Neonatal cardiomyopathy in mice homozygous for the Arg403Gln mutation in the α cardiac myosin heavy chain gene

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Introduction

Familial hypertrophic cardiomyopathy (FHC) is inherited as an autosomal dominant trait that is characterized by unexplained hypertrophy of the ventricular myocardium with histologic evidence of myocyte and myofibrillar disarray. Myocyte loss and replacement fibrosis are often prominent features, which can contribute to arrhythmias and altered myocardial hemodynamics in FHC. Molecular genetic studies have demonstrated that FHC is caused by mutations in genes encoding cardiac sarcomere proteins; these defects are responsible for the disease phenotype. Myocardial fibrosis and cell viability are hallmarks of FHC. Myocytolysis, cardiomyocyte death, and cell viability are hallmarks of FHC.

Heterozygous mice bearing an Arg403Gln missense mutation in the α cardiac myosin heavy chain gene (α-MHC403/−) exhibit the histopathologic features of human familial hypertrophic cardiomyopathy. Surprisingly, homozygous α-MHC403/403 mice die by postnatal day 8. Here we report that neonatal lethality is caused by a fulminant dilated cardiomyopathy characterized by myocyte dysfunction and loss. Heart tissues from neonatal wild-type and α-MHC403/403 mice demonstrate equivalent switching of MHC isoforms; α isoforms in each increase from 30% at birth to 70% by day 6.

Cardiac dimensions and function, studied for the first time in neonatal mice by high frequency (45 MHz) echocardiography, were normal at birth. Between days 4 and 6, α-MHC403/403 mice developed a rapidly progressive cardiomyopathy with left ventricular dilation, wall thinning, and reduced systolic contraction. Histopathology revealed myocardial necrosis with dystrophic calcification. Electron microscopy showed normal architecture intermixed with focal myofibrillar disarray. We conclude that 45-MHz echocardiography is an excellent tool for assessing cardiac physiology in neonatal mice and that the concentration of Gln403 α cardiac MHC in myocytes influences both cell function and cell viability. We speculate that variable incorporation of mutant and normal MHC into sarcomeres of heterozygotes may account for focal myocyte death in familial hypertrophic cardiomyopathy.

Table 1
Echocardiographic data for wild-type (+/+) and cardiac MHC mutant mice on postnatal days 0, 4, and 6

<table>
<thead>
<tr>
<th>Mouse genotype</th>
<th>Day 0</th>
<th>Day 4</th>
<th>Day 6</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>(n = 6)</td>
<td>(n = 4)</td>
<td>(n = 11)</td>
</tr>
<tr>
<td>Wild-type (+/+)</td>
<td>1.38 ± 0.23</td>
<td>2.37 ± 0.52</td>
<td>3.34 ± 0.83</td>
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<tr>
<td>Heart rate (bpm)</td>
<td>268 ± 54</td>
<td>314 ± 49</td>
<td>304 ± 41</td>
</tr>
<tr>
<td>LVDD (mm)</td>
<td>1.37 ± 0.08</td>
<td>1.66 ± 0.17</td>
<td>1.78 ± 0.26</td>
</tr>
<tr>
<td>LVSD (mm)</td>
<td>0.91 ± 0.08</td>
<td>0.98 ± 0.10</td>
<td>0.98 ± 0.12</td>
</tr>
<tr>
<td>LVFS (%)</td>
<td>34 ± 4</td>
<td>41 ± 6</td>
<td>45 ± 8</td>
</tr>
<tr>
<td>LVWT (mm)</td>
<td>0.21 ± 0.02</td>
<td>0.21 ± 0.03</td>
<td>0.28 ± 0.05</td>
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</table>

Abbreviations as explained in text.

MHC403/+, and wild-type (strain 129/Black Swiss) mice were studied between postnatal day 0 (birth) and day 6. Mouse genotype was determined by restriction enzyme digestion of PCR-amplified tail DNA. All mice were maintained according to protocols approved by the Institutional Animal Care and Use Committees of Harvard Medical School and New York University Medical Center.

