Molecular basis of the C-terminal tail-to-tail assembly of the sarcomeric filament protein myomesin

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Abstract

Sarcomeric filament proteins display extraordinary properties in terms of protein length and mechanical elasticity, requiring specific anchoring and assembly mechanisms. In order to establish the molecular basis of terminal filament assembly, we have selected the sarcomeric M-band protein myomesin as a prototypic filament model. The crystal structure of the myomesin C-terminus, comprising a tandem array of two lg domains My12 and My13, reveals a dimeric end-to-end filament of 14.3 nm length. Although the two domains share the same fold, an unexpected rearrangement of one β -strand reveals how they are evolved into unrelated functions, terminal filament assembly (My13) and filament propagation (My12). The two domains are connected by a six-turn α -helix, of which two turns are void of any interactions with other protein parts. Thus, the overall structure of the assembled myomesin C-terminus resembles a three body beads-on-the-string model with potentially elastic properties. We predict that the found My12-helix-My13 domain topology may provide a structural template for the filament architecture of the entire C-terminal Ig domain array My9-My13 of myomesin.

Keywords: myomesin / muscle sarcomere / M band / filament / protein assembly.

Subject Category: Structural Biology / Cell & Tissue Architecture

Introduction

The sarcomeric units of skeletal and cardiac muscles contain one of the most complex multi-protein assemblies characterized to date. Many of these protein components, which function either as filaments or filament bridges, are composed of long arrays of immunoglobulin-like (Ig) and fibronectin type III (Fn) domains (Kenny et al., 1999; Li et al., 2002; Tskhovrebova and Trinick, 2003; Williams et al., 2003). Since many of these proteins actively contribute to the force generating apparatus of muscle cells, they need to have elastic properties that support reversible conformational changes in response to these forces. As a prerequisite, these filament systems need to be tightly assembled and anchored within stable subcellular structures, such as the Z-disk and M-band of the sarcomeric unit. These processes frequently take place at one or both termini of the respective sequences.

The giant muscle protein titin is a prototype protein for investigations aimed at unraveling the underlying molecular mechanisms for filament anchoring, assembly and elasticity (Lange et al., 2006; Sotomayor and Schulten, 2007; Watanabe et al., 2002). The molecular basis of the N-terminal telethoninmediated assembly of titin in the Z-disk has been recently elucidated (Zou et al., 2006). However, it is unlikely that this unique arrangement could serve as a model for the terminal assembly of other sarcomeric filaments. Biochemical data indicated that the C-terminal M-band portion of titin is crosslinked with myosin

and myomesin filaments by yet unknown molecular mechanisms (Obermann et al., 1996). The potential importance of these interactions has been underlined by recent studies on targeting the C-terminal M-band region of titin in embryonic stem cells or animals, revealing extensive defects in myofibrillogenesis and in the maturation and maintenance of intact sarcomeres (Weinert et al., 2006).

In order to unravel the general molecular principles in the assembly of long sarcomeric proteins, we have selected the M-band protein myomesin as a molecular model. Myomesin and M-protein resemble the giant muscle protein titin in terms of modular domain composition and structure (Labeit et al., 1997; Vinkemeier et al., 1993). The C-terminal regions of both proteins are composed of five modular immunoglobulin-like (Ig) domains interspersed at regular sequence intervals. While myomesin is generally found in all vertebrate striated muscle types, M-protein is specifically expressed during late embryonic heart development and appears to be absent in adult slow-twitch muscles, indicating that these two homologous proteins exhibit different functions (Obermann et al., 1996). An early model of the M-band displayed myomesin as a filament bridge connecting myosin with titin (Obermann et al., 1996). However, only recently has it been demonstrated that the C-terminal domain My13 is involved in myomesin homodimerization, supporting a functional role of the protein in the cross-linking of neighboring myosin filaments across the sarcomeric M-band (Lange et al., 2005). The assembly of C-terminal myomesin has been shown to be direct.

We have been able to crystallize a myomesin fragment that comprises the two Cterminal Ig domains My12 and My13. The crystal structure reveals how the myomesin C-terminus is end-to-end assembled and forms a molecular filament of 14.3 nm in length. The central dimeric interface is formed by β -strand A' of My13, leading to an intermolecular, antiparallel β -sheet arrangement. In the preceding domain My12, the equivalent β -strand associates with a different β sheet and is involved in filament propagation rather than assembly. The two Ig domains of each myomesin molecule are connected by a long α -helix, rendering an overall architecture of the assembled myomesin C-terminus that can be associated with a three beads-on-the-string model. The overall domain arrangement is different from other known sarcomeric filaments and could become a prototype model for future studies to unravel filament assembly, anchoring and elasticity.

Results

Overall structure of the C-terminal filament assembly of myomesin

We determined the crystal structure of the C-terminal domains My12My13 (residues 1459-1667) to a resolution of 2.24 Å (**Table 1, Figure 1)**. Experimental phases were obtained by the multiple anomalous dispersion (MAD) method, using a seleno-methionine substituted version of the protein. The structure comprises a dimeric assembly of two myomesin molecules, of which all except the N-terminal five residues of the myomesin fragment used for crystallization were interpreted. In addition, 248 ordered solvent molecules were found in the electron density.

The structure reveals an end-to-end dimeric assembly of the two My12My13 molecules, generating a filament-like shape with overall dimensions of 14.3 nm x 5.0 nm x 4.1 nm (**Figure 1**). The two My13 domains form an extensive antiparallel dimer interface, generating a central two-domain body within the overall architecture. In contrast, the two My12 domains are located at the two tips of the structure. Each of them is connected with the My13 domain of the same molecule by a long α -helix that is partly void of any interactions with the two adjacent My12 and My13 domains. Thus, an overall arrangement of three rigid bodies My12, My13/My13 and My12 is generated, resembling a model of three beads-on-the-string (**Figure 1B**).

Region variance analysis (Schneider, 2002) confirms this interpretation by identifying two hinges with limited flexibility in symmetric positions, situated at the C-termini of the two My12My13 connecting helices (**Figure 1C**). Comparison of relative domain/domain arrangements (Bork et al., 1996) reveals an almost identical tilt angle, with a domain rotation along the filament axis of 81° and 82° for the two My12My13 molecules, respectively. In contrast, the two twist angles, which define the angles of the axes of the two individual Ig domains, are 95° and 124°, revealing a substantial difference of 29° (**Figure 1C**). Because of these tilt angles, the arrangement of the domains My12 and My13 is far from being linear.

