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Structure of the F-actin-tropomyosin complex

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Abstract

Filamentous actin (F-actin) is the major protein of muscle thin filaments, and actin microfilaments are the main component of the eukaryotic cytoskeleton. Mutations in different actin isoforms lead to early-onset autosomal dominant non-syndromic hearing loss¹, familial thoracic aortic aneurysms and dissections², and multiple variations of myopathies³. In striated muscle fibres, the binding of myosin motors to actin filaments is mainly regulated by tropomyosin and troponin^{4,5}. Tropomyosin also binds to F-actin in smooth muscle and in non-muscle cells and stabilizes and regulates the filaments there in the absence of troponin⁶. Although crystal structures for monomeric actin (G-actin) are available⁷, a high-resolution structure of F-actin is still missing, hampering our understanding of how disease-causing mutations affect the function of thin muscle filaments and microfilaments. Here we report the three-dimensional structure of F-actin at a resolution of 3.7 ångstroms in complex with tropomyosin at a resolution of 6.5 ångstroms, determined by electron cryomicroscopy. The structure reveals that the D-loop is ordered and acts as a central region for hydrophobic and electrostatic interactions that stabilize the F-actin filament. We clearly identify the density corresponding to ADP and Mg^{2+} and explain the possible effect of prominent disease-causing mutants. A comparison of F-actin with G-actin reveals the conformational changes during filament formation and identifies the D-loop as their key mediator.

Supplementary Information is available in the online version of the paper.

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Author Contributions J.v.d.E. and S.R. designed the project. M.M. and D.J.M. provided protein samples. W.L. and D.J.M. provided information on tropomyosin and actin. J.v.d.E. performed research, analysed the data, and prepared figures. J.v.d.E. and P.A.P. improved image processing of helical specimens. S.R. managed the project and wrote the manuscript. All authors discussed the results and commented on the manuscript.

Author Information The coordinates for the electron microscope structure has been deposited in the Electron Microscopy Data Bank under accession code EMD-6124. Coordinates of F-actin in complex with tropomyosin have been deposited in the Protein Data Bank with accession number 3J8A. The authors declare no competing financial interests. Readers are welcome to comment on the online version of the paper.

Online Content Methods, along with any additional Extended Data display items and Source Data, are available in the online version of the paper; references unique to these sections appear only in the online paper.

We also confirm that negatively charged tropomyosin interacts with a positively charged groove on F-actin. Comparison of the position of tropomyosin in F-actin–tropomyosin with its position in our previously determined actin–tropomyosin–myosin structure⁸ reveals a myosin-induced transition of tropomyosin. Our results allow us to understand the role of individual mutations in the genesis of actin- and tropomyosin-related diseases and will serve as a strong foundation for the targeted development of drugs.

> To determine the structure of F-actin is inherently difficult because of its flexibility and its resistance to crystallization. Therefore, the only structural models of F-actin so far have been determined either from medium-resolution electron cryomicroscopy (cryo-EM) maps^{9–13} or by interpreting X-ray fibre diffraction data¹⁴, which has certain limitations. Using a direct electron detector and drift correction and by improving the image processing of helical specimens (see Methods), we have determined the structure of F-actin in complex with tropomyosin at an average resolution of 3.7 Å for F-actin and 6.5 Å for tropomyosin using cryo-EM (Fig. 1a, Extended Data Fig. 1a, b, 2a, Supplementary Video 1). During refinement the helical parameters-that is, the rise per subunit and the azimuthal rotation-were estimated to be 27.5 Å and 166.4°, respectively (see Methods). The side-chain densities of most actin residues were clearly resolved (Extended Data Fig. 3, Supplementary Video 2) and allowed us to build an atomic model of F-actin (Fig. 1b, Extended Data Fig. 3). The first four residues of the amino terminus and the last four residues of the carboxy terminus were not resolved (Extended Data Fig. 2b-d), indicating that these regions are disordered in the filament. However, we could clearly identify density corresponding to ADP and the coordinated cation, which is most probably Mg^{2+} (Fig. 1b, Extended Data Fig. 3a).

> The overall organization of F-actin is similar to that described in previous structures and models^{10,14}. However, given the superior resolution of our structure, we could clearly identify many salt bridges and therefore directly reveal intra- and intermolecular interactions of the F-actin filament in detail (Extended Data Fig. 4a).

F-actin is composed of two long-pitch helical strands. Interactions between actin subunits of the same strand and the opposing strand-the so-called intrastrand and interstrand interactions, respectively-stabilize the F-actin filament (Fig. 2, Extended Data Figs 4 and 5). Intrastrand contacts are mediated by subdomains SD2 and SD4 of one part of the actin strand with the SD3 of the adjacent part of the actin strand (Fig. 2a). Besides several salt bridges between the edges of SD4 and SD3 (Fig. 2b), the major site of interaction is between the D-loop and the bottom of the β -sheet of SD3 (Fig. 2c–f, Extended Data Fig. 4b–e). The D-loop encloses tyrosine 169 of the neighbouring subunit, resembling a lock-and-key interaction (Fig. 2c). In addition, adjacent residues fit snugly into the groove formed by regions next to the D-loop around isoleucine 64 (Fig. 2d) and a prominent hydrophobic patch in the D-loop interacts with a hydrophobic groove on the neighbouring F-actin subunit (Fig. 2e, f). Thus, the intrastrand interface at this position is large and is stabilized not only by electrostatic and hydrophobic interactions but also by geometric surface complementarity. As expected, mutations at this site (which, for instance, induce charges at hydrophobic interfaces) hamper F-actin polymerization and stability (Extended Data Fig. 4d, e) and can lead to different forms of nemaline myopathy^{3,15} (Extended Data Fig. 4a).

In contrast to most G-actin structural models, in which the D-loop is not resolved owing to its high flexibility, the D-loop is well defined in our structure (Extended Data Fig. 3b), corroborating its importance in mediating actin–actin interaction. It also contrasts with results of medium-resolution cryo-EM data¹² and disulphide cross-linking studies¹⁶ that suggested the presence of multiple D-loop conformational states. In addition, these and other studies^{17,18} suggested a close interaction of the D-loop with the C terminus of actin. Although the C terminus is located next to the D-loop, it is not resolved in the cryo-EM density (Extended Data Fig. 2c), indicating that the interaction between these two regions is either transient or structurally flexible.

Our structural determination also gives a possible explanation for the recently described Factin disassembly by Mical, a multidomain cytosolic actin regulator¹⁹. Mical posttranslationally oxidizes methionine 44 within the D-loop of actin²⁰, leading to the less flexible and more hydrophilic sulfoxide. The increased polarity could lead to a weakening of the hydrophobic actin–actin interaction (Extended Data Fig. 4d) and consequently to a Mical-induced disassembly of the F-actin filament.

The F-actin structure confirms that the so-called hydrophobic plug⁹ connects three F-actin subunits with each other (Fig. 2a). However, as suggested previously¹⁰, this interstrand interaction is formed by salt bridges (Fig. 2a), not by hydrophobic interactions (Extended Data Fig. 5a–e). A large part of the plug is negatively charged on the surface, electrostatically interacting with a positively charged patch on the opposing actin, with histidine 173 and lysine 284 at its centre (Extended Data Fig. 5c–e). Another possible salt bridge connects two neighbouring actins at lysine 113 of SD1 and glutamate 195 of SD4 (Extended Data Fig. 5f). However, the interface atthis position is quite small and not surrounded by other prominent electrostatic or hydrophobic interactions. In general, the two strands of F-actin are held together by only a few contacts, and a large empty space is maintained between the strands. This is probably an important requisite for the flexibility of F-actin filaments.

