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Alternative N-terminal regions of *Drosophila* myosin heavy chain II regulate communication of the purine binding loop with the essential light chain

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**ABSTRACT** We investigated the biochemical and biophysical properties of one of the four alternative exon-encoded regions within the *Drosophila* myosin catalytic domain. This region is encoded by alternative exons 3a and 3b and includes part of the N-terminal β-barrel. Chimeric myosin constructs (IFI-3a and EMB-3b) were generated by exchanging the exon 3-encoded areas between native slow embryonic body wall (EMB) and fast indirect flight muscle myosin isoforms (IFI). We found that this exchange alters the kinetic properties of the myosin S1 head. The ADP release rate (*kD*) in the absence of actin is completely reversed for each chimera compared to the native isoforms. Steady-state data also suggest a reciprocal shift, with basal and actin-activated ATPase activity of IFI-3a showing reduced values compared to wild-type IFI, whereas for EMB-3b these values are increased compared to wild-type EMB. In the presence of actin, ADP affinity (*KAD*) is unchanged for IFI-3a, compared to IFI, but ADP-affinity for EMB-3b is increased, compared to EMB, and shifted towards IFI values. ATP-induced dissociation of acto-S1 (*K1k2*) is reduced for both exon 3 chimeras. Homology modeling, combined with a recently reported crystal structure for *Drosophila* EMB, indicate that the exon 3 encoded region in the myosin head is part of the communication pathway between the nucleotide binding pocket (purine-binding loop) and the essential light chain, emphasizing an important role for this variable N-terminal domain in regulating acto-myosin cross-bridge kinetics, in particular with respect to the force-sensing properties of myosin isoforms.

**Introduction**

Muscle myosin isoforms display a large variety in kinetic properties and force production, despite their sequences being highly conserved. The various isoforms of *Drosophila melanogaster* muscle myosin heavy chain (MHC) are encoded by a single gene (*Mhc*) and expressed using alternative splicing (1). The first four of the six alternative exon sets (exons 3, 7, 9 and 11) encode regions located in the myosin head domain (2) and the use of alternative domains in the myosin head allows for fine-tuning of myosin properties (see Figure 1A). Two native myosin isoforms that differ in
Myosin alternative N-terminal domains influence kinetics

all four alternative regions in the head domain display very different kinetic and mechanical properties. The EMB (embryonic) myosin isoform is found in the embryonic body wall muscle, which is used for slow locomotion of the larvae, whereas the IFI (indirect flight muscle isoform) myosin is present in the muscle that can generate very high wing beat frequencies and enables flight. Transgenic expression of EMB in the indirect flight muscle resulted in loss of flight ability (3) and subsequent studies using isolated muscle fibers and/or myosin proteins confirmed the striking differences in kinetic and mechanical properties between IFI and EMB myosin isoforms (4–6). Exchange of the variable regions between IFI and EMB has been used as a strategy to estimate the effect of each alternative domain on muscle myosin kinetics and mechanics. Here we focus on the variable region near the N-terminus of MHC, encoded by exon 3. Two alternative regions, encoded by exons 3a and 3b, are expressed in Drosophila myosin and their sequences are shown in Figure 1B. EMB contains the region encoded by exon 3a, while the exon 3b encoded sequence is normally expressed within IFI. Previous work found that exchange of the exon 3 regions between IFI and EMB significantly changes the steady-state kinetic properties of both Drosophila muscle myosin isoforms. Inserting the exon 3a area into IFI resulted in significantly reduced ATPase rates and $V_{\text{max}}$ for IFI-3a (compared to IFI), whereas inserting the exon 3b region into EMB had surprisingly little effect on ATPase and $V_{\text{max}}$ of EMB-3b (compared to EMB) (7). The same study showed that in vitro actin sliding velocity was increased for EMB-3b, although not restored to IFI levels, and unaltered for IFI-3a, compared to wild type.

Exchange of the exon 3 area also affects the mechanical properties of the indirect flight muscle. Flight ability of IFI-3a Drosophila is slightly decreased, compared to wild-type, whereas introduction of the exon 3b region in an embryonic background does not restore flight ability for EMB-3b Drosophila (7). A follow-up study found that IFI-3a Drosophila showed a reduction in both maximum power generation ($P_{\text{max}}$) and optimal frequency for power production ($f_{\text{max}}$), whereas for EMB-3b Drosophila an increase in both $P_{\text{max}}$ and $f_{\text{max}}$ was found (8). Based on these results it was suggested that the exon 3 region can influence at least two steps of the cross-bridge cycle independently, thereby fine-tuning myosin muscle kinetics for optimal force generation. To understand the biochemical kinetics of various steps in the cross-bridge cycle that are influenced by the exon 3 domain, we have now performed steady-state and transient kinetics measurements using the exon 3 chimeric myosin S1 isoforms (IFI-3a and EMB-3b). Our data show that exchange of this variable N-terminal area significantly alters the kinetic properties of the myosin S1 head. The catalytic activity $k_{\text{cat}}$ of the myosin S1 samples generated from all full-length molecules (IFI, EMB, EMB-3b, and IFI-3a) followed the trend of activities so far observed for the full-length molecules. Interestingly, the ADP release rate ($k_{\text{diss}}$) in the absence of actin is completely reversed for both chimeras, compared to their wild-type backbones. In the presence of actin, the ADP affinity ($K_{\text{AD}}$) is unchanged for IFI-3a, whereas ADP-affinity for EMB-3b is increased and shifted towards IFI values. ATP-induced dissociation of acto-S1 ($K_{\text{AD}}k_{\text{diss}}$) is reduced for both exon 3 chimeras. Detailed analysis of a recently published crystal structure for EMB (9) combined with homology modeling indicate that the exon 3 encoded area is part of the communication pathway between the nucleotide binding pocket (purine-binding loop) and the essential light chain, emphasizing an important role for exon 3 in regulating actomyosin cross-bridge kinetics, in particular with respect to the force-sensing properties of the myosin isoforms.

Results

ATP-induced dissociation of acto-S1 ($K_{\text{AD}}k_{\text{diss}}$) is reduced for both exon 3 mutants

The ATP-induced dissociation of the acto-S1 complex was measured as described previously (10) using the flash photolysis method. Plotting the observed rate constant $k_{\text{obs}}$ versus ATP concentration allows the apparent second-order rate constant $K_{\text{AD}}k_{\text{diss}}$ to be determined from the slope of the graph. Example traces of the measured light scattering data are shown in Figure 2A for EMB-3b after ATP release. The analyzed data are depicted in
Figure 2B for the two exon 3 chimeras, together with IFI and EMB wild type (see also Table 1). The measured $K{\cdot}k_\text{obs}$ value of EMB-3b ($0.71\pm 0.12 \, \mu \text{M}^{-1} \text{s}^{-1}$) is significantly lower compared to EMB ($p<0.007$, Table 1) but not significantly different from the value previously recorded for IFI ($0.75\pm 0.08 \, \mu \text{M}^{-1} \text{s}^{-1}$). For IFI-3a, a significantly lower $K{\cdot}k_\text{obs}$ value ($0.66\pm 0.08 \, \mu \text{M}^{-1} \text{s}^{-1}$) compared to IFI was recorded using an unpaired t test ($p<0.04$, Table 1), although this difference did not reach statistical significance using a more stringent one-way Welch’s ANOVA that compared all four constructs. Thus, introduction of the IFI version of the exon 3 region into EMB shifted the ATP-induced dissociation of EMB-3b towards IFI values, whereas the presence of the EMB version of the exon 3 region did not shift IFI-3a towards wildtype EMB levels.

