Two Drosophila Myosin Transducer Mutants with Distinct Cardiomyopathies Have Divergent ADP and Actin Affinities*

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Two Drosophila myosin II point mutations (D45 and Mhc⁴) generate Drosophila cardiac phenotypes that are similar to dilated or restrictive human cardiomyopathies. Our homology models suggest that the mutations (A261T in D45, G200D in Mhc⁵) could stabilize (D45) or destabilize (Mhc⁵) loop 1 of myosin, a region known to influence ADP release. To gain insight into the molecular mechanism that causes the cardiomyopathic phenotypes to develop, we determined whether the kinetic properties of the mutant molecules have been altered. We used myosin subfragment 1 (S1) carrying either of the two mutations (S1 A261T and S1 G200D) from the indirect flight muscles of Drosophila. The kinetic data show that the two point mutations have an opposite effect on the enzymatic activity of S1. S1 A261T is less active (reduced ATPase, higher ADP affinity for S1 and actomyosin subfragment 1 (actin-S1), and reduced ATP-induced dissociation of actin-S1), whereas S1 G200D shows increased enzymatic activity (enhanced ATPase, reduced ADP affinity for both S1 and actin-S1). The opposite changes in the myosin properties are consistent with the induced cardiac phenotypes for S1 A261T (dilated) and S1 G200D (restrictive). Our results provide novel insights into the molecular mechanisms that cause different cardiomyopathy phenotypes for these mutants. In addition, we report that S1 A261T weakens the affinity of S1-ADP for actin, whereas S1 G200D increases it. This may account for the suppression (A261T) or enhancement (G200D) of the skeletal muscle hypercontraction phenotype induced by the troponin I held-up² mutation in Drosophila.

Myosins are a large family of actin-based molecular motor proteins, of which at least 35 different classes are known (1). They show a wide variety of different mechanical activities, such as muscle contraction, phagocytosis, cell motility, tension maintenance, and vesicle transport (2). Striated muscle myosin is a class II molecular motor and consists of two myosin heavy chains (MHC), two essential light chains, and two regulatory light chains. Muscle contraction results from cyclic interactions between myosin II motors and actin filaments, also known as the cross-bridge cycle. These force-generating interactions are driven by the hydrolysis of ATP at the myosin active site (3). Mutations in human myosin II domains have been associated with both skeletal and cardiac myopathies (4).

Drosophila melanogaster has been described as a useful model system to study cardiomyopathies (5, 6). A recent study of myosin heavy chain mutations in Drosophila has shown that two mutations in the myosin motor domain generate defects in the Drosophila heart. The mutation known as D45⁴ (point mutation A261T) has a phenotype similar to human dilated cardiomyopathy (DCM), whereas Mhc⁵ (point mutation G200D) has a restrictive cardiomyopathy phenotype (RCM, a subset of hypertrophic cardiomyopathies (HCMs)) (7). The two myosin mutations identified are localized in the transducer region (Fig. 1), which is encoded by constitutive exons of the alternatively spliced Mhc transcript. D45 (A261T) is located on one of the β-strands (β̄) close to the β-bulge between β6 and β7, whereas Mhc⁵ (G200D) is at the N-terminal side of loop 1 (using chicken skeletal myosin residue numbers as introduced by Kronert et al. (8); the analogous Drosophila residues are Gly²⁰⁰ and Ala²⁵⁰). The transducer region is thought to integrate the structural changes from different parts of the motor domain and includes the seven-stranded β-sheet plus various flexible elements such as loop 1, the β-bulge between β6 and β7, the HO-linker, and the nucleotide-sensing elements such as the phosphate binding loop (P-loop), switch-1, and switch-2 (11). The seven-stranded β-sheet becomes distorted when myosin switches between the rigor-like and post-rigor states, and this distortion is facilitated by the flexible elements.

In an earlier study, D45 and Mhc⁵ were shown to interact with a mutation in troponin I (held-up² (hidp²)) that causes hypercontractility of the indirect flight muscle (IFM) (8). D45 is a suppressor of the TnI mutation (A116V) in the IFM, and in Drosophila hearts, it leads to a DCM-like phenotype in the presence of normal troponin I, whereas Mhc⁵ leads to RCM in the

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⁴ The two mutants used throughout this study are: Mhc⁴, which indicates the myosin heavy chain mutation G200D; and D45, which indicates the myosin heavy chain mutation A261T.