Pathology. Fixed hearts were examined grossly and then bisected transversely at the midventricular level before histologic examination. Both apical and basal halves were embedded in paraffin or glycol methacrylate plastic (JB-4, Polysciences Inc., Warrington, Pennsylvania, USA), sectioned from the cut surfaces at ~5 μm and 2 μm, respectively, and stained for light microscopy with hematoxylin and eosin (for overall morphology). Selected paraffin sections were also stained with Mason’s trichrome stain (for collagen) and Von Kossa’s reagent (for calcium phosphates). Myocardial necrosis and calcification were graded as follows: 0, not present; 1+, mild, comprising a single small focus of necrosis and/or calcification; 2+, moderate, multifocal-to-confluent foci; and 3+, severe, large, confluent clusters and/or transmural involvement. Histologic specimens were further counterstained with methyl green for morphologic analysis of myocyte nuclei.

The myocardial tissue processed for transmission electron microscopy was fixed in 2.5% glutaraldehyde and 2% paraformaldehyde in cacodylate buffer at pH 7.4, postfixed in 2% osmium tetroxide, dehydrated in propylene oxide, and embedded in Poly/Bed 812 (Polysciences Inc.). Sections were cut at 60 nm, stained with lead citrate and uranyl acetate, and examined with a JEOL-100CX transmission electron microscope (JEOL USA Inc., Cranford, New Jersey, USA) at an accelerating voltage of 80 kV.

Echocardiography. Transthoracic echocardiography was performed without anesthesia and with mice lightly restrained in a supine position, at ambient temperatures >25°C, using a Humphrey Ultrasound Biomicroscope (model 840; Humphrey Instruments, San Leandro, California, USA). Two-dimensional images were obtained using a mechanically scanned, focused transducer (gently positioned over a layer of acoustic gel on the anterior chest to avoid pressure that might cause bradycardia) that operated at a nominal frequency of 45 MHz. The spatial resolution of the echocardiographic system was 30 μm in the axial (depth) direction and 65 μm in the lateral (horizontal) direction; penetration depth was 4–5 mm. Two-dimensional 5 × 5-mm, 8-bit grayscale video images were acquired at a scan rate of 8 Hz, with a gain setting of 80 dB and a time gain correction of 5 dB/mm.

Continuous-wave Doppler. Continuous-wave (CW) Doppler interrogation was performed immediately after two-dimensional echocardiographic imaging. The CW Doppler probe was manipulated at a constant angle on the left anterior chest wall until maximal blood velocity signals were obtained. A high-frequency Doppler ultrasound system using two air-backed lithium niobate transducers operating at an ultrasound frequency of 43 MHz (6) was used. The two transducers were aligned to obtain overlapping ultrasound beams (angle of intersection is 30°), the intersection of which defines the Doppler sample volume. Transducers were fabricated with diameters between 1 and 1.5 mm, resulting in Doppler sample volumes between 0.5 and 1.5 mm³; CW Doppler data were recorded onto digital audio tapes and analyzed on a PC computer in the form of a spectrogram (6). Conversion between Doppler shift frequency and blood velocity was made assuming a speed of sound in blood of 1,580 m/s (7) to obtain a conversion factor of 0.02 (mm/s)/Hz.
Data analysis. Left ventricular (LV) diameters, areas, and wall thicknesses were obtained from cross-sectional short-axis views. Diastolic and systolic LV dimensions were measured from six consecutive cardiac cycles. To account for the low frame rates and lack of synchronization of image acquisition with the cardiac cycle, LV diastolic diameter (LVDD) was calculated from the mean of the three maximal diastolic measurements; LV systolic diameter (LVSD) was calculated from the mean of the three minimal systolic measurements. Left ventricular wall thickness (LVWT) was derived from the mean of three minimum diastolic wall thickness measurements. The coefficient of variation for each set of measurements was expressed as the SD/mean. Left ventricular fractional shortening (LVFS) was expressed as a percentage, using the formula (LVDD – LVSD) / LVDD · 100. Heart rates were calculated using the time interval between successive waveforms on CW Doppler tracings. Measurements were made without prior knowledge of genotype. Inter- and intraobserver variability was assessed by comparison of independent measurements of LV dimensions by two observers, and on two different occasions, by one observer.

Protein assays. α-MHC403/403 and wild-type mouse hearts were harvested at postnatal days 0, 2, 4, and 6. Ventricles were isolated, washed in PBS, blotted on filter paper, and immediately frozen in liquid N2. Tissues were homogenized in 1 ml of ice-cold inhibiting buffer (8) containing 50 mM KH2PO4, 70 mM NaF, and 5 mM EDTA, with protease inhibitors (5 μg/ml antipain, 10 μg/ml leupeptin, 5 μg/ml pepstatin A, 43 μg/ml phenylmethylsulfonyl fluoride, 5 mM EGTA, and 0.1 μM sodium orthovanadate), and centrifuged at 15,000 g for 5 min at 4°C. The pellet was incubated on ice for 30 min in inhibiting buffer plus 1% Triton X-100 plus protease inhibitors, and then centrifuged at 5,000 g for 5 min. Detergent-extracted myofibrils were resuspended in inhibiting buffers.