**ADP-affinity ($K_{AD}$) is unchanged for IFI-3a but increased for EMB-3b compared to wildtype**

The ADP-affinity for S1 in the presence of actin, described by the equilibrium dissociation constant $K_{AD}$, was determined according to established methods (5). ATP-induced dissociation of acto-S1 was measured in the presence of increasing amounts of ADP. Figure 2C shows example traces of the light scattering data for EMB-3b, recorded with increasing ADP concentration. The amplitude of the signal drops as the ADP concentration increases. From the light scattering signals, $k_\text{obs}$ can be obtained by fitting to a single exponential and plotting $k_\text{obs}$ versus ADP-concentration, which allows $K_{AD}$ to be estimated. The data show that the measured value for the embryonic chimera EMB-3b ($K_{AD} = 496 \pm 79 \, \mu \text{M}$) is significantly lower, as assessed by t test, compared to the wild-type EMB value ($K_{AD} = 587 \pm 48 \, \mu \text{M}$), suggesting a tighter ADP-binding affinity for EMB-3b (Figure 2C), although this did not reach statistical significance using Welch’s one-way ANOVA. The measured value of $K_{AD}$ for IFI-3a ($K_{AD} = 409\pm 18 \, \mu \text{M}$) is not significantly different from the wild-type IFI value (see Table 1).

**The ADP-release rate of S1 ($k_D$) is reversed for both IFI-3a and EMB-3b chimeras**

To estimate the rate constant of ADP dissociation from S1 in the absence of actin, the change in fluorescence of a coumarin-labelled ADP analog (eda-deac ADP, (11) was measured upon displacement of eda-deac ADP (cmADP) by ATP-binding to S1 (12). A single laser flash released 15-20 µM ATP from cATP (100 µM) and the fluorescence change resulting from cmADP release was recorded (Figure 3). The fluorescence signal could be fitted to a single exponential, which gives the ADP-release rate ($k_D$). The small increase in fluorescence at the start of the transient is an artefact resulting from the laser flash which is difficult to eliminate entirely when working with low fluorescence signals as used here. Note the direction of the change is the same in both transients.

The two exon 3 chimeras, IFI-3a and EMB-3b, show an almost complete reversal of their $k_D$ values compared to their wild-type counterparts (Table 1). Insertion of the embryonic exon 3a area into IFI results in an ADP release rate typically found in wild-type EMB ($k_D = 2.1 \pm 0.41 \, \text{s}^{-1}$), whereas the introduction of the fast IFI exon 3b region into EMB increases the ADP-release rate of EMB3b to wild-type IFI levels ($k_D = 7.0 \pm 1.3 \, \text{s}^{-1}$). Reversal of $k_D$ rates after exon exchange was previously found for one of the other variable domains, the relay loop encoded by exon 9, as introduction of the exon 9a domain into IFI reduced the ADP-release rate to embryonic levels, whereas the substitution of the exon 9b region into EMB increased the ADP-release rate to IFI-levels (10).

The directionality of the observed fluorescence traces is variable and can even be reversed as seen here for EMB-3b. The fluorescence trace of IFI-3a shows a signal decrease after release of cmADP, similar to what was seen previously for IFI and EMB (5). However, EMB-3b fluorescence increases upon cmADP release and this reversal of the fluorescence signal has been observed in other studies. We previously reported a similar reversal of fluorescence change for a series of probes and myosin isoforms (12) and also for other chimeric Drosophila myosin isoforms after exchange of exon 7 (13) and exchange of exon 9 (10). We currently have no explanation of why this occurs, except that this must reflect a change in the environment of the fluorescent probe, that will require analysis of a highresolution structure to resolve.
Catalytic activity is reduced for IFI-3a and increased for EMB-3b

Basal and actin-activated ATPase activity of wild-type IFI S1 and the two exon 3 mutants were measured (see Figure 4 and Table 2). IFI-3a S1 showed a significant reduction in basal Ca-ATPase (4.92±1.05 s$^{-1}$) compared to wild-type IFI S1 (9.79±1.39 s$^{-1}$), whereas CaATPase activity of EMB-3b S1 (5.16±2.01 s$^{-1}$) is significantly increased compared to values for wild-type EMB (1.83±0.08 s$^{-1}$) (Figure 4A and Table 2). The same trends were seen for basal Mg-ATPase activity, as well as actin-activated ATPase values ($V_{\text{max}}$), although differences did not reach statistical significance (Table 2 and Figure 4B/C). While previous CaATPase, Mg-ATPase and $V_{\text{max}}$ measurements for full-length myosin (7) showed significantly reduced values for IFI-3a compared to IFI wildtype, the full-length protein showed an increase only in Ca-ATPase levels for EMB-3b compared to EMB.

Using flash photolysis, the turnover number ($k_{\text{cat}}$) was measured at a fixed actin concentration for S1 of both exon 3 exchange mutants and compared with IFI wild type (Figure 4D). The time taken to hydrolyze all released ATP ($t_{\text{cat}}$) was estimated from the time at which the dissociation reaction was 50% complete ($t_{\text{diss}}$) to the time for 50% recovery of light scattering ($t_{\text{max}}$). This time period ($t_{\text{cat}}$) is linearly dependent upon the amount of released ATP (see Figure 4D). The steady-state rate of ATP hydrolysis is the inverse of the slope (steady-state rate = [ATP]/$t_{\text{cat}}$), allowing the catalytic activity ($k_{\text{cat}}$) to be determined. For IFI a steady-state rate of 0.104 (μM s$^{-1}$) was found, resulting in a $k_{\text{cat}}$ of 0.18 s$^{-1}$. This value is essentially the same as the $k_{\text{cat}}$ value reported previously for IFI S1 using the same experimental setup ($k_{\text{cat}}$ = 0.17±0.006 s$^{-1}$) (10).

The catalytic activity of EMB-3b ($k_{\text{cat}}$ = 0.021±0.009 s$^{-1}$) is nearly identical to the value measured previously for EMB (0.028 s$^{-1}$), whereas the $k_{\text{cat}}$ value for IFI-3a (0.057±0.003 s$^{-1}$) shows a decrease compared to IFI (10) (see Table 1). Previously reported $V_{\text{max}}$ values for full length myosin constructs showed similar behavior (7), with unchanged $V_{\text{max}}$ values for EMB-3b compared to EMB, and a 2.36-fold decrease in $V_{\text{max}}$ for IFI-3a compared to IFI.

Homology models and general description of the exon 3 encoded area

Sequence alignment of various myosin isoforms show that the exon 3 area is highly variable, with only five residues that are fully conserved: Asp$^{90}$, Asn$^{105}$, Arg$^{109}$, Ile$^{115}$ and Tyr$^{116}$ (Figure S1). The recently published crystal structure of embryonic Drosophila myosin in the rigor-like conformation (PDB 5W1A) shows that the exon 3 region (residues 69-116) starts with two β-strands (β4:69-73 and β5:77-79), which form part of the SH3-like domain found in many other myosins (Figure 5A). The β4 and β5 strands are followed by two short helices (HA: 83-85 and HB: 91-93) and a longer helix C (residues 98-111). The Cterminal residues of the exon 3 area are at the start of the next β-strand (S1B, Figure 5B) which forms the first β-strand of the central seven-stranded β-sheet in the myosin head.

Conserved residue Asp$^{90}$ is located between exon 3 helix A and helix B, residues Asn$^{105}$ and Arg$^{109}$ are located on the same side of helix C and Ile$^{115}$ and Tyr$^{116}$ are part of the first β-strand of the central 7-stranded β-sheet, a conserved feature of all myosins. The EMB crystal structure shows that Asp$^{90}$ forms a strong salt-bridge with Arg$^{148}$ and that the latter also forms a hydrogen-bond with Gly$^{120}$, located between the first two β-strands of the central 7stranded β-sheet (Figure 5B). Conserved residue Asn$^{105}$ forms hydrogen bonds with the backbone oxygen of three residues close to Asp$^{90}$ (Lys$^{87}$, Ile/Ala$^{88}$ and Met$^{91}$), whereas conserved residue Arg$^{109}$ contacts Tyr$^{116}$.