⁵ The abbreviations used are: DCM, dilated cardiomyopathy; HCM, hypertrophic cardiomyopathy; RCM, restrictive cardiomyopathy; S1, myosin subfragment 1; actin-S1, actomyosin subfragment 1; CATP, caged ATP; cADP, caged ADP; eda-deac-ATP, 3′-O-(2-(7-diethylamino-coumarin-3-carboxamido)ethyl)carbamoyl-ATP; eda-deac-ADP, 3′-O-(2-(7-diethylamino-coumarin-3-carboxamido)ethyl)carbamoyl-ADP; IFI, indirect flight muscle isoform; IFM, indirect flight muscle; Tn, troponin; TnI, troponin I; EMB, embryonic myosin S1.
heart and is lethal in combination with \( hdp^2 \). This suggests the likelihood of opposite effects of the two myosin mutations in regard to TnI interaction and that such differential activity may lead to the observed alternative cardiomyopathies.

Although a large number of human cardiomyopathies have been linked with specific mutations in sarcomeric proteins, including myosins, the causal relationship is not well understood. For mutations in the thin filament proteins tropomyosin (Tm) and troponin (Tn), there is an emerging paradigm that mutations that increase calcium sensitivity of the thin filament regulation of contraction result in HCM, whereas a decrease in calcium sensitivity is associated with DCM (9, 10). Because the \( hdp^2 \) TnI mutation leads to over-contraction, the myosin mutation may reduce calcium sensitivity (\( D45^- \)) to compensate for the TnI mutation or increase it further (\( Mhc^5^- \)) to cause lethality. Indeed, \( Drosophila \) D45 myosin has reduced ATPase activity and reduced \( \text{in vitro} \) motility, whereas \( Mhc^5 \) myosin displays an increase in both ATPase activity and \( \text{in vitro} \) motility (7). It was argued that this is consistent with \( D45^- \) suppressing the TnI mutation and \( Mhc^5^- \) enhancing it. However, the molecular mechanism for this interaction remains to be explored.

In this study, we used myosin subfragment 1 (S1) to investigate the transient and steady-state kinetics of the two transducer mutants of myosin S1 that suppress (A261T) or enhance (G200D) the TnI defect (A116V). Because the \( hdp^2 \) TnI mutation leads to over-contraction, the myosin mutation may reduce calcium sensitivity (\( D45^- \)) to compensate for the TnI mutation or increase it further (\( Mhc^5^- \)) to cause lethality. Indeed, \( Drosophila \) D45 myosin has reduced ATPase activity and reduced \( \text{in vitro} \) motility, whereas \( Mhc^5 \) myosin displays an increase in both ATPase activity and \( \text{in vitro} \) motility (7). It was argued that this is consistent with \( D45^- \) suppressing the TnI mutation and \( Mhc^5^- \) enhancing it. However, the molecular mechanism for this interaction remains to be explored.

In this study, we used myosin subfragment 1 (S1) to investigate the transient and steady-state kinetics of the two transducer mutants of myosin S1 that suppress (A261T) or enhance (G200D) the TnI defect (A116V). We will use S1\(^{A261T}\) and S1\(^{G200D}\) nomenclature for the two mutants throughout this study, and the properties and nomenclature of these mutants are summarized in Table 1. The kinetic data show that the two point mutations have an opposite effect on the enzymatic activity of S1 myosin as compared with wild type, resulting in a less active myosin (S1\(^{A261T}\)) or more active myosin (S1\(^{G200D}\)). These observations are consistent with the induced cardiac phenotypes for the two mutants. In addition, we report that the two mutations change the affinity of S1-ADP for actin in opposite directions and that these changes can account for the differential interaction of the two mutations with the TnI \( hdp^2 \) mutation in \( Drosophila \).

**TABLE 1**

| Point mutation | Mhc\(^5^-\) | D45^- |
|----------------|-------------|-------|
| Location in transducer | G200D | A261T |
| Interaction with flightless TnI (A116V) \( \text{held-up}^2 \) allele | Lethal | Rescued |
| \( Drosophila \) cardiac phenotype | RCM | DCM |