As previously described^{9,10,14}, F-actin is more flattened than is G-actin because of a rotation of SD1 and particularly SD2 (Fig. 3a, Supplementary Video 3). New salt bridges between SD1, SD4 and SD2 stabilize the new interface between the opposing subdomains at the nucleotide-binding cleft (Fig. 3b–f, Extended Data Fig. 6a).

The coordination of ADP and the divalent cation in the nucleotide-binding cleft by nearby residues is very similar to that in G-actin (Fig. 3b, Extended Data Fig. 6b). However, glutamine 137, which has been shown to play a central role in ATP hydrolysis²¹, is clearly moved closer to ADP, coordinating not only the cation (as in G-actin) but also the nucleotide β -phosphate (Extended Data Fig. 6c, d, Supplementary Video 3). Interestingly, the presence of ATP instead of ADP in the nucleotide-binding site would be sterically unfavourable (Extended Data Fig. 6d, Supplementary Video 3), suggesting that a different intermediate conformation exists for F-actin–ATP. High-resolution crystal structures of nonvertebrate actin showed that glutamine 137 positions the nucleophilic water that attacks the γ -phosphate of ATP²². Therefore, as previously suggested¹⁴, the shorter distance of glutamine 137 to the γ -phosphate probably induces ATP hydrolysis and then afterwards the

cation takes the position of the γ -phosphate in the finally resulting ADP-binding-state (Extended Data Fig. 6e, Supplementary Video 3). A mutation of glutamine 137 connected to nemaline myopathy²³ further highlights the importance of this residue in the proper function of F-actin (Extended Data Fig. 6f).

The comparison of the structures of F-actin with that of G-actin (PDB accession codes 3EL2 and 1J6Z), which represent the start and end points of filament formation, allows us to describe the possible conformational changes during polymerization (Extended Data Fig. 7, Supplementary Discussion).

The F-actin filaments were completely decorated by tropomyosin (Fig. 1a, Extended Data Fig. 1a, b, i). Although tropomyosin consists of seven pseudo-repeating units, applying the helical symmetry of F-actin for the reconstruction of the F-actin–tropomyosin complex results in a reduced resolution of tropomyosin and the overlap region of the N and C termini is not visible in the map (see Methods). However, densities corresponding to the helices of the coiled-coil were well resolved (Fig. 1a, b) and allowed us to fit a previous chimeric model of tropomyosin derived from X-ray, nuclear magnetic resonance and molecular dynamics simulation data²⁴ and to determine the register of the chains of tropomyosin. Together with the atomic resolution of F-actin, this allows us to describe the interaction between actin and tropomyosin in greater detail and with higher accuracy than previous studies.

Tropomyosin, which is mainly negatively charged on its surface, interacts with a positively charged groove on F-actin, confirming previous predictions (Fig. 4a, b, Extended Data Fig. 8a). In particular, lysines 326 and 328 on F-actin strongly interact with the different pseudo-repeats of tropomyosin (Fig. 4c). The distance between tropomyosin and actin varies depending on the position along the filament between 38 Å and 40 Å. The distance would allow for direct interactions of opposing residues (Fig. 4c) and is similar to the distance in the M-state⁸ and to what was previously suggested for the other states of tropomyosin^{25,26}. The interactions between tropomyosin and F-actin probably determine the position of tropomyosin on F-actin in the Apo-state (A-state), a term we recently introduced to describe the position of tropomyosin in the absence of troponin, myosin or Ca²⁺ (ref. 27).

The position of tropomyosin in the A-state in our structure (Fig. 4a, Extended Data Fig. 8b) is close to that described by Sousa *et al.*¹³, but differs from a previous positioning based on negative stain data and molecular dynamics simulations²⁴. To determine the source of this discrepancy, we first repeated the negative stain work (Extended Data Fig. 1c–j) and indeed reproduced the different position (Extended Data Fig. 8c). However, when we cross-linked the sample before staining, tropomyosin was in the same position as in our cryo-EM structure (Extended Data Fig. 1c–h, j, Extended Data Fig. 8d–g). This shows that negative staining rather than cryo-preparation (the sample is frozen quickly in liquid ethane without ice crystal formation) results in a change of the tropomyosin position on F-actin. The relatively low pH of the stain (pH 4) probably induces repulsions at the F-actin–tropomyosin interface and forces tropomyosin to change its position (Extended Data Fig. 8h). In any case, the change in position depending on preparative conditions confirms the striking ability of tropomyosin to translocate azimuthally across the actin filament at low energy cost²⁷.

In comparison to the previously assumed position of tropomyosin in the A-state, the new position blocks only minor regions of the myosin-binding site. This supports our previous model of myosin binding to actin filaments in the absence of troponin⁸ even more strongly. If the current position of tropomyosin is the default state for tropomyosin, then only loop 4 and the cardiomyopathy loop of myosin would be sterically hindered from binding to the F-actin filament (Extended Data Fig. 9a–d, Supplementary Video 4) and probably allows for a weak binding of myosin, especially with its lower 50-kDa domain. Actin-induced cleft closure of myosin would then move tropomyosin to its M-state⁸ (Extended Data Fig. 9b–g, Supplementary Video 4).

Comparing the position of tropomyosin in the new A-state with the M-state(Extended Data Fig. 9h), we find two possible transitions between the states. Either tropomyosin rolls on the F-actin filament by an azimuthal rotation of about 16° and a left-handed rotation of about 70° or a right-handed rotation of about 110° along its own axis (Extended Data Fig. 9i, j) or tropomyosin shifts by about 37 Å (half the length of a pseudo-repeat) along the actin filament (Extended Data Fig. 9k, 1) or a mixture of the two effects occurs. The distance of tropomyosin to the filament axis of F-actin is not altered during the transition. Gestalt-binding and rigidity of tropomyosin²⁸ support a shifting mechanism, whereas the spatial limitations inside the sarcomere make a shift of 37 Å rather unlikely. However, direct localization of the N/C-terminal overlap of tropomyosin will be needed to discriminate between the two possibilities.

In summary, the structure of the F-actin-tropomyosin complex shows how F-actin filaments are stabilized in health and destabilized in certain diseases. Our results provide the structural framework for further experiments towards resolving the molecular details of actin polymerization and its interaction with actin-regulating proteins such as tropomyosin.

METHODS

Protein expression and purification

F-actin was isolated from rabbit skeletal muscle acetone powder and purified with several polymerization and depolymerization steps as described previously²⁹. The composition of the G-actin buffer was 5 mM Tris pH 7.5, 1 mM DTT, 0.2 mM CaCl₂. Recombinant α -tropomyosin (*Mus musculus*, alpha-1 chain, Uniprot-ID P58771) with an alanine-serine extension was purified from *Escherichia coli*, based on the protocol of ref. 30. The α -tropomyosin isoform we used is abundant in smooth and striated muscles. The actin–tropomyosin complex was prepared by mixing F-actin with tropomyosin initially at a molar ratio of 7:1 in filament buffer (5 mM Tris-HCl pH 7.5, 1 mM DTT, 100 mM KCl, and 2 mM MgCl₂). The final concentration of tropomyosin for either negatively stained or frozen specimens was then adjusted empirically to obtain complete decoration and only few unbound tropomyosins in the background. For the cross-linking studies between F-actin and tropomyosin, samples were prepared in filament buffer containing 5 mM HEPES (pH 7.5).