The available crystal structure of embryonic Drosophila myosin in the rigor-like conformation (PDB 5W1A) was used as a template to generate a homology model for IFI myosin. Overlay of the IFI model and the EMB crystal structure showed that the backbone topology for both is very similar (rmsd of 1.59 ˚A). Inspecting the homology models using scallop myosin templates that represent conformations at different steps of the mechanochemical cycle shows that some of these conserved interactions are maintained throughout the cycle, whereas others are lost (see Discussion for details).
**Exon 3 interacts with the nucleotide binding site**

The kinetic data show that exchange of the exon 3 area alters the nucleotide binding properties of *Drosophila* myosins. A large body of research has established that nucleotide binding and processing in the active site of myosin involves four highly conserved loops: P-loop (GESGAGKT), switch 1 (AKTxxN(ND)NSSR), switch 2 (DIxGFE) and the purine binding loop (NPxxxxxy) (14). The exon 3 region (residues 69-116) could potentially alter nucleotide binding properties by interacting with any of these highly conserved loops. Inspection of the EMB crystal structure (Figure 6A) shows that the purine binding loop (residues 127-135) interacts with the exon 3 region; for example, exon 3 residue Tyr110 forms a hydrogen bond with purine binding loop residue Asn127 (Figure 6B). Asn127 also has a hydrogen bond with the backbone carbonyl of P-loop residue Gly182. These contacts suggest a role for the exon 3 region of the myosin head in regulating nucleotide binding and release via interaction with the purine binding loop.

Homology models derived from scallop myosin structures at various states of the crossbridge cycle were generated for both IFI and EMB and used for further analysis, since the EMB crystal structure is only available for a rigor-like configuration that has no nucleotide bound. The homology models confirm the interaction between the purine-binding loop and the exon 3 area, and suggest this interaction depends on the myosin state involved (see Figure S2 for a detailed description of these interactions for different myosin conformations). However, the interactions between the purine binding loop and the exon 3 area are very similar for IFI and EMB (see Figure S2/S3 for details).

The EMB crystal structure shows that the exon 3 region has no direct contacts with any of the other variable domains in the myosin head. However, the SH1-SH2 helix is wedged between the exon 3 and exon 9 (relay loop) regions and makes contacts with both variable domains. Since Drosophila EMB and IFI share the same SH1-SH2 sequence, the two variable regions could potentially interact differently with this element, thereby altering the myosin properties. However, detailed analysis showed that contacts between exon 3 residues and the highly conserved SH1-SH2 region in the myosin head were also found to be very similar for both chimeras (see Figure S4 for details). In summary, the conserved interaction of the exon 3 region with both the nucleotide binding loop and the SH1-SH2 area suggest that alternative versions of exon 3 differentially affect myosin properties via another mechanism.

**Exon 3 interaction with the myosin essential light chain**

Part of the exon 3 area is embedded in the SH3 element, a β-barrel found at the N-terminus of the MHC. In vertebrates this SH3 element is thought to interact with the extended N-terminus of the myosin essential light chain (ELC) in the presence of actin, thereby modulating myosin ATPase kinetics (15). The EMB crystal structure does not show any direct contacts between the exon 3 region and the ELC, as the ELC of *Drosophila* lacks this N-terminal extension (16). However, two ELC residues (Glu104 and His107) form salt-bridge/Hbond contacts with two MHC residues located outside the SH3 domain (Arg25 and Ser29). The latter two are part of a small helix near the N-terminus of the MHC. Using the EMB crystal structure (Figure 6A, C). Another residue located on this small helix (Asp27) forms a strong salt-bridge with exon 3 residue Lys87 (Figure 6C). Lys87 is part of a network of highly conserved interactions in the exon 3 area, as it makes contacts with conserved residues Arg109 and Asn105 (see Figure 5B). Lys87 (backbone oxygen) also interacts with the sidechain of helix NT residue Arg24 (Figure 6C). Therefore, the exon 3 region is likely to be part of a communication pathway between the essential light chain and the purine binding loop, which also involves helix NT.

Homology models for IFI show a similar picture when using the EMB crystal structure 5W1A as a template, with no direct ELC-exon 3 region contacts, but indirect contacts via helix NT (Figure 7A). Homology models representing other states in the crossbridge cycle indicate that for IFI, helix NT maintains contacts with the exon 3 region throughout the cycle, whereas for EMB these contacts are lost. For instance, the pre-power stroke state of IFI (using 1QVI as template)
shows interactions between exon 3 residue Asn$^{112}$ and helix NT residue Arg$^{24}$ (Figure 7B). Exon 3 residue 112 is a variable residue, Ala$^{112}$ for EMB, and therefore the pre-power stroke state of EMB lacks this exon 3 contact with helix NT. The post-power stroke state of IFI (using 1KK8 as template) maintains the Asn$^{112}$-Arg$^{24}$ contact and has additional interactions between the sidechain of Arg$^{85}$ (helix NT) and the backbone oxygen of exon 3 residues Pro$^{83}$ and Lys$^{84}$ (Figure 7C). For EMB the post-power stroke state again does not show any contacts between exon 3 and helix NT, indicating the interaction between the exon 3 region and helix NT is fundamentally different between the IFI and EMB isoforms.

**Discussion**

The use of alternative domains in the myosin head is thought to play a role in finetuning the kinetic and mechanical properties of myosin isoforms (2), but the exact mechanisms whereby alternative structures affect these properties are not well understood. Herein we investigate the structure-function relationship of the area encoded by exon 3, a variable domain that forms part of the SH3-fold found near the N-terminus of the myosin head. Our results show that exchange of the exon 3 region between the indirect flight isoform (IFI) and the embryonic isoform (EMB) alters various kinetic properties of the generated myosin S1 chimeras (IFI-3a and EMB-3b). The most profound effect is on the ADP-release rates in the absence of actin, as the $k_D$ values for the chimeras are completely reversed, compared to their wildtype counterparts. In the presence of actin, the ADP affinity for IFI-3a is not affected compared to wild-type IFI. However, the ADP affinity of EMB-3b is significantly tighter, compared to wild-type EMB and shifts towards IFI values. ATP-induced dissociation is also affected after exchange of the exon 3 area between IFI and EMB, although the effect is less dramatic compared to ADP-release. Introduction of exon 3b into EMB shifted the ATP-induced dissociation towards IFI values, however, the insertion of the exon 3a region into an IFI background did not shift the ATP-induced dissociation towards wild-type EMB levels. The steady-state kinetic parameters measured for the IFI, EMB, and exon 3 chimeric S1 isoforms show a similar pattern for $k_{cat}$, with values of IFI-3a shifting towards the EMB donor isoform, but values for EMB-3b being very similar to EMB. While this pattern holds for IFI vs. IFI-3a values for $V_{max}$ and basal Ca- and Mg-ATPase, increases in these parameters were seen for EMB-3b relative to EMB.