**EXPERIMENTAL PROCEDURES**

Myosin Isolation and Generation of Its S1—Myosin was isolated from IFM of less than 1-day-old wild-type and D45(A261T) or Mhc\(^5\)(G200D) mutant flies as described previously (12). The original procedure has been modified so that all dilutions were carried out with water containing 10 mM DTT and all solution volumes were reduced to 75% (7). Digestion of the S1 subfragment was carried out with chymotrypsin as described previously (13–15) with the following modifications. The final myosin pellet obtained after centrifugation was resuspended in digestion buffer (120 mM NaCl, 20 mM Na\(_2\)PO\(_4\), pH 7.0, 1 mM EDTA, and 20 mM DTT). Samples were incubated on ice for \(~6\) min. The myosin solution was then placed in a \(20^\circ\)C water bath for 5 min for temperature equilibration. The addition of \( \alpha \)-chymotrypsin (0.2 mg/ml final concentration from 10 mg/ml stock solution) to the myosin solution was carried out in the \(20^\circ\)C water bath followed by incubation for 6 min. To quench the reaction, 1.5 mM (final concentration) of phenylmethylsulfonyl fluoride (PMSF) was added. To pellet undigested myosin or myosin rod, the sample was immediately cen-
trifuged at 68,000 rpm (250,000 × g, TLA 100.3 rotor) for 30 min in a Beckman ultracentrifuge. The supernatant was removed and diluted 10-fold with low salt buffer (30 mM KCl, 5 mM MgCl₂, 20 mM MOPS, pH 7.0, and 4 mM DTT) and incubated for 30 min on ice. Any remaining uncut myosin or rod contamination was eliminated by centrifugation at 68,000 rpm for 20 min. For S1 concentration, the samples were centrifuged at 12,000 × g in a Sorvall MC 12 V at 4°C using a Millipore Ultrafree 0.5-μm centrifugal Biomax-5 filter with a 5-kDa cut-off. The final volume of the supernatant was 40–60 μl. S1 concentration was determined from A₂₈₀ (0.75 cm⁻¹ for 1 mg of S1 per ml) and a molecular mass of 115 kDa. A single band of S1 was detected on SDS polyacrylamide gels for all samples. All steady-state kinetic measurements were performed with freshly prepared S1, and transient kinetics experiments were performed within 1 week after S1 preparation.

G- and F-actin Preparation—G-actin was isolated from acetone powder of chicken skeletal muscle as described previously (1–4). After multiple cycles of polymerization–depolymerization, soluble G-actin was obtained after dialysis against 2 mM Tris-Cl (pH 8), 0.2 mM ATP, 2 mM CaCl₂, and 1 mM EDTA and quantified spectrophotometrically using an extinction coefficient of 0.62 cm⁻¹ (A₃₄₀ – A₂₈₀ nm) for 1 mg of actin ml⁻¹. To obtain a G-actin solution of 150–250 μM, the G-actin solution was concentrated using a Millipore Biomax-5 filter as described above. F-actin was prepared by adding 1 volume of 10% polymerization buffer (50 mM Tris-Cl, pH 8, 0.5 mM KCl, 20 mM MgCl₂, and 10 mM EDTA) to 9 volumes of G-actin. Actin was stored on ice at 4°C and used within 1 week of preparation for actin-activated Mg²⁺-ATPase assays.

Steady-state ATPase Activity—ATPase activities were determined using [γ-³²P]ATP as described previously in detail for full-length myosin (7). Ca²⁺ ATPase was carried out as described above. Basal Mg²⁺ ATPase was carried out as described above, but basal Mg²⁺ ATPase activities obtained in the presence of 500 nM S1 and 10 mM ADP with 1 μM actin (1–3 μM S1, and 500 μM cATP, eda-deac-ADP dissociation experiments were also performed in this low salt buffer and contained 4 μM S1, 10 μM eda-deac-ATP (source of eda-deac-ADP), and 100 μM cATP (S1A²⁶¹T) or caged ADP (cADP) (S1G²⁰⁰D₂).

Analysis of Transient Kinetic Data—The following equation was derived from the interaction of actin and S1 with ATP and ADP shown in Scheme 1 and was used to determine Kₐᵢₜ.

\[
k_{obs} = K_{k_{10}}[ATP](1 + [ADP]/K_{AD})
\]

where \( k_{obs} \) is the observed rate constant for the ATP-induced dissociation of actin-S1; \( k_{k_{10}} \) is the second-order rate constant for ATP binding to actin-S1; and \( K_{AD} \) is the equilibrium dissociation constant for the binding of ADP to actin-S1. The equation \( k_{rel} = k_{obs}/k_{0} \) was used to determine the relative rate constant (\( k_{rel} \)) shown in Fig. 5, where \( k_{0} \) is the value when [ADP] = 0.

Co-sedimentation Assay—Co-sedimentation assays were done as described previously with some minor modifications (14). S1 (100 nm) was incubated with increasing actin concentrations (0–500 nM) in low salt buffer (30 mM KCl, 5 mM MgCl₂, 20 mM MOPS, pH 7.0, and 4 mM DTT) with a total volume of 100 μl and then centrifuged at 400,000 × g for 30 min at low temperature (4°C) to determine \( K_{A} \). Measurements were repeated in the presence of 10 mM ATP with 500 nM S1 and increasing actin concentrations (0–5 μM) to determine \( K_{DA} \).

Homology Models—Homology models of the wild-type indirect flight muscle isoform (IFI) and the two point mutants S1A²⁶¹T and S1G²⁰⁰D₂ were built as described using SWISS-MODEL (19). When aligning the IFI myosin head sequence with class II scallop myosin, the sequence identity is 60 ± 1%, enabling us to build well resolved homology models (20). Crystal structures of the various states of scallop myosin during the cross-bridge cycle were chosen to generate three-dimensional homology models of IFI, S1A²⁶¹T, and S1G²⁰⁰D₂: the near-rigor state of myosin (Protein Data Bank (PDB) code 1sr6), the post-

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**Scheme 1:** The interaction of S1 with actin, ATP and ADP. M, A, T, and D symbolize S1, actin, ATP, and ADP, respectively.
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rigor actin-detached state (PDB code 1kk8) that contains ADP-BeF₃, the pre-powerstroke state (PDB code 1qvi) that contains ADP-VO₄, and a novel conformation that contains partially bound ADP-SO₄ (PDB code 1s5g). For regions of insertions or deletions in the target-template alignment, SWISS-MODEL constructs an ensemble of fragments compatible with the neighboring stems using constraint space programming or a loop library (20, 21).

RESULTS

Catalytic Activity Is Reduced for S1¹₂₆₁T and Increased for S1²₀₀₀D — The basal and actin-activated Mg²⁺-ATPase activities of wild-type myosin S1 and the two point mutants (S1¹₂₆₁T and S1²₀₀₀D) were measured (Fig. 2), and the results are summarized in Table 2. The Tnl hdp² suppressor mutant S1¹₂₆₁T showed a more than 2-fold reduction in basal Mg²⁺-ATPase activity (0.04 s⁻¹), a nearly 3-fold reduction in basal Ca²⁺-ATPase (1.97 s⁻¹), and a 10-fold reduction in V_max (0.23 s⁻¹) as compared with wild type (0.09, 5.33, and 2.54 s⁻¹, respectively) (difference significance at <0.001 in each case). In contrast, the hdp² enhancer mutant S1²₀₀₀D showed nearly a 2-fold increase in basal Mg²⁺-ATPase (0.14 s⁻¹) and no significant change in basal Ca²⁺-ATPase or V_max.

The earlier ATPase measurements for full-length myosin (7) cannot be compared directly with the S1 ATPases described here because the two assays use different salt concentrations and the filamentous two-headed full-length myosin has a different affinity for actin than the soluble S1. However, the relative effects of the mutations on the S1 and the full-length myosin basal Mg²⁺-ATPases and the V_max are similar (full-length myosin basal Mg²⁺-ATPase, 0.2, 0.48, and 0.11 s⁻¹, and V_max of 1.8, 1.5, and 0.69 s⁻¹ for wild type, G200D, and A261T, respectively). The Kₘ for actin with the S1 constructs of either S1¹₂₆₁T (Kₘ = 2.15 μM) or S1²₀₀₀D (Kₘ = 3.79 μM) showed a decrease relative to wild type (Kₘ = 5.77 μM), which was not seen previously with full-length myosin. The ratio V_max/Kₘ, referred to as the catalytic efficiency, shows a significant (p < 0.01) increase for S1²₀₀₀D (0.70 μM⁻¹s⁻¹) as compared with wild type (0.45 μM⁻¹s⁻¹), whereas for S1¹₂₆₁T, this ratio decreases 4-fold (0.11 μM⁻¹s⁻¹, significance p < 0.001). Using flash photolysis, the turnover number (k_cat) was measured at a fixed actin concentration (2 μM) for the point mutants S1²₀₀₀D and S1¹₂₆₁T and compared with wild type (Fig. 3). The catalytic activity of S1¹₂₆₁T (k_cat = 0.064 s⁻¹) is reduced as compared with wild type (0.172 s⁻¹), whereas k_cat of S1²₀₀₀D (0.210 s⁻¹) is increased. This demonstrates that the differences seen in the lower ionic strength ATPase assay above remain at the higher salt concentrations (30 mM KCl) used for the transient kinetic measurements.

FIGURE 2. Actin-stimulated ATPase activity of wild-type IFI (A), S1²₀₀₀D (B), and S1¹₂₆₁T (C). Basal Mg²⁺-ATPase and actin-stimulated Mg²⁺-ATPase data were obtained in the absence or in the presence of increasing concentrations of actin as described under “Experimental Procedures.” To obtain Kₘ and V_max basal Mg-ATPase values were subtracted from actin-activated ATPase values, and data points were fit to a hyperbola. Statistical significance of data comparisons are shown in Table 2.

TABLE 2

Steady-state kinetic parameters measured for wild-type, G200D and A261T Drosophila myosin S1

| Myosin S1 | Basal Ca-ATPase V_max | Kₘ | V_max/Kₘ |
|-----------|----------------------|-----|----------|
|           | s⁻¹                  | μM  | μM⁻¹s⁻¹  |
| Wild-type IFI | 5.33 ± 0.62 | 0.09 ± 0.01 | 2.54 ± 0.29 | 0.45 ± 0.09 |
| S¹₂₆₁T(Mic³) | 5.36 ± 0.27 | 0.14 ± 0.01 | 2.60 ± 0.18 | 0.70 ± 0.08 |
| S¹₂₆₁T(D45) | 1.97 ± 0.46 | 0.04 ± 0.01 | 0.23 ± 0.04 | 2.15 ± 0.73 |

*p < 0.001 determined by Student’s t test as compared with IFI.
* p < 0.01 determined by Student’s t test as compared with IFI.
ATP-induced Dissociation of Actin-S1 Is Slower for S1^{A261T} but Unaltered for S1^{G200D}—The second-order rate constant governing the dissociation of acto-S1 by ATP ($K_{12}$; see Scheme 1) was measured as described previously (14). Briefly, 1 μM actin and 1–3 μM S1 were preincubated and then dissociated by ATP released from cATP. In the presence of apyrase, a single sample was subjected to a series of laser flashes decreasing in intensity to release varying concentrations of ATP. The rate of change in light scattering was recorded for each laser flash and could be well described by a single exponential at each ATP concentration (Fig. 4, A and B). The slope of a graph of $k_{obs}$ against ATP concentration determines the apparent second-order rate constant $K_{AD}$ for the dissociation of acto-S1 by ATP (Fig. 4C). As compared with the value for wild type (0.75 μM$^{-1}$ s$^{-1}$), S1^{A261T} showed a reduced ATP-induced dissociation rate (0.47 μM$^{-1}$ s$^{-1}$), whereas the rate measured for S1^{G200D} (0.64 μM$^{-1}$ s$^{-1}$) was similar to wild type (Table 3).

Affinity of Actin-S1 for ADP Is Weak for S1^{G200D} but Strong for S1^{A261T}—The ADP affinity for S1 in the presence of actin, defined by the equilibrium dissociation constant $K_{AD}$, was determined as described previously (14). In the presence of increasing concentrations of ADP, the ATP-induced dissociation of acto-S1 was measured using light scattering, and plotting the relative $k_{obs}$ ($k_{rel}$) versus ADP concentration allows the measurement of the ADP affinity $K_{AD}$ for acto-S1 (Fig. 5 and Scheme 1). The $K_{AD}$ of S1^{A261T} ($K_{AD} = 293$ μM) is reduced by almost 30% as compared with wild type ($K_{AD} = 409$ μM), whereas for S1^{G200D}, the $K_{AD}$ value is increased ($K_{AD} = 604$ μM) (Fig. 5 and Table 3). Thus the ADP affinity of the two point mutants is affected in opposing ways with S1^{G200D} having a weaker ADP affinity and S1^{A261T} displaying a stronger ADP affinity as compared with wild type.

ADP Release from S1 ($k_{rel}$) Is Fast for S1^{G200D} but Slow for S1^{A261T}—To estimate the rate constant of ADP dissociation from S1 in the absence of actin, the change in fluorescence of coumarin-labeled ADP (eda-deac-ADP) was measured upon displacement of eda-deac-ADP by ATP binding to S1. It was shown previously that this coumarin-labeled analog has very similar kinetic properties as compared with ADP (17). Measuring the ADP release rate ($k_{rel}$) for wild type and S1^{A261T} was relatively straightforward. A single laser flash released 15–20 μM ATP from cATP (100 μM), and the fluorescence change resulting from eda-deac-ADP release was well described by a single exponential function, as shown in Fig. 6A. The observed rate constant for ADP release ($k_{rel} = 41$ s$^{-1}$) was slower as compared with wild type ($k_{rel} = 7.3$ s$^{-1}$).

For S1^{G200D}, a different approach was needed because it was noted that the fluorescence signal of the sample containing S1^{G200D} increased on adding cATP, indicating that cATP could displace eda-deac-ADP for this mutant. This displacement of eda-deac-ADP by cATP was not seen for wild-type S1, indicat-
TABLE 3

| Myosin S1          |  \(K_{\text{K}_{\text{AD}}}\)  |  \(K_{\text{AD}}\)  |  \(k_{\text{AD}}\)  |  \(k_{\text{AD}}^1\)  |  \(K_{\text{AD}/KD}}\)  |  \(K_{\text{AD}}\)  |
|---------------------|-----------------|-----------------|----------------|----------------|----------------|----------------|
| Wild-type IFI       | 0.75 ± 0.08 s^{-1} | 409 ± 26 s^{-1} | 4090          | 7.5 ± 1.3      | 7.5           | 54            |
| S1(G200D)          | 0.63 ± 0.09 s^{-1} | 604 ± 157 s^{-1} | 6040          | 71 ± 19         | 71            | 8.5           |
| S1(A261T)          | 0.47 ± 0.08 s^{-1} | 293 ± 61 s^{-1} | 2930          | 4.1 ± 0.3       | 4.1           | 71            |
| S1(G200D)          | 0.47 ± 0.08 s^{-1} | 293 ± 61 s^{-1} | 2930          | 4.1 ± 0.3       | 4.1           | 71            |

* Data are estimated from  \(K_{\text{AD}}\) assuming an association rate constant of 10^{7} M^{-1} s^{-1}.  
* Data are estimated from  \(k_{\text{AD}}\) assuming an association rate constant of 10^{6} M^{-1} s^{-1}.  
* Data are from Miller et al. (14).  
* \(p < 0.05\) determined by Student’s t test as compared with IFI.  
* \(p < 0.01\) determined by Student’s t test as compared with IFI.

FIGURE 5. The affinity of ADP (\(K_{\text{AD}}\)) for actin S1. The dissociation of actin-S1 was induced by ATP in the presence of ADP ranging from 0 to 1200 \(\mu\text{M}\). The light-scattering traces were fitted with single exponentials to determine the \(k_{\text{obs}}\). Hyperbolic plots of the \(k_{\text{obs}}(k_{\text{obs}}/k_{0})\) versus ADP concentration were fitted with an equation derived from Scheme 1 (18) to determine \(K_{\text{AD}}\). The fits yielded a value of 293 ± 61 \(\mu\text{M}\) for S1(G200D) (A) and 604 ± 157 \(\mu\text{M}\) for S1(G200D) (B), as compared with 409 ± 26 \(\mu\text{M}\) for wild-type IFI (C).

FIGURE 6. Rate of eda-deac-ADP dissociation (\(k_{\text{AD}}\)) from Drosophila point mutant S1(A261T) (A) and S1(G200D) (B). A fluorescence signal after releasing ATP (15 \(\mu\text{M}\)) from caged ATP (100 \(\mu\text{M}\)) by a single laser flash displacing the eda-deac-ADP bound to S1(A261T) (A) and S1(G200D) (B). Fitting with a single exponential resulted in \(k_{\text{AD}} = 4.1 ± 0.3 s^{-1}\). B, fluorescence signal after release of ADP (7 \(\mu\text{M}\)) from cADP (1000 \(\mu\text{M}\)) by a single laser flash displacing eda-deac-ADP from S1(G200D). Fitting with a single exponential resulted in \(k_{\text{AD}} = 71 ± 19 s^{-1}\).

Affinity of S1 for Actin in the Presence of ADP Is Weak for S1(A261T) but Strong for S1(G200D)—The measurement of the affinity of S1 and actin-S1 for ADP (\(K_{\text{AD}}\)) allows us to calculate the thermodynamic coupling constant (\(K_{\text{AD}}/K_{\text{D}}\)) for the interaction between the actin and ADP sites on S1. This ratio of two independent values carries a large error and includes the assumption that \(K_{\text{AD}}\) has the same value for all constructs. The values listed in Table 3 are shown for illustration, acknowledging the error. The numbers do show a 6-fold reduction for S1(G200D) in the coupling (from 54 for wild type to 8.5 for the mutation). In contrast, A261T has a 30% increase in coupling (\(K_{\text{AD}}/K_{\text{D}} = 71\)), which is not very different from the 20% error of the \(K_{\text{AD}}\) value. Here we can use the values to predict the effect of ADP on the affinity of S1 for actin for the two mutants. In our earlier work (14), we estimated that the affinity of wild-type S1 for actin (\(K_{\text{a}}\)) was ±10 nm. Assuming that the affinity of S1 for
Homology Models Suggest a Role for Loop 1—Homology models were built of wild type and mutants S1G200D and S1A261T to help understand the mechanism of action of the mutated residues. Fig. 8 shows part of the transducer area of the homology structures representing the pre-powerstroke state, the differences between the two transducer mutants, our results provide new insights into the molecular mechanisms that cause different phenotypes for these mutants on their own and in combination with the held-up2 TnI mutation.

Summary
The affinity of S1 for actin was characterized with the co-sedimentation method and in particular, loop 1. The model of S1A261T suggests a helical structure of loop 1 for wild type, which is disrupted in S1G200D. The bound nucleotide is shown as solid spheres, and switch 1 (SW-I) and the phosphate binding loop (P-loop) are also indicated. The altered structure of loop 1 may affect its flexibility and ultimately change ADP release rate for the mutants, with faster release for S1G200D (less ordered loop 1 structure) and slower release for S1A261T (more rigid loop 1 structure). FIGURE 8. Homology models of S1G200D (A) and S1A261T (B) suggest a role for loop 1. A, overlay of the loop 1 area of homology models of wild-type S1 (Wt) and S1G200D (built using pre-powerstroke scallop myosin (PDB code 1qvi) as template) showing a helical structure of loop 1 for wild type, which is disrupted in S1G200D. The bound nucleotide is shown as solid spheres, and switch 1 (SW-I) and the phosphate binding loop (P-loop) are also indicated. B, loop 1 area of S1A261T shows a wild type-like helical structure for loop 1. The conserved loop 1 residue Lys203 can interact with Thr261, which is located on β′. The altered structure of loop 1 may affect its flexibility and ultimately change ADP release rate for the mutants, with faster release for S1G200D (less ordered loop 1 structure) and slower release for S1A261T (more rigid loop 1 structure).
Point Mutants of Drosophila Muscle Myosin

There is an emerging consensus that in the case of HCM (of which RCM is a subset), mutant contractile proteins result in increased contractility or motor function and are usually associated with an increase in the calcium sensitivity of contraction. In contrast, mutations that result in DCM yield decreased contractility or motor function that is often associated with a decrease in calcium sensitivity (9, 23–27). Our results are consistent with the induced cardiac phenotypes because actin-S1A261T has decreased activity and results in a DCM phenotype (dilated), whereas actin-S1G200D results in increased activity, and this mutation causes an RCM phenotype (restrictive). Our data also show for the first time that the affinity of S1-ADP for actin is affected in opposite ways with the RCM mutant having very tight actin affinity and the DCM mutant showing weaker actin affinity as compared with wild type. Because these particular mutations affect the transducer region, which provides a communication pathway between the actin and nucleotide binding sites (11), the changes in transducer structure might be predicted to have effects on the actin affinity of S1 in the presence of ADP. However, note that although residues Ala261 and Gly200 and the transducer region are common to both the cardiac and the IFI myosin isoforms, there are differences in other domains of the myosin head between the IFI and cardiac myosin isoforms (28). Thus the IFI isoform can model the heart isoforms, but is not identical.

