is the cross-over with that of wild-type at \(\sim\text{Ca}50\). As already discussed, this feature may be manifested as changes in relaxation dynamics (Fig. 4). The mechanism for decreased cooperativity in mutant muscle is not understood at present, but it would not be surprising if it turns out to be a direct consequence of the structural alteration (Arg403Gln) in the myosin heavy chain.

While our findings reveal novel alterations of calcium cycling in FHC mutant muscle, the underlying mechanisms remain to be explored. Given the nature of the primary defect, these alterations may well be secondary to altered mechanics due to defective myofilament structure. We can clearly conclude that, for whatever reason, \(\text{Ca}^2+\) mobilization is upregulated in FHC myocardium. These changes in \(\text{Ca}^2+\) regulation may have numerous potential sequelae yet to be characterized. Given what we know about the key role of \([\text{Ca}^2+]\) in the pathogenesis of cardiac arrhythmias (22), it seems probable that the higher incidence of sudden cardiac death in FHC with Arg403Gln mutation arises from the alterations in calcium cycling, rather than directly from the myofilament defect. In this respect, it would be very interesting to investigate whether changes in calcium cycling are less marked in muscles containing more benign mutations in myosin, e.g., Val606Met.

Acknowledgments

Supported by National Institutes of Health grant R01 HL-44065 (to E. Marbán).