**Correlation between $K_k$, $k_{AD}$ and motility**

It is generally accepted that ADP release is rate-limiting for slow sarcomere myosin isoforms (17). In contrast, for faster myosins, ATP-induced dissociation can be ratelimiting, depending on the particular conditions used (18) (19). Plotting the measured ADP affinity ($K_{AD}$) for EMB, EMB-3b, IFI and IFI3a against previously reported in vitro motility values, with and without the actin-binding protein troponymosin (Tpm) (7), shows a decrease in velocity with increasing $K_{AD}$ values (lower ADP affinity) (see Figure S5 and Table 3). This implies that ADP release is not ratelimiting for these *Drosophila* myosin isoforms, as the actin sliding velocity is higher for isoforms with higher ADP affinity. This agrees with previous biochemical studies that reported ADP affinity measured in solution did not correlate with actin sliding velocity of other *Drosophila* myosin isoforms (5, 10, 13). Mechanical studies on isolated *Drosophila* muscle fibers also found no correlation between ADP affinity and frequency of maximum work production (4, 8, 20, 21). Plotting in vitro motility versus ATP-induced dissociation ($K_{k_{cat}}$) for IFI, EMB and the two exon 3 chimeras, shows reduced motility with increasing $K_{k_{cat}}$ values, although the correlation is not particularly strong ($R^2 = 0.86$, Figure S5).

**ADP release ($k_{AD}$) is faster than ATP-induced dissociation of AM ($k_{dis}$)**

In order to estimate if either ADP release or ATP-induced dissociation could be rate limiting for these *Drosophila* myosins, the theoretical rate-limiting step $k_{min}$ was calculated using the equation $k_{min} = V/d$, assuming a working stroke ($d$) of 5 nm (18) and using the published motility data for velocity in the presence ($V^*$) and absence ($V$) of tropomyosin (Tm) (7). The calculated values for the ratelimiting step $k_{min}$ are listed in Table 3.
together with the motility data and estimated values for ADP release ($k_{AD}$) and ATP-induced dissociation ($k_{diss}$). The values associated with ATP-induced dissociation ($k_{diss,2}$) are within 2600-3600 s$^{-1}$, whereas values estimated for ADP release ($k_{AD}$) are within 4000-5900 s$^{-1}$. Thus, both ATP-induced dissociation and ADP release are significantly faster than the rate limiting step $k_{min}$ (800-1200 s$^{-1}$). Therefore, it seems unlikely that either ADP-release or ATP-induced dissociation is rate-limiting under these conditions.

At lower ATP concentrations, one predicts that $k_{diss}$ could be limiting the velocity for these myosins, in particular for IFI, which has an unusual low affinity for ATP, compared to EMB (20). At 1 mM ATP concentration, $k_{diss,1}$ is around 1300-1800 s$^{-1}$, close to the values calculated for $k_{min}$, indicating that at such low ATP concentrations $k_{diss}$ can indeed be ratelimiting for these myosins. At saturating ATP levels, another step in the cross-bridge cycle could become rate-limiting, for instance phosphate release (IFI) or an isomerization step before ADP release (EMB) as proposed by Swank et al. (20). However, EPR studies of Drosophila myosins, investigating the isomerization step between the open to closed conformation of the nucleotide binding pocket, found that the pocket is predominantly closed for both IFI and EMB with ADP bound. Therefore, this isomerization step is unlikely to control velocity for these fast myosin isoforms (22). Overall, our calculations suggest that ADP-release is too fast to be rate-limiting, but ATP-induced dissociation could be ratelimiting at lower [ATP].

**Biological implications**

Homology modeling, combined with the crystal structure for Drosophila EMB, indicate that the exon 3 encoded region in the myosin head is part of a communication pathway between the nucleotide binding pocket (purine-binding loop) and the essential light chain via helix NT, emphasizing an important role for this variable N-terminal domain in regulating acto-myosin cross-bridge kinetics. IFI maintains the interaction between helix NT and exon 3 during the cross-bridge cycle for different conformational states, whereas for EMB these interactions are lost. The presence of helix NT-exon 3 contacts throughout the cross-bridge cycle allows the IFI isoform to rapidly communicate any conformational change from the nucleotide binding site towards the ELC and vice versa, whereas for the EMB isoform this process is likely to be slower, due to the loss of exon 3 – helix NT contacts.

The presence of additional stabilizing interactions between the exon 3 region and helix NT for IFI throughout the cross-bridge cycle might be expected to stabilize the structure and make transitions between states more difficult thus reducing the reaction rates of its chemical-mechanical cycle compared to EMB, which is opposite to experimental data. In contrast to these expectations, the increased stability may restrict the number of conformational states the myosin head can adopt and actually improve its function. In this regard, a recent paper by Schmid and Hugel (23) explores this concept and introduces the idea of ‘conformational confinement of proteins’, in which the authors argue that restricting conformational states enhances the function of a protein "by limiting the non-productive degrees of freedom". For myosin IFI, the presence of helix NT-exon 3 contacts throughout the cross-bridge cycle may therefore ensure that the conformational states of myosin are confined to those that enhance its function.

If the exon 3 region plays an important role in the communication pathway between the essential light chain and the purine binding loop, via helix NT, any mutations along this route are expected to have a serious effect on the biological function of myosin. An extensive literature search (24) revealed that various cardiomyopathy mutations have been reported along this pathway, involving the exon 3 region, helix NT and the essential light chain. Mutations have been reported for exon 3 residues Pro$^{81}$, Ala$^{100}$, and Tyr$^{115}$ (the equivalent exon 3 encoded residues in Drosophila are Pro$^{82}$, Ser$^{101}$/Cys$^{101}$ and Tyr$^{116}$). Tyr$^{115}$ can interact with the purine binding loop (Tyr$^{116}$ in Figure S2 B/C) and therefore a mutation of Tyr$^{115}$ is expected to alter this interaction and affect the nucleotide binding properties of myosin. The helix NT equivalent in human βmyosin heavy chain also contains a cardiomyopathy site, Ala$^{26}$ (Glu$^{27}$ in both Drosophila EMB and IFI myosin) which could disrupt the signal transfer between the essential...
light chain and the purine binding loop. The essential light chain residue Arg\textsuperscript{154} (Gln\textsuperscript{106} equivalent in \textit{Drosophila}) can result in serious cardiomyopathy when mutated into His (25). Gln\textsuperscript{106} is close to ELC residues Glu\textsuperscript{104} and His\textsuperscript{107}, which both interact with the NT helix (Figure 6C). Based on kinetic measurements, and molecular modeling, we propose that these mutations alter myosin function by disrupting the relay pathway between the nucleotide binding pocket and the essential light chain. A recent structure of beta-cardiac myosin (PDB: 6FSA) shows a strong contact (salt-bridge) between exon 3 residue Asp\textsuperscript{85} and helix NT residue Arg\textsuperscript{29}, in addition to contacts between exon 3 residue Tyr\textsuperscript{115} and purine binding loop residue Tyr\textsuperscript{134} (26), strongly supporting our hypothesis that the exon 3 area is part of the relay pathway towards the purine binding loop via helix NT.

Disruption of the above-mentioned pathway does not only affect the cross-bridge kinetics but is also expected to alter the mechanical properties of myosin. The N-terminal region (NTR) of myosin has been identified as an important element in tuning the mechanical properties of myosin-I (27). Exchange of the NTR from Myo-1b, a highly tension-sensitive motor, onto Myo-1c, which is less tension-sensitive, converts Myo-1c into a highly tension-sensitive motor, resulting in sensitivity to forces < 2 pN. Overlay of the crystal structures of Myo-1b (PDB: 4L79)(28) and the EMB crystal structure (PDB 5W1A) shows a remarkable similarity between the secondary structure elements of the N-terminal region of EMB and Myo-1b, except for the SH3 domain of EMB (Figure 8). The NTR of Myo1b and a small helix immediately following the NTR are located near helix A and helix B of EMB, whereas the longer helix C of EMB nearly overlaps with an equivalent longer helix of Myo-1b (see Figure 8A/B). Overlay of EMB with Myo-1c shows a similar picture (Figure 8C) with the NTR and the first two helices of Myo-1c following the backbone topology of the exon 3 area secondary structure elements (helix A, B and C). The similarity between the Nterminals of EMB and the Myo-1b/Myo-1c structures suggests a similar functional role, implicating the exon 3 area as a mechanosensing element.