Myofibrillar proteins were separated by SDS-PAGE. Proteins were solubilized and denatured in sample preparation buffer containing 4% SDS, 0.23 M β-mercaptoethanol, and 0.2 M Tris-HCl (pH 6.5) at 100°C for 5 min. Protein concentration was determined using the Bradford protein assay. Proteins were separated by PAGE in the presence of 0.1% SDS at constant current (50 mA, for 30 h) on 6% polyacrylamide gels with recirculated, cooled (15°C) running buffers. After electrophoresis, the gels were silver-stained and dried. The relative proportions of α and β cardiac MHC isoforms and a high molecular weight protein reference standard were quantified using densitometry.

Statistical analysis. The significance of differences in survival of α-MHC403/403, α-MHC403/+, and wild-type mice was assessed by the one-group χ2 test. Echocardiographic measurements made in the three groups of mice were compared by one-factor ANOVA and Student’s t test. Relationships between continuous variables were characterized using linear regression analysis. Comparison of inter- and intraobserver measurements were determined using linear regression analysis and Bland and Altman’s “limits of agreement” (9), in which acceptable reproducibility requires ≥95% of the data to be within ± 2 SD of the mean difference. Data are expressed as mean ± SD. P < 0.05 was considered significant.

Results

Survival analysis. MHC403/403 mice were produced by mating heterozygous α-MHC403/+ mice. At birth, the ratio of α-MHC403/403/α-MHC403/+ wild-type mice approximated...
inflammation but often was accompanied by secondary dystrophic calcification (Fig. 2d). By postnatal day 6, myocardial necrosis was present in 100% (n = 15) of α-MHC<sup>403/403</sup> mice; in 14 animals, this was moderate to severe (grades 2+/3+). Dystrophic calcification was present in 87% (n = 13); in nine hearts, this was moderate to severe. No mice had calcification in the absence of necrosis.

Electron microscopy of α-MHC<sup>403/403</sup> hearts demonstrated normal sarcomere structure (Fig. 3), with alignment of myofibrils along the long axis of the myocytes and the presence of intact Z bands throughout the cell length. Although some sections showed focal myofibrillar disarray, most were indistinguishable from sections derived from α-MHC<sup>403/4</sup> or wild-type mice.

Echocardiographic studies were performed in 97 mice between ages 0 and 6 days without procedural complications or deaths. Image acquisition was completed on each mouse within 10–15 minutes. Cross-sectional LV images were obtained in a short-axis view in each animal. The right ventricle (RV) was visualized in the majority of studies. The LA was observed only when pathological dilation was present; the RA was not observed in any study.

Good correlations were obtained between two observers for LV cavity dimensions: LVDD (r = 0.88, P < 0.001), LV diameters and areas. Intraobserver correlations for all LV parameters were high: LVDD (r = 0.98, P < 0.001), LVDD = 0.85, LVSD (r = 0.98, P < 0.001), LV systolic area (LVSA, r = 0.55, P < 0.001), LVFS (r = 0.74, P < 0.001). Left ventricular wall thickness measurements demonstrated greater interobserver variability (r = 0.30, P = 0.007) than did LV diameters and areas. Intraobserver correlations for all LV parameters were high: LVDD (r = 0.98, P < 0.001), LVDD (r = 0.98, P < 0.001), LVSD (r = 0.97, P < 0.001), LVFS (r = 0.96, P < 0.001), LVFS (r = 0.96, P < 0.001), LVWT (r = 0.50, P = 0.001). All LV parameters were within, or close to, the 95% limits of agreement (range 92%–98%). The coefficient of variation for measurements made by a single observer on one occasion was <5% for LV diameters and areas and <7% for LV wall thickness.

No significant differences were observed in cardiac morphology or size between wild-type and α-MHC<sup>403/4</sup> mice during days 0–6 (Table 1). Heart rates were also equivalent but notably lower (~50%) than those observed in conscious adult animals (Table 1 and data not shown).

Echocardiographic parameters of α-MHC<sup>403/4</sup> mouse hearts were indistinguishable from those of littermates at birth (Table 1). Increases in LV chamber dimensions (LVDD and LVSD) and LV wall thinning were evident by postnatal day 4. Contractile function was preserved in three of four α-MHC<sup>403/4</sup> mice but was reduced in four mice (LVFS = 19%, 23%, 26%, and 31%). These parameters tended to worsen by postnatal day 6, at which time all α-MHC<sup>403/4</sup> mice exhibited a global reduction of systolic contraction and/or LV dilation. Echocardiographic studies on postnatal day 6 demonstrated biventricular failure in 50% of α-MHC<sup>403/4</sup> mice: LA, LV, and RV dilatation and pericardial effusion (Fig. 4) were present. Echo-dense material detected in the LA of these animals was caused by thrombus in the dilated LA cavity, with foci of necrosis in the LA myocardium (Fig. 5).

The extent of myocardial necrosis in α-MHC<sup>403/4</sup> mouse hearts correlated with chamber dimensions and function (LVDD, P < 0.001; LVFS, P < 0.001). By postnatal

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Figure 4
Two-dimensional images obtained by 45-MHz echocardiography demonstrating the left ventricle (LV) and right ventricle (RV) in a short-axis view in diastole (a and d) and systole (b and e). Normal cardiac size and function at postnatal day 6 in a α-MHC<sup>403/4</sup> mouse (a and b). The LV is dilated and thin-walled with reduced systolic contraction in an α-MHC<sup>403/4</sup> mouse (d and e). In addition, RV dilation, LA dilation, and a pericardial effusion (d, arrow) were identified. Echocardiographic measurements (c) were as follows: transverse LV diameter (arrow), LV area (dashed lines), and LV wall thickness (asterisk). (f) Bidirectional blood flow velocity pattern on CW Doppler interrogation of the left hemithorax, consistent with transmural LV inflow (positive velocity) and aortic outflow (negative velocity). Scale bar, 1 mm. LV, left ventricle; RV, right ventricle; LA, left atrium.
day 4, myocardial necrosis was present in all α-MHC403/403 mice with reduced LVFS (n = 4) and in two of the three mice with normal LVFS (necrosis grades 1+ and 3+, respectively). On day 6, myocardial necrosis was present in all mice with reduced LVFS (n = 4) and in two mice with normal LVFS (necrosis grades 2+ and 3+, respectively).

Cardiac MHC isoforms. The total amount of cardiac MHC in the myofibrillar fractions was assessed by comparison to a high molecular weight protein; the total amount of cardiac MHC was the same in myofibrillar fractions from wild-type and α-MHC403/403 mouse hearts (data not shown). Levels of α and β cardiac MHC isoforms were assessed in wild-type and α-MHC403/403 mice by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) (Fig. 6). In wild-type mice, ventricular levels of β cardiac MHC exceeded α cardiac MHC at day 0 (β = 66%, α = 34%); by day 6, the cardiac MHC isoform switch was virtually complete, with a predominance of the α isoform (β = 27%, α = 73%). This pattern of expression was not significantly altered in α-MHC403/403 mice: day 0, β-MHC = 73%, α-MHC = 27%; day 6, β-MHC = 30%, α-MHC = 70%.

Discussion
Cardiac structure and function of neonatal α-MHC403/403 mice demonstrate that homozygous mice bearing the Arg403Gln mutation develop a rapidly progressive, lethal cardiomyopathy. Unlike the cardiomyopathy observed in heterozygous α-MHC403/403 mice, systolic dysfunction, myocardial necrosis, and dystrophic calcification are the predominant features of the α-MHC403/403 pathology. These data demonstrate the technical feasibility of in vivo cardiac assessment of neonatal mice and indicate that myocyte dysfunction and loss may be important in the pathophysiology of human FHC.