Divergent β -sheet arrangements in domains My12 and My13 support distinct functions

Both the My12 and My13 domains belong to the I-set category of the Ig-fold, which has previously been found in several Ig domains of the protein titin (Marino et al., 2005; Muller et al., 2007; Zou et al., 2006). Apart from local deviations in distinct loop regions, the structures of these domains are generally closely related (Marino et al., 2005; Muller et al., 2007). However, when the two My12 and My13 Ig domains are superimposed onto each other (Krissinel and Henrick, 2004), their coordinates differ by 1.92 Å (first molecule) and 1.76 Å (second molecule) r.m.s. deviation, thus revealing substantial structural differences (**Figure 2**). The resulting structure-based alignment comprises only 21% identical

amino acids in matching positions (**Figure 2A**). Direct structural comparison reveals that the N-termini of the two sequences cannot be aligned. Of note, those residues of the My12 domain (1461-1464) that approximately match β -strand A of My13 appear to be in a β -strand-like conformation, but their geometry is too irregular to be recognized as a β -strand. In addition, β -strand G of My13 is four residues longer than that in My12. There are also local conformational differences in loops BC, DE and FG, not allowing a structure-based alignment of the respective sequence segments. Collectively, these structural deviations lead to substantial differences of the overall shapes of the two domains (**Figure 3**), providing the molecular basis of divergent structural functions of My12 and My13 within the observed end-to-end filament assembly and propagation.

Analysis of the secondary structure topology of the two myomesin Ig domains reveals that β -strand A' in My12 is associated with a different Ig β -sheet (A'BED) than the equivalent β -strand in My13, which participates in the canonical I-set β sheet A'GFCC' (**Figure 2B**). However, no obvious reason for these structural differences could be discerned, especially considering the similarity in sequence specific interactions involving residues from β -strand A' in both domains (not shown). The only noticeable difference in My13 is the interaction between Thr1583 from β -strand A' and Ser1662 from β -strand G (**Figure 2A**). In contrast, Ser1470 from My12, which is equivalent to Thr1583 from My13, interacts with one of the main chain carbonyls of the My12My13 connecting helix (**Figure 4A**). Therefore, the structural swap of strand A' is most likely caused by the additional

interface of the My12 domain with the My12My13 connecting α -helix (details below). To date, this unusual β -sheet topology has only been described for the growth factor-binding d5 domain of several Trk receptors (Ultsch et al., 1999), but has not been found for any sarcomeric protein with long arrays of lg domains.

The role of the My12My13 connecting helix in C-terminal myomesin filament formation

The arrangement of the My12 and My13 lg domains in the C-terminal myomesin filament is defined by the connecting six-turn α -helix (1549-1570) (**Figures 1**, **2B).** The helix is amphiphilic over its entire length. The hydrophobic face of its Nterminal part contains an array of aromatic residues (Tyr1551, Tyr1555, Phe1558), followed by one charged residue (Lys1562). The side chains of these residues are at the same face of the helical wheel and provide the framework of an interface with residues from the My12 domain, which is in the order of 540 $Å^2$ (Figures 3A and 4A). In particular, Tyr1551 binds into a deep cleft of the My12 domain, which is formed by several residues from the swapped β -strand A' and the FG loop. The only additional polar interactions, observed in this interface, are two hydrogen bonds, connecting Lys1562 and Glu1520 from the EF loop, and the main chain carbonyl group of Ala1554 and the hydroxyl group of Ser1470 from the swapped β -strand A' of the My12 domain. The interface of the Nterminal part of the helix and the My12 domain is virtually identical in each of the two myomesin molecules, demonstrating a rigid arrangement.

In contrast, the C-terminal part of this connecting α-helix is not involved in lateral interface interactions with any of the two adjacent Ig domains, thus providing an exposed spring-like appearance of this part of the helix. Remarkably, the polar side chains of helix residues Gln1559, Arg1560, Gln1563, Lys1569 and Asn1570 point to the solvent without any interactions to other protein residues. However, an extensive network of capping interactions occurs at the C-terminus of the My12My13 connecting helix (**Figure 4B**). Critical contributions are provided by residues Arg1571, Arg1573, Trp1597, and Asn1650 from My13. Arg1573 is the central residue of a negative-positive-negative hydrogen bond network, involving Glu1568 from the C-terminus of the My12My13 connecting helix. The nature of the My13/helix interface provides a molecular rational for the observed structural deviations in the overall architecture of the myomesin dimer, in contrast to the rigid My12/helix interface (**Figure 1C**).

The molecular basis of the end-to-end myomesin filament assembly

The central dimeric assembly of the C-terminal myomesin filament is defined by a symmetric interface, formed by extensive interactions between the two My13 domains (**Figure 5**). The interface covers an area of about 710 Å² per molecule, corresponding to 12% of the entire My13 surface. It can be basically divided in two parts. The first interaction site is formed by an intermolecular, antiparallel β-sheet with C'CFGA'-A'GFCC' topology. The regular β-sheet hydrogen bond

pattern extends over residues 1581-1583 from both My13 domains and is symmetrically flanked by a pair of hydrogen bonds, involving the side chain of Asp1580. In addition, one ordered solvent molecule is found in the very centre of the intermolecular β -sheet. The second part of the interface is formed by equivalent residues of β -strands A and B from both My13 domains. The β strands of the second interaction site are, however, not properly oriented to support regular β -sheet interactions. Instead, four symmetrical pairs of separate hydrogen bond pairs are formed: two involving interactions between the Cterminus of β -strand A (Gly1577) and residues from β -strand B (Asn1589, Asn1591); one between residues of β -strand B (Asn1589, Thr1593); and a flanking hydrogen bond by Asp1580 and Lys1588. Moreover, this part of the interface is highly solvated, with a total of 2 x 5 ordered solvent pairs observed in the X-ray structure. The symmetric arrangement of solvent molecules in this interface is remarkable, considering the limited resolution of the structure and the lack of any non-crystallographic symmetry constraints or restraints imposed upon the structure during structure refinement.