Recent crystal structures of cardiac myosin with the myosin activator omecamtiv mecarbil (OM) bound also hint at the possibility that the exon 3 area is involved in fine-tuning the mechano-sensing properties of myosin (29, 30). OM is a selective, small-molecule cardiac myosin activator that binds to the myosin head domain and can increase the power output of the cardiac muscle (31). OM is currently in clinical trials for the treatment of heart failure. Two cardiac myosin binding sites have been reported for OM (29, 30), one of which shows the OM binding site in a narrow cleft between the Nterminal domain and lower 50K domain (29). Three out of six N-terminal domain residues that interact with OM are located in the exon 3 area (A\textsuperscript{91}, M\textsuperscript{92}, L\textsuperscript{96}), whereas the other three residues are just outside the exon 3 area on the loop between the first (S1β) and second β-strand (S2β) of the seven-stranded β-sheet (S\textsuperscript{118}, G\textsuperscript{119} and F\textsuperscript{121}). Since OM can alter the power output of myosin, the involvement of exon 3 encoded residues in this process is another indication of this region’s potential in fine-tuning mechanical properties of myosin isoforms.

In conclusion, we find that the alternatively expressed domain encoded by exon 3 in muscle myosin is critical for optimal myosin performance. Various steps in the cross-bridge cycle are affected by exchange of the exon 3 regions, in particular ADP release and ATP binding. Based on the recently reported EMB crystal structure, combined with the homology modelling presented here, we propose that this exon 3 area is part of the communication pathway between the nucleotide binding pocket (purine-binding loop) and the essential light chain and plays an important role in modulating acto-myosin cross-bridge kinetics and loaddependent mechanics.

**Experimental Procedures**

**Generation of subfragment-1 (S1) from isolated full-length myosin**

Myosin was isolated from the indirect flight muscles of 140 wild-type (\textit{PwMhc2} transgenics), 200 EMB, 140 IFI-3a or 300 EMB-3b transgenic flies (those expressing the IFI, EMB, EMB-3b or IFI-3a myosin isoforms in the
indirect flight muscles, respectively) as previously described (6). The production of S1 by α-chymotrypsin digestion was carried out using a method based on Silva et al. (32) with the following modifications: the final myosin pellet was dissolved in digestion buffer (120 mM NaCl, 20 mM Na₂PO₄, pH 7.0, 1 mM EDTA, and 4 mM DTT). The myosin was incubated at 20 °C for 5 min to equilibrate, and then incubated with 0.2 mg/ml α-chymotrypsin for 6 min. To quench the reaction, phenylmethylsulfonyl fluoride (PMSF) was added to a final concentration of 1.5 mM. The reaction was subsequently centrifuged at 60,000 rpm (TLA 100.3 rotor) for 20 min in a Beckman ultracentrifuge to pellet the undigested myosin and myosin rods. The supernatant containing the S1 was removed and diluted to 1 ml with low salt buffer (30 mM KCl, 15 mM MgCl₂, 20 mM MOPS, pH 7.0 and 4 mM DTT). To concentrate the S1, the samples were centrifuged at 15,000 rpm in a Sorvall MC 12V microcentrifuge at 4 °C using a Millipore Ultrafree 0.5 µm centrifugal filter with a 5,000 kDa cut-off. The final volume of the supernatant was 30–40 µl, containing an S1 concentration of roughly 1.5 – 2 µg/[l]. S1 concentration was determined from the absorbance at 280 nm (E¹% = 0.73 cm⁻¹) and the molecular mass of 115 kDa.

Flash photolysis

Because of the small amounts of protein available, flash photolysis was used to measure the kinetics of the mutant Drosophila myosin S1 (5, 33). The ATP-induced dissociation of the acto-S1 complex was followed by measuring changes in the light scattering signal. The dissociation of nucleotide from S1 alone was detected by changes in fluorescence using fluorescently labelled analogues. In brief, the 20 µl sample was held in a quartz cuvette and ATP was released by a single 5 ns flash at 353 nm from a neodymiumyttrium-aluminum-garnet laser (Surelite I-10, 70 mJ maximum power) along the vertical axis of the cell at a rate of 90 s⁻¹ to start the reaction.

Both absorbance (to determine the ATP concentration) and light scattering (to monitor the acto-S1 complex) or fluorescence (to determine cmADP release) were measured from the cuvette simultaneously following the laser flash. White light > 389 nm was introduced to the sample from a 100-watt halogen lamp and the change in the amount of light scattered 90° was monitored after each flash. The absorbance at 405 nm was measured with a monochromator to determine the amount of ATP liberated from caged ATP during each laser flash. Coumarin fluorescence changes were detected by monitoring the emission through a 455 nm cutoff filter after excitation at 434 nm (75-watt xenon/mercury lamp).

All light scattering experiments were conducted in a low salt buffer (pH 7.0: 30 mM KCl, 5 mM MgCl₂, 20 mM MOPS and 4 mM DTT) with 1 µM actin, 1-3 µM S1, 500 µM cATP, 10 mM DTT and either apyrase (2 units/ml: ATP-induced dissociation of acto-S1) or ADP (various concentrations) and a glucosehexokinase system (0.03 units/ml hexokinase, 1 mM glucose and 100 µM Ap₃A: Kₐd determination). Each sample was subjected to multiple laser flashes. During Kₐd determination, ADP and cATP were added after each flash. The cmADP dissociation experiments were also performed in this low salt buffer and contained 4 µM S1, 10 µM cmATP (source of cmADP) and 100 µM cATP. For the determination of k₄, the acto-S1 sample, incubated without apyrase, was irradiated by a series of laser pulses of different intensities, which released a range of ATP concentrations. The time taken to hydrolyze all of the ATP (t₄cat) was estimated from the time at which the dissociation reaction was 50% complete (t₄diss) to the time for 50% recovery of the light scattering (t₄att) (33).

Analysis of the kinetic data

Equation 1 was derived from the interaction of actin and S1 with ATP and ADP shown in Scheme 1 and was used to determine Kₐd.

\[
[k_{\text{obs}} = K_k k_{2} ([\text{ATP}]/(1 + [\text{ADP}]/K_{\text{AD}})) \]  

(1)

where \(k_{\text{obs}}\) is the observed rate constant for the ATP-induced dissociation of acto-S1: \(K_k k_{2}\) is the second-order rate constant for ATP binding to acto-S1; \(K_{\text{AD}}\) is the equilibrium dissociation constant for the binding of ADP to acto-S1. The equation \(k_{\text{cat}} = k_{\text{obs}}/k_{0}\) was used to determine the relative rate constant (\(k_{\text{cat}}\)) shown in Figure 2D, where \(k_{0}\) is the value when [ADP] = 0. Plotting \(t_{\text{cat}}\) versus ATP concentration allows one to
estimate the steady-state rate of ATP hydrolysis according to Equation 2:

\[
\text{steady-state rate} = \frac{[\text{ATP}]}{t_{\text{cat}}} \quad (2)
\]

From which one can determine the \( k_{\text{cat}} \) (Equation 3):

\[
k_{\text{cat}} = \frac{\text{steady-state rate}}{[S1]} \quad (3)
\]

Values reported for flash photolysis studies are mean ± S.D. based on a minimum of 3 preparations, except \( k_{\text{cat}} \) which is based on two preparations and therefore \( k_{\text{cat}} \) is not included in the statistical analysis that was carried out by unpaired t tests and Welch's correction, with significant differences assumed at \( p<0.05 \). Significance was also assessed using one-way Welch’s ANOVA.