Preparation for negative-stain electron microscopy

To stabilize and straighten the F-actin filaments (this is important for image processing), we decorated them with tropomyosin. Actin–tropomyosin complexes were directly prepared before the negative staining procedure. 4 μ l of sample was applied on a freshly glow-discharged copper grid (Agar Scientific; G2400C) with an additional thin and continuous carbon film. After incubation for 60 s on the grid the sample was blotted using filter paper (Whatman no. 4), washed twice with filament buffer and stained with 0.75% uranyl formate (pH 4.0) for 30 s. For cross-linking studies, after incubation for 60 s of the sample solution on the grid and one washing step, a droplet of filament buffer with 0.25% glutardialdehyde was applied for 30 min on the grid before negative staining to avoid putative pH-induced artefacts. All images were taken with a JEOL JEM-1400 electron microscope equipped with a LaB₆ cathode at an operation voltage of 120 kV. Micrographs were recorded manually with a 4,000 × 4,000 pixel complementary metal–oxide–semiconductor (CMOS) TemCam F416 (TVIPS) camera under minimal dose conditions.

Image processing of negatively stained filaments

All micrographs had a pixel size of 2.32 Å per pixel and filaments were boxed with a boxing distance of 13 pixels (overlap ~90%) and a total box size of 128 pixels. We analysed 8,371 segments from 40 images of bare F-actin, 27,926 segments from 111 images of F-actin decorated with tropomyosin, and 27,011 segments from 81 images of F-actin decorated with tropomyosin and cross-linked with glutardialdehyde. Subsequently, all segment stacks were aligned and classified independently using reference-free alignment and *k*-means classification procedures as implemented in SPARX³¹ (Extended Data Fig. 1c–h, j). During the helical refinement and reconstruction using the *helicon* package implemented in SPARX³¹, a cylinder filled with Gaussian noise was used as an initial three-dimensional template. The position of tropomyosin–bare F-actin and glutardialdehyde–bare F-actin were calculated with the 'vop subtract' command in Chimera³² (Extended Data Fig. 8c–g).

Grid preparation and image acquisition for cryo-EM

A 1.5-µl sample was applied to a glow-discharged holey carbon grid (C-flats 2/1, Protochips), incubated for 10 s and manually blotted for 3 s from the backside with filter paper (Whatman no. 5), before vitrification by plunging the grid into liquid ethane using a Cp3 (Gatan). Screening for the best sample and blotting conditions was performed on a JEOL JEM 3200FSC electron microscope equipped with a field emission gun and operated at a voltage of 200 kV. The omega in-column energy filter of the microscope was used to estimate the best ice conditions (~40–60 nm thickness). Finally, a data set was taken with a spherical-aberration (Cs)-corrected FEI Titan Krios transmission electron microscope equipped with a X-field emission gun and operated at a voltage of 300 kV. Images were recorded with a back-thinned $4k \times 4k$ FEI Falcon 2 direct detection camera under minimal dose conditions using the automatic data collection software EPU (FEI). Within each selected grid-hole, three different positions were imaged, each with a total exposure of 1 s and a frame time of 55 ms. Seven frames from 85 ms to 475 ms with a total dose of 14.6 electrons per Å² and one total average with an electron dose of 30.7 electrons per Å² were used for image processing. The defocus range of the data set was $0.8-2.6 \mu m$. The pixel size was 1.14 Å per pixel, based on the reflection of gold at the same magnification. Since the inaccuracy of pixel size determination is about 2% with this method, we used the well-defined rise per subunit of F-actin (27.6 Å) from fibre diffraction studies^{14,33} instead to calibrate the pixel size, which was then 1.12 Å per pixel.

Image processing of the cryo-EM data set

As a first quality control, all 1,311 frame averages were manually inspected and only 689 images with high filament quality (Extended Data Fig. 1a) were used for further processing. Frames were aligned and afterwards summed up using motion correction³⁴. Filaments were manually selected with helixboxer in SPARX. A total of 109,242 segments from 7,854 filaments were extracted with a box size of 256 pixels and a boxing distance of 29 pixels (overlap ~ 90%). Thus, the approximate distance between them (~32 Å) slightly exceeded the rise of the helical assembly of actin ($\sim 27-28$ Å), as required by the *helicon* design. The defocus and astigmatism and the accuracy of both were determined by cter³⁵ in SPARX. All segments were aligned and classified using reference-free alignment and k-means classification procedures implemented in SPARX (Extended Data Fig. 1b, i). For the initial reconstruction, the data set was binned twice to a pixel size of 2.24 Å and a box size of 128 pixels. We used a cylinder filled with Gaussian noise as a three-dimensional initial template for the helical refinement and reconstruction in the helicon. After adjusting the pixel size as described above, the helical symmetry parameters converged to a rise per subunit of 27.5 Å and an azimuthal rotation of 166.4°. Initially, the entire F-actin-tropomyosin map was used as the reference during the first rounds of refinement, using the helical symmetry parameters of F-actin. However, tropomyosin spans over seven actin subunits and therefore has a symmetry different to that of actin. Nevertheless, because it is divided into seven pseudorepeating units, each of which binds to a successive actin subunit along F-actin, the symmetry of F-actin could be used for reconstructions of actin-tropomyosin complexes of up to 6 Å resolution, accepting that the overlap region of the N and C termini of tropomyosin would not be discernible in these maps. Once the refined map reached a resolution of 6.5 Å, as determined by the $FSC_{0.5}$ criterion³⁶, we masked out tropomyosin in the reference volume before continuing with high-resolution refinements of F-actin. In the last refinement iteration, we used only filaments derived from summed up frames with a drift of less than 7 Å, as estimated in the motion correction³⁴ step and with a sufficient content of high-resolution information, as determined by cter³⁵ in SPARX. This yielded a subset of 74,228 segments for the final map, which corresponded to ~120,000 asymmetric subunits. The segments' statistics are given in Extended Data Table 1. Since an electron microscope map of a filament is truncated at the edges of the cubic volume of the box (here 256^3 voxels), a mask with Gaussian edges must be applied for resolution estimation to minimize artefacts. Therefore, Fourier shell correlation (FSC) analysis was performed within the central area of the volume, resulting in a nominal resolution of 3.7 Å (FSC $_{0.5}$ criterion³⁶) for the F-actin electron density map (Extended Data Fig. 2a). The map of F-actin was then sharpened using a negative b factor of -50 Å^2 and filtered to its nominal resolution. Finally, the tropomyosin map filtered to 6.5 Å was merged with the final F-actin map to obtain a map of the entire F-actin-tropomyosin complex.

We also tried to process tropomyosin separately from F-actin. After alignment of the filaments based on F-actin, we masked out tropomyosin. To find the N- or C-terminal overlap and guarantee the proper alignment of tropomyosin, we used very large boxes to allow tropomyosin to shift by almost its full length and tried to align the molecules. However, this approach did not work. We suspect that the pseudo-repeats and even the N/C-terminal overlaps are too similar to each other (at least in cryo-EM raw images), resulting in an improper alignment. We are therefore investigating labelling strategies to solve this problem.