Validation of the molecular architecture of the My12My13 dimer

Although the described myomesin dimeric assembly is the most prominent within the crystal form used for structure determination, other dimeric assemblies can also be generated in the crystal lattice (**Figure 6**). To validate the dimeric arrangement in solution, small angle X-ray scattering (SAXS) was employed. The scattering pattern computed from the My13-mediated dimer matches the experimental SAXS data with a χ^2 value of 1.58, while other potential dimeric assemblies generated from the crystal lattice yielded much poorer fits. Therefore, the SAXS analysis confirms the presence of an end-to-end filament dimer of the myomesin C-terminus that is assembled by the terminal My13 domains.

Further, we mutated two residues (Asp1580 and Lys1588) that form a direct hydrogen bond within the My13 dimerization interface (Figure 5) into residues with opposite charges (Asp1580, Lys1588). Since these two residues make critical contributions to the myomesin dimer interface, we expected that charge reversal of one would diminish or even abolish the entire binding interface. Size exclusion chromatography indeed indicated that both mutants elute with an apparent molecular weight of a monomer (**Figure 7A**), in contrast to wild-type myomesin. Moreover, SAXS data collected from the two mutants are most compatible with a monomeric association state (Figure 7B-C, Table 2). In addition to confirming the crystal structure of My12My13 dimer, the gel filtration and SAXS data obtained with the mutants reveal that the Asp1580-Lys1588 hydrogen bond within the My13 dimer interface is essential for myomesin homodimerization. The analysis of these mutants also implies that the formation of the antiparallel β -sheet by β -strands A' from each molecule is not sufficient for constitutive dimer assembly.

It is noteworthy mentioning that in comparison to the K1588D mutant, the SAXS distance distribution function of D1580K is more skewed and suggests the presence of a residual fraction of dimer assembly (**Table 2**). However, given that the same mutant also shows an additional shoulder in the gel filtration (**Figure 7A**), we cannot rule out that the effect may be due to another, unknown abnormality of the solution properties of this mutant.

In an attempt to test for the dimerization ability of the different combinations of the myomesin C-terminus (wild-type, D1580K, K1588D) we also carried out pulldown experiments using polyhistidine-tagged versions of the three variants (**Figure 7D**). For all combinations with predicted charge repulsion in the interface (wild-type/D1580K; wild-type/K1588D; D1580K/D1580K; K1588D/K1588D), dimer binding was found to be either largely diminished or completely abolished. Intriguingly, the double mutant D1580K/K1588D was able to partially dimerize, indicating that the dual charge reversal in residues Asp1580 and Lys1588 may allow partial restoration of the My13 dimer interface.

Discussion

Ig-domain mediated filament assembly and propagation

The architecture of the C-terminal myomesin My12My13 dimer reveals the molecular basis of a novel type of terminal filament assembly and propagation by a sarcomeric model filament. Our structural findings are consistent with previous yeast two-hybrid interaction and biochemical data, demonstrating direct Cterminal assembly of myomesin *via* the My13 domain (Lange et al., 2005). However, taking into account that more than one half of a total of 13 myomesin domain modules consist of Ig domains, including two (My2, My3) within the Nterminal region and an array of five Ig domains (My9-My13) from the C-terminus of myomesin, the molecular origin of the specific ability of My13 for dimeric assembly remained elusive. Specific My13-mediated myomesin assembly also raises an important question about the underlying molecular parameters of sorting, avoiding unwanted fortuitous assembly with other Ig domains. Moreover, although myomesin filament cross-linking was considered to be similar to α actinin links (Lange et al., 2005), there have been no data available, to date, providing a molecular model of myomesin filament propagation from the Cterminal assembly.

The crystal structure of the C-terminal myomesin dimer reveals how two domains that both belong to the same I-set Ig domain category perform different functions within the context of a large structural protein, leading filament assembly (My13) and propagation (My12). In order to detect specific structural differences within the two Ig domains My12 and My13, we made use of previously defined structural markers of this fold (Chothia and Jones, 1997). One of the specific

features of the I-set family of Ig domains is the presence of an additional β -strand A'. In the canonical I-set arrangement, this strand interacts with the second β -sheet of A'GFC topology while the preceding β -strand A associates with the first Ig domain β -sheet of ABED topology.

Structural comparison of the My12 and My13 lg domains has identified the Nterminal sequence segment as the most divergent region (**Figure 2**). The difference is highlighted by an unexpected swapped association of β -strand A' in the myomesin My12 domain, using the My13 arrangement and other canonical Iset lg domains from titin (Marino et al., 2005; Muller et al., 2007) as reference. The different β -sheet association of this strand in domains My12 and My13 leads to distinct structural functions of residues from β -strand A'. While in My12, the strand is critically involved in the formation of an *intramolecular* interface with the My12My13 connecting α -helix, in My13 the same strand forms an *intermolecular* antiparallel β -sheet arrangement over six residues with the equivalent strand of the second myomesin My13 domain (**Figure 4, 5B**).

However, since this My13/My13 β -sheet is formed by main chain atom interactions, this type of interface on its own may not be sufficient to avoid fortuitous β -sheet assembly with other I-set Ig domains from myomesin or different sarcomeric filament proteins. The structure of the My12My13 dimer and biochemical validation data demonstrate that additional sequence specific

interactions, originating from residues on β-strands A and B, are essential for the formation of the My13 dimerization interface (**Figure 4B**). For instance, abolishment of the charged hydrogen bond by Asp1580 and Lys1588 impairs My13-mediated terminal myomesin dimer assembly (**Figure 7**).

In contrast, the swapped β -sheet association of β -strand A' in My12 does not support the formation of a My13-like dimer interface by the My12 domain. Thus, our present data demonstrate how specific structural rearrangements in domains with an identical fold can lead to different functional readouts, either My13mediated myomesin filament assembly or the formation of a specific helix interface by My12 that provides the structural basis of filament propagation within the My12My13 dimer. Moreover, comparison of the sequences of My12 and further preceding Ig domains from myomesin (My9, My10, My11) does not reveal any conservation of residues involved in the My13 dimer interface (data not shown), thus making it unlikely that any these domains may have the ability to form a My13-like dimer interface.