**Homology modelling**

Three-dimensional homology models were generated for the *Drosophila* IFI myosin motor domain and exon 3 chimeras using the SWISS-MODEL automatic comparative protein modeling server (35–37). The embryonic myosin crystal structure (PDB 5W1A) was used as a template to build a homology model of the IFI isoform, after pairwise alignment of the primary sequences of the *Drosophila* IFI and EMB S1 domains (96% conserved). To represent various states from the cross-bridge cycle, scallop myosin structures were also used as templates: PDB 1KK8 (actindetached state), 1QVI (pre-power stroke state) and 1S5G (near-rigor ADP-bound state) using the CLUSTALW alignment protocol. The alignments were submitted to the alignment interface of SWISS-MODEL. The chosen templates allowed us to generate 3D homology models of the IFI and EMB myosin heads and of the chimeras obtained after swapping the exon 3 region (IFI-3a and EMB-3a).

All data described in the manuscript are either contained within the manuscript or are to be shared upon request to corresponding author.

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this work, Aileen Knowles (San Diego State University) for help with Ca-ATPase data, Corey Dambacher, Jennifer Suggs and Floyd Sarsoza (San Diego State University) for dissecting indirect flight muscles and preparing S1 samples and Sam Lynn (University of Kent) for excellent technical support.

**Conflict of interest:** The authors declare that they have no conflicts of interest with the contents of this article.

**Author contributions:** MB performed flash photolysis and homology modelling work. KH did the S1 protein purification and ATPase assays. SB and MG designed the research. MB, KH, SB and MG contributed to data analysis and writing of the manuscript. All authors approved of the final manuscript before submission.

**References:**


**Abbreviations and nomenclature:**

- **MHC**: Myosin heavy chain
- **EMB**: Embryonic muscle myosin isoform in *Drosophila*
- **IFI**: Adult indirect flight muscle isoform of myosin
- **IFM**: Indirect flight muscle
- **ELC**: Essential light chain
- **myosinS1**: Myosin proteolytic subfragment-1
- **cmADP**: coumarin-ADP
- **cATP**: caged ATP
- **Tpm**: tropomyosin
- **NTR**: N-terminal region
- **OM**: omecamtiv mecarbil
- **PMSF**: phenylmethylsulfonyl fluoride
Figure 1: (A) Structure of the embryonic myosin isoform (EMB) in the rigor-like conformation, with the location of the four variable regions in the myosin head indicated: exon 3 region (green), exon 7 region (purple), exon 9 region (dark red) and exon 11 region (light brown). The essential light chain (ELC) is shown in dark-grey (the coordinates of the EMB crystal structure (PDB: 5W1A) were used to generate this image). (B) Alternative sequences encoded by the exon 3a and 3b regions, with nonconservative differences denoted by asterisks: 3a is found in the slow EMB isoform and 3b is present in the fast IFI isoform.
Myosin alternative N-terminal domains influence kinetics

Figure 2: ATP-induced dissociation ($K_{1+k_2}$) and ADP affinity ($K_{AD}$) of the two myosin S1 exon 3 chimeras. (A) Example of light scattering traces for acto-S1 dissociation with EMB-3b S1 (B) The second order rate constant ($K_{1+k_2}$) for the ATP-induced dissociation of S1 from actin is determined from a linear fit to the plot of the $k_{obs}$ vs. [ATP] (see Experimental Procedures). The linear fits yielded values of $0.71 \pm 0.12 \mu M^{-1} s^{-1}$ for EMB-3b (●) as compared to $0.91 \pm 0.13 \mu M^{-1} s^{-1}$ for EMB (○). For IFI-3a the linear fits yielded mean values of $0.66 \pm 0.08 \mu M^{-1} s^{-1}$ (■) as compared to $0.75 \pm 0.08 \mu M^{-1} s^{-1}$ for IFI S1 (□). (C) Light scattering traces for acto-S1 dissociation with EMB-3b S1 at various ADP concentrations. (D) Comparison of the affinity of ADP for acto-S1 ($K_{AD}$) for EMB and EMB-3b, with the relative $k_{obs}$ ($k_{rel}$) values shown. Hyperbolic fits resulted in $K_{AD}$ values of $587 \pm 48 \mu M$ for EMB (open circles) and $496 \pm 79 \mu M$ for EMB-3b S1 (closed circles).
Myosin alternative N-terminal domains influence kinetics

Figure 3: Rate of ADP release (k_D) from Drosophila S1 isoforms. The rate constant for cm.ADP dissociation (k_D) from S1 in the absence of actin was determined using flash photolysis. After release of ATP (15 µM) from caged-ATP (100 µM), a fluorescent ADP analogue (eda-deac ADP) bound to S1 was displaced by ATP. The change in fluorescence upon release of eda-deac ADP from S1 was used to determine k_D. Exchange of either the exon 3a or 3b domain resulted in a complete reversal of the ADP release rate (see also Table 1). The dissociation rate for EMB-3b (k_D: 7.0 s⁻¹) is not significantly different to IFI (k_D: 7.5 s⁻¹) whereas that of IFI-3a (k_D: 2.1 s⁻¹) is similar to EMB (k_D 1.8 s⁻¹).

Figure 4: Steady-state ATPase activity of IFI, EMB and exon 3 chimeric S1 isoforms. (A) Basal Ca-ATPase activity, (B) basal Mg-ATPase, (C) actin-activated Mg-ATPase activity (V_max) and (D) the turnover number (k_cat) for acto-S1 were determined as described in Experimental Procedures. Notations above histograms indicate the level of statistically significant differences (*, p < 0.05; **, p < 0.01; ***, p < 0.001; ****, p < 0.0001, ns, not statistically significant). Significant differences were assumed for p < 0.05.
Figure 5: Location and structure of exon 3 domain (green) within the myosin heavy chain N-terminus. (A) Secondary structure elements encoded by exon 3 (residues 69-116) include β4 (69-73), β5 (77-79), helix A (HA,83-85), helix B (HB, 91-93), helix C (HC, 98 -111) and the first β-strand of the seven-stranded β-sheet S1β (starts at 115, indicated with an arrow). The full SH3 element is also depicted (β1:Lys36-Glu43, β2: Glu45-Lys56, β3: Ile59-Gln65). (B) Conserved interactions within exon 3 regions involve residues Asp⁹⁰, Asn¹⁰⁵, Arg¹⁰⁹ and Tyr¹¹⁶. Asn¹⁰⁵ forms hydrogen bonds with backbone oxygens of three residues close to Asp⁹⁰ (Lys⁸⁷, Ile/Ala⁸⁸ and Met⁹¹) whereas Arg¹⁰⁹ forms H-bonds with Tyr¹¹⁶.
Figure 6: The exon 3 area is part of the communication pathway between the essential light chain and the purine binding loop. (A) Overview of elements in the communication pathway between the ELC (blue) and purine binding loop (127-135, orange), shown for rigor-like EMB myosin S1 (PDB: 5W1A). In addition to the exon 3 area (69-116, green), the small N-terminal helix (21-30, red) is also involved in signal transfer from the ELC towards the nucleotide binding pocket. (B) Detailed view of purine-binding loop interactions with the exon 3 area and the P-loop (PDB: 5W1A). Figure 6B is reused and extended in the supplementary section to demonstrate that interaction of the exon 3 region with the purine binding loop depends on conformational state of the myosin head (Figures S2/S3). (C) Close up of N-terminal helix interactions with the ELC (Glu$^{104}$ and His$^{107}$) and the exon 3 region (Lys$^{87}$) (PDB: 5W1A).
**Figure 7: Exon 3 interaction with helix NT is distinct between IFI and EMB and depends on conformational state.** Comparison of helix NT interaction with exon 3 residues and the ELC throughout the crossbridge cycle for IFI and EMB myosin isoforms, with exon 3 area (residues 69-116) in green, the NT-helix (residues 21-30) in red and the essential light chain (ELC) in blue. (A) Near rigor state: EMB crystal structure (PDB 5W1A) and corresponding IFI homology model show similar interaction of helix NT with exon 3 region and ELC (B) Pre-power stroke state: interaction of helix NT (Arg24) with the exon 3 area is present for IFI (Asn112) whereas for EMB (Ala112) this interaction is absent (PDB template 1QVI). (C) Post-power stroke state: IFI maintains contacts between the sidechains of variable exon 3 residue Asn112 and helix NT residue Arg25. Additional contacts are found for exon 3 residues Pro83 and Lys84 (backbone oxygen) with helix NT residue Arg25(sidechain). No exon 3 - helix NT contacts were found for EMB (PDB template 1KK8).
Myosin alternative N-terminal domains influence kinetics

