

Mechanoenzymatics of titin kinase

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Biological responses to mechanical stress require strain-sensing molecules, whose mechanically induced conformational changes are relayed to signaling cascades mediating changes in cell and tissue properties. In vertebrate muscle, the giant elastic protein titin is involved in strain sensing via its C-terminal kinase domain (TK) at the sarcomeric M-band and contributes to the adaptation of muscle in response to changes in mechanical strain. TK is regulated in a unique dual autoinhibition mechanism by a C-terminal regulatory tail, blocking the ATP binding site, and tyrosine autoinhibition of the catalytic base. For access to the ATP binding site and phosphorylation of the autoinhibitory tyrosine, the C-terminal autoinhibitory tail needs to be removed. Here, we use AFM-based single-molecule force spectroscopy, molecular dynamics simulations, and enzymatics to study the conformational changes during strain-induced activation of human TK. We show that mechanical strain activates ATP binding before unfolding of the structural titin domains, and that TK can thus act as a biological force sensor. Furthermore, we identify the steps in which the autoinhibition of TK is mechanically relieved at low forces, leading to binding of the cosubstrate ATP and priming the enzyme for subsequent autophosphorylation and substrate turnover.

atomic force microscopy | force-probe molecular dynamics simulation | muscle signaling | protein kinase regulation | single-molecule force spectroscopy

Mechanical activity and adaptive responses to changes in work load in muscle are tightly linked, but the mechanosensors triggering the sweeping adaptive changes seen *in vivo* are as yet poorly understood on the molecular level. In the vertebrate muscle sarcomere, titin serves as a molecular ruler for sarcomere assembly and is responsible for resting elasticity of muscle (1, 2) (Fig. 1A). At the M-band, titin contains a serine/threonine protein kinase domain (TK) (Fig. 1B) (3, 4). TK is regulated in a dual autoinhibition mechanism by a C-terminal regulatory tail, blocking the ATP binding site, and tyrosine autoinhibition of the catalytic base by tyrosine-170 (5). For access to the ATP binding site and the autoinhibitory tyrosine, the C-terminal autoinhibitory tail must be removed.

In most autoinhibited kinases, the relief of intramolecular autoinhibition is essentially a partial unfolding event of the autoinhibited conformation, driven by ligand binding or post-translational modification. Although TK activity can be moderately stimulated by calmodulin when tyrosine phosphorylation is mimicked, calmodulin or other calcium binding proteins are unable on their own to activate it (5). Because titin is firmly embedded in the contractile machinery (Fig. 1A), its conformation and function can readily be affected by mechanical forces (1, 6). The M-band, being much more compliant than the Z-disk (7, 8), is ideally placed as a strain sensor (9, 10). Because the M-band lattice is deformed only during active contraction, it is optimal for detecting the actual workload on the myofibril (10). Force-probe molecular dynamics simulations of the mechanical properties of TK suggested that kinase activation might be possible by mechanical forces (11). Indeed, a mechanosensitive signaling complex (signalosome) was identified that interacts with an open conformation of TK, and by controlling protein turnover and

muscle gene transcription (12) seems to contribute to the adaptation of muscle in response to changes in mechanical strain. The importance of TK in maintaining the turnover of muscle proteins is highlighted by a point mutation in the human kinase domain that causes a myopathy with failure of load-dependent protein turnover (12).

Two recent reports on single-molecule force spectroscopy of titin kinase and *C. elegans* giant muscle protein kinase (13, 14) showed that these giant muscle protein kinases can unfold in a stepwise fashion, as predicted (11). To be strain regulated, association of the cytoskeletal lattice with the N- and C-terminal ends of the kinase domain is required. Titin is firmly integrated into the M-band lattice by interactions with obscurin, obscurin-like 1, and myomesin, which form a ternary complex at its C terminus (15). Because of their I-band (16) or broad A-band localization (17), the nematode giant muscle protein kinases have been implicated in contraction regulation. It is therefore as yet unclear whether the invertebrate giant protein kinases are similarly integrated into the cytoskeleton, follow the same activation pathways, and serve analogous functions as the M-band-associated TK.

However, experimental proof of direct mechanical activation, rather than simply partial unfolding, is lacking not only for titin kinase, but for all biological force sensors in muscle (2). Can mechanical force really induce a catalytically competent kinase conformation that will be able to bind substrates? The complex protein composition of the sarcomere M-band precludes an unequivocal experimental answer. Studying single molecules in