The My12My13 structure displays the propagation of single myomesin filaments, originating from the C-terminal My13 end-to-end assembly. Our data are in agreement with previous *in vivo* and biochemical data that point to the presence of single myomesin filament strands rather than oligomeric rod assemblies (Lange et al., 2005; Obermann et al., 1996). In contrast, even though Ig domain-mediated terminal assemblies have been demonstrated in filamentous proteins

like titin and different filamin isoforms (McCoy et al., 1999; Popowicz et al., 2004; Pudas et al., 2005; Zou et al., 2006) it is unlikely or unknown whether the antiparallel arrangement of the assembly site propagates throughout the remaining filament of these proteins. Based on structural data of human filamin C, a flexible hinge region near the dimerization module was proposed (Pudas et al., 2005). This model attempted to reconcile the observed antiparallel assembly of the C-terminal domain observed in the crystal structure with previous electron microscopy data that supported a parallel rod arrangement. However, in the Cterminal myomesin filament the structural rigidity of the Ig domain connecting α helical linker does not allow comparable bending, which could lead to a reversal of the overall filament orientation. The concept of single filament propagation for myomesin is also unrelated to the type of filament formation in α -actinin, which is oligometric and formed by α -helical coiled coil domains instead of lg domains (Djinovic-Carugo et al., 1999). Interestingly, the individual domains of the α actinin rod R1-R4 are also connected by α -helical linkers, which are, however, short and not exposed to the external solvent over several residues.

Does the My12My13 connecting helix function as an elastic spring?

Intensive research on the molecular parameters leading to passive elasticity of the giant muscle protein titin has revealed that the flexible "PEVK" regions within the I-band segment give rise to most of the protein's elastic properties, while the involvement of folded Ig or Fn domains remains controversial (Preetha et al., 2005; Tskhovrebova et al., 2005). Similar to titin, a recent model also postulated a molecular requirement for elastic properties in myomesin, in order to build flexible bridges to the titin and myosin filaments under substantial external force conditions *in vivo* (Agarkova and Perriard, 2005). Recent data, using singlemolecule force spectroscopy and transmission electron microscopy, demonstrated that the Embryonic Heart (EH)-segment within the central part of the EH-isoform of myomesin may act as an entropic spring (Bertoncini et al., 2005; Schoenauer et al., 2005). However, the entire C-terminal Ig domain array My9-My13 of myomesin, in the absence of high-resolution structural data, has not been investigated in terms of potential elastic properties of the protein, to date.

Nonetheless, the structure of the dimeric My12My13 filament provides an unexpected model to test molecular elasticity of the C-terminal part of myomesin. In contrast to other known Ig domain arrays with available 3D structures (Mrosek et al., 2007; Muller et al., 2007; Zou et al., 2006), domains My12 and My13 of myomesin are linked by an α -helix, of which about one half is void of any interactions with other protein residues of the myomesin C-terminus (**Figure 8**). This finding is remarkable considering that fully exposed α -helices are not commonly found as domain connecting elements in soluble proteins. Indeed, they are like a mechanical coiled spring, universally used in natural processes requiring elasticity (Chouaieb et al., 2006).

In contrast to amphiphilic or hydrophobic helices, which are generally either embedded in side-wise interactions with other protein residues or are assembled into coiled coils, the mechanical stability of a structurally separate helix, like that observed for the C-terminal part of the My12My13 connecting helix, is solely determined by the internal α -helical hydrogen bond pattern. Previous measurements revealed that only low mechanical forces below 30 pN are required to disrupt the internal hydrogen bond pattern (Choe and Sun, 2005; Root et al., 2006). Previous experimental and computational data, investigating the mechanical stability of the cytoskeletal protein spectrin are consistent with our hypotheses, as they demonstrate a distinct and reversible stretching-out pattern of α -helical linker regions under low external force conditions (Law et al., 2003; Rief et al., 1999). Considering the exposure of myomesin to external forces within sarcomeres (Agarkova and Perriard, 2005), the mechanic stability of such helical linker may be inferior to that of domain/domain arrangements investigated previously or comparable to the measured elastic properties of the EH-region (Bertoncini et al., 2005; Schoenauer et al., 2005). Similar investigations for the Cterminal myomesin filament, taking advantage of the structural knowledge of the domain/domain linkers in My12My13, are eagerly awaited. However, such studies will require an extension of the present analysis of the C-terminal My12My13 fragment into a larger array of C-terminal myomesin Ig domains.

My12My13 as model for a repetitive filament architecture in myomesin

Using a suite of secondary structure prediction programs (Rost et al., 2004), the length and the position of the My12My13-connecting helix can be accurately predicted with high confidence values (**Figure 8C**). Extending our analysis over the entire C-terminal half of the myomesin sequence, spanning five Ig domains (My9-My13), we predict a repetitive pattern of α -helices at basically identical sequence intervals of about 110 residues (**Figure 8C**), also with high confidence values. The predicted helices are consistently located in sequence segments between myomesin Ig-domains My9-My13. The prediction is supported by circular dichroism spectra of the myomesin fragments My12My13 and My9-My13 (**Figure 8C**). Both constructs show an significant α -helical content of about 20%. We are not aware about a similar pattern reported for any other sarcomeric protein.

Thus, the above findings support the emergence of an attractive filament model of a repetitive array of Ig domain/linker repeats, which may form the molecular basis of the filament architecture of myomesin over long distances (Lange et al., 2005; Obermann et al., 1996). Previous immunoelectron microscopy data found that the N-terminal part of myomesin is located at or close to the symmetric M4 and M4' lines of the sarcomeric M band, while domains of the C-terminal Ig domain array of myomesin are located between the two lines (Obermann et al., 1996). Recent biochemical data (Lange et al., 2005) and our present data, demonstrating C-terminal dimeric assembly of myomesin, therefore suggest that the C-terminal half of myomesin, spanning domains My9-My13, functions as a

bridging filament. Such filament would allow to connect protein components with myomesin interaction sites that have been found in the M4/M4' lines, such as creatine kinase (Hornemann et al., 2003) and titin (Obermann et al., 1997), across the central M line through the C-terminal myomesin filament.

