ARTICLE

Cardiomyopathic mutations in essential light chain reveal mechanisms regulating the super relaxed state of myosin

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In this study, we assessed the super relaxed (SRX) state of myosin and sarcomeric protein phosphorylation in two pathological models of cardiomyopathy and in a near-physiological model of cardiac hypertrophy. The cardiomyopathy models differ in disease progression and severity and express the hypertrophic (HCM-A57G) or restrictive (RCM-E143K) mutations in the human ventricular myosin essential light chain (ELC), which is encoded by the MYL3 gene. Their effects were compared with near-physiological heart remodeling, represented by the N-terminally truncated ELC (Δ43 ELC mice), and with nonmutated human ventricular WT-ELC mice. The HCM-A57G and RCM-E143K mutations had antagonistic effects on the ATP-dependent myosin energetic states, with HCM-A57G cross-bridges fostering the disordered relaxed (DRX) state and the RCM-E143K model favoring the energy-conserving SRX state. The HCM-A57G model promoted the switch from the SRX to DRX state and showed an ∼40% increase in myosin regulatory light chain (RLC) phosphorylation compared with the RLC of normal WT-ELC myocardium. On the contrary, the RCM-E143K-associated stabilization of the SRX state was accompanied by an approximately twofold lower level of myosin RLC phosphorylation compared with the RLC of WT-ELC. Upregulation of RLC phosphorylation was also observed in Δ43 versus WT-ELC hearts, and the Δ43 myosin favored the energy-saving SRX conformation. The two disease variants also differently affected the duration of force transients, with shorter (HCM-A57G) or longer (RCM-E143K) transients measured in electrically stimulated papillary muscles from these pathological models, while no changes were displayed by Δ43 fibers. We propose that the N terminus of ELC (N-ELC), which is missing in the hearts of Δ43 mice, works as an energetic switch promoting the SRX-to-DRX transition and contributing to the regulation of myosin RLC phosphorylation in full-length ELC mice by facilitating or sterically blocking RLC phosphorylation in HCM-A57G and RCM-E143K hearts, respectively.

Introduction

Cardiac muscle contraction results from the ATP-coupled interactions between myosin cross-bridges and actin/troponin-tropomyosin (Tm-Tn) filaments. It is regulated by Ca2+ ions and controlled by myosin filament-based mechanisms, including phosphorylation of myosin regulatory light chain (RLC) and myosin binding protein C (MyBP-C; Brunello et al., 2020). Myosin binds ATP and actin in the motor domain, and its lever arm region amplifies small conformational changes generated in the catalytic domain into the large movements needed to produce force and muscle contraction (Rayment et al., 1993; Geeves and Holmes, 2005). The myosin lever arm is structurally supported by myosin RLC and essential light chain (ELC), two important myosin subunits that bind to their respective IQ motifs within the myosin heavy chain lever arm region. During muscle relaxation, myosin head domains of paired myosin heavy chains appear to form an interacting-head motif that structurally characterizes two different myosin states, a disordered relaxed (DRX) state, in which myosin heads protrude into the interfilament space, and a super relaxed (SRX) state, in which they interact with the myosin backbone and each other (Alamo et al., 2017). Compared with DRX, the SRX state is characterized by low ATP turnover rates, typical of an energy-conserving state that serves as a modulator of energy utilization in cardiac muscle (Hooijman et al., 2011; McNamara et al., 2015). The normal

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cardiac relaxation phase is highly significant for normal heart function, and the disruption of the SRX state of myosin has been thought to increase the possibility for inefficient myocardial performance and poor clinical outcomes (Toepfer et al., 2020).

The myosin ELC constitutes an essential part of the lever arm structure, but it also plays a role in the contractile force development and the ATP-dependent kinetics of actin-myosin interactions (VanBuren et al., 1994; Timson, 2003; Miller et al., 2005; Kazmierczak et al., 2009). A functionally important domain of the cardiac ELC is its 43-amino-acid-long N-terminal extension (N-ELC) shown by many to directly interact with the C-terminal region of actin (Trayer et al., 1987; Milligan et al., 1990; Aydt et al., 2007; Kazmierczak et al., 2009; Wang et al., 2018). We previously reported that transgenic mice expressing the truncated N-ELC (Δ43 mice) display near-physiological cardiac remodeling with an increase in left ventricular (LV) wall thickness but no changes in cardiac morphology or function or in myofilament contraction/relaxation parameters are observed (Sitbon et al., 2020). On the other hand, pathological hypertrophy due to hypertrophic cardiomyopathy (HCM) or restrictive cardiomyopathy (RCM) is characterized by interstitial fibrosis, myofilament disarray, reduced LV cavity, and impaired heart’s relaxation phase and can lead to diastolic dysfunction and sudden cardiac death (Seidman and Seidman, 1998; Spirito et al., 2000; Arad et al., 2002; Ip et al., 2013). RCM is additionally manifested by biaxial enlargement, increased stiffness of the LV wall with no increase in wall thickness (Kushwaha et al., 1997; Yuan et al., 2017). Both HCM and RCM can result from genetic mutations in major sarcomeric proteins such as β-myosin heavy chain, cardiac MyBP-C, titin, actin, Tm, TnT, Tni, TnC, and both myosin RLC and ELC.

In this report, we focus on the A57G and E143K mutations in the ventricular ELC, the products of the mutated MYL3 gene that were shown by population studies to cause human HCM or RCM, respectively (Lee et al., 2001; Olson et al., 2002; Caleshu et al., 2011). We previously reported that transgenic mice expressing the A57G ELC mutation (HCM-A57G model) exhibited characteristic HCM remodeling that included increased LV wall thickness, fibrosis, and diastolic abnormalities manifested by prolongation of isovolumetric relaxation time (Sitbon et al., 2020). Skinned and intact papillary muscle (PM) experiments demonstrated increased Ca$^{2+}$ sensitivity of force, decreased maximal tension, and delayed Ca$^{2+}$ transients in HCM-A57G mice (Kazmierczak et al., 2013; Sitbon et al., 2020). Likewise, transgenic mice expressing the E143K ELC mutation (RCM-E143K model) displayed a typical RCM phenotype with a decrease in LV inner diameter and increased LV stiffness, fibrosis, and diastolic dysfunction, as judged by prolongation of isovolumetric relaxation time and an increase in Ca$^{2+}$ sensitivity of force (Yuan et al., 2017). Interestingly, in contrast to HCM-A57G mice, RCM-E143K animals exhibited decreased cardiac output and stroke work in vivo (Yuan et al., 2017) and downregulation of myosin power and a reduced force–velocity relationship in the in vitro motility assay (Wang et al., 2018).

These in vivo and in vitro functional characteristics of ELC transgenic animal models built the scientific premise of the current investigation aimed at the comprehensive assessment of the bioenergetic signatures present in HCM-A57G, RCM-E143K, and Δ43 mice. We first focused on the ATP-dependent interaction of myosin and actin assessed in skinned and intact PMs to provide insight into the energetic states of myosin and cross-bridge kinetics at the level of myofilaments. Specifically, we looked at the mechanism by which HCM and RCM ELC mutations modulate the SRX-to-DRX transition and how myosin energetic states in the hearts of HCM-A57G, RCM-E143K, and Δ43 mice correlate with myofilament protein phosphorylation. Because RLC-mediated regulation of myosin motor activity via myosin light chain kinase–dependent RLC phosphorylation has long been acknowledged to be critical for heart function in health and disease (reviewed in Yadav and Szczesna-Cordary, 2017), the emphasis was put on testing the myosin RLC phosphorylation in the hearts of mice. We thoroughly analyzed the interplay between myosin energetic states, SRX↔DRX equilibrium, and the degree of myosin RLC phosphorylation in all investigated ELC models.

Finally, because of our recently published proteomic data on HCM-A57G and Δ43 relative to WT-ELC hearts (Gomes et al., 2015) and on RCM-E143K versus WT-ELC mice (Yuan et al., 2017) that revealed significant differences between the models in biogenesis, cellular organization, and biological regulation, we also investigated ELC mutation-dependent mitochondrial and metabolic remodeling, oxidative phosphorylation (OXPHOS), and ATP respiration.