Figure 8: Overlay of N-terminal regions of EMB and Myo-1b or Myo-1c. (A-B) EMB (grey) with the exon 3 area (green) and the N-terminal helix (NT) shown in red, Myo-1b (purple) and Myo-1c (yellow). Note the similar orientation of exon 3 secondary structure elements (helix B and C) of EMB with respect to the N-terminal region (NTR) of Myo-1b and/or Myo-1c in both nearrigor and pre-power stroke state. (A) Overlay of crystal structures of Myo-1b (4L79) with EMB (rigorlike, 5W1A). (B) Overlay of Myo-1b (6C1D) and Myo-1c (4BYF) with EMB homology model (prepower stroke, 1QVI template). (C) Sequence alignment of N-terminal regions of EMB and Myo-1b showing conserved secondary structure elements (HB and HC) for the exon-3 encoded region (EMB) and the NTR (Myo-1b).

Table 1: Kinetic parameters measured for myosin S1 isoforms and the exon 3 chimeras using flash photolysis. Values are mean ± S.D. based on a minimum of 3 preparations except $k_{cat}$ (n=2). $K_1k_{+2}$ is the second order rate constant for ATP-induced dissociation of acto-S1. $K_D$ and $K_{AD}$ are dissociation equilibrium constants determined by division of the dissociation rate constant by the association rate constant (e.g. $K_D = k_D / k_{+2}$). $k_D$ and $k_{AD}$ are the ADP dissociation rate constants in the absence and presence of actin, respectively. $K_{AD} / K_D$ is the thermodynamic coupling constant describing the relationship between actin and ADP affinities. $k_{cat}$ is the catalytic activity.
Myosin alternative N-terminal domains influence kinetics

Table 2: Steady-state kinetic parameters measured for wild-type IFI, IFI-3a, EMB and EMB-3b Drosophila myosin S1. Values are mean ± S.D. based on a minimum of four preparations (except for EMB Ca-ATPase, which is 3).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>IFI†</th>
<th>EMB†</th>
<th>IFI-3a</th>
<th>EMB-3b</th>
</tr>
</thead>
<tbody>
<tr>
<td>( K_{i}k_{+2} ) (( \mu M \cdot s^{-1} ))</td>
<td>0.75±0.08</td>
<td>0.91±0.13</td>
<td>0.66±0.08(^a)</td>
<td>0.71±0.12(^b)</td>
</tr>
<tr>
<td>( K_{AD} ) (( \mu M ))</td>
<td>409±26</td>
<td>587±48</td>
<td>409±18</td>
<td>496±79(^c)</td>
</tr>
<tr>
<td>( k_{AD} ) (s(^{-1}))</td>
<td>4090</td>
<td>5870</td>
<td>4090</td>
<td>4960</td>
</tr>
<tr>
<td>( K_{D} ) (( \mu M ))(^e)</td>
<td>7.5</td>
<td>1.8</td>
<td>2.1</td>
<td>7.0</td>
</tr>
<tr>
<td>( k_{D} ) (s(^{-1}))</td>
<td>7.5±1.3</td>
<td>1.8±0.3</td>
<td>2.1±0.41(^f)</td>
<td>7.0±1.3(^g)</td>
</tr>
<tr>
<td>( K_{AD}/K_{D} )</td>
<td>54.5</td>
<td>326</td>
<td>194</td>
<td>70.9</td>
</tr>
<tr>
<td>( k_{cat} ) (s(^{-1}))</td>
<td>0.170±0.006(^i)</td>
<td>0.028±0.006(^i)</td>
<td>0.057±0.003</td>
<td>0.021±0.009</td>
</tr>
</tbody>
</table>

\(^a\) Statistically different compared to IFI, p<0.04 by t test, but did not reach statistical significance using Welch’s one-way ANOVA.
\(^b\) Statistically different compared to EMB, p<0.007.
\(^c\) Statistically different compared to EMB, p<0.04 by t test, but did not reach statistical significance using Welch’s one-way ANOVA.
\(^d\) Data are estimated from \( K_{AD} \) assuming an association rate constant of 10\(^7\) M\(^{-1}\) s\(^{-1}\).
\(^e\) Data are estimated from \( k_{D} \) assuming an association rate constant of 10\(^6\) M\(^{-1}\) s\(^{-1}\).
\(^f\) Statistically different compared to IFI, p<0.0001.
\(^g\) Statistically different compared to EMB, p<0.0001.
\(^h\) Actin is not at saturating conditions.
\(^i\) Data from (10)
Myosin alternative N-terminal domains influence kinetics

<table>
<thead>
<tr>
<th>Myosin S1</th>
<th>Basal Ca-ATPase (s⁻¹)</th>
<th>Basal Mg-ATPase (s⁻¹)</th>
<th>Vₘₐₓ (s⁻¹)</th>
<th>K (μM) m</th>
</tr>
</thead>
<tbody>
<tr>
<td>IFI</td>
<td>9.79±1.39</td>
<td>0.092±0.035</td>
<td>1.25±0.29</td>
<td>3.26± 2.36</td>
</tr>
<tr>
<td>IFI-3a</td>
<td>4.92±1.05</td>
<td>0.066± 0.047</td>
<td>1.07±0.26</td>
<td>2.38±0.91</td>
</tr>
<tr>
<td>EMB-3b</td>
<td>5.16±2.01</td>
<td>0.053±0.039</td>
<td>0.83±0.37</td>
<td>2.95±2.95</td>
</tr>
<tr>
<td>EMB</td>
<td>1.83±0.08</td>
<td>0.016±0.002†</td>
<td>0.67±0.06†</td>
<td>2.54±0.46†</td>
</tr>
</tbody>
</table>

*Data from (10)

a Statistically different compared to IFI, p < 0.0001  
b Statistically different compared to EMB, p < 0.05
Myosin alternative N-terminal domains influence kinetics

<table>
<thead>
<tr>
<th>Isoform</th>
<th>Velocity (-Tpm) (μm/s)①</th>
<th>Velocity (+Tpm) (μm/s)①</th>
<th>$K_A D$ (μM)</th>
<th>$K_{i+k+2}$ (μM·s⁻¹)</th>
<th>$k_{AD}$ (s⁻¹)</th>
<th>$k_{min,5}$ (s⁻¹)</th>
<th>$k_{diss,2}$ (s⁻¹)</th>
<th>$k_{diss,1}$ (s⁻¹)</th>
</tr>
</thead>
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<tr>
<td>IFI-3a</td>
<td>6.5</td>
<td>6.2</td>
<td>409</td>
<td>0.66</td>
<td>1.32</td>
<td>4090</td>
<td>1300</td>
<td>1240</td>
</tr>
<tr>
<td>IFI</td>
<td>6.4</td>
<td>5.7</td>
<td>409b</td>
<td>0.75②</td>
<td>1.50</td>
<td>4090</td>
<td>1280</td>
<td>1140</td>
</tr>
<tr>
<td>EMB-3b</td>
<td>3.8</td>
<td>5.2</td>
<td>496</td>
<td>0.71</td>
<td>1.42</td>
<td>4960</td>
<td>760</td>
<td>1040</td>
</tr>
<tr>
<td>EMB</td>
<td>0.7</td>
<td>4.0</td>
<td>587b</td>
<td>0.91②</td>
<td>1.82</td>
<td>5870</td>
<td>140</td>
<td>800</td>
</tr>
</tbody>
</table>