A repetitive domain structure for My9-My13 is expected to lead to a dimeric myomesin end-to-end filament with a total of 4 x 2 lg domain/helix modules (My9, My10, My11, My12) and one central My13 dimerization unit. If we estimate each domain module to be in the order of 6 nm in length, using the My12My13 crystal structure and the SAXS distribution function as reference (**Figures 1 and 7C**), such myomesin My9-My13 filament would be about 55 nm long, assuming a fully extended arrangement of adjacent domains. However, since our structural data indicate a filament with twist angles that would considerably deviate from a linear arrangement (**Figure 1**), such a filament may approach the estimated M4/M4' distance of 44 nm in the relaxed sarcomere state.

At present, the lack of available experimental data of the C-terminal myomesin Ig domain array beyond the My12My13 fragment prevents further speculations. In order to confirm the molecular filament architecture of the entire C-terminal Ig array of myomesin, therefore, a combination of high resolution structural biology methods and solution methods, allowing molecular imaging *in vitro* and *in vivo*, will be required. Should the presence of an array of repetitive Ig domains-helix linker elements be experimentally confirmed, the C-terminal myomesin filament

could become a unique model system to study molecular mechanisms of protein elasticity mediated by long distance filament cross-linkers under physiological conditions.

Experimental Procedures

Protein expression and purification

The DNA sequence (Q6H969 HUMAN) encoding for myomesin domains My12My13 (residues 1459-1667) was amplified by PCR from existing constructs (Lange et al., 2005), using primers to generate XhoI and MluI sites at the 5' and 3' ends, respectively. The DNA fragments were digested and inserted into a modified pET22b(+) vector (Novagen) carrying an N-terminal polyhistidine-tag sequence and a TEV protease cleavage site. The My12My13 fragment and the versions of its two single point mutants D1580K and K1588D were overexpressed in the E. coli strain BL21(DE3) and induced with 1 mM IPTG. For production of the seleno-L-methionine (SeMet)-incorporated My12My13 domain construct, the corresponding expression vector was used to transform the methionine auxotroph *E. coli* strain B834(DE3). A preculture grown in LB broth medium was washed with M9 minimal medium and resuspended in 1.5 l of medium A (M9 supplied with 20 mM D-(+)-glucose, 1 mM MgCl₂, 0.3 mM CaCl₂, $4 \,\mu\text{M}$ biotin, 2.7 μM thiamine) and trace element solution (0.3 mM FeCl₃·6H₂O, 0.06 mM ZnCl₂, 0.008 mM CuCl₂·4H₂O, 0.004 mM CoCl₂·6H₂O, 0.016 mM

H₃BO₃, 0.0007 mM MnCl₂·6H₂O). The cells were grown at 310K for 3 hours (starvation step) prior to adding the L-amino acids mixture (40 μ g ml⁻¹ of each amino acid, except methionine) and SeMet (60 μ g ml⁻¹). When the culture entered the logarithmic phase of growth, it was induced with 1 mM IPTG, and grown overnight. All proteins were purified from crude cell extracts using Ni-NTA affinity chromatography, followed by removal of the hexahistidine-tag by TEV protease. The proteins were further purified by size exclusion chromatography, using a Superdex 200 column (16/60 or 10/30), equilibrated with 25 mM Tris/HCl, pH 7.5, and 150 mM NaCl. The *wild type and Se-Met* version of My12My13 eluted with an apparent molecular weight of approximately 48 kDa, matching the molecular weight of a dimer, while the single point mutants were eluted as monomers.

X-ray structure determination and interpretation

The myomesin My12My13 fragment was crystallized by the hanging-drop vapor diffusion technique. 1 μ l of about 10 mg·ml⁻¹ of protein solution was mixed with 1 μ l of 14% (w/v) PEG 20000, 0.1 M Bis-Tris, pH 7.5, and 0.16 M ammonium acetate, and suspended above a reservoir of the same solution. Crystals of a trigonal shape appeared the next day and grew to an optimal size of about 200 x 200 x 50 μ m³ within one week. They were identified to belong to space group C222₁, containing two copies of the molecule per asymmetric unit, with a solvent content of 56.5% (Matthews, 1968).

A multiple wavelength anomalous dispersion (MAD) experiment using the SeMet protein was carried out on the tunable wiggler beamline BW7A (EMBL/DESY, Hamburg, Germany) and a native data set was collected on the bending magnet beamline X11 (EMBL/DESY, Hamburg, Germany). Prior to data collection, the crystals were immersed in the mother liquid containing 18% [v/v] MPD, and flash cooled at 100 K under the cryostream attached to the goniometer. All data sets were integrated, scaled, and merged using the HKL suite (Otwinowski, 1997). The expected two selenium atoms were located using the heavy atom search routine of the CNS program suite (Brunger et al., 1998). These positions were refined; initial phases of the protein were calculated to 3.1 Å resolution. and density modification was applied, improving the figure of merit from 0.455 to 0.898 as defined in CNS (Brunger et al., 1998). The electron density map was of sufficient quality to fit the titin module M5 NMR model (PDB code: 1NCT) by phased molecular replacement (Murshudov et al., 1997) at the two positions of the domain My13. The remaining model was built manually using the program 'O' (Jones et al., 1991). Further refinement using molecular dynamics (CNS) resulted in a map where the entire protein model, except for the first five disordered residues, could be built. Solvent molecules were added by ARP/wARP (Perrakis et al., 2001) and the final cycles of refinement were carried out through REFMAC5, using TLS refinement of rigid groups (Winn et al., 2001). The data collection and refinement statistics are reported in **Table 1**.

A structure-based sequence alignment was computed by SSM (Krissinel and Henrick, 2004). Protein/protein interfaces were determined with the program AREAIMOL of the CCP4 suite (Collaborative Computational Project, 1994). Secondary structural elements were determined by PROCHECK (Laskowski et al., 1993).

Small angle X-ray scattering analysis

Measurements were carried out at the beamline X33 (EMBL/DESY, Hamburg, Germany). Each sample was exposed to X-rays of wavelength λ =1.5 Å for 3 minutes, and the signal was recorded on a MAR345 IP detector. The measurements were carried out at 290 K using a sample-detector distance of 2.7 m, with a momentum transfer range of 0.10 nm⁻¹ < s < 4.5 nm⁻¹ (here, s = 4π $\sin(\theta)/\lambda$ where 20 is the scattering angle). All myomesin samples were measured at, at least, two different concentrations with intermittent buffer solution (25 mM Tris/HCl, pH 7.5, and 150 mM NaCl). To monitor radiation damage, two consecutive 2-minute exposures at the highest protein concentration were compared. The data were processed using standard procedures, corrected for buffer contribution, and extrapolated to infinite dilution using the program PRIMUS (Konarev et al., 2003). The radius of gyration R_q and forward scattering I(0), the maximum particle dimension D_{max} , and the distance distribution function p(r) were evaluated using the program GNOM (Svergun, 1992). The molecular masses of the different constructs were calculated by comparing with the

reference BSA samples. The scattering patterns from the high-resolution models were calculated using the program CRYSOL (Svergun et al., 1995). The best fits in terms of mixtures of monomers and dimers for My12My13 mutants were computed using the program OLIGOMER (Konarev et al., 2003).

The discrepancy of the experimental data was calculated as:

$$\chi^{2} = \frac{1}{N-1} \sum_{j} \left[\frac{I_{\exp}(s_{j}) - cI_{calc}(s_{j})}{\sigma(s_{j})} \right]^{2}$$

where *N* is the number of experimental points, *c* is a scaling factor, and $I_{calc}(s)$ and $\sigma(s_j)$ are the calculated intensity and the experimental error at the momentum transfer s_j , respectively. The SAXS data statistics are summarized in **Table 2.**

Circular dichroism measurements

The myomesin fragments My12My13 and My9-My13 were dialyzed against 30 mM potassium phosphate, pH 7.5. The concentration of each sample was determined by measuring the absorbance of the proteins, which were diluted with 8M urea. Far-UV circular dichroism (CD) spectra were obtained on station 12.1 at the Synchrotron Radiation Source, Daresbury Laboratory, UK. Experiments were performed at a temperature 277 K and a wavelength range of 280-175 nm. The

spectra were analyzed with the CONTIN procedure (Provencher and Glockner, 1981).

Biochemical characterization

GFP-labeled myomesin My9-My13 constructs (residues 1134-1685) were generated via PCR and subcloned into the pEGFP-C1 vector (Clontech). Sitedirected mutagenesis to produce the D1580K and the K1588D mutants was carried out as per the Quickchange protocol (Stratagene). All constructs and mutants were verified by sequencing. For the pull-down assay, Cos-1 cells were transfected with the GFP constructs using Escort IV (Sigma), and expression was measured after 2 days. Cells were lysed in IP-buffer (150 mM NaCl, 10 mM Tris-HCI, pH 7.9, 1 mM DTT, 0.2% NP-40, 1x Roche Protease Inhibitor Cocktail), sonicated briefly, and centrifuged at 4°C to separate the soluble and insoluble cell fractions. The soluble Cos-lysate fraction was incubated with His-tagged My12My13(wild type), My12My13(D1580K), or My12My13(K1588D) bound to Ni-NTA beads (Qiagen) for 2 hours on ice. Beads were washed three times with icecold IP buffer and samples were processed for SDS-PAGE followed by immunoblotting. Visualization of GFP-tagged proteins was done using the monoclonal mouse anti-GFP antibody (Roche) followed by a horseradish peroxidase coupled rabbit anti-mouse antibody (DAKO).

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Accession Numbers

The coordinates and the structure factors were deposited in the Protein Data Bank with accession code 2R15.

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Nature, **439**, 229-233.

Figure Legends

Figure 1. Overall structure of the dimeric filament assembly of the Cterminal myomesin fragment My12My13. (A), Ribbon representation in two different orientations, defining the three rigid body segments: the central $(My13)_{2}$ assembly, and the two distal motifs, each comprising the My12 domain and the My12My13 connecting α -helix. The two molecules of the dimeric assembly are shown in different colors (orange-light orange / blue-light blue). The dimension of the overall structure and the three rigid body segments are indicated. Three specific interface regions, including My12/helix (A), My13/helix (B) and My13/My13 (C1, C2) are framed (right panel). The interfaces are further illustrated in Figures 4 and 5. (B), Representative σ_A weighted $2F_o$ - F_c electron density map contoured at 1σ , showing the sequence segments around residues Asp1580 and Lys1588, which form a hydrogen bond in the My13 dimer interface (cp. Figure 4) and which have been used for biochemical validation of the structure (cp. Figure 7). The structure is shown in atom-type colors, following the scheme of panel A for carbon atoms. (C) Superposition of the two myomesin molecules, using the coordinates of the My13 domains as common basis. It reveals a 29° difference of the twist angle that determines the relative rotation of My12 and My13 with respect to each other (Bork et al., 1996). The hinge is located at the C-terminus of the My12My13 connecting helix (green).

Figure 2: Sequence and structural relations of domains My12 and My13. (A) Structure based sequence alignment. Residue pairs that structurally match are shown in capitals, the remaining ones are in small characters. Those residues that are involved either in interfaces with the My12My13 connecting α -helix or in the My13 dimer assembly are highlighted in green and orange, respectively. The secondary structural elements are shown below the respective sequences of My12 and My13 (E, β -strand; H, α -helix). The coloring of structural elements indicates association with the first β -sheet (cyan, ABED) and second β -sheet (magenta, A'GFCC'). Please note that β -strand A is missing in My12, and β strand A' of My12 has swapped its β -sheet association, taking the canonical I-set arrangement of My13 as reference (Chothia and Jones, 1997). The My12My13 connecting α -helix is colored in green. The approximate consensus positions of the secondary structural elements, common to My12 and My13, are labeled on top of the alignment. (B), Ribbon presentation of the structural modules, My12, My13 and the My12My13 connecting α -helix. The color codes are as in panel (A), except for the My12My13 connecting helix, in which only the part of the ribbon presenting fully exposed residues is colored in green. The six turns of the helix are numbered.

Figure 3: Semitransparent surface presentation of My12 and My13. The interface areas with the My12My13 connecting helix and the My13 dimerization interface are colored in green and blue, respectively, demonstrating overlapping interface areas in My12 and My13. The ribbon of each domain is in grey, except

 β -strand A' (yellow) that swaps in My12 in terms of β -sheet association. Those structural elements that are involved in interface surfaces are labeled.

Figure 4: (A,B) My12/helix and (C) My13/helix interface. Residues of the My12My13 connecting α -helix are in green. Those side chains that are involved in one of these interfaces are shown and labeled. Hydrogen bonds are displayed by dashed lines. For the My12/helix interface, a surface presentation (panel A) is also shown to indicate the complementarity of the My12 surface to interact with residues from the N-terminal part of the My12y13 connecting helix. The two interfaces are labeled "A" and "B" in Figure 1A.

Figure 5: My13 dimerization interface. (A), Overview, showing the My13 domain assembly in two different orientations. Upper panel, A'-A' interface; lower panel, A/B-A/B interface. (B), Zoom. Left panel, A'-A' interface by intermolecular β -sheet interactions of β -strands A' of the two myomesin molecules. Right panel, A/B-A/B interface formed by interactions of several residues from β -strands A and B. The terminal residues of the sequence segments from β -strands A, A' and B are labeled. Side chains of those residues that are involved in interface interactions are shown. Hydrogen bonds are indicated by dashed lines. Ordered solvent molecules are displayed by red spheres. The color code has been adopted from Figure 1. Labels, referring to the second myomesin molecule of the dimer are indicated with asterisks. The two interfaces relate to those areas that are labeled C1 and C2 in Figure 1A.

Figure 6. SAXS characterization and validation of the dimeric My12My13 arrangement. (A), Distance distribution plots, computed for three possible Cterminal myomesin dimer arrangements (model-1, blue; model-2, brown; model-3, green) that can be generated from the crystal lattice. (B), Ribbon representations of the three models examined, using the colors of panel A for one molecule of each dimer. The inlet table lists some of the crystallographic model parameters and the agreement with the experimental SAXS data. Comparison with the experimental SAXS curve of *wt*My12My13 (cp. Figure 7, Table 2) demonstrates that the best fit is calculated for model-1, which serves as the basis of structural interpretation. Note that the distance distribution of this model contains an additional maximum at about 6 nm (arrow), reflecting the approximate distance between domains My12 and My13.

Figure 7: Validation of the My13 mediated dimer interface. (A), Size exclusion chromatography of *wt*My12My13 (blue), My12My13(D1580K) (orange), and My12My13(K1588D) (red), indicating that the interaction Asp1580-Lys1588 is essential for dimer formation. (B), Experimental SAXS data of *wt*My12My13 (blue), My12My13(D1580K) (orange), and My12My13(K1588D) (red) are shown by data points. They confirm that the interaction D1580-K1588 is essential for dimer formation (cp. Figure 4). The calculated curves of the My12-My13 crystal structure (dimer, blue; one protomer, black) are provided for comparison. (C), Distance distributions calculated from the scattering curves in panel B. For one of

the mutants (D1580K) the SAXS data, as indicated by the R_g and the p(r) functions, reveal residual dimer formation in the order of 20% (**Table 2**). (D), Pull down assays, using (i) wtMy12My13, (ii) My12My13(D1580K), (iii) My12My13(K1588D), indicating that charge reversal of either Asp1580 or Lys1588 impairs My12My13 dimer formation. In contrast, charge reversal in both residues selected for site-directed mutagenesis (D1580K, K1588D) partially restores the ability of myomesin to dimerize.

Figure 8: Prediction of a myomesin filament beads-on-the-string model, consisting of 2 x 5 lg domains that are connected by α -helical linkers and end-to-end C-terminal assembly. (A), Beads-on-the-string presentation of the crystal structure of the My12My13 dimeric assembly of the myomesin C-terminus (cp. Figure 1). (B), Schematic model of the My9-My13 C-terminal filament, in which adjacent domains are connected by α -helical linkers. (C), Prediction of α helical segments at repetitive sequence intervals, interspersed myomesin domains My9, My10, My11, My12 and My13, using PredictProtein (Rost et al., 2004). The starting residue number, the sequence interval with respect to the previous predicted α -helical segment and the sequence of each predicted α -helix segment are presented. Residues that are predicted to be α -helical by PROF (for details, see PredictProtein) are shown in capital characters, those that are predicted with a confidence level of at least 82% are shown by small characters. The predicted helix length, using the two categories is indicated (second number in parentheses). For comparison, the experimentally determined My12My13

connecting helix is highlighted in green. (D), Circular dichroism curves of My12My13 (blue) and My9-My13 (magenta). The estimates for secondary structural elements for My12My13 and My9-My13 are: helix, 0.23/0.19; strand, 0.26/0.36; turn, 0.18/0.13; unordered 0.33/0.34.

| Omerical | Native | Se-Met substituted | | |
|---|-------------|--------------------|-------------------|-------------|
| Crystal | | | | |
| Data collection | | | | |
| Space group | | C | C222 ₁ | |
| | 61.50 | 61.58 | | |
| Cell dimensions <i>a</i> , <i>b</i> , <i>c</i> (Å) | 87.87 | 88.49 | | |
| | 113.0 | 203.58 | | |
| | | Peak | Inflection | Remote |
| Wavelength | 0.8126 | 0.98214 | 0.98264 | 0.91837 |
| Desclution $(Å)$ | 20.0-2.24 | 15.0-3.06 | 15.0-3.06 | 15.0-3.20 |
| Resolution (A) | (2.32-2.24) | (3.11-3.06) | (3.11-3.06) | (3.25-3.20) |
| R _{merge} ^a (%) | 7.6 (38.2) | 4.8 (12.8) | 5.5 (16.8) | 6.6 (14.1) |
| <1/0 <i>1</i> > | 17.8 (3.7) | 21.6 (5.4) | 17.4 (3.9) | 13.9 (4.3) |
| Completeness (%) | 95.7 (93.2) | 94.4 (89.8) | 94.2 (66.8) | 95.8 (77.5) |
| Redundancy | 4.9 | 2.9 | 2.7 | 3.1 |
| Refinement | | | | |
| Resolution (Å) | 20.0-2.24 | | | |
| No. reflections | 25,014 | | | |
| R _{cryst} ^b / R _{free} | 19.7/22.3 | | | |
| No. atoms | | | | |
| Protein | 3,489 | | | |
| Glycerol | 24 | | | |
| Solvent | 247 | | | |
| Rms Δ bonds ^c (Å) | 0.012 | | | |
| Rms Δ angles ^c (°) | 1.380 | | | |
| Mean <i>B</i> factor (Å ²) | 44.7 | | | |
| Main-Chain Dihedral Angles (%) | | | | |
| Most favored | 90.6 | | | |
| Allowed | 9.1 | | | |
| Generously allowed | 0.3 | | | |

Table I. X-ray structure determination

Parentheses devote values for the highest resolution shell.

^a $R_{merge} = \sum |I - \langle I \rangle | / \sum I$; I, intensity.

^b $R_{cryst} = \sum |F_o - F_{calc}| / \sum F_o$; F_o , observed structure-factor amplitude; F_{calc} , calculated structure-factor amplitude.

^c Root-mean-square deviations from ideal values.

| Sample | My1213 | My1213 D1580K | My1213 K1588D |
|--|-----------------|------------------|------------------|
| Theoretical molecular mass (kDa) ^a | 23.4 | 23.4 | 23.4 |
| Experimental molecular mass (kDa) ^b | 47 ± 4 | 21 ± 2 | 20 ± 2 |
| Maximum size, d _{max} X-ray model (nm) ^c | 14.1 | 10.0 | 10.0 |
| Exp. maximum size, d_{max} Gnom (nm) ^d | 14±1.5 | 13±1.5 | 9±1 |
| Radius of gyration, R_{g} X-ray model (nm) ^d | 3.93 | 2.79 | 2.79 |
| Experimental radius of gyration R_g (nm) ^d | 4.04 ± 0.08 | 3.30 ± 0.05 | 2.77 ± 0.05 |
| Model fit, χ against raw data (monomer) | 5.4 | 1.16 | 0.90 |
| Model fit, χ against raw data (dimer) | 1.58 | 2.80 | 8.6 |
| Monomer/dimer fraction (%) ^e | 7/93 | 78/22 | 100/0 |

Table 2. SAXS data collection statistics of the myomesin My12My13 fragment and itsmutants D1580K and K1588D

^a Molecular mass of the monomeric construct calculated from the sequence.

^b Molecular mass from comparison with reference solutions of bovine serum albumin.

^c For the D1580K, K1588D mutants: Calculations against chain A of the crystallographic model.

^d Maximum particle dimension and radius of gyration calculated by indirect transformation of the scattering data using GNOM.

^e The volume fractions of are calculated by the best fits of the experimental data by the curves computed from monomeric and dimeric constructs. The accuracy of the estimate is about 5 %.



Rigid body 2 B С Rigid body 1 Rigid body 3 My12 N/ -5.0 nm------—4.1 nm—→

Figure 1, Pinotsis et al.





chain A



-





29 deg

Α

Sec.Str. Topology <-A> <-A'--> <--B---> My12 1461 --LSATDLKiqs---taeGIQLYSFVTYyvEDL Sec. Str. (My12) EEEE EEEEEEE My13 1571 raRV<mark>LGGLPdvvtiq</mark>eg<mark>kALNLT</mark>CNVWG<mark>d</mark>-PPP EEEEEEE EEEE EEEEE Sec. Str. (My13)

| Sec.Str. | . Toporogy | <- |
|----------|------------|-----|
| Myo12 | 1513 | EQI |
| Sec. Sti | c. (My12) | EF |
| Myo13 | 1628 | RTA |
| Sec. Sti | c. (My13) | EF |

B)



Figure 2, Pinotsis et al.



My12

My13

Sec.Str. Topology <--E--> <---F--> <----G----> IWLQINEPTPNDKGKYVMELFDGkTGHQKTVDLSGqaydeayaefqrlkqaaiaekn HHH EEEEEEE EEEEE A<mark>Y</mark>FTINGVSTADSGKYGLVVKNK<mark>y</mark>GSET<mark>SD</mark>FTVSV<mark>f</mark>ipe-----------HHH EEEEEEEE EEEEEEEEEEEE EEEEE





My12My13 connecting helix

ЕЕЕЕЕЕЕ НННННННННННННННННН

| | <c-></c-> | <c'></c'> | <d-></d-> |
|---|-----------|-----------|----------------------------------|
| L | KVNWSHN | IGSAIRY | SDRVKTGVTG-E |
| | EEEEEE | ΕE | EEEEE |
| > | EVSWLKN | IEKALAÇ | <u>DHCNLKFEAgR</u> |
| | EEEEE | EEE | EEEEEE |



A)



Figure 3, Pinotsis et al.



A) My12/helix interface A





Figure 4, Pinotsis et al.

B) My13/helix interface B

A)



B)

Figure 5, Pinotsis et al.





Model

Interface

Symmetry op

Dimer formation

Interface size

Maximum dimen

Radius of gyrat

Model fit

Figure 6, Pinotsis et al.

| m) 12 | | | No. |
|------------|-------|---------------------|---------------|
| | 1 | 2 | 3 |
| e | My13 | My12My13 helix | My12 |
| perator | x,y,z | -x+1/2,y-1/2,-z+1/2 | -x+1,y,-z+1/2 |
| n [chain] | A/B | A/B | A/A |
| e [Ų] | 668 | 815 | 626 |
| nsion [nm] | 14.3 | 8.1 | 17.5 |
| tion [nm] | 3.93 | 2.90 | 4.77 |
| χ^2 | 1.58 | 8.0 | 3.0 |



B





(ii)

Figure 7, Pinotsis et al.







My9

Length: 10(12)

My10 1335(+110) GDVFKKLQKEAEFQRQEWIRK Helix prediction hHHHHHHHHHHHHHHHHHHHHHHH Length: 19(21)

My11 1438(+103) KSRLKLVDEAFKELMMEVCKKIA Helix prediction hhhhhHHHHHHHHHHHHHHHHHHHH Length: 17(23)

My12 1548(+110) SQAYDEAYAEFQRLKQAAIAEKN Helix prediction -hнннннннннннннннн Length: 21(22)



Figure 8, Pinotsis et al.

1225 EEELKRLLALSH

Helix prediction HHHHHHHHHHHhh

wavelength (nm)