**Materials and methods**

**Transgenic mice**

All animal procedures and experiments were performed following the “Guide for the Care and Use of Laboratory Animals” (National Institutes of Health publication 85-23, revised 2011). All the protocols were approved by the Institutional Animal Care and Use Committee at the University of Miami Miller School of Medicine (protocol #18-110), which has an Animal Welfare Assurance on file with the Office of Laboratory Animal Welfare, National Institutes of Health (assurance number #A-3224-01, approved through November 30, 2019). We are registered with US Department of Agriculture Animal and Plant Health Inspection Service (registration #58-R-007, approved through December 3, 2023). We have full accreditation with the Association for Assessment and Accreditation of Laboratory Animal Care (site 001069, latest effective date, November 8, 2016). Mice were euthanized through CO$_2$ inhalation followed by cervical dislocation.

We have previously generated transgenic mice models expressing the mutated human cardiac ELC in HCM-A57G, RCM-E143K, and physiological-like model Δ43 mice as well as transgenic WT mice expressing the human ventricular ELC (Kazmierczak et al., 2009; Muthu et al., 2011; Yuan et al., 2017). Multiple crosses of transgenic ELC mice with B6SJL/F1 mice were performed before the animals were used for experiments. For this study, two lines of WT (L1, 76%; and L4, 71%), two of A57G (L1, 80%; and L5, 74%), and two of Δ43 (L8, 34–39%; and L9, 35–40%) were used. We also used L2 of E143K mice expressing 55% of mutant ELC protein (Yuan et al., 2017). All experiments were performed on the hearts of 8 ± 1.5-mo-old male and female mice.
Skinned PM experiments and the SRX state of myosin

N-methylanthraniloyl (mant)-ATP assays were performed on skinned PM fibers from HCM-A57G, RCM-E143K, Δ43, and WT mice using the IonOptix instrumentation as described in Yadav et al. (2019a). Fibers were incubated in a rigor solution containing 250 µM mant-ATP, and decay in fluorescence versus time curves were collected when mant-ATP was rapidly exchanged with 4 mM nonlabeled ATP. Decay isotherms were fitted to a two-state exponential equation, and the P1 and P2 and T1 and T2 parameters were calculated to describe the proportion of myosin heads in the SRX (P2) or DRX (P1) states and assess their respective T2 and T1 lifetimes (in seconds; Yadav et al., 2019a; Sitbon et al., 2020).

Intact PM fiber mechanics and force/calcium transients

Intact PM fiber studies were performed according to protocols described previously (Yuan et al., 2018; Sitbon et al., 2020). Briefly, intact PMs were carefully dissected from the LVS of mice in preoxygenated (95% O2, 5% CO2) 2,3-butanedione monoxime (BDM)-Krebs-Ringer solution and incubated in BDM-Krebs-Ringer buffer containing 50 µM Fura-2 AM (Life Technologies) at room temperature for 1.5 hr in the dark, followed by several washes in BDM-Krebs-Ringer solution to remove excess Fura-2 AM. Fibers were then mounted in a flow chamber between a hook connected to a length-controlled micromanipulator and a force transducer. Preoxygenated Krebs-Ringer solution (95% O2, 5% CO2, pH 7.4) was perfused into the chamber at a flow rate of ∼2 ml/min. The temperature in the muscle chamber was kept constant at 25°C by a heat exchanger at the inflow line and a circulating water bath. The muscle was stretched to an optimum length at which the developed tension was maximal during an isometric twitch. Sarcomere length, monitored by a camera, was maintained at ∼2.1 µm for another 10–15 min to allow the force development to stabilize. The stimulus was applied through platinum hooks, and strength was adjusted to 20% above the force development to stabilize. The stimulus was applied (PS726 and P0044; Sigma-Aldrich), 5 mM DTT, and 1 µl/ml protease inhibitor cocktail (P8340; Sigma-Aldrich) and homogenized in a Mixer-Mill MM301. The homogenate was centrifuged for 4 min at 8,000 × g, and the supernatant was discarded. After centrifugation, the pellets were left on ice for 2 min. This step was repeated three times with CMF buffer without Triton X-100. The pellets were then resuspended in the CMF buffer, and the samples were subsequently mixed at a 1:1 ratio with Laemmli buffer and 5% β-mercaptoethanol and heated at 100°C for 5 min. Samples were loaded onto 12% or 4–20% gradient gels (Mini-Protein TGX; Bio-Rad), and phosphoproteins were detected using Pro-Q Diamond/ Sypro system (Invitrogen, Thermo Fisher Scientific). Total protein was also detected using Coomassie staining as described previously (Kazmierczak et al., 2014). RLC and cMyBP-C phosphorylation was tested by Western blots and protein-specific antibodies. To detect phosphorylated RLC, ∼30 µg of protein per lane was loaded into 15% SDS-PAGE, and the membranes were probed with phospho-specific RLCmas antibody (gift from Dr. Neal Epstein, National Institutes of Health, Bethesda, MD), which recognizes the phosphorylated form of the RLC and does not react with nonphosphorylated RLC. Total RLC protein was detected with rabbit polyclonal CT-1 antibody, produced in this laboratory (Szczesna-Cordary et al., 2005), which served as a loading control. Phosphorylation of cMyBP-C was tested at Ser282. Approximately 20 µg of protein per lane was loaded into 12% SDS-PAGE, and the membranes were probed with phospho-specific cMyBP-Cser282 antibody (gift from Dr. Sakhthivel Sadayappan, University of Cincinnati, Cincinnati, OH). Goat polyclonal antibody was used to detect total cMyBP-C. Myofilament protein phosphorylation was calculated based on the density of bands of phosphorylated protein/total protein. The bands were analyzed using ImageJ software (National Institutes of Health).

Analysis of sarcomeric protein phosphorylation

Phosphorylation of cardiac myosin RLC, TnI, and cMyBP-C in all investigated animal models was measured in cardiac myofibrils (CMFs) isolated from six to eight hearts per group as described previously (Kazmierczak et al., 2013). Specifically, 4 M and 3 F (7.5–10-mo-old) WT, 3 M and 5 F (8–10-mo-old) HCM-A57G, 3 M and 3 F (6–9-mo-old) RCM-E143K, and 4 M and 2 F (7.5–10-mo-old) Δ43 mice were used. After euthanasia, the hearts were excised and placed in liquid nitrogen. Before the experiment, the tissue was thawed in CMF buffer consisting of 5 mM NaH2PO4, 5 mM NaHPO4, pH 7.0, 0.1 mM NaCl, 5 mM MgCl2, 0.5 mM EGTA, 5 mM ATP, 5 mM microcystin, 0.1% Triton X-100, 10 µl/ml of phosphatase inhibitor cocktails 2 and 3 (P5726 and P0044; Sigma-Aldrich), 5 mM DTT, and 1 µl/ml protease inhibitor cocktail (P8340; Sigma-Aldrich) and homogenized in a Mixer-Mill MM301. The homogenate was centrifuged for 4 min at 8,000 × g, and the supernatant was discarded. After centrifugation, the pellets were left on ice for 2 min. This step was repeated three times with CMF buffer without Triton X-100. The pellets were then resuspended in the CMF buffer, and the samples were subsequently mixed at a 1:1 ratio with Laemmli buffer and 5% β-mercaptoethanol and heated at 100°C for 5 min. Samples were loaded onto 12% or 4–20% gradient gels (Mini-Protein TGX; Bio-Rad), and phosphoproteins were detected using Pro-Q Diamond/Sypro system (Invitrogen, Thermo Fisher Scientific). Total protein was also detected using Coomassie staining as described previously (Kazmierczak et al., 2014). RLC and cMyBP-C phosphorylation was tested by Western blots and protein-specific antibodies. To detect phosphorylated RLC, ∼30 µg of protein per lane was loaded into 15% SDS-PAGE, and the membranes were probed with phospho-specific RLCmas antibody (gift from Dr. Neal Epstein, National Institutes of Health, Bethesda, MD), which recognizes the phosphorylated form of the RLC and does not react with nonphosphorylated RLC. Total RLC protein was detected with rabbit polyclonal CT-1 antibody, produced in this laboratory (Szczesna-Cordary et al., 2005), which served as a loading control. Phosphorylation of cMyBP-C was tested at Ser282. Approximately 20 µg of protein per lane was loaded into 12% SDS-PAGE, and the membranes were probed with phospho-specific cMyBP-Cser282 antibody (gift from Dr. Sakhthivel Sadayappan, University of Cincinnati, Cincinnati, OH). Goat polyclonal antibody was used to detect total cMyBP-C. Myofilament protein phosphorylation was calculated based on the density of bands of phosphorylated protein/total protein. The bands were analyzed using ImageJ software (National Institutes of Health).

Statistical analyses

All values are shown as mean ± SD. Differences between multiple groups were assessed using one-way ANOVA followed by Tukey’s multiple comparison test. To detect statistical differences between the genotypes that were sex-dependent, two-way ANOVA followed by Tukey’s or Sidak’s multiple comparison test was used with significance defined as P < 0.05 (GraphPad Prism v7.05).

Online supplemental material

In the supplemental text (at the end of the PDF), the Materials and methods section addresses experimental procedures regarding OXPHOS complexes, enzymatic activities, and mitochondrial respiration. The Results section provides assessment of calcium transients presented in Fig. S1 and Table S1; mitochondrial...
remodeling in HCM-A57G, RCM-E143K, and Δ43 versus WT-ELC mice (mitochondrial protein expression [Fig. S2], enzymatic activities [Fig. S3], and cellular respiration and oxygen consumption rates [OCRs; Fig. S4 and Table S2]).

Results

Energetic remodeling in HCM-A57G, RCM-E143K, and Δ43 ELC mice at the level of myofilaments

SRX state of myosin

The cardiac relaxation phase is critically important for normal heart function, and disruption of SRX ↔ DRX equilibrium has been postulated to increase the risk for inefficient heart performance that can lead to adverse clinical outcomes (Toepfer et al., 2020). Both the HCM-A57G and RCM-E143K ELC mutations have been shown to cause disease-specific abnormalities in patients (Lee et al., 2001; Olson et al., 2002; Caleshu et al., 2011) and in transgenic mice (Kazmierczak et al., 2013; Yuan et al., 2017), but little is understood about the mechanistic basis by which these mutations confer distinct in vivo cardiac function. To learn about the effect of ELC mutations on SRX ↔ DRX equilibrium and the proportion of myosin heads existing in the SRX (P2) versus DRX (P1) conformations, we performed ATP binding/exchange assays in skinned PM fibers from HCM-A57G and RCM-E143K mice. The effects due to these two pathological mutations were compared with WT-ELC mice and to Δ43 animals that manifest a time-dependent hypertrophic growth with no obvious abnormalities in heart function (Muthu et al., 2011). Fibers were incubated in a solution containing 250 µM mant-ATP, and decay fluorescence curves as a function of time were collected during a rapid mant-ATP exchange with 4 mM non-labeled ATP. Averaged fluorescence decay curves were fitted to a double-exponential equation (Yadav et al., 2019a; Sitbon et al., 2020):

\[ F = 1 - P1[1 - \exp(-t/T1)] - P2[1 - \exp(-t/T2)] \]

where P1 and P2 (in %) are the amplitudes of fast and slow phases of fluorescence intensity decay, indicating percentage of heads in the DRX versus SRX state, respectively. Averaged fluorescence decay curves and their simulated single-exponential traces for the fast and slow phases of fluorescence decay for fibers from all genotypes are shown in Fig. 1A, Fig. 1B and Table 1 compare the fluorescence intensity decay isotherms and depict the fraction of heads existing in either the DRX or SRX state, and Fig. 1C analyzes the effect of ELC mutations on SRX ↔ DRX equilibrium separately for male and female mice. To complete the comparison between all genotypes studied, previously reported SRX data for HCM-A57G and Δ43 mice (Sitbon et al., 2020) were included in the analysis of the newly acquired results on HCM-A57G and Δ43 mice as well as RCM-E143K and WT mice (Fig. 1 and Table 1).

The HCM-A57G mutation significantly reduced the SRX state of myosin and promoted the SRX-to-DRX transition (Fig. 1A and B), usually associated with greater ATP consumption (Alamo et al., 2016). The ratio of myosin heads existing in the DRX to SRX (P1:P2) states was 56:42 for WT-ELC and changed to 67:30 in HCM-A57G mice (Table 1). The RCM-E143K and Δ43 models promoted the SRX conformation and exhibited a significantly larger proportion of myosin heads occupying the SRX (P2) state compared with P2 observed in HCM-A57G mice (Fig. 1B and Table 1). Interestingly, male mice of all genotypes demonstrated a more pronounced phenotype compared with female counterparts (Fig. 1C), indicating sexual dimorphism in the effect of the ELC mutation on SRX ↔ DRX equilibrium. No changes between the genotypes were noted in T1 or T2 lifetimes (Table 1).

Overall, these results explain the opposing effects of two pathological ELC variants as the results of a disturbance in SRX ↔ DRX equilibrium and underscore the importance of studying the energetic states of myosin at the level of myofilaments. The HCM-A57G mutation promoted the SRX-to-DRX transition that mechanistically signifies the HCM clinical phenotype of hypercontractility, while the RCM-E143K mutation favored the energy-saving SRX state. Similar to RCM-E143K, the near-physiological Δ43 model also stabilized the sequestered energy-saving conformation. Larger SRX populations may be cardioprotective in normal Δ43 myocardium and during times of stress in cardiomyopathic RCM-E143K mice. The data on Δ43 mice favoring the SRX conformation suggests a likely involvement of the N-ELC (missing in Δ43 mice) in regulation of the SRX-to-DRX transition in full-length ELC mutant mice. The results also revealed sex dimorphism among the ELC models and the dominant phenotype of male mice.

Kinetics of force development

Our previous in vivo investigations demonstrated major differences in function between the pathological HCM-A57G and RCM-E143K models and suggested distinct mechanisms by which they influence cardiac function. In contrast to HCM-A57G (Kazmierczak et al., 2013), RCM-E143K animals exhibited decreased cardiac output and stroke work in vivo (Yuan et al., 2017). These differences most likely originated from different effects of mutations on myosin motor power, with A57G upregulating and E143K downregulating myosin power as observed in the in vitro motility assay (Wang et al., 2018). Interestingly, despite such different in vivo and in vitro behaviors due to these two ELC variants, the mutations resulted in a common phenotype of diastolic dysfunction in mice (Yuan et al., 2017; Sitbon et al., 2020) and humans (Lee et al., 2001; Olson et al., 2002; Caleshu et al., 2011).

To understand the origin of physiological differences between the HCM-A57G and RCM-E143K models, we compared their ability to produce force in electrically stimulated intact PMs from these mice. Unlike the RCM-E143K model that has not been characterized before, HCM-A57G and Δ43 were tested for their effects on the kinetics of the force and calcium transients during twitch contractions (Sitbon et al., 2020). To compare the effect of all ELC mutations on the kinetics of force/calcium transients and relate the results to the distribution of myosin energetic states in these models, the data previously reported for HCM-A57G and Δ43 mice (Sitbon et al., 2020) were merged with the new results on RCM-E143K animals and compared with sex- and age-matched WT controls (Fig. 2). We observed no differences in the effects of the two genotypes on maximal force (in millinewtons) or the peak force/cross-section area (in...
millinewtons per square millimeter; Fig. 2 A and Table 2). However, differences between the two pathogenic models were noted in force relaxation kinetics, with \( t_{25}, t_{50}, \) and \( t_{75} \) times (in milliseconds) at 25, 50, and 75% peak force to baseline being significantly longer in RCM-E143K compared with HCM-A57G and WT fibers (Fig. 2 B and Table 2). Interestingly, as in SRX state experiments, male mice presented with a more pronounced functional phenotype than female counterparts (Fig. 2 B). Despite differences in force transient kinetics, both pathological models, HCM-A57G and RCM-E143K, displayed delayed calcium transients compared with WT mice, indicating diastolic abnormalities in these mice (Fig. S1 and Table S1). The finding of longer versus shorter force transients observed for RCM-E143K and HCM-A57G models, respectively, are consistent with the SRX results showing slower cross-bridge kinetics and stabilization of the sequestered SRX conformation in RCM-E143K mice versus faster cross-bridge relaxation kinetics and larger DRX populations in the HCM-A57G mice.

### Protein phosphorylation in HCM-A57G, RCM-E143K, and Δ43 ELC mice

The structure of cardiac thick and thin filaments is known to undergo dynamic changes produced by posttranslational modifications such as altered phosphorylations of myosin RLC, cMyBP-C, or TnI that may lead to alterations in energy utilization and cardiac output (Moss and Fitzsimons, 2010). To test this idea, we assessed the phosphorylation status of myofilament proteins in the hearts of female and male WT, HCM-A57G, RCM-E143K, and Δ43 mice. The phosphorylation of myosin RLC, cMyBP-C, and TnI was determined in CMFs prepared from the hearts of mice that were run on SDS-PAGE and stained with ProQ/Coomassie (Fig. 3) or blotted with protein-specific antibodies (Fig. 4).

ProQ/Coomassie data revealed a significant upregulation of myosin RLC phosphorylation in HCM-A57G compared with the RLC of WT-ELC (\( P < 0.05 \)) and RCM-E143K (\( P < 0.001 \)) mice (Fig. 3). A significantly higher RLC phosphorylation level was also observed in Δ43 CMFs compared with RCM-E143K hearts.
No changes in cMyBP-C or TnI phosphorylation were noted in HCM-A57G, RCM-E143K, or Δ43 CMFs compared with WT-ELC (Fig. 3). A more quantitative analysis of RLC and MyBP-CSer282 phosphorylation assessed with Western blotting and protein-specific antibodies (Fig. 4) showed ~40% upregulation of myosin RLC phosphorylation in HCM-A57G versus WT-ELC mice (P < 0.01) and ~2.7-fold greater RLC phosphorylation in HCM-A57G versus RCM-E143K (P < 0.0001) CMFs. No changes in cMyBP-C phosphorylation seen by ProQ/Coomassie (Fig. 3) were further confirmed by Western blotting using cMyBP-CSer282 specific antibodies (Fig. 4). On the other hand, an approximately twofold lower level of myosin RLC phosphorylation was observed in RCM-E143K versus WT-ELC mice (P < 0.01) and an ~2.6-fold lower level when compared with the RLC of Δ43 (P < 0.0001) CMFs. Remarkably, upregulation and downregulation of myosin RLC phosphorylation seen in HCM-A57G and RCM-E143K hearts, respectively, were correlated with an increase in DRX populations in HCM-A57G mice and a larger SRX population in RCM-E143K animals (Fig. 1), further highlighting the consistency in phenotypic differences between the two pathological models.

Upregulation of RLC phosphorylation was also observed in Δ43 versus WT-ELC hearts (Fig. 4), which corresponded to larger SRX populations and presumably greater energy conservation in this model (Fig. 1). These data imply that the N-ELC that is missing in the hearts of Δ43 mice not only binds to actin during muscle contraction (Timson et al., 1998) but may also interact with myosin RLC and modulate its phosphorylation in

![Graph](image-url)

Table 1. The SRX state of myosin measured by mant-ATP/ATP chase assays in skinned PMs from the hearts of HCM-A57G, RCM-E143K, Δ43, and WT-ELC mice

| Parameter | WT     | HCM-A57G | Δ43   | RCM-E143K |
|-----------|--------|----------|-------|-----------|
| P1 (%)    | 55.8 ± 3.8 | 67.4 ± 8.4a | 48.5 ± 9.5b | 48.3 ± 8.4c |
| P2 (%)    | 41.5 ± 3.8 | 29.5 ± 8.4a | 49.4 ± 8.7b | 49.6 ± 7.4c |
| T1 (s)    | 8.2 ± 4.3  | 6.2 ± 3.4  | 4.9 ± 3.5  | 9.0 ± 5.2  |
| T2 (s)    | 182.8 ± 126.3 | 109.0 ± 71.5 | 197.3 ± 108.7 | 186.3 ± 123.9 |

Number of animals

|       | WT (5 M, 6 F) | HCM-A57G (4 M, 4 F) | Δ43 (5 M, 5 F) | RCM-E143K (4 M, 3 F) |

Data are mean ± SD. P1 and P2 represent the percentage of myosin cross-bridges in the DRX and SRX states, respectively; T1 and T2 depict their respective lifetimes. P values were calculated with one-way ANOVA with Tukey's multiple comparison test.

*P < 0.05 for mutant versus WT-ELC.

aP < 0.05 for Δ43 versus A57G.

bP < 0.05 for E143K versus A57G.

![Graph](image-url)

Figure 2. Force transients in intact PMs from 8–10-mo-old HCM-A57G (5 M and 3 F), Δ43 (3 M and 3 F), and RCM-E143K (3 M and 5 F) compared with WT-ELC (7 M and 4 F). The data for A57G and Δ43 animals were taken, with permission, from Sitbon et al. (2020). (A) Left: Normalized force transients for WT-ELC (blue), HCM-A57G (green), Δ43 (red), and RCM-E143K (black) mice. Right: Force (in millinewtons per square millimeter) was calculated as a ratio of peak force to physiological cross-sectional area (see Materials and methods) and plotted as a function of genotype and sex. (B) Time points t25, t50, and t75 depicting the time (in milliseconds) from peak force to 25%, 50%, and 75% baseline for male and female mice of all genotypes. Open symbols depict female mice, and closed symbols depict male mice. Data are mean ± SD of male or female mice analyzed by two-way ANOVA followed by Tukey’s or Sidak’s multiple comparison test. *, P < 0.05 for mutant versus WT; $, P < 0.05 for A57G versus E143K; and &, P < 0.05 for A57G versus Δ43.

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Transgenic HCM-A57G, RCM-E143K, and Δ43 ELC mice

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NDUFB8 (Fig. S2 B). Decreased expression of NDUFS4 may be indicative of a decreased ability of RCM-E143K to properly use NADH to fuel the mitochondrial electron transport chain, leading to less ATP production in RCM-E143K hearts, while increased expression of NDUFB8 in HCM-A57G myocardium may suggest some abnormalities in CI function due to HCM.

We also studied the expression of complex IV (CIV; cytochrome c oxidase [COX]), which contributes to the proton gradient across the inner mitochondrial membrane to drive ATP synthesis (Bourens and Barrientos, 2017). Except for small but significant alterations in Cox5b expression observed in female HCM-A57G mice (lower) versus female RCM-E143K animals (higher; Fig. S2 C), no differences in any of the CIV subunits were observed. Compared with HCM-A57G mice, the Δ43 ELC model of near-physiological hypertrophic remodeling showed a significant upregulation of hexokinase 2 (HK2), an enzyme involved in glucose utilization pathways (Fig. S2 B). Similar to alterations in CI/CIV expression, the differences in HK2 expression between HCM-A57G and Δ43 mice were also observed in females (Fig. S2 C).

We then investigated the enzymatic activities of the OXPHOS complexes, as any alterations in CI or CIV activities could be indicative of defective mitochondrial respiration, uncoupled proton gradient, and altered ATP production (Diaz, 2010). Spectrophotometric assessment of heart homogenates from HCM-A57G, RCM-E143K, and Δ43 relative to WT mice showed no changes of CI/CIV enzymatic activities in any of the investigated models (Fig. S3 A). Interestingly, analysis of sex differences in CI activity between the genotypes revealed that females of the pathological HCM-A57G model had significantly higher activity compared with Δ43 females (Fig. S3 B).

Finally, oxygen consumption profiles were determined in mitochondria isolated from HCM-A57G, RCM-E143K, and Δ43 hearts relative to WT hearts in the presence of specific substrates and inhibitors (Fig. S4 A). Determination of basal OCR or state II respiration using pyruvate and malate as substrates showed that OCR rates were similar among all genotypes (Fig. S4 B and Table S2), and no differences were noted between females and males of all genotypes (Fig. S4 C). No changes were observed in ADP-dependent state III respiration, state IVo and state IIIu respiration, or respiratory control ratio (RCR), calculated as state III/IIATP to state IVo ratio (Rogers et al., 2011), and delineating the mitochondrial quality control (Khalifa et al., 2017; Fig. S4). Overall, these data indicated that no mitochondrial respiratory dysfunction occurred in the hearts of investigated ELC models and that the hearts of mice were supplied with enough ATP to sustain their contractile function.

### Discussion

Cardiac muscle contraction depends on the ATP-driven cyclical interaction of myosin with thin filaments comprising actin and Tm-Tn (Grees and Holmes, 2005). The myosin head or crossbridge is the molecular motor of the heart and contains a motor domain with actin-activated MgATPase activity and the lever-arm region where both myosin ELC and RLC are attached to their respective myosin heavy chain IQ motifs (Rayment et al., 1993). The outstanding question in myosin ELC biology regards...
the function of the cardiac-specific 43-amino-acid-long N-ELC, comprising a lysine-rich actin-binding region that can bridge the ELC core of the myosin head with the actin filament (reviewed in Yadav et al., 2019b). Based on extensive research of myosin ELC function in the heart, we hypothesized that the cardiac N-ELC works as a molecular linker and/or energetic switch of the actin-myosin interaction that can regulate the SRX-to-DRX transition and the proportion of myosin heads that exist in either of these two energetically different states (Fig. 5).

In DRX, the myosin heads protrude into the interfilament space, but they are restricted from binding to actin, while in the SRX state, myosin heads are neatly ordered along the thick filament axis but cycle very slowly at a correspondingly low ATP turnover rate (Hooijman et al., 2011; McNamara et al., 2015). Because variations in the size of the myosin SRX population are considered central to modulating sarcomeric force production and energy utilization in cardiac muscle, we aimed to explore the mechanism by which the two pathological models of cardiomyopathy, HCM-A57G and RCM-E143K, as well as the structural Δ43 ELC mutation, affect the stability of the sequestered state and SRX→DRX equilibrium and whether these processes can be regulated by N-ELC and myosin RLC phosphorylation (Fig. 5).

Figure 3. Myosin RLC, cMyBP-C, and TnI phosphorylation in ventricular CMFs purified from the hearts of WT, HCM-A57G, RCM-E143K, and Δ43 mice. The hearts of 4 M and 3 F (7.5–10-mo-old) WT-ELC, 3 M and 5 F (8–10-mo-old) A57G, 3 M and 3 F (6–9-mo-old) E143K, and 4 M and 2 F (7.5–10-mo-old) Δ43 mice were used. (A) SDS-PAGE and sarcomeric protein phosphorylation monitored by ProQ/Coomassie. (B) Quantification of myosin RLC, cMyBP-C, and TnI phosphorylation in CMFs from all genotypes. Open symbols depict female mice and closed symbols depict male mice. Data are mean ± SD and were analyzed using one-way ANOVA with Tukey’s comparison test with significance depicted as *, P < 0.05 between mutant versus WT; $$$, P < 0.001 for A57G versus E143K; and &, P < 0.05 for E143K versus Δ43 mice. Abbreviations: +P-RLC, phosphorylated form of mouse ventricular RLC; +P-MyBP-C, phosphorylated form of mouse cMyBP-C; +P-TnI, phosphorylated form of TnI; F/A, F-actin; ELC mse, mouse ventricular essential light chain; ELC hu, human transgenic ELC; Δ43 ELC, N-terminally truncated ELC protein standard.

Myosin energetic states and cross-bridge kinetics in HCM-A57G, RCM-E143K, and Δ43 ELC mouse models
Mant-ATP chase assays revealed that the two models, HCM-A57G and RCM-E143K, exhibited antagonistic effects on SRX→DRX equilibrium, with HCM-A57G favoring the DRX state and RCM-E143K favoring the energy-saving SRX state (Fig. 1). Interestingly, fluorescence decay isotherms of RCM-E143K were similar to those produced by Δ43 animals, with a larger number of heads occupying the SRX state (P2) compared with the SRX (P2) of the HCM-A57G model (Table 1). The fact that the structural Δ43 ELC mutation that brings about the physiological-like hypertrophy phenotype in mice also promotes the formation of SRX suggests that the N-ELC, which is lacking in Δ43 mice, may be involved in regulating the myosin sequestered state and the SRX-to-DRX transition (Fig. 5). All models demonstrated sex dimorphism, showing greater effects and significant differences in myosin energetic states in male mice (Fig. 1 C).

Force production and energy use within the sarcomere are intricately regulated by the SRX mechanism (Schmid and Toepfer, 2021). We observed a strong correlation between myosin energetic states and the kinetics of force transients measured in electrically stimulated LV PMs from mice (Fig. 2). Faster
relaxation rates calculated in HCM-A57G fibers were related to the SRX-to-DRX transition, while the stabilization of the SRX state correlated with slower force transients measured in RCM-E143K samples compared with WT controls (Table 2). While both pathological variants demonstrated significant differences in comparison to WT, no changes in force transient kinetics were observed in Δ43 versus WT-ELC mice. Notably, in agreement with myosin SRX/DRX experiments, the kinetics of force transients were significantly different between male mice, indicating sexual dimorphism (Fig. 2 B). Considering their distinct clinical phenotypes, and despite showing differences in force transient kinetics, both models of ELC-related cardiomyopathy

Figure 4. Myosin RLC and cMyBP-CSer282 phosphorylation in myofibrils from heart ventricles of WT-ELC, HCM-A57G, RCM-E143K, and Δ43 mice detected by Western blotting. (A) SDS-PAGE and sarcomeric protein phosphorylation monitored by Western blots and RLC-specific (left) or cMyBP-C-specific (right) antibodies (Ab). Phosphorylated forms of the mouse RLC and cMyBP-C (upper panels) were detected with +P-RLC MSE or +P-MyBP-CSer282 antibodies, while total proteins were assessed with RLC tot (CT-1) or MyBP-C tot antibodies (lower panels). (B) Quantification of myosin RLC and cMyBP-CSer282 phosphorylation in CMFs purified from WT-ELC (4 M, 3 F), HCM-A57G (3 M, 5 F), RCM-E143K (3 M, 3 F), and Δ43 (4 M, 2 F) mice. Open symbols depict female mice and closed symbols depict male mice. Data are mean ± SD and were analyzed using one-way ANOVA with Tukey’s comparison test. Significance was depicted as *, P < 0.05 and **, P < 0.01 between mutant versus WT-ELC; $$$$, P < 0.0001 for HCM-A57G versus RCM-E143K; and &&&&, P < 0.0001 for RCM-E143K versus Δ43 mice.

Figure 5. Schematic representation of the antagonistic mechanism by which the HCM-linked A57G and RCM-associated E143K mutations in cardiac myosin ELC regulate the SRX state of myosin and myosin RLC phosphorylation. The HCM-A57G mutation promotes the SRX-to-DRX transition, and this process is coupled with a ~40% increase in myosin RLC phosphorylation compared with the level of RLC phosphorylation in WT-ELC myocardium. The RCM-E143K model favors the SRX conformation, and its RLC phosphorylation is approximately twofold less than in WT-ELC hearts, as judged by Western blotting and RLC-specific antibodies. Mechanistically, it is proposed that N-ELC, which is missing in the Δ43 model, regulates the SRX-to-DRX transition and contributes to sterically blocking (RCM-E143K) or facilitating (HCM-A57G) myosin RLC phosphorylation. Fig. 5 was created with https://BioRender.com.
displayed delayed calcium transients compared with WT-ELC mice (Fig. S1 and Table S1), suggesting the potential for diastolic dysfunction. Taken together, the A57G and E143K ELC mutations that were shown to cause human HCM (Lee et al., 2001) or RCM (Olson et al., 2002; Caleshu et al., 2011), respectively, presented clear differences in myosin energetic states and force transients kinetics (Figs. 1 and 2). They showed an upregulation (HCM-A57G) or downregulation (RCM-E143K) of myosin power in vitro (Wang et al., 2018) and exhibited opposite effects on cardiac output in vivo (Kazmierczak et al., 2013; Yuan et al., 2017).

Myosin RLC phosphorylation is upregulated in HCM-A57G and ΔA43 hearts and downregulated in RCM-E143K hearts

Myosin-based regulation of cardiac muscle is increasingly recognized as supplementing actin-based mechanisms in controlling the strength and time course of contraction (Brunello et al., 2020). The RLC-mediated modulation of myosin motor activity via myosin light chain kinase–dependent RLC phosphorylation has long been known to be crucial for normal heart function and was also shown to be perturbed in heart disease (Scruggs et al., 2009; Chang et al., 2015). The myocardium containing dephosphorylated myosin was shown to have limited ability to produce force and maintain cardiac function at physiological levels (Muthu et al., 2012; Yuan et al., 2015), suggesting myosin RLC phosphorylation as a promising target for the development of novel therapies (Yadav et al., 2019c). Structurally, RLC phosphorylation is proposed to control the myosin head conformation by disrupting a compact OFF conformation of myosin heads, which is equivalent to an energy-saving SRX state (Fig. 5). Disruption of the OFF state and promoting the DRX cross-bridge conformation is linked to potentiated contractility by increasing the population of myosin heads that are available for interaction with actin and force production in muscle (Jung et al., 2008; Kampourakis et al., 2016; Alamo et al., 2017).

Comparison of the results from the two cardiomyopathy models strongly suggests that RLC phosphorylation can regulate SRX ↔ DRX equilibrium in cardiac muscle and the proportion of force-producing myosin heads. Myosin RLC phosphorylation promoted the SRX-to-DRX redistribution in HCM-A57G myocardium, while the absence of RLC phosphorylation in RCM-E143K mice contributed to stabilization of the sequestered SRX state (Fig. 5). The interplay between the level of myosin RLC phosphorylation and the energetic states of myosin further explains the mechanisms of phenotypic differences between the HCM-A57G and RCM-E143K models. Upregulation of RLC phosphorylation was also observed in ΔA43 versus WT hearts and, as we showed before (Sitbon et al., 2020) and in this study, the ΔA43 myosin favored the energy-conserving SRX conformation. One can speculate that the N-ELC that is truncated in the hearts of ΔA43 mice can assume different conformations in the full-length ELC models to either promote or sterically hinder myosin RLC phosphorylation in HCM-A57G or RCM-E143K mice, respectively (Fig. 5).

The interrelation between myosin N-ELC and RLC phosphorylation has been addressed before: the Burghardt group, using single-molecule approaches and quantum dot motility assays elegantly demonstrated that N-ELC is involved in stimulus frequency modulation to affect power output (Wang et al., 2016) and that this process is highly regulated by RLC phosphorylation (Wang et al., 2014). After RLC phosphorylation, which increases at higher stimulus frequencies, an 8-nm step was the dominant myosin step size resulting in a significant gain in the average step size and leading to increased work produced per ATPase cycle (Wang et al., 2014). Of note, in our recent work, we determined that the deletion of N-ELC in ΔA43 mice increased the prevalence of 5-nm step sizes with coincidental loss of the 8-nm step as compared with native full-length ELC (Wang et al., 2016; Wang et al., 2018). Ensemble motility and single myosin mechanical characteristics were also consistent with the notion that HCM-A57G mutation impairs N-ELC–actin binding, while RCM-E143K mutation leads to weakening of myosin lever arm stability (Wang et al., 2018). High-resolution structural approaches (e.g., x-ray crystallography and in silico analysis; Robert-Paganin et al., 2018) coupled with time-resolved biophysical assessments (e.g., Guhathakurta et al., 2017) of the ELC and RLC conformational transitions associated with HCM-A57G and RCM-E143K mutations in myosin ELC ± RLC phosphorylation would provide a still greater understanding of the effects of these mutations on the prevalence of the sequestered state of myosin and how the mutations modify myosin power and the number of heads available for force production.

Mitochondrial energy supply is not affected by myosin ELC mutations

Cardiac function is highly dependent on ample energy supply, and the regulation of heart muscle contraction and relaxation relies heavily on mitochondria, which produce energy in the form of ~6 kg of ATP daily (Hall et al., 2014; Ritterhoff and Tian, 2017). Most of the cell’s ATP is produced via the OXPHOS system, comprising five multi-subunit complexes embedded in the mitochondrial inner membrane. Oxidative metabolism in mitochondria provides most of the energy consumed by the heart, and the inability to generate and/or transfer energy may trigger abnormalities in contractile function (Tian et al., 2019). Considering the clear differences between the ELC mutations in the myosin energetic states and force transient kinetics, it was somewhat surprising not to observe any significant alterations in the OXPHOS complexes or mitochondrial respiration.

In summary, our study revealed that under normal conditions, the overall respiratory function assessed in all investigated ELC models was normal, and the hearts of HCM-A57G, RCM-E143K, and ΔA43 mice were supplied with enough ATP to sustain their contractile function. However, the minute differences in CI subunit expression noted between the two pathological models (Fig. S2) may become more evident under stress conditions (Garnier et al., 2003; Dai et al., 2012) that may exacerbate mutation-dependent deficits in energy transfer and mitochondrial function.

Significance and innovation

Our study contributes new information on myosin energetic states associated with hypertrophic (A57G) and restrictive (E143K) cardiomyopathy phenotypes and provides insight into the interplay between myosin SRX and DRX conformations and the phosphorylation of myosin RLC (Fig. 5). The hypercontractility of
that results from the SRX-to-DRX transition in the HCM-A57G myocardium is also associated with increased phosphorylation of the myosin RLC, while stabilization of the SRX energy-conserving state of cross-bridges in RCM-E143K hearts is related to the downregulation of myosin RLC phosphorylation. We observed a strong correlation between the increased or decreased degree of RLC phosphorylation in HCM-A57G and RCM-E143K mice, respectively, with their enhanced or depressed in vivo function monitored by echocardiography (Kazmierczak et al., 2013; Yuan et al., 2017) and upregulation or downregulation of myosin power in the in vitro motility assay (Wang et al., 2018).

We propose a new regulatory function for N-ELC, which is missing in Δ43 mice, as being involved in the modulation of myosin energetic states and myosin RLC phosphorylation in full-length ELC mutant mice (Fig. 5). The N-ELC is presumed to adopt different conformations relative to myosin heavy chain and myosin RLC to either facilitate (HCM-A57G) or sterically block (RCM-E143K) myosin RLC phosphorylation and is considered a key regulator of SRX→DRX equilibrium in these two pathogenic mouse models of cardiomyopathy (Fig. 5). Taken together, our findings show that both mechanisms, RLC phosphorylation and N-ELC, could be crucial targets in therapeutic interventions for ELC-associated cardiomyopathy.

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**Supplemental material**

**Materials and methods**

**OXPHOS complexes: immunoblotting of metabolic proteins**

Heart homogenates from each group of mice were rinsed with cold PBS, homogenized for 1-2 min at 30 Hz in a Mixer-mill MM301 (Retsch USA) in buffer containing 75 mM sucrose, 0.22 mM sorbitol, 1 mM EGTA, 10 mM HCl, pH 7.4, and 10 μl/ml protease inhibitor cocktail (P8340; Sigma-Aldrich), prepared for SDS-PAGE using Laemmli buffer, and heated for 20 min at 67°C. 5 mg of protein was loaded on 4-20% Tris-glycine polyacrylamide precast Stain-Free Gels (456-8096; Bio-Rad) and transferred to polyvinylidene fluoride membrane using the Turbo blot system from Bio-Rad. Membranes were blocked in 5% milk in PBST and blotted with specific primary antibodies and then with horseradish peroxidase-conjugated secondary antibodies (Cell Signaling). The signal was detected using chemiluminescence using ECL Plus reagent (GE Healthcare) and captured in the ChemiDoc imaging system (Bio-Rad). Optical densities were quantified using ImageJ software. The following antibodies were used: peroxisome proliferator-activated receptor-γ coactivator 1-α (PGC-1α, ab54481; Abcam); NADH-ubiquinone oxidoreductase subunit S4 (NDUFS4, MS104; Mitoscience), subunit B8 (NDUFB8, ab10242; Abcam), and subunit A9 (NDUFA9, ab14713; Abcam); COX subunit 1 (Cox1, ab14705; Abcam) and subunit 5b (Cox5b, ab14705; Abcam); hexokinase isoform 1 (HK1, 2867; Cell Signaling) and isoform 2 (HK2, 2024; Cell Signaling); and carnitine palmitoyltransferase 1A (CPT1A, ab176320; Abcam). All primary antibodies were used at a 1:1,000 dilution and secondary antibodies at a 1:2,000 dilution. The signal intensity of the immunoreactive bands was calculated using ImageJ, and values were normalized to total protein and relative to WT and visualized by stain-free technology (Bio-Rad).

**Assessment of enzymatic activities**

The activities of CI, CIV, and citrate synthase (CS) were measured spectrophotometrically using heart homogenates from WT, HCM-A57G, RCM-E143K, and Δ43 mice as described previously, with some modifications (Barrientos et al., 2009; Spinazzi et al., 2012; Iommarini et al., 2020). Briefly, CI activity was measured by the oxidation of NADH at 340 nm over time in a plate reader (Sinergy 1; A57G, RCM-E143K, and Δ43G mice as described previously, with some modifications (Barrientos et al., 2009; Spinazzi et al., 2012; Iommarini et al., 2020). Transgenic HCM-A57G, RCM-E143K, and Δ43 ELC mice https://doi.org/10.1085/jgp.202012801)

**Mitochondria isolation**

Mitochondria were isolated from mouse hearts as described in Bharadwaj et al. (2015), with some modifications. After euthanasia, hearts were harvested and washed three times with DPBS to remove excess blood. Hearts were incubated for 5-10 min with a solution of pCa8 + 30 mM BDM on ice, muscles were chopped and incubated in 1.5 ml of Chappel-Perry buffer I (CPI; 100 mM KCl, 50 mM MOPS, 1 mM EDTA, 5 mM MgSO4, and 1 mM ATP adjusted at pH 7.4) + 0.2 mg of subtilisin A (P5380; Sigma-Aldrich) in a final volume of 200 μl (Barrientos et al., 2009; Iommarini et al., 2020). Enzymatic activities of CI, CIV, and CS were calculated as specific activity in millimoles per minute per milligram of protein and presented relative to WT.

**Seahorse XFp-based mitochondrial respiration**

We assessed OCR in isolated mitochondria using the Seahorse XFp analyzer (Agilent). Briefly, isolated mitochondrial fractions from heart tissue (~2 μg per well) were attached to the bottom of an XFp Cell Culture Microplate (S7802A; Agilent; 8 wells) by
centrifuging the plate at 2,000 × g for 20 min at 4°C. Mitochondria were viewed under a microscope at 20× magnification to ensure consistent adherence to the bottom of the wells. CI-driven respiration was initiated with 2 mM malate (46940-U; Sigma-Aldrich) and 10 mM pyruvate (P2256; Sigma-Aldrich) dissolved in MAS 1× buffer and pH adjusted to 7.4 at 37°C. Respiration was measured in picomoles of O₂ per minute per micromgram of mitochondria in the presence of CI substrates (state II or basal respiration), followed by the addition of 2 mM ADP (A2754; Sigma-Aldrich) leading to mitochondrial respiration state IIIADP. The addition of 10 µM oligomycin (O4876; Sigma-Aldrich), a complex V inhibitor, was used to induce mitochondrial respiration state IVo, followed by the addition of 10 µM carbonyl cyanide-p-trifluoromethoxyphenyl-hydrazone (FCCP; C2920; Sigma-Aldrich), a mitochondrial uncoupler, to promote maximal respiration (state IIIu). Finally, 5 µM antimycin A (A8674; Sigma-Aldrich) and 5 µM rotenone (R8875; Sigma-Aldrich), inhibitors of CIII and CI, respectively, were added to shut down respiration (state IV) as described in Rogers et al. (2011). Each drug was loaded in the corresponding port, and the cartridge was placed in the instrument to allow for proper calibration and ejection of each drug at specific times. All reagents listed above were dissolved in MAS 1× buffer. The RCR was calculated using the ratio between state IIIADP and state IVo (Rogers et al., 2011).

Results

Calcium transients in intact PM fibers
Simultaneously with the force transients, we also measured the rates of [Ca²⁺] transients in PM fibers from RCM-E143K versus WT-ELC mice using Ca²⁺ indicator, Fura 2-AM (Fig. S1 and Table S1). The data for calcium transients in HCM-A57G and Δ43 animals were taken, with permission, from Sitbon et al. (2020) and plotted together with RCM-E143K and WT (Fig. S1). Compared with WT-ELC fibers, the [Ca²⁺] transients were significantly delayed in the RCM-E143K model as manifested by increased relaxation times (in milliseconds) of t₅₀, t₅₀, and t₇₅ (Table S1). As reported earlier (Sitbon et al., 2020), increased t₅₀ was also observed in HCM-A57G versus WT-ELC mice (Table S1), indicating that both pathological phenotypes share diastolic abnormalities. In conclusion, delayed relaxation times of force (Fig. 2 and Table 2) and calcium (Fig. S1 and Table S1) transients in PMs from RCM-E143K versus WT-ELC hearts were indicative of diastolic dysfunction, the phenotype associated with the E143K mutation in humans (Olson et al., 2002; Caleshu et al., 2011) and observed in RCM-E143K mice (Yuan et al., 2017).

Mitochondrial remodeling

Mitochondrial protein expression CI (NADH:ubiquinone oxidoreductase) is a key membrane protein complex in mitochondria that catalyzes the oxidation of NADH in the mitochondrial matrix, and any alterations in CI subunits may cause a wide variety of defects in mitochondrial respiration (Kang et al., 2018). Analysis of Western blots shown in Fig. S2 A revealed that the RCM-E143K mutant decreased the expression of CI-NDUFS4 subunit compared with WT-ELC (Fig. S2 B), and this deficiency might be indicative of a decreased ability of the RCMI43K model to properly use NADH and resulting in less ATP production. Interestingly, the significant difference in CI-NDUFS4 was observed in female RCM-E143K versus female WT mice (Fig. S2 C). RCM-E143K female mice also showed a significant decrease in NDUFS4 expression compared with female HCM-A57G mice (Fig. S2 C), further highlighting the phenotypic differences between the two pathological models. The HCM-A57G model also showed increased expression of the beta subunit of NADH dehydrogenase, NDUFB8, compared with WT-ELC mice (Fig. S2 B), and this change could be indicative of HCM-associated abnormalities in CI function. However, the differences between HCM-A57G and WT mice were not significant in either female or male mice (Fig. S2 C), diminishing the importance of these small changes observed in NDUFB8 expression in HCM-A57G versus WT hearts (Fig. S2 B).

COX or CIV is a copper-heme oxidase that couples transfer of electrons from cytochrome c to oxygen with proton extrusion across the inner membrane that contributes to the proton gradient across the membrane to drive ATP synthesis (Bourrens and Barrientos, 2017). As Cox5b is an integral part of CIV and is required for the stability and assembly of COX, any alterations in its expression can modulate CIV activity (Galati et al., 2009). However, the difference in Cox5b expression between the two pathological models was observed only in female HCM-A57G (lower) versus RCM-E143K (higher) mice (Fig. S2 C).