The application of genetic engineering to study the molecular basis of cardiovascular structure and function has posed an important research challenge: assessment of cardiac physiology in small rodents. We demonstrate that 45-MHz echocardiography is a powerful tool for assessing in vivo cardiac function despite the extremely small size and high beating rates of neonatal mouse hearts. An important finding from these studies is a developmental profile of normal neonatal cardiac function during the first week of life. Neonatal mice had slower heart rates and reduced contractile function when compared with adult mice. Although LVFS increases (from 34% to 45%) during the first week of life, neonatal heart rates (~300 beats per minute [bpm]) remain slow. Increases in LVFS are probably due to altered cardiac MHC isoform expression that occurs in the first postnatal week (Fig. 6 and ref. 10), a hypothesis supported by previous in vitro and in vivo investigations. Myocytes expressing β cardiac MHC have lower myosin adenosine triphosphatase (ATPase) activity and lower velocity of fiber shortening than do myocytes that express α cardiac MHC (11–16). Furthermore, the LVFS values (34%–36%) observed in neonatal mice during the first few days after birth were equivalent to values reported in the embryonic mouse at E12.5–E13.5 (5) and in other species, such as humans (17–19) and rabbits (20), in which the β-MHC isoform predominates. In contrast, the LVFS observed after one week (45%) was similar to that reported previously in adult mice (21–23) and in other small rodents, such as adult rats (24), in which the α cardiac MHC isoform predominates.

Unlike wild-type and heterozygous littersmates, an increase in systolic contractility did not occur in α-MHC403/403 mouse hearts, and progressive LV dilation ensued. Given that the α/β cardiac MHC isoform ratio at postnatal day 6 in α-MHC403/403 mice was similar to that observed in wild-type hearts, the relatively lower mean LVFS could not be attributed to failure of the β-to-α isoform switch nor to upregulation of the β isoform in response to the LV functional deficit. The myocardial contractile dysfunction observed in α-MHC403/403 mouse...
hearts is caused by the Arg403Gln mutation in α cardiac MHC. Residue 403 is situated in the head of the MHC at the base of a loop that interacts with actin (25), and *in vitro* studies (26, 27) have demonstrated the Arg403Gln substitution alters the kinetics of the actin–myosin interaction by reducing actin translocating activity and actin-associated ATPase. Reductions in actin translocation can result in a lowered force/stiffness ratio and depressed velocity of shortening (28, 29), thereby producing global contractile dysfunction and LV dilation in neonatal α-MHC403/403 mice.

Although the mechanism by which mutant MHC produces myocyte necrosis remains to be elucidated, calcium is a likely participant. Heightened calcium sensitivity in heterozygous α-MHC403/4 hearts (30) and prolonged calcium transients have been observed in ventricular muscle preparations from animal models and patients with FHC (31); both suggest a propensity for calcium overload. However, the gross morphologic calcification in α-MHC403/403 heart sections (Fig. 2d) probably reflects a different process; its rapid appearance in preexisting necrotic foci is most consistent with accelerated dystrophic calcification (32) in young animals, in which bone mineralization mechanisms are active (33, 34). Left ventricular contractile dysfunction and dilation could promote myocardial ischemia and subsequent necrosis by reducing coronary artery perfusion, either directly through LV pump failure or indirectly through catecholamine-induced vasospasm (35). However, the distribution of necrotic lesions in α-MHC403/403 hearts are atypical for coronary ischemia, in which pathology is positioned along vascular beds, and observed heart rates were not consistent with increased catecholamine levels.

These data have important ramifications for understanding the histopathology in human FHC. Affected individuals with this disorder are almost always heterozygous, with one mutant sarcomere protein gene allele and one normal allele (1). However, individuals with homozygous sarcomere protein gene mutations have been observed very infrequently (36, and our unpublished data). Cardiac histopathology in heterozygous individuals typically shows marked variation in the distribution and severity of findings; foci with marked myofibrillar disarray, myocyte death, and replacement fibrosis are juxtaposed to normal myocardium. On the basis of the studies of α-MHC403/403 mice, we speculate that variable incorporation of mutant α cardiac MHC into sarcomeres may account for variability of cell viability in FHC. This is consistent with a “poison polypeptide” effect: levels above a critical threshold for mutant peptide (or sarcomere dysfunction) may cause the myocyte to die. Patchy myocyte loss probably contributes to arrhythmias in FHC; more global involvement results in the dilated phase of this disease. Myocyte loss has also been implicated in the progression of ventricular dysfunction and remodeling in cardiomyopathies associated with chronic ischemia, myocardial infarction, pressure overload hypertrophy, hypertension, chronic tachycardia, pacing, and aging (37–42). Hence, elucidation of the cellular events triggered by sarcomere dysfunction in the α-MHC403/403 mouse should provide insights into myocyte necrosis and LV dilation in many cardiomyopathies.

**Acknowledgments**

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