Model building and refinement of F-actin

The central F-actin subunit (chain A) shows all available contacts to adjacent chains (B, C, D, E) and was therefore used for further structural analysis. In a first step, five central subunits (chains A-F) of an F-actin model (PDB accession code 4A7N; ref. 8) was rigidbody fitted into the electron microscope map, using 'Fit in Map' and the map was initially segmented with 'Split Map' in Chimera^{32,37}. Homology modelling was performed using MODELLER³⁸ and the resulting model of the central subunit was then flexibly fitted into its corresponding part of the density using DireX³⁹. Finally, the flexibly fitted subunit was used to create a starting F-actin model of five adjacent subunits (hereafter referred to as the 'pentamer') for further model building. The electron density was converted to structure factors with the CCP4 program suite⁴⁰. The model and the map were then used for real space refinement and model building in COOT⁴¹. After a single iteration of model building of the central F-actin subunit, its part of the map was further used for an initial automatic reciprocal space refinement using PHENIX.refine⁴². Since this part was less well resolved in the N terminus and the C terminus, and model building was difficult for a part of the D-loop, we deleted amino acids 1-4, 42-50 and 372-375 from the model. Owing to the high reliability of the experimental phases of the cryo-EM map, we were able to apply experimental phase restraints during refinement in PHENIX. After refinement the output model was again manually refined in COOT using the original cryo-EM map to minimize model bias. Rebuilding of the N or C terminus was not possible even in the refined map (Extended Data Fig. 2c, d). However, after refinement in PHENIX the definition of the Dloop improved, so ultimately it could be manually rebuilt ab initio.

After refinement of the subunit had converged and model building was completed, the resulting subunit was used to assemble the F-actin pentamer and the respective full map of the central pentamer was then segmented. The molecular model and the respective original density were then used as input for PHENIX.refine. Again, we alternated manual model building and refinement steps. Finally, the resulting model was validated using MOLPROBITY⁴³ (see data statistics in Extended Data Table 1). For further structural analysis we used the final model and the cryo-EM map of the actin pentamer.

Structure analysis and visualization

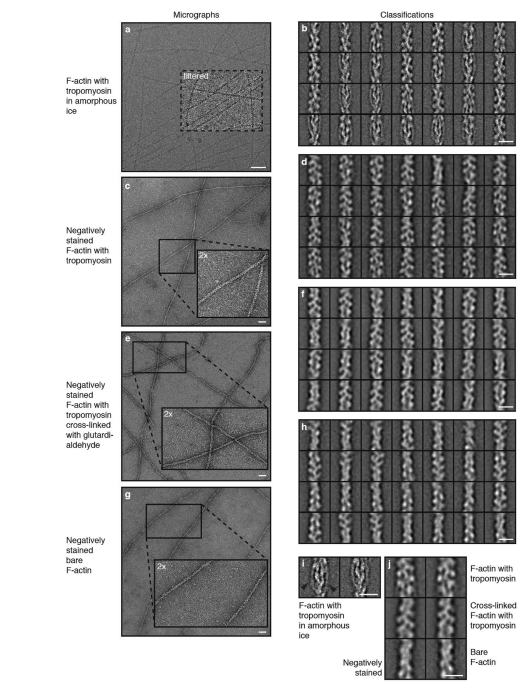
To describe the interaction between F-actin and tropomyosin, a chimaeric molecular model of tropomyosin⁴⁴ was initially rigid-body fitted into the cryo-EM density of F-actin– tropomyosin. Subsequently, the respective density of tropomyosin was extracted as described above using Chimera. The molecular model of tropomyosin was then flexibly

fitted into the resulting density using DireX³⁹. Furthermore, during flexible fitting, in order to preserve its electrostatically preferred position on F-actin we fixed tropomyosin in its vertical position on the filament. Atomic clashes and stereochemistry were repaired using PHENIX.geometry_minimization⁴². Since the tropomyosin N/C-terminal overlap is not fully understood⁴⁵, we reduced the model to the five central pseudo-repeats. Owing to the limited resolution of the cryo-EM density in the region of tropomyosin, we avoided interpretation of tropomyosin on the single-amino-acid level.

We used the refined actin pentamer and the modelled tropomyosin to reconstruct a full filament for visualization of the actin–tropomyosin interaction (Fig. 3). The filament was protonated using $H++^{46}$ at different pH values and the electrostatic coulomb potential of the filament surface was calculated to range from -10 kcal mol⁻¹ to +10 kcal mol⁻¹ in Chimera (Fig. 4b, Extended Data Fig. 8h). For visualization of the hydrophobicity per amino acid residue, we used 'Define attribute' in Chimera and generated amino-acid-specific scores⁴⁷. Point mutations were introduced via COOT or Chimera to show charge-induced repulsions and changes in hydrophobicity. The HGMD⁴⁸ library was browsed to find mutations in regions of interest.

For detailed analyses of intra- and intermolecular interactions in F-actin, we applied PISA tools⁴⁹ and distance measurements in Chimera. The G- to F-actin transition within a monomer/subunit was visualized and interpreted with the 'Morph Conformations' in Chimera. Furthermore, the hinge region and the global rotation angle were detected with the DynDom Sever⁵⁰ using an atomic model of G-actin (PDB accession code 1J6Z; ref. 51) and one subunit of our F-actin model (Fig. 3).

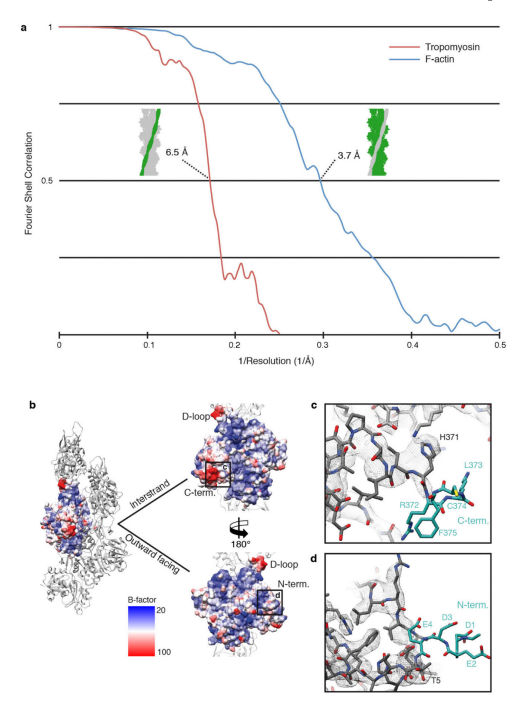
Extended Data



Extended Data Figure 1. Micrographs and classifications of different data sets

a–**h**, Representative digital micrographs and corresponding representative two-dimensional class averages of F-actin decorated with tropomyosin in amorphous ice (in total 300 class averages of randomly chosen 40,000 phase-flipped segments of in total 109,242 segments from 689 images) (**a**, **b**), negatively stained (in total 300 class averages of 27,926 segments from 111 images) (**c**, **d**), negatively stained after cross-linking with glutardialdehyde (in total 300 class averages of 27,011 segments from 81 images) (**e**, **f**), and a micrograph of

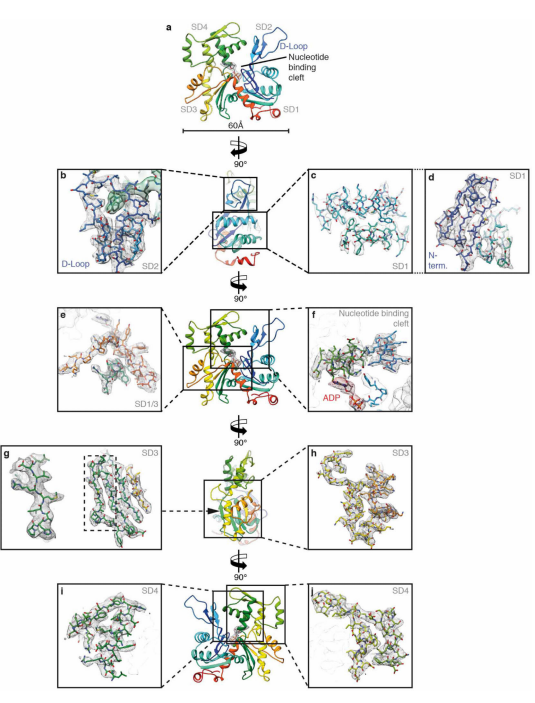
negatively stained bare F-actin (in total 100 class averages of 8,371 segments from 40 images) (\mathbf{g} , \mathbf{h}). Some areas of the micrographs are either filtered or enlarged, as indicated in the figures. Scale bars, 50 nm. Each class average ('Classifications') contains 130–200 (cryo data set) or 70–90 (negatively stained data sets) single segments. Scale bars, 10 nm. A boxed region of the digital micrograph in \mathbf{a} was band-pass filtered to allow a better visualization of the filaments. Insets in \mathbf{c} , \mathbf{e} and \mathbf{g} show 2× magnified regions of the digital micrographs. \mathbf{i} , Two of the class averages depicted in \mathbf{b} that show a clear tropomyosin density. The tropomyosin density is indicated by black arrows. Scale bar, 10 nm. \mathbf{j} , Comparison of representative class averages of the three negative-stain data sets. Class averages of the data sets with tropomyosin (top panels) show additional density and a larger diameter than bare F-actin (bottom panel). Scale bar, 10 nm. 8,371 segments from 40 images of bare F-actin, 27,926 segments from 111 images of F-actin decorated with tropomyosin, and 27,011 segments from 81 images of F-actin decorated with tropomyosin and cross-linked with glutardialdehyde.



Extended Data Figure 2. Resolution of the F-actin-tropomyosin complex

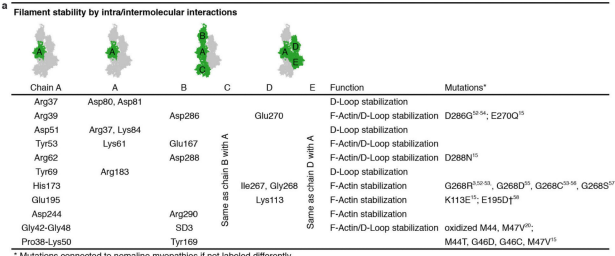
a, FSC curves of different areas of interest by masking (see Methods). The resolution of tropomyosin was estimated using the twice down-sampled data set (2.24 Å per pixel). The FSC_{0.5} criterion indicates that the tropomyosin electron density map has a resolution of 6.5 Å. The resolution of the final F-actin electron density map is estimated at a resolution of 3.7 Å (FSC_{0.5}). **b**, Surfaces of F-actin with B-factors (high is red, low is blue; the B-factor indicates the true static or dynamic mobility of an atom) estimated by the reciprocal space refinement in PHENIX⁴². A side view as well as top views on the inward- and outward-

facing surfaces, that is, facing the adjacent F-actin subunit inside the filament and oriented to the periphery, respectively, are shown. **c**, **d**, Putative structures of the C and N termini (cyan), respectively. Electron density is missing in these regions.



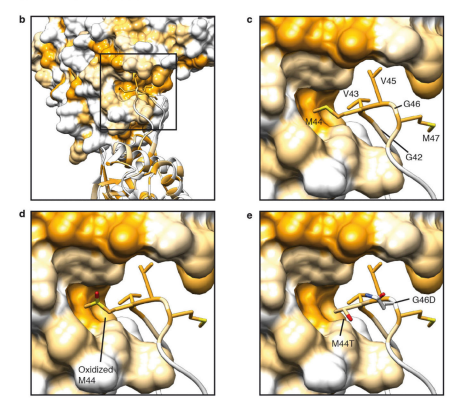
Extended Data Figure 3. Representative regions of the F-actin electron density map a, Overview of the atomic model of an F-actin subunit rainbow-coloured from the N terminus (blue) to the C terminus (red). ADP and the coordinated cation, probably Mg²⁺, are depicted inside their corresponding electron densities. **b–d**, Side view. Interface between the

D-loop (SD2) and the SD3 of the adjacent actin subunit (b), outer two helices of SD1 (c), and inner β -sheet with N terminus (d). e, f, Front view. Connection of SD1 and SD3 (e) and nucleotide binding cleft (f). g, h, Side view. Inner β -sheet of SD3 with one highlighted strand (g) and outer part of SD3 (h). i, j, Back view. SD4 separated in two parts.



* Mutations connected to nemaline myopathies if not labeled differently

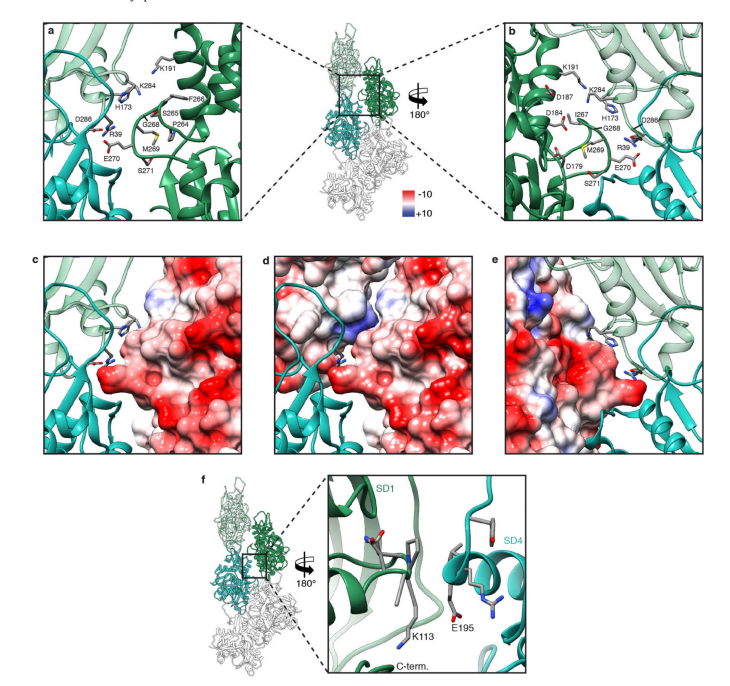
† Mutation connected to neuromuscular disorders



Extended Data Figure 4. Overview of inter- and intrastrand interactions and hydrophobic Dloop

a, Table of identified residues that are involved in intra- and intermolecular interactions and known mutations. b, Overview of the D-loop bound to the hydrophobic cleft in SD3 of the

F-actin molecule on top. Surfaces and residues are coloured from high (yellow) to low (white) hydrophobicity. **c**, Involved hydrophobic residues of the D-loop. **d**, **e**, Modifications in the D-loop, like oxidation of methionine 44 (ref. 20) (**c**) or mutations of methionine 44 and glycine 46 (M44T, G46G)^{15,20} (**d**) change the polarity or insert charges and consequently weaken the hydrophobic interactions. Thus, this destabilization of the intrastrand contact is connected to anomalous actin filament assembly and nemaline myopathies^{3,15,20,52–58}.



Extended Data Figure 5. Interstrand and intrastrand F-actin interactions

a–**e**, The interface at the plug involves three residues (R39, E270, D286) that form salt bridges and mediate not only one interstrand contact but also one intrastrand contact. In addition, the orientation of residues 264–269 result in a negatively charged patch that electrostatically interacts with positively charged residues on the opposing actin. **a**, **b**, Front and back view of the interface at the plug, respectively. **c**–**e**, Surface representations (front views in **c** and **d**, back view in **e**) depicting the Coulomb potential, indicating that the interaction of the upper region of the plug with adjacent inter- or intrastrand molecules is mediated by electrostatic interactions. **f**, Another interstrand contact is formed by residues 110–115 of SD1 and residues 191–199 of SD4 of the adjacent actin. However, no prominent electrostatic or hydrophobic interactions could be identified at this interface.

a	Intramolecular	interaction by	salt bridges and	ADP/Cation	coordination in	the nucleotide	binding cle

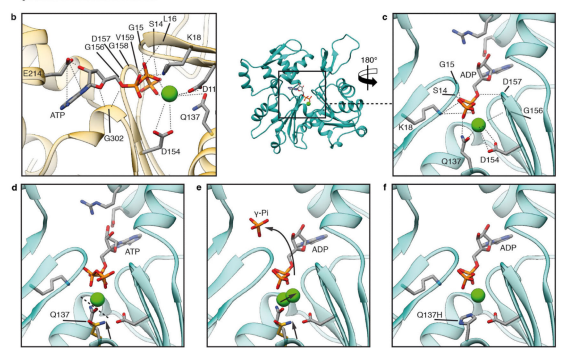
Chain A	А	Function	Mutations*
Tyr69	Arg183	Cleft stabilization	R183C ^{53-54,60} ; R183G ^{53,56} , R183L ¹⁵ ; R183S ³
His73	Asp179	Cleft stabilization	H73N ¹⁵ ; H73L, H73R, D179N, D179H ³ ; H73R ⁶¹ ; D179G ⁶²
Asp157	Arg183	Cleft stabilization	R183C ^{53-54,60} ; R183G ^{53,56} , R183L ¹⁵ ; R183S ³
Asp187	Arg206	Cleft stabilization	
Asp207	Arg210	Cleft stabilization	E207D ⁵⁷
Asp211	Arg210	Cleft stabilization	
Lys213	Tyr306	Cleft stabilization	
Tyr337	Lys18, Lys336	Cleft stabilization	K336E ⁴⁸³ ; K336I ³
Ser14, Gly15	ADP	ADP binding	G15R‡ ⁵²⁻⁵⁴ , G15D§ ⁶⁴
Lys18	ADP	ADP binding	
Gln137	ADP	ADP binding	Q137H ²³
Gly156, Asp157	ADP	ADP binding	
Arg210	ADP	ADP binding	
Glu214	ADP	ADP binding	
Gly302	ADP	ADP binding	
Gln137	Ion	Ion coordination	Q137H ²³
Asp154	Ion	Ion coordination	
Gly156	lon	Ion coordination	

* Mutations connected to nemaline myopathies if not labeled differently

† Mutation connected to nemaline myopathies and hypertrophic cardiomyopathies

‡ Mutation connected to actin myopathies

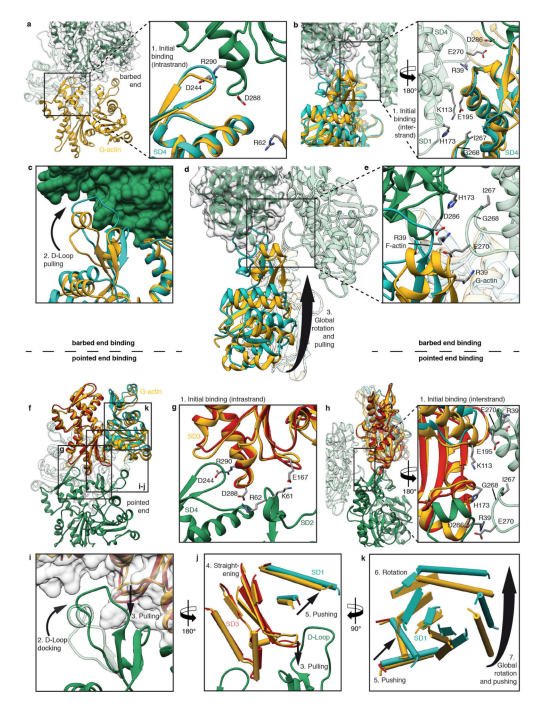
§ Mutation connected to fetal akinesia



Extended Data Figure 6. Nucleotide binding site and intramolecular interactions

a, Table of identified intramolecular interactions that result in a stabilization of the nucleotide binding cleft, coordination of ADP and a divalent cation. Data are from refs 3, 15, 23, 53, 54, 56, 57 and 60–64. **b**, Coordination of ADP and Ca^{2+} in the nucleotide binding cleft in G-actin (PDB accession code 3EL2; ref. 59). c-e, Back view of the nucleotide binding cleft of F-actin (cyan) with bound ADP-Mg²⁺ or ADP-Ca²⁺ (c), ATP-Ca²⁺ (d, relative position taken from PDB accession code 3EL2; ref. 59) and conformational changes between the G-actin-ATP and F-actin-ADP state (e). Glutamine 137 is moved

closer to ADP, coordinating not only the cation (as in G-actin) but also the nucleotide β phosphate (**c**). The presence of ATP instead of ADP in the nucleotide-binding site would be sterically unfavourable, suggesting that a different intermediate conformation exists for Factin-ATP (**d**). The shorter distance of glutamine 137 to the γ -phosphate probably induces ATP hydrolysis and then afterwards the cation takes the position of the γ -phosphate in the ADP-state (**e**). For comparison the position of glutamine 137 in G-actin is shown in yellow and the transition from G-actin to F-actin is depicted by arrows. **f**, Mutation of glutamine 137 to histidine results in hampered coordination of the ion and the nucleotide and is connected to nemaline myopathies²³.