**Notes:**

- ① From (7) Tpm is tropomyosin. * indicates parameters determined in the presence of tropomyosin.
- ② From (5)
- ③ Value of $K_{i+k+2}$ corrected for inhibition of the ATP reaction by cATP (correction factor 2, (18)).
- ④ Assumes $k_{+.AD} = 10^7$ M⁻¹·s⁻¹ and $k_{AD} = K_{AD} · k_{+.AD}$. * Assumes $k_{diss} = K_{i+k+2} · [ATP]$ and [ATP] is 2 mM (ATP concentration used in motility assays).
- ⑤ Assumes $k_{diss} = K_{i+k+2} · [ATP]$ and [ATP] is 1 mM.
Scheme 1: The interaction of S1 with actin, ATP and ADP. M, A, T and D symbolize myosin S1, actin, ATP and ADP, respectively.

\[
\begin{align*}
AM + T & \xrightleftharpoons[K_{\text{AD}}]{K_{1}} A.M.T. \xrightarrow[k_{+2}]{k_{-2}} A-M.T \rightarrow A + MT \\
+D & \uparrow K_{\text{AD}} \downarrow
\end{align*}
\]

A.M.D.
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Alternative N-terminal regions of *Drosophila* myosin heavy chain II regulate communication of the purine binding loop with the essential light chain

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Running title: *Myosin alternative N-terminal domains influence kinetics*

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Figure S1: Sequence alignment of the N-termini of various myosins. Conserved residues in the exon 3-encoded region are labelled explicitly by rectangles. “*” indicates that the residues are identical in all sequences in the alignment. “:” represents conserved substitutions, whereas “.” indicates semiconserved substitutions. The corresponding human β-myosin cardiomyopathy sites in the exon 3-encoded domains (Pro81, Ala100 and Tyr115) and the short N-terminal helix (Ala26) are indicated with a ▲.

Interaction between the exon 3 region and the purine binding loop depends on conformational state of the myosin head (Figures S2/S3).
The rigor-like EMB crystal structure (5W1A, Figures S2A/S3A) has contacts between the exon 3 area (Tyr<sub>110</sub> and Lys<sub>113</sub>) and the purine binding loop (Asn<sub>127</sub> and Tyr<sub>132</sub>). In the pre-power stroke conformation (EMB-1QVI, figure S2B/S3B), the EMB homology structure maintains contacts between the exon 3 area (Tyr<sub>110</sub> and Tyr<sub>116</sub>) and the purine binding loop (Pro<sub>128</sub> and Tyr<sub>135</sub>). Homology models of the ADP-bound near rigor state (1S5G template, figure S2D/S3D) show that all direct contacts between the exon 3 area and the purine binding loop are lost, whereas in the post-power stroke conformation (EMB-1KK8, figure S2C/S3C) contacts are maintained between exon 3 (Lys<sup>113</sup>, Tyr<sup>116</sup>) and the purine binding loop (Tyr<sup>132</sup>, Tyr<sup>135</sup>). For IFI very similar contacts are found between exon 3 and the purine binding loop (see Figure S3 for summary of interactions). Taken together, the crystal structures and homology models suggest that exon 3 is involved in regulating the conformation of the purine binding loop, as the myosin head goes through the various conformational states during the crossbridge cycle. However, the interactions between the purine binding loop and the exon 3 area are very similar for IFI and EMB.

Figure S2: Overview of exon 3 region – purine-binding loop contacts throughout the cross-bridge cycle for EMB. (A) In the near-rigor state exon 3 residue Tyr<sub>110</sub> interacts with the purine binding loop (PDB: 5W1A). (B) In the pre-power stroke state exon 3 residues Tyr<sub>110</sub> and Tyr<sub>116</sub> both interact with the purine-binding loop (1QVI used as template). (C) In the post-power stroke state exon 3 residues Lys<sub>113</sub> and Tyr<sub>116</sub> contact the purine-binding loop (1KK8 used as template). (D) In the ADP-bound near-rigor state no contacts are seen between exon 3 and the purine-binding loop (1S5G used as template).
Figure S3 Overview of purine binding loop contacts for EMB and IFI. The purine binding loop is shown at the top of each panel (residues 127-135). Interacting EMB residues are shown below in black, IFI residues are in red. Shaded residues are in the exon 3 area. (A) **rigor-like state** using 5W1A EMB crystal structure as template for IFI. (B) **pre-power stroke state** using 1QVI as template for both EMB and IFI (C) **post-power stroke state** using 1KK8 as template (D) **ADP-bound near-rigor state** using 1S5G as template. In addition to contacts with exon 3 residues (Tyr110, Lys113 and Tyr116), the purine binding loop interacts with the P-loop (Gly182, Glu187) and the bound nucleotide (ADP).

**Interactions of the SH1-SH2 helix with the exon 3 region are conserved for IFI and EMB.**

The EMB crystal structure in the rigor-like state (5W1A) shows that the exon 3 region has no direct contacts with any of the other variable domains in the myosin head (Figure 1A). However, the SH1SH2 helix is wedged between the exon 3 and exon 9 (relay loop) regions and makes contacts with both variable domains. Since *Drosophila* EMB and IFI share the same SH1-SH2 sequence, the two variable regions could potentially interact differently with this element, thereby altering the myosin properties. An overview of interactions between SH1-SH2 and the exon 3/9 regions is summarized in Figure S4 (see below). In addition to conserved exon 3 residues L97 and N98, two variable residues between EMB and IFI (A92/S92 and M94/L94) interact with the SH1-SH2 region in the rigor-like state (left panel) and the pre-power stroke state (middle panel). However, for both residues it is the backbone oxygen that is involved in the contacts with the SH1-SH2 element, and thus not expected to significantly change the exon 3 – SH1-SH2 interaction. Overall for the three conformational states of the myosin molecule.
investigated here (near-rigor, pre-power stroke and post-power stroke state), the homology models show very similar interactions between the two Drosophila myosin isoforms, indicating that the interaction of exon 3 with SH1-SH2 is highly conserved for EMB and IFI.

**Figure S4: Interactions of the SH1-SH2 helix with the exon 3 region are conserved for IFI and EMB.** Residues of SH1-SH2 and the exon 9-encoded relay loop are shown in black for both EMB and IFI. Exon 3 residues are shown in black (EMB) and red (IFI). Left panel: Rigor-like state using the EMB crystal structure (5W1A) as template for IFI. Middle panel: pre-power stroke state for IFI and EMB homology models based on scallop crystal structure (1QVI). Right panel: Post-power stroke state of IFI and EMB homology models based on scallop crystal structure (1KK8).
Figure S5: Comparison of kinetic data with in vitro motility (IVM) as a function of $K_{AD}$ or $K_{1k_{2}}$ for Drosophila myosin isoforms IFI, IFI-3a, EMB and EMB-3b. Left two panels: $K_{AD}$ correlation with motility; linear fits give slopes of $-32 \pm 1$ ($R^2=0.99$) and $-88 \pm 15$ ($R^2=0.94$) in the absence (-Tpm) and presence (+Tpm) of tropomyosin, respectively. Right two panels: $K_{1k_{2}}$ correlation with motility; slopes of $-0.03 \pm 0.01$ ($R^2=0.72$) and $-0.11\pm0.03$ ($R^2=0.86$) without and with tropomyosin respectively were determined. IVM data is from Swank et al 2003.

References for supplementary materials: