Hypertrophic and Dilated Cardiomyopathy-Associated Troponin T Mutations R130C and ΔK210 Oppositely Affect Length-Dependent Calcium Sensitivity of Force Generation

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Length-dependent activation of calcium-dependent myocardial force generation provides the basis for the Frank-Starling mechanism. To directly compare the effects of mutations associated with hypertrophic cardiomyopathy and dilated cardiomyopathy, the native troponin complex in skinned trabecular fibers of guinea pigs was exchanged with recombinant heterotrimeric, human, cardiac troponin complexes containing different human cardiac troponin T subunits (hcTnT): hypertrophic cardiomyopathy-associated hcTnT\textsuperscript{R130C}, dilated cardiomyopathy-associated hcTnT\textsuperscript{ΔK210} or the wild type hcTnT (hcTnT\textsuperscript{WT}) serving as control. Force-calcium relations of exchanged fibers were explored at short fiber length defined as 110% of slack length (L\textsubscript{0}) and long fiber length defined as 125% of L\textsubscript{0} (1.25 L\textsubscript{0}). At short fiber length (1.1 L\textsubscript{0}), calcium sensitivity of force generation expressed by \(−\log [Ca^{2+}]\) required for half-maximum force generation (pCa\textsubscript{50}) was highest for the hypertrophic cardiomyopathy-associated mutation R130C (5.657 ± 0.019), intermediate for the wild type control (5.580 ± 0.028) and lowest for the dilated cardiomyopathy-associated mutation ΔK210 (5.325 ± 0.038). Lengthening fibers from 1.1 L\textsubscript{0} to 1.25 L\textsubscript{0} increased calcium sensitivity in fibers containing hcTnT\textsuperscript{R130C} (delta-pCa\textsubscript{50} = +0.030 ± 0.010), did not alter calcium sensitivity in the wild type control (delta-pCa\textsubscript{50} = −0.001 ± 0.010), and decreased calcium sensitivity in fibers containing hcTnT\textsuperscript{ΔK210} (delta-pCa\textsubscript{50} = −0.034 ± 0.013). Length-dependent activation indicated by the delta-pCa\textsubscript{50} was highly significantly (\(P < 0.001\)) different between the two mutations. We hypothesize that primary effects of mutations on length-dependent activation contribute to the development of the diverging phenotypes in hypertrophic and dilated cardiomyopathy.

Keywords: contractility, length dependent activation, thin filament regulation, cardiomyopathy, sarcomere length, force generation, troponin, calcium sensitivity
INTRODUCTION

The Frank-Starling law describes the intrinsic ability of the heart ventricle to adapt the systolic stroke volume to the previous diastolic filling. One main reason for this ability is the increased calcium sensitivity of the stretched myocardium, reviewed in Hanft et al. (2008), de Tombe et al. (2010) and Ait Mou et al. (2015) and also termed length-dependent activation (LDA). LDA has been attributed to different mechanisms intrinsic to the sarcomere, involving changes in filament lattice spacing (Hanft et al., 2008), stretch-dependent Ca$^{2+}$ regulation of troponin (Arteaga et al., 2000; Konhilas et al., 2003; Korte et al., 2012; Zhang et al., 2017), ordering of myosin head orientation (Farman et al., 2011), strain-sensing in titin (Millman and Irving, 1988; Ait Mou et al., 2015; Ait-Mou et al., 2016; Linke, 2018), and the communication between these mechanisms (Ait-Mou et al., 2016; Zhang et al., 2017).

Mutations in proteins of the sarcomere are associated with hypertrophic cardiomyopathy (HCM) and dilated cardiomyopathy (DCM). HCM and DCM mutations can occur in the same protein like in human cardiac troponin T (hcTnT) (Kamisago et al., 2000; Marston and Hodgkinson, 2001; Song et al., 2005; Lu et al., 2013). Cardiac troponin T (cTnT) transfers the Ca$^{2+}$ to the troponin C (cTnC) to allow the cardiac sarcomere, involving changes in filament lattice spacing (Hanft et al., 2008), stretch-dependent Ca$^{2+}$ regulation of troponin (Arteaga et al., 2000; Konhilas et al., 2003; Korte et al., 2012; Zhang et al., 2017), ordering of myosin head orientation (Farman et al., 2011), strain-sensing in titin (Millman and Irving, 1988; Ait Mou et al., 2015; Ait-Mou et al., 2016; Linke, 2018), and the communication between these mechanisms (Ait-Mou et al., 2016; Zhang et al., 2017).

MATERIALS AND METHODS

Skinned Fiber Preparation and Mechanical Setup

Skinned fibers were dissected from the left ventricular papillary muscles of the guinea pig (Stehle et al., 2002) and stored for up to 60 h at 0°C in skinning solution containing 5 mM KH$_2$PO$_4$, 5 mM Na-acetate, 3 mM magnesium acetate, 5 mM K$_2$EGTA, 3 mM Na$_2$MgATP, 47 mM sodium creatine phosphate, 2 mM dithiothreitol (DTT), 0.2 mM 4-(2-aminoethyl) benzenesulfonyl-fluoride (AEBSF), 10 µM leupeptin, 10 µM antipain, 5 mg/l aprotinin. Skinned fibers were mounted in skinning solution in the mechanical setup between a force transducer (KG7A) with bridge-amplifier DUBAM 7C (Scientific Instruments, Heidelberg, Germany) and a fixed clamp. After mounting, fibers were stretched by 10% of their slack length $L_0$ to $1.1 L_0$.

Troponin Exchange

The three subunits of hcTn, i.e., hcTnC, hcTnT, and hcTnI were separately expressed in Escherichia coli and isolated as described previously (Kruiger et al., 2003). For the exchange of the endogenous guinea pig cTn for the exogenous recombinant human cTn, the fibers were incubated in the mechanical setup at 10°C for 15 min in exchange buffer (in mmol/L): 132 NaCl, 5 KCl, 1 MgCl$_2$, 10 Tris, 5 EGTA, 1 NaN$_3$, pH 7.1 (20°C) followed by incubation in the same buffer containing in addition 3 mg/ml hcTn for 180 min at 20°C (Neulen et al., 2007).

The exchange of the endogenous guinea pig cTn (gcTn) for the exogenous hcTn was probed by 12.5% SDS-PAGE and visualizing proteins by Commassie-R250-staining (Solzin et al., 2007). Guinea pig cTn (gcTn) contains one more amino acid and migrates less than hcTnI on the gel (Supplementary Figure S1 in Supplementary Material). Exchange efficiency was defined by the ratio of hcTnI intensity per total intensity of hcTnI and gcTnI and quantified using Phoretix-1 as illustrated in Supplementary Figure S1.

Force-pCa Relations

Force-pCa relations were measured using mixtures of Ca$^{2+}$ buffered activating and relaxing solutions containing 3 mM (CaCl$_2$)$_2$K$_2$EGTA (activating solution, pCa 4.7) or 3 mM K$_4$Cl$_2$EGTA (relaxing solution, pCa 7), 10 mM imidazole, 10 mM Na$_2$MgATP, 3 mM MgCl$_2$, 32.7 mM sodium creatine phosphate, 2 mM DTT, pH 7.0, µ = 178 mM. To ensure saturation of free Ca$^{2+}$ concentration at all conditions, an extra activation solution (pCa 4.28) was prepared by adding 3 mL 60 mM CaCl$_2$ per 100 mL activating solution. Experimental temperature was 10°C.
Force–pCa relations were fitted by sigmoidal Hill equation: 
\[ F_{\text{norm}} = 1 + 10^{\frac{\text{pCa}_{50} - \text{pCa}}{n_H}} \]
where \( F_{\text{norm}} \) is the force at pCa, \( \text{pCa} = -\log ([\text{Ca}^{2+}]/M \text{ normalized to maximum force at pCa}) \). 4.28, \( \text{pCa}_{50} \) is the pCa at which \( F_{\text{norm}} = 0.5 \), and \( n_H \) is the Hill coefficient indicating the slope of the force–pCa relation.

### Statistical Analysis

Two-way repeated measures analysis of variance (Two-way RM ANOVA) was performed under GraphPad Prism 4 to test the effects of two factors, the effect of hcTnT-type (hcTnTR130C, hcTnWT, and hcTnTAK210), and the effect of fiber length (1.1 \( L_0 \) and 1.25 \( L_0 \)) on each analyzed parameter. Data was subject-matched (fiber-matched) for analyzing the effect of fiber length on parameters that were measured in each individual fiber first at 1.1 \( L_0 \) and then at 1.25 \( L_0 \). Subject matching was highly indicated by \( P < 0.0001 \) for each parameter. Significant hcTnT-type-fiber length interaction (\( P < 0.05 \)) in the two-way RM ANOVA indicated dissimilar change of the parameter among the three hcTnT-types. To probe the cause for significant interaction, post hoc analysis was performed using Tukey’s multiple comparison test yielding the \( P \)-values indicated in the results by * for \( P < 0.05 \), ** for \( P < 0.01 \), and *** for \( P < 0.001 \). When length affected the parameter with no significant interaction, the significance for length changing the parameter indicated by the subject-matched delta values of the parameter being significantly different from zero was analyzed by Bonferroni post-tests and indicated by # for \( P < 0.05 \), ## for \( P < 0.01 \), and ### for \( P < 0.001 \). All parameter values are given as mean ± SEM (standard error of the mean) of \( n \) fibers exchanged for each hcTnT-type.

### RESULTS

#### Control of Troponin Exchange

The endogenous troponin complex in the left ventricular skinned fibers from guinea pig was exchanged by exogenous recombinant human cardiac heterotrimeric troponin complex (hcTn) containing the hcTnC and hcTn wild type subunits and either hcTnWT, hcTnR130C or hcTnTAK210. The exchange in the fibers of the exogenous hcTn complexes for the endogenous cTn was tested by preparing three samples for each type of recombinant hcTn exchange. Each sample contained two fibers that were subjected to the exchange protocol in the chamber of the mechanical setup under the same conditions as performed for the mechanical measurements and each sample was then quantified for the relative amounts of endogenous and exogenous hcTn (Supplementary Figure S1). The exchange efficiency defined by the ratio of hcTn per total cTn (sum of endogenous gcTnI and exogenous hcTnI) was 46 ± 2% for the exchange done with hcTn containing the hcTnWT, 44 ± 1% for the one containing the hcTnR130C and 48 ± 2% for the one containing the hcTnTAK210 (mean ± SEM of each \( n = 3 \)). There were no significant differences in the efficiencies for the three exchanges. Similar exchange efficiencies have been reported in a previous in vitro study of the hcTnTAK210 mutation using exchange of recombinant hcTn for endogenous cTn in permeabilized rabbit cardiac muscle fibers (Morimoto et al., 2002).

#### Biomechanical Measurements

For the force measurements, fibers were prepared from 7 guinea pig hearts. 18 fibers were exchanged for hcTn containing HCM-associated hcTnTR130C, 19 fibers contained the hcTnWT control and 19 fibers contained the DCM-associated hcTnTAK210. Figure 1A shows the resting tension (\( F_{\text{REST}} \)) of fibers exchanged with the three different hcTnT at short fiber length (110% of slack length, 1.1 \( L_0 \)) and after stretching them to long fiber length (125% of slack length, 1.25 \( L_0 \)). Although the statistical analysis by two-way RM ANOVA indicated strong significant increase of \( F_{\text{PASS}} \) by stretch (\( P < 0.0001 \)) for each hcTnT-type by Bonferroni post-test (see ### in Figure 1A), there was no significant interaction (\( P = 0.37 \)) between the effects of fiber length and hcTnT-type on \( F_{\text{PASS}} \) (Figure 1A and Table 1). No interaction indicates similar passive mechanical properties of fibers containing the three different types of hcTnT. Similarly to passive tension, two-way RM ANOVA revealed no interaction between the effects of fiber length and hcTnT-type on the maximum tension (\( F_{\text{MAX}} \)) during contraction (Figure 1B and Table 1). \( F_{\text{MAX}} \) was strongly significantly increased (\( P < 0.0001 \)) by stretching the fibers. Bonferroni post-tests confirmed the significant effect of stretch on \( F_{\text{MAX}} \) for each hcTnT-type. The results for \( F_{\text{MAX}} \) indicate that the mutations do not alter the maximum force-generating capacity or its length dependence. The values of \( F_{\text{PASS}} \) and \( F_{\text{MAX}} \) are summarized in Supplementary Table S1 (see Supplementary Material) and their statistical analysis in Table 2.

Along with resting tension and maximum tension, the full force–pCa relations were determined before and after lengthening fibers from 1.1 \( L_0 \) to 1.25 \( L_0 \). Figure 1C illustrates the average force–pCa relations of the three groups of different hcTn-exchanged fibers at short (1.1 \( L_0 \), circles) and long fiber length (1.25 \( L_0 \), triangles). At short fiber length, the relation of the hcTnTR130C-exchanged fibers is shifted to the left compared to the relation of the hcTnTWT control, i.e., to higher pCa values or lower [Ca2+]0. This leftward shift is slightly enhanced upon lengthening the fibers from 1.1 \( L_0 \) to 1.25 \( L_0 \). Opposite to the HCM mutation, the relation of the fibers containing the DCM-associated hcTnTAK210 is slightly shifted to the right compared to the relation of the control fibers containing the hcTnTWT (Figure 1C). The basal calcium desensitization by the DCM mutation observed at short fiber length is further enhanced by lengthening the fibers to 1.25 \( L_0 \).

To test for statistical significant differences in calcium-dependent force generation among the three types of hcTn exchange and among the two lengths, the force–pCa relation plotted for each individual fiber at each length was fitted by the sigmoidal Hill function (see section “Materials and Methods”) to quantify the mean and variation of pCa50 and \( n_H \) as indicators for calcium sensitivity and cooperativity of the calcium-dependent force generation, respectively. Two-way RM ANOVA revealed a highly significant effect (\( P < 0.0001 \)) of the hcTnT-type on the pCa50 (Figure 2A, Table 2, and Supplementary Table S1). Most important, there was high interaction (\( P = 0.0009 \)) of the
FIGURE 1 | Resting, maximum and calcium-dependent force generation of skinned fibers exchanged for human hcTn containing HCM-associated hcTnT\textsuperscript{R130C} (n = 18 fibers), hcTnT\textsuperscript{WT} (n = 19 fibers), or DCM-associated hcTnT\textsuperscript{1K210} (n = 19 fibers). (A) Resting tension measured in relaxing solution (pCa 7) at short (1.1 $L_0$) and long (1.25 $L_0$) fiber length. (B) Maximum tension measured at pCa 4.28. (C) Force-pCa relations at short fiber length (1.1 $L_0$) and long fiber length (1.25 $L_0$). Normalized force is scaled as percentage from 0% for resting tension to 100% for maximum tension. ### indicates highly significant different to 1.1 $L_0$ in paired Bonferroni post tests.

TABLE 1 | Changes in resting tension, maximum tension, pCa\textsubscript{50} and $n_H$ induced by lengthening fibers from 1.1 to 1.25 $L_0$.

|                      | hcTnT\textsuperscript{R130C} | hcTnT\textsuperscript{WT} | hcTnT\textsuperscript{1K210} |
|----------------------|-------------------------------|----------------------------|-------------------------------|
| Delta-$F_{\text{REST}}$ (mN/mm$^2$) | +4.89 ± 0.94$^{***}$      | +3.28 ± 0.39$^{***}$      | +4.09 ± 0.70$^{***}$      |
| Delta-$F_{\text{MAX}}$ (mN/mm$^2$) | +2.56 ± 0.56$^{***}$      | +1.85 ± 0.35$^{***}$      | +2.01 ± 0.50$^{***}$      |
| Delta-$pC_{a50}$     | −0.030 ± 0.010$^{***}$     | −0.001 ± 0.010           | −0.034 ± 0.013$^g$       |
| $n_H$                | −0.06 ± 0.12                | −0.02 ± 0.07             | +0.22 ± 0.14              |

Values present mean ± SEM of subject-matched parameter changes, i.e., the parameter changes of the individual fibers exchanged for hcTn containing different hcTnT (hcTnT\textsuperscript{R130C}: n = 18, hcTnT\textsuperscript{WT}: n = 19, hcTnT\textsuperscript{1K210}: n = 19). Significant change by lengthening: $^aP < 0.05$, $^{**}P < 0.01$, $^{***}P < 0.001$. Significant different to hcTnT\textsuperscript{1K210}: $^{***}P < 0.001$.

Effects of hcTnT-type and fiber length on the pCa\textsubscript{50} indicating dissimilar lengthening-induced change of pCa\textsubscript{50} (delta-pCa\textsubscript{50}) among the fibers containing the three different hcTnT-types. In contrast to the strong interaction found for the pCa\textsubscript{50}, there is no interaction ($P = 0.18$), no effect of fiber length ($P = 0.37$) and no effect of hcTnT-type ($P = 0.47$) on the Hill coefficient $n_H$ in the two-way RM ANOVA (Figure 2B, Table 2, and Supplementary Table S1).

For further analysis of the effect of hcTnT-type on the pCa\textsubscript{50}, the data sets were separated for either short (1.1 $L_0$) or long (1.25 $L_0$) fiber length and post hoc tested by one-way ANOVA which confirmed the significant effect of hcTnT-type on the pCa\textsubscript{50} for each of the two lengths ($P < 0.001$). Tukey’s multiple comparison revealed that the DCM mutant highly significantly ($P < 0.001$) decreases Ca sensitivity at both lengths when either compared to the wild type or to the HCM mutant, whereas the HCM mutant increases Ca sensitivity only at long ($P < 0.05$) but not at short fiber length compared to the wild type (Figure 2A and Supplementary Table S1), thus lengthening manifested in an effect of the HCM mutant compared to wild type.

Post hoc analysis for searching the reason of the high interaction of hcTnT-type and length effects on the pCa\textsubscript{50}
by Tukey’s multiple comparison yielded highly significant (P < 0.001) different delta-\(pC_{50}\) of fibers containing HCM-associated hcTnT\textsuperscript{R130C} compared to fibers containing DCM-associated hcTnT\textsuperscript{AK210} (Figure 2C and Tables 1, 2). The 95% confidence interval of the delta-\(pC_{50}\) of fibers containing hcTnT\textsuperscript{R130C} was fully positive (+0.002 to +0.058) whereas that of fibers containing hcTnT\textsuperscript{AK210} was fully negative (−0.062 to −0.007). The significant difference (P < 0.05) of the intervals from zero is also reflected by the corresponding Bonferroni post-tests (* marks in Figure 2C) and indicates that lengthening induced calcium sensitization in fibers containing the HCM mutation whereas lengthening caused calcium desensitization in fibers containing the DCM mutation. The delta-\(pC_{50}\) of fibers containing wild type hcTnT is in-between the delta-\(pC_{50}\) of the fibers containing the mutants and not significantly different to each mutant.

To analyze the effects of hcTnT and length on the calcium-dependent force at sub-maximally activating [\(Ca^{2+}\)], the normalized force at each \(pC\) was tested by two-way RM ANOVA. Strong significant interaction of hcTnT-type and fiber length effects on normalized force were found for \(pC\) 6.07 (P = 0.0002), \(pC\) 5.88 (P < 0.0001), \(pC\) 5.77 (P = 0.0006), \(pC\) 5.65 (P = 0.0022), \(pC\) 5.51 (P = 0.0016), and \(pC\) 5.32 (P = 0.006) indicating dissimilarity of lengthening-induced change of normalized force (delta-\(F_{\text{norm}}\)) for at least one hcTnT-type at the respective \(pC\) (Table 2). The delta-\(F_{\text{norm}}\) values at submaximal activating \(pC\) are plotted in Figure 2D. Post-tests confirmed that at each of the above \(pC\), fibers containing the HCM and the DCM mutation differ significantly by at least P < 0.01 in their delta-\(F_{\text{norm}}\) (Table 2). At \(pC\) 5.88, all three hcTnT-types differed significantly in delta-\(F_{\text{norm}}\) (P < 0.01 for HCM versus WT, P < 0.05 for DCM versus WT, and P < 0.001 for HCM versus DCM) (Figure 2D and Table 2). In summary, stretching fibers containing the HCM mutation increased calcium sensitivity (\(pC_{50}\)) and \(F_{\text{norm}}\) whereas stretching fibers containing the DCM mutation decreased calcium sensitivity and \(F_{\text{norm}}\).

**DISCUSSION**

**Basic Effects of the hcTnT Mutations on Calcium Sensitivity**

The two mutations R130C and ΔK210 in hcTnT have been associated with autosomal dominant inherited hypertrophic or dilated cardiomyopathy, respectively (Kamisago et al., 2000; Song et al., 2005; Wang et al., 2007). The basal effects of the HCM-associated mutation R130C and of the DCM-associated mutation ΔK210 on calcium sensitivity found in this study resemble the most common phenotype of HCM-mutations and DCM-associated mutations on calcium sensitivity. They are in general agreement with numerous previous studies of HCM- and DCM-associated mutations in hcTnT reporting increase of calcium sensitivity by HCM- and decrease of calcium sensitivity by DCM-associated mutations (Morimoto et al., 2002; Lu et al., 2003, 2013; Venkatraman et al., 2003; Mirza et al., 2005; Robinson et al., 2007; Messer et al., 2016).

The functional consequences of the ΔK210 mutation in hcTnT have been extensively analyzed using in vitro, ex vivo, and in vivo models (Morimoto et al., 2002; Robinson et al., 2002; Venkatraman et al., 2003; Mirza et al., 2005; Du et al., 2007; Robinson et al., 2007; Sfichi-Duke et al., 2010) whereas to the best of our knowledge, there is no functional study for the R130C mutation. A study of knock-in mice expressing cTnT\textsuperscript{ΔK210} showed that the decrease of calcium sensitivity was higher in homozygous than in heterozygous cTnT\textsuperscript{ΔK210} mice indicating that the calcium desensitization by this mutation increased with the relative amount of mutant protein (Du et al., 2007). The exchange efficiencies in this study of 44–48% are in a good range to mimic the typical co-expression of mutant

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### Table 2: P-values obtained in statistical analysis of the parameters. \(F_{\text{pc}}\)'s correspond to the normalized force values at the respective \(pC\).

| Parameter          | Interaction | hcTnT-type | Fiber length | Subject match | Tukey’s multiple comparison | Bonferroni post-test |
|--------------------|-------------|------------|--------------|---------------|----------------------------|---------------------|
| \(F_{\text{REST}}\) | 0.37        | 0.78       | ***          | ***           | n.d.                       | HCM vs. WT          |
| \(F_{\text{ACT}}\) | 0.55        | 0.41       | ***          | ***           | n.d.                       | HCM vs. DCM         |
| \(pC_{50}\)        | ***         | 0.81       | ***          | –             | HCM vs. WT                 | DCM vs. WT          |
| \(n_{\text{v}}\)   | 0.18        | 0.37       | 0.47         | ***           | n.d.                       | HCM                 |
| \(F_{\text{pC56.25}}\) | 0.44      | 0.35       | 0.25         | ***           | n.d.                       | WT                  |
| \(F_{\text{pC56.07}}\) | ***      | **         | 0.67         | ***           | HCM vs. WT                 | DCM                 |
| \(F_{\text{pC56.88}}\) | ***      | ***        | 0.68         | ***           | HCM vs. WT                 |                     |
| \(F_{\text{pC55.77}}\) | ***      | ***        | 0.53         | ***           | HCM vs. WT                 |                     |
| \(F_{\text{pC55.65}}\) | ***      | ***        | 0.50         | ***           | HCM vs. WT                 |                     |
| \(F_{\text{pC55.51}}\) | ***      | ***        | 0.63         | ***           | HCM vs. WT                 |                     |
| \(F_{\text{pC55.32}}\) | ***      | ***        | 0.11         | ***           | HCM vs. WT                 |                     |
| \(F_{\text{pC55.06}}\) | 0.97      | ***        | 0.33         | ***           | n.d.                       |                     |
| \(F_{\text{pC54.7}}\) | 0.27      | ***        | 0.86         | ***           | n.d.                       |                     |

Two-way RM ANOVA and Tukey’s multiple comparison post-tests were used to test for significant differences among hcTnT-types: *P < 0.05, **P < 0.01, ***P < 0.001. n.d., not determined because of insignificant interaction (P > 0.05). Bonferroni post-tests indicate significant length-dependent changes of parameters for the hcTnT-type: *P < 0.05, **P < 0.01, ***P < 0.001; ↑ marks increase, ↓ marks decrease of parameter; – indicates not significant (P > 0.05).
and wild type hcTnT protein in the heterozygous allelic patients albeit the relative amount of the expression of mutant protein in patients can substantially differ from the theoretical value of 50% (Tripathi et al., 2011). The significant lower calcium sensitivity in absence of significant differences of maximum force and cooperativity of calcium-dependent force generation of fibers containing hcTnT<sup>1K210</sup> compared to hcTnT<sup>WT</sup> found in this study resemble the previously reported effects on these parameters found between fibers isolated from heterozygous cTnT<sup>ΔK210</sup> knock-in and wild type mice (Du et al., 2007). Thus, the <i>in vitro</i> exchange of hcTn in cardiac fibers performed in this study qualitatively reproduces the basic functional phenotype found in the <i>ex vivo</i> fiber approach (Du et al., 2007). The suitability of the <i>in vitro</i> approach to mimic primary effects of mutations on calcium sensitivity together with their diverging basal effects on calcium sensitivity at short fiber length provides a promising starting point for studying their effects on LDA.

**Limitations of the Present Study**

An unexpected result in our study was the lack of lengthening-induced change of calcium sensitivity (delta-pCa<sub>50</sub> = 0) and the low increase of $F_{MAX}$ (+10%) in fibers exchanged with hcTn wild type. Most likely the exchange for the recombinant human cardiac troponin complex in the fibers does not restore the LDA of the native cTn. Low LDA might partly result from the dephosphorylated state of cTnI in recombinant cTn (Konhilas et al., 2003). However, previous studies examining effects of cTnT mutation on LDA by exchange of recombinant gcTn into guinea pig cardiac fibers reported delta-pCa<sub>50</sub> values of +0.1 and increase of $F_{MAX}$ by 62–65% after wild type exchange (Reda and Chandra, 2018, 2019). Reda and Chandra (2018, 2019) used recombinant cTn consisting of the guinea pig isoforms while we used recombinant cTn consisting of the human isoforms. Thus, the lacking length-dependent change of pCa<sub>50</sub> and the low increase of $F_{MAX}$ for the wild type control in our study might
result from the species-specific difference in the cTn. We chose the human isoforms because the aim of our study was to compare the effects of cTnT mutations related to human cardiomyopathy.

To the best of our knowledge, no functional study of the HCM-associated mutation hCTnT^{R130C} exists so far. We found no difference in calcium sensitivity compared to wild type at short fiber length for this mutation but significant higher calcium sensitivity at long fiber length. Thus, screening the effect of this mutation on calcium sensitivity under basal conditions only would have been negative because the calcium sensitization only became evident under stretch. However, this finding underlines the positive effect of this HCM mutation on LDA.

Finally, we choose rather simple protocol for working at same relative fiber length instead of measuring and adjusting sarcomere length prior activation. Slack length ($L_0$) were measured prior to hCTn exchange, and biomechanical parameters determined after hCTn exchange at 1.1- and 1.25-fold of that $L_0$ measured prior exchange. Therefore, all fibers should have similar sarcomere length at $L_0$ independent of the type of hCTnT. Furthermore, the three different types of hCTnT-exchanged fibers exhibited similar passive and maximum tension and therefore likely adopted similar sarcomere lengths before and during calcium activation in the mechanical experiments.

### Effect of hCTnT Mutations on LDA

The primary aim of the study was to test if HCM and DCM-associated mutations in the human troponin complex exert different effects on the length dependence of mechanical parameters, in particular of calcium sensitivity reflecting LDA. As two-way RM ANOVA analysis indicated no difference in the effects of lengthening on resting and maximum tension for the three type of hCTnT, the HCM-associated R130C and the DCM-associated ΔK210 mutation seem not to alter the basal inhibitory and the maximum regulatory capacity of hCTn. Lengthening changed calcium sensitivity in opposite direction for the two mutations as indicated by the opposite signs of their 95% confidence intervals and the highly significant difference ($P < 0.001$) for their delta-pCa, indicating calcium sensitization by the HCM-associated and calcium desensitization by the DCM-associated mutation. LDA has been also associated to changes in the lattice spacing (McDonald and Moss, 1995; Fuchs and Smith, 2001) and lattice spacing depends on filament charge according to the Donnan potential (Millman and Irving, 1988). Since R130C and ΔK210 both lead to loss of a positive charge in hCTnT, their opposite effects on LDA cannot be explained by filament charge. Instead they likely reflect specific effects of the two sites, 130 and 210 on hCTnT, in the modulation of LDA.

Consistent with our finding of calcium desensitization by the DCM-associated ΔK210 mutation being augmented under stretch, the DCM-associated mutation R174W decreases calcium sensitivity and attenuates the sarcomere length-dependent increase of calcium sensitivity in guinea pig cardiac fibers (Reda and Chandra, 2019). Whether this applies for all DCM-associated mutations in hCTnT needs to be tested in future studies. In any case, it is definitive from previous studies that the mechanism does not apply for all HCM-associated mutations in thin filament proteins and that not all mutations increasing calcium sensitivity enhance LDA. The hypertrophic cardiomyopathy-associated mutation F87L in the central region of cTnT enhances calcium sensitivity but attenuates LDA (Reda and Chandra, 2018) and the RCM-associated mutation hCTnI^{R145W} does not affect LDA although it strongly increases calcium sensitivity (Dvornikov et al., 2016). Thus, modifications of different sites in cTnT might exert either positive or negative effects on LDA. Probing the effects of further mutations in Tm and cTn subunits on LDA provides a promising approach to map protein domains involved in LDA for understanding how these proteins integrate the length and the calcium signal for modulating myocardial contraction.

### Possible Contribution of LDA in the Diverging Phenotype of DCM and HCM

Several hypotheses have been formulated to explain the diverging heart phenotypes in HCM versus DCM manifesting from specific mutations within the same protein like cTnT: (1) mutations directly affecting calcium sensitivity (Robinson et al., 2002, 2007), (2) mutations affecting EC coupling or Ca$^{2+}$ homeostasis (Tardiff et al., 2015; Crocini et al., 2016), and (3) mutations interfering with the effect of posttranslational modifications on calcium sensitivity (Sfichi-Duke et al., 2010; Memo et al., 2013; Messer et al., 2016). The systemic development of each of the two diseases in the human is even more complex and highly variable (Maron et al., 2012; Deranek et al., 2019). Studies of human samples revealed that hCTnI is hypo-phosphorylated in myocardial samples from HCM and DCM patients compared to control samples from donor hearts (Hamdani et al., 2008; Sequeira et al., 2013, 2015). It is known that phosphorylation of cTnI by PKA increases lengthening-induced calcium sensitization by increasing the delta-pCa (Konhilas et al., 2003). The lower phosphorylation of cTnI in human patients and animal models for cardiomyopathies compared to control samples could therefore complicate the elucidation of the direct effect of the mutation on LDA. For example, low phosphorylation of hCTnI in HCM-associated patients might reduce LDA and prevent detection of possible increase of LDA by the mutation itself. The simple approach of exchanging recombinant cTn in guinea pig cardiac fibers, like in previous (Mickelson and Chandra, 2017; Reda and Chandra, 2018, 2019) and this study does not include this complication. The direct comparison of HCM and DCM mutant hCTnI found in our study supports the hypothesis that if one excludes posttranslational modulation of myofilaments proteins, HCM- and DCM-associated mutations can increase and decrease LDA, respectively.

Although, our study is in agreement with previous functional studies of the ΔK210 mutation and the prevalent disposition of HCM mutations increasing and DCM mutations decreasing calcium sensitivity, the definite reasons why the ΔK210 mutation results in DCM and the R130C mutation in HCM remain elusive. While typical features of HCM are increased wall thickness, cardiomyocyte disarray, fibrosis and impaired diastolic filling, DCM is characterized by enlarged ventricles, reduced ventricular wall thickness to volume ratio and impaired systolic contraction, i.e., reduced ejection fraction (Garfinkel et al., 2018;...
An interesting hypothesis is that the primary, acute effects of mutations on LDA, in the long term, might contribute to the directionality of the diverging histological and morphological phenotypes of HCM and DCM. Chronically enhanced response to stretch could contribute to strain imbalance of cardiomyocytes, cardiomyocyte disarray and wall thickening in HCM, while impaired contractile response to stretch might lead to overstretched thin cardiomyocytes and enlarged ventricles in DCM. Certainly, the primary effects of the mutations on LDA found in this study are counteractive and not compensatory mechanisms for the primary diastolic dysfunction in HCM and systolic dysfunction in DCM. Regarding the finding that the Frank–Starling mechanism is also impaired in the late-stage heart failure (Schwinger et al., 1994), the effect of the DCM mutation on impairing LDA is expected to be detrimental.

**DATA AVAILABILITY STATEMENT**

The datasets generated for this study are available on request to the corresponding author.

**ETHICS STATEMENT**

Guinea pig were killed for subsequent removal of the heart according to the guidelines approved by the ethical committee of the Landesamt für Natur, Umwelt und Verbraucherschutz Nordrhein-Westfalen (LANUV NRW Leibnizstraße 10 D-45659 Recklinghausen Germany).

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**SUPPLEMENTARY MATERIAL**

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**Conflict of Interest:** The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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