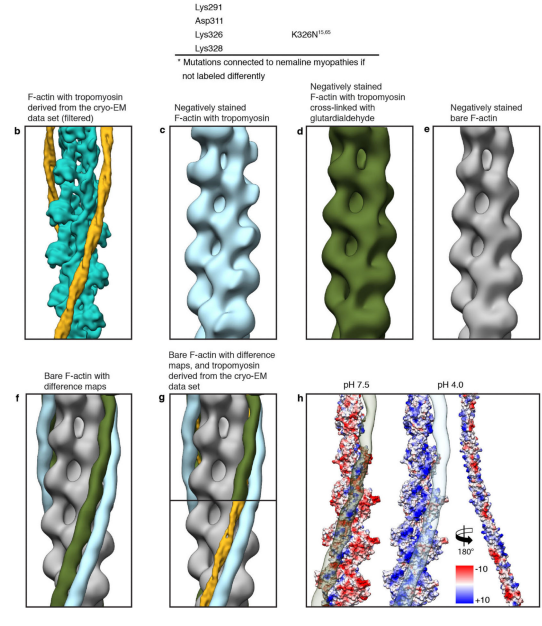


Extended Data Figure 7. Model of barbed-end and pointed-end binding and G- to F-actin transition based on a comparison of start point (G-actin) and end point (F-actin)
a, b, Binding of new G-actin (yellow, PDB accession code 3EL2; ref. 59) at the barbed end of F-actin (green) is initiated by the intrastrand binding of SD4 of G-actin to SD3 of F-actin (a, side view) and the interstrand binding of SD4 and SD1 (b, front view), respectively. G-actin is overlaid with a subunit of the structure found in F-actin (cyan). The main interstrand contacts are already present at the start of the transition and thereby guide the binding G-actin to its correct position, determining the symmetry of the filament. c, After initial binding the D-loop is trapped in the hydrophobic cleft of SD3 of F-actin and pulls on SD2.

F-actin is depicted in surface representation (green). d, Finally, the SD2 and concomitantly SD1 are rotated and the final F-actin conformation is stabilized by various intramolecular interactions (see Figs 2c-f and 3c-f, Extended Data Fig. 6a). e, Owing to the relatively large distance from the SD1 and SD2 subdomains of the newly bound G-actin to the subunit of the opposite strand, the only influence of the opposite strand on the binding of G-actin is at the docking position of the D-loop. The transition of R39 illustrates well the docking of the D-loop. The residue is depicted in both states (G-actin and F-actin). f-h, Binding of new Gactin (yellow, PDB accession code 3EL2; ref. 59) at the pointed end of F-actin (green) is initiated by initial intrastrand binding (g) of SD3 of G-actin to SD4 of F-actin at the pointed end and interstrand binding of SD4 to SD3 (h), respectively. Again, the main interstrand contacts are available before the transition of G-actin (yellow) to F-actin (SD1 and SD2 in cyan, SD3 and SD4 are red). i, During binding to F-actin the transition from G-actin to Factin is initiated by an induced fit of the F-actin D-loop to the hydrophobic cleft of the newly bound G-actin. This leads to a pulling down of the central β -sheet of SD3 of G-actin. j, The β -sheet is thereby straightened and pushes up two adjacent helices of SD1. **k**, The slight dislocation of these helices is transmitted to other regions of SD1 and thereby emphasized. This leads to a global rotation of SD1, which results in a considerable rotation of SD2 by an angle of 20° and a closure of the nucleotide binding cleft (Fig. 3; see Supplementary Video 3).

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Tropomyosin binding to F-actin in A-state

Mutations*

R147K¹⁵

a

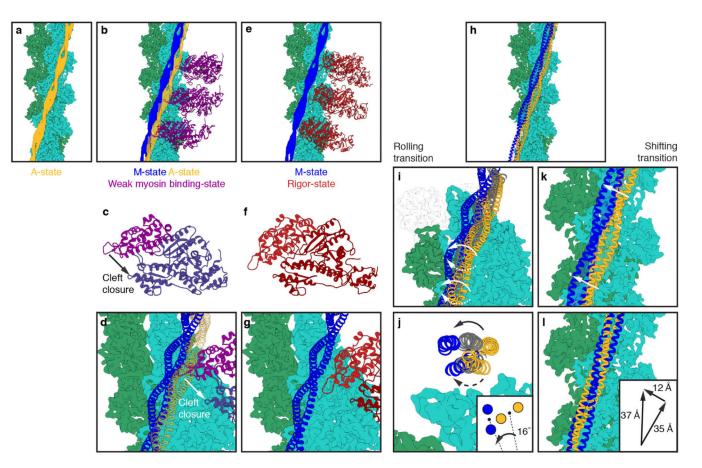
Actin

Arg147 Lys215

Extended Data Figure 8. Tropomyosin binding and comparison of reconstructions regarding the tropomyosin position on F-actin

a, Table showing putative residues of F-actin involved in tropomyosin binding and known mutations. **b**–**e**, Reconstructions of F-actin decorated with tropomyosin calculated from: the cryo-EM data set filtered to 15 Å (**b**), from a negatively stained data set (**c**), from a negatively stained data set after cross-linking with glutardialdehyde (**d**), and from negatively stained bare F-actin (**e**). **f**, By calculating a difference map between tropomyosin–bare F-actin (blue) and glutardialdehyde–bare F-actin (green), differences in the tropomyosin

position on bare F-actin (grey) are visualized. **g**, Overlay of difference maps showing that the position of tropomyosin in the cryo-EM reconstruction (yellow) corresponds to the tropomyosin position of the negatively stained data set with the cross-linked complex (green). **h**, Surface of F-actin and tropomyosin (pseudo-repeats 2–6) with the electrostatic Coulomb potentials at pH 7.5 and pH 4 (see also Fig. 4b). Tropomyosin was rotated by 180° and shifted to the right to allow a better view of the F-actin–tropomyosin interface. Difference maps of the glutardialdehyde–bare F-actin and the tropomyosin–bare F-actin map are shown on the F-actin surface at pH 7.5 (left) and pH 4.0 (middle), respectively^{15,65}.



Extended Data Figure 9. Model of tropomyosin transition on F-actin during myosin binding a, Cryo-EM structure of the F-actin–tropomyosin complex with tropomyosin in the A-state. Tropomyosin (yellow, A-state), F-actin (green). **b**–**d**, Initial weak binding of myosin (magenta, PDB accession code 1LKX; ref. 66) to the F-actin–tropomyosin filament in the absence of troponin. Most of the myosin binding sites on F-actin are not occupied by tropomyosin and only loop 4 and the cardiomyopathy loop are sterically hindered from binding to the F-actin filament (**b**). Actin-induced closure of the 50 kDa cleft of myosin (**c**) results in a strong binding of myosin and tropomyosin moves to its M-state position (blue) (**d**). **e**–**g**, Actin–tropomyosin–myosin complex in the rigor state (PDB accession code 4A7H; ref. 8). Myosin is shown in red. **h**–**l**, There are two possible ways for the transition of tropomyosin from the A-state to the M-state. Tropomyosin either rolls (**i**, **j**) or slides (**k**, **l**) from one to the other position. Rolling would involve an azimuthal rotation of ~16° with

respect to the F-actin axis (inset of **j**) and a left-handed rotation of ~70° (indicated by solid arrows) or a right-handed rotation of ~110° (indicated by dotted arrows) with respect to its own axis (**i**, **j**). Sliding would imply not only an azimuthal shift of ~12 Å (indicated by white arrows in **k**), but also a tremendous shift of a half-tropomyosin repeat (that is, ~35 Å) along the F-actin filament (**k**, **l**). The radius to the filament axis would be preserved in both situations. The inset of **l** depicts the vectors for a shifting transition of tropomyosin: an azimuthal and longitudinal shift of 12 Å and 35 Å, resulting in an overall shift of 37 Å.

Extended Data Table 1

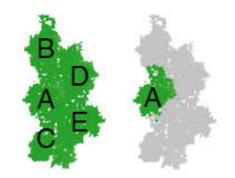
Data collection and refinement statistics

Data collection			
	Magnification	×122,270	
	Defocus range (µm)	0.8-2.6	
	Voltage (kV)	300	
	Mircoscope	Titan Krios (Cs corrected)	
	Camera	Falcon 2 (4k DED)	
	Frame recording time (s)	0.085-0.475	
	Number of frames	7	
	Electron dose (e ⁻ Å ⁻²)	14.6	
	Pixel size (Å)	1.12	
Particle statistics			
	Filaments *	7,854 (5,239)	
	Box size (px)	256	
	Boxing distance (px)	29	
	Rise (Å)	27.5	
	Azimuthal rotation (°)	166.4	
	Segments *	109,242 (74,228)	
	Asymmetric subunits *	~180,000 (~120,000)	
		B D C E	A
Model composition		ABCDE	A
	Non-hydrogen atoms	14,450	2,8

 Non-hydrogen atoms
 14,450
 2,890

 Protein residues
 1,835
 367

 Ligand (ADP Mg²⁺)
 140
 28



Model composition	Chain ID	ABCDE	Α
Refinement			
	Resolution used for refinement (Å)	95.2-3.7	
	Map sharpening B-factor (Å ²)	-50	
	Average B factor (Å ²)	55.40	53.40
	R factor ^{$\dot{\tau}$}	0.277	
	Cross-Resolution function [≠]	0.81	0.83
Rms deviations			
	Bonds (Å)	0.004	0.005
	Angles (°)	1.10	1.12
Validation			
	Clashscore, all atoms	19.99	18.32
	Poor rotamers (%)	1.23	1.29
Ramachandran plot			
	Favored (%)	87.20	87.40
	Outliers (%)	1.37	1.37

Refinement statistics are shown after the last step of the refinement of the F-actin pentamer. In addition, relevant values of the central chain A are shown. Rms, root mean squared.

*Number in parentheses is count after sorting the dataset regarding micrograph quality

 \hat{T} R factor = $\Sigma |F_{obs}| - |F_{calc}| / \Sigma |F_{obs}|$

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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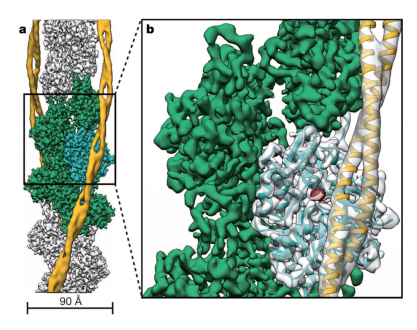


Figure 1. Cryo-EM structure of F-actin decorated with tropomyosin

a, Full cryo-EM reconstruction of F-actin (grey, with five central subunits in green and one subunit in cyan) decorated with tropomyosin (yellow). **b**, Close-up view of **a** with the atomic and molecular model of an F-actin subunit (cyan) and tropomyosin (yellow) and their corresponding densities, respectively. The density corresponding to ADP is depicted in red.

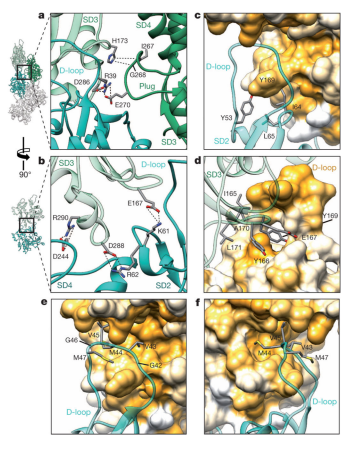


Figure 2. Filament stability by intrastrand and interstrand interactions

a–**f**, Neighbouring F-actin subunits stabilize the F-actin filament by interaction through salt bridges (**a**, **b**) and by hydrophobic interactions (**c**–**f**). The central F-actin subunit is depicted in cyan, and adjacent subunits are shown in shades of green. Surfaces are coloured from high (yellow) to low (white) hydrophobicity. Oxygen is coloured red, nitrogen is blue, carbon is grey and hydrogen is colourless. Interactions between amino acids are highlighted with dotted lines. **a**, Interstrand and intrastrand salt bridges of three actin subunits involving the plug. **b**, Several intrastrand salt bridges at the actin–actin interface. **c**–**f**, Front view (**c**, **e**), back view (**d**) and side view (**f**) of the D-loop interacting with the SD3 of the neighbouring intrastrand F-actin subunit. The D-loop wraps around Y169 of the neighbouring subunit and other residues snugly fit into the groove formed by regions adjacent to the D-loop (**c**, **d**), resembling a lock-and-key interaction. In addition, a prominent hydrophobic patch in the D-loop interacts with a hydrophobic groove on the neighbouring F-actin subunit (**e**, **f**).

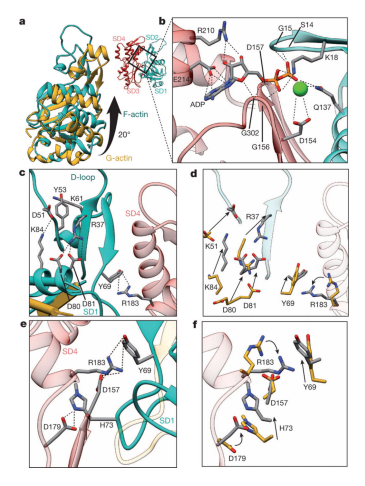


Figure 3. G-actin to F-actin transition

a, A global rotation of SD1 and SD2 leads to a flattening of G-actin (yellow) during transition to F-actin (cyan). **b**, Coordination of ADP and a cation in the nucleotide binding cleft in F-actin (SD1–SD2 cyan, SD3–SD4 red). The cation is most probably Mg²⁺. However, Ca²⁺ cannot be excluded as the coordinated cation because the actin was purified in Ca²⁺-containing buffer. **c**–**f**, The conformation of F-actin is finally stabilized by intramolecular interactions between SD2, SD1 and SD4 (**c**, **e**). Most of the residues involved in the stabilization of the F-actin conformation show a considerable movement during the transition between G-actin and F-actin (**d**, **f**). **a**, G-actin with defined D-loop for better visualization; PDB accession code 1J6Z. **c**–**f**, ATP-bound G-actin; PDB accession code 3EL2.

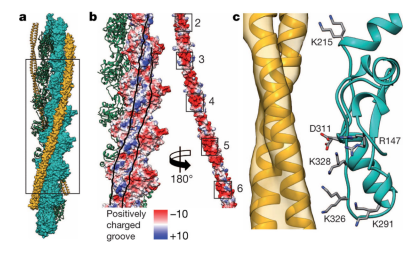


Figure 4. F-actin interaction with tropomyosin

a, Structural overview of an F-actin filament (green and cyan) decorated with tropomyosin (yellow). Half of the filament is shown in surface representation. The N and C termini of F-actin are included in this model. **b**, Surface of F-actin and tropomyosin (pseudo-repeats 2–6) with the electrostatic Coulomb potentials ranging from -10 kcal mol⁻¹ to +10 kcal mol⁻¹ at pH 7.5. Tropomyosin was rotated by 180° and shifted to the right to allow a better view on the F-actin–tropomyosin interface, which is delimited by lines drawn onto the surfaces. The overall negatively charged tropomyosin interacts with a positively charged groove on F-actin. **c**, Several charged residues of actin are within distances that would make it possible to interact with tropomyosin via putative salt bridges. Different rotamers (orientations of a residue) of the same residue are shown to indicate how F-actin subunits could adjust to the surfaces of different tropomyosin pseudo-repeats.