Compared with HCM-A57G mice, the Δ43 ELC model showed a significant upregulation of hexokinase 2 (HK2), an enzyme involved in glucose utilization pathways (Fig. S2 B). However, the difference between the genotypes was significant only in female mice (Fig. S2 C). There was also a significant difference in HK2 between female HCM-A57G (lower) and RCM-E143K (higher) models (Fig. S2 C). As HK2 catalyzes the phosphorylation of glucose to glucose-6-phosphate in the heart, its higher expression in Δ43 versus HCM-A57G (Fig. S2 B) may be suggestive of higher reliance on glucose for energy provision in Δ43 versus HCM-A57G hearts (Ritterhoff and Tian, 2017). Overexpression of HK2 has also been found to confer cellular protection against H₂O₂ in cardiomyocytes and prevent maladaptive cardiac hypertrophy in vivo (Roberts and Miyamoto, 2015). It is therefore intuitive to see the upregulation of HK2 in hypertrophied hearts of the near-physiological Δ43 model compared with the pathological HCM-A57G phenotype. Finally, analysis of PGC-1α, considered a “master regulator” of mitochondrial biogenesis (Scarpulla, 2008; Rowe et al., 2010), showed no changes in any of the ELC mice (Fig. S2 B).
**Enzymatic activities** Oxidative metabolism in mitochondria provides most of the energy consumed by the heart, and the inability to generate and transfer energy has been considered to underlie abnormal contractile function (Tian et al., 2019). Any alterations in CI activity may cause a wide variety of defects in mitochondrial respiration, and alterations in CIV activity may lead to uncoupled proton gradient and altered ATP production (Diaz, 2010). Spectrophotometric assessment of enzymatic activities of CI and CIV in heart homogenates from HCM-A57G, Δ43, and RCM-E143K mice relative to WT mice is presented in Fig. S3. As CS is considered a marker of mitochondrial mass (Spinazzi et al., 2012; O’Brien et al., 2017), the activities of CI and CIV were normalized to CS and expressed relative to WT-ELC samples (Fig. S3 A). Overall, no effects on CI or CIV activities were observed for any of the investigated models. Despite the significant decrease in the expression of NDUF5 subunit of CI by the RCM-E143K model compared with WT mice (Fig. S2 B), no alterations in CI or CS/CS activities were monitored (Fig. S3 A). Interestingly, analysis of sex differences in CI activity between the genotypes revealed that female HCM-A57G mice had significantly higher activity compared with female Δ43 mice (Fig. S3 B). As for CI, the activity of CIV was not altered by any investigated ELC mutations compared with WT controls, and no effect of sex on CI and CIV activities was observed for any of the groups (Fig. S3 A).

**Cellular respiration and OCR** We then examined the CI-driven mitochondrial respiration in all ELC models compared with WT-ELC mice (Fig. S4), using heart-isolated mitochondria as described previously (Bharadwaj et al., 2015). Mitochondrial respiration or OCRs were recorded as a function of [O2] versus time and expressed as picomoles of O2 per minute per microgram of mitochondria. Fig. S4 A illustrates average traces of oxygen consumption profiles in WT, HCM-A57G, Δ43, and RCM-E143K hearts assessed in the presence of specific substrates and inhibitors. Determination of basal OCR or state II respiration using pyruvate and malate as substrates showed that OCRs were similar among all genotypes (Fig. S4 A and Table S2), and no differences were noted between female and male mice (Fig. S4 B). The same was true for state III respiration, defined as ADP-dependent respiration (IIIADP), which is controlled by the activity of ATP turnover and substrate oxidation rates (Brand and Nicholls, 2011). When challenged with oligomycin inhibiting complex V and preventing the conversion of ADP to ATP, no changes in state IVo respiration among all genotypes and/or sexes were observed (Fig. S4 B). The maximal respiration induced by FCCP, a mitochondrial uncoupler, showed no changes in state IIIu respiration, indicating no dysfunction in respiratory chain components, substrate translocases, or dehydrogenases. When mitochondrial respiration was inhibited with a combination of antimycin and rotenone (state IV), a decrease in changes in state IIIu respiration, indicating no dysfunction in respiratory chain components, substrate translocases, or dehydrogenases. When mitochondrial respiration was inhibited with a combination of antimycin and rotenone (state IV), a decrease in

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**Figure S1.** Calcium transients in intact PMs from HCM-A57G, RCM-E143K, and Δ43 versus WT-ELC hearts. Data for HCM-A57G and Δ43 were taken, with permission, from Sitbon et al. (2020). (A) Normalized calcium transients for HCM-A57G (green), RCM-E143K (black), Δ43 (red), and WT (blue) mice. (B) Time points t25, t50, and t75 depicting the time (in milliseconds) from peak [Ca2+] to 25%, 50%, and 75% baseline for all four genotypes. Open symbols depict female mice and closed symbols depict male mice. The hearts of n = 8 (5 M, 3 F) HCM-A57G, n = 6 (3 M, 3 F) Δ43, n = 8 (3 M, 5 F) RCM-E143K, and n = 15 (10 M, 5 F) WT-ELC mice were tested. Values are mean ± SD and were analyzed by one-way ANOVA with Tukey’s multiple comparisons test. Significance was depicted as *P < 0.05 for mutant versus WT-ELC; and &, P < 0.05 for RCM-E143K versus Δ43. Note delayed relaxation times for HCM-A57G (t25) and RCM-E143K (t25, t50, and t75) compared with WT-ELC mice.

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Sitbon et al., 2020; O’Brien et al., 2017; Spinazzi et al., 2012; Diaz, 2010; Brand and Nicholls, 2011; Spinazzi et al., 2012; Tian et al., 2019; Sitbon et al. (2020).
Figure S2. Assessment of mitochondrial protein expression in heart homogenates from 7–11-mo-old HCM-A57G, Δ43, and RCM-E143K relative to WT-ELC mice. (A) Representative Western blots of mitochondrial and metabolism proteins in heart homogenates of mice. (B and C) Expression of CI, CIV, glycolysis, fatty acid subunits (HK1, HK2, and CPT1A), and peroxisome proliferator-activated receptor γ coactivator 1-α (PGC-1α; PPAR-γ coactivator and master regulator of mitochondrial biogenesis) for all four genotypes (B), and separately for female versus male mice (C). Values are mean ± SD of n = 8 (5 M, 3 F) WT-ELC; n = 8 (4 M, 4 F) HCM-A57G; n = 8 (4 M, 4 F) Δ43; and n = 8 (4 M, 4 F) RCM-E143K animals with significance depicted as *, P < 0.05 for mutant versus WT or between genders. $, P < 0.05 for HCM-A57G versus RCM-E143K; and &, P < 0.05 between HCM-A57G versus Δ43 analyzed by one-way ANOVA with Tukey’s multiple comparison test (B) and two-way ANOVA with two nominal variables (genotype, sex) followed by Tukey’s multiple comparison test (C). Optical density was assessed with ImageJ. Each protein band was normalized to total protein content assessed for each blot before the primary antibody was applied. HK, hexokinase (glucose metabolism); NDUFS4, NDUFA9, and NDUFB8, CI subunits (NADH dehydrogenase); Cox5b and Cox1, CIV subunits; CPT1A, carnitine palmitoyltransferase 1A (fatty acid metabolism).

Figure S3. Enzymatic activities in the hearts of ELC mice. (A and B) Spectrophotometric assessment of enzymatic activities in heart homogenates from HCM-A57G (3 M, 3 F), Δ43 (3 M, 3 F), and RCM-E143K (4 M, 2 F) relative to WT (3 M, 3 F) mice (A), and separated by sex (B). Values are mean ± SD and analyzed using two-way ANOVA with Tukey’s or Sidak’s multiple comparisons test with significance depicted as &, P < 0.05 between HCM-A57G versus Δ43. Open symbols depict female mice and closed symbols depict male mice.
Provided online are two tables. Table S1 shows calcium transients in intact PMs from RCM-E143K versus WT mice. Table S2 shows CI-driven mitochondrial respiratory states in isolated mitochondria from the hearts of WT-ELC, HCM-A57G, RCM-E143K, and Δ43 mice using Seahorse XFp analyzer.