Can our results help explain the interaction between the mutant myosins characterized here and the TnI held-up2 mutation? The regulation of muscle contraction by troponin relies, at one level, on a competition between TnI and myosin for binding sites on actin. In the absence of calcium, TnI binds actin more tightly than myosin; in the presence of calcium, the TnI interaction is weakened, and now myosin binds more tightly. In vitro motility assays of regulated thin filaments showed that the hdp2 mutant troponin I caused activation of motility at lower calcium concentrations than wild type (29, 30). If the held-up2 phenotype is caused by the A116V TnI mutation weakening the inhibitory interaction of TnI with actin, then a mutation in myosin that weakens its interaction with actin could bring the two affinities back into balance. The AM-ADP complex is expected to be a significant force-holding state in the cross-bridge cycle. Alterations in the affinity of AM-ADP for actin could alter the ability of the AM-ADP state to compete with TnI for binding to actin. Our estimate of KDA for S1A261T (weaker actin binding than wild-type IFI) is consistent with this prediction and with this mutation operating as a suppressor of the held-up2 phenotype. In contrast, S1G200D has a higher affinity of S1-ADP for actin, and the G200D mutation is expected to be able to activate more readily than wild-type myosin. In addition to the direct effect of competition between TnI and myosin, a more strongly bound myosin would, through the cooperativity in the thin filament, enhance the affinity of the Tn complex for calcium (31). This, in combination with held-up2, will result in activation at very low calcium concentrations. This can explain the lethality observed when these two mutations are placed within the same organism, where muscles essential for viability are constitutively activated. These characteristics of the G200D activating more readily and A261T activating less well are also consistent with alterations in calcium sensitivity associated with the DCM and RCM cardiac defects.

Another characteristic change produced in myosin by the two mutations is the change in the in vitro motility. The velocity was enhanced by 16% for G200D and reduced by 55% for A261T (7). The velocity at which myosin moves actin in a motility assay and the maximum speed of muscle fiber shortening are normally considered to be limited by the rate constant for cross-bridge detachment after completion of the working stroke (32–34). For fast myosins, cross-bridge detachment is usually limited by ATP binding, whereas for slower myosins, ADP release is rate-limiting. However, for Drosophila IFM fibers at saturating ATP (20 μM), the rate-limiting step of fiber shortening is inorganic phosphate release (35). Because IFM fibers have a very weak affinity for ATP, at the ATP concentration used for the motility assays (2 mM), ATP binding could be rate-limiting instead of phosphate release. However, in vitro motility data are collected under unloaded conditions, whereas fiber data represent loaded conditions, which could result in different rate-limiting steps. For the IFM, as for fast mammalian muscle, the ADP release is expected to be much too fast to limit the velocity, and this remains true for the two mutant myosins. Here we report that the second-order rate constant (K1k+2) for ATP-induced dissociation of actin-S1A261T is reduced to about 60% of the value of the IFI. This is similar to the 55% reduction in the velocity of in vitro motility assays. In contrast, there is no significant difference between the rate constant for ATP-induced dissociation of actin-S1G200D and wild type and only a 16% difference in the in vitro motility velocity.

The homology models suggest a possible loop 1-based mechanism for how the two mutations affect myosin kinetics in opposite ways. The rate constant of ADP release (kAD and hence K1k+2) has been correlated with the size and flexibility of loop 1, with the larger and more flexible loops giving higher release rate constants (36). Loop 1 is also one of the key players in transducer region function, and its importance has been emphasized by Houdusse and co-workers (11) who called the transducer region “loop 1 tuning elements.” Although the surface loop 1 is not usually seen in myosin crystal structures because of its inherent flexibility, our homology models do predict differences in the apparent flexibility of the loop in the three myosins. For wild type, going from near-rigor to post-rigor to pre-powerstroke states, the models predict an increase in the probability of a helical structure, implying less flexibility for the loop in the pre-powerstroke state. A similar trend is seen for the three structures of each of the two mutants, with A261T tending to have more helix in loop 1 in each case and G200D less. The models suggest that for wild-type myosin in the most stable (pre-powerstroke) state, loop 1 is largely an α-helix that starts at Gly200. For S1G200D, the introduction of the larger Asp side chain appears to disrupt the start of the helical structure, resulting in a more flexible loop that may lead to faster ADP release rates. In the case of S1A261T, loop 1 may become less flexible due to the interaction between Lys203 (in loop 1) and Thr261, resulting in slower ADP release.

In summary, this study of the two myosin point mutations reveals characteristics that are compatible with the observed IFM and cardiac phenotypes. In addition, the changes in the
two mutant S1s provide an explanation of the ability of A261T to suppress the held-up² phenotype as well as the lethality of the G200D mutation in combination with held-up². The opposing biochemical defects we uncovered provide insight into how mutations within the same gene can result in disparate mutant muscle phenotypes.

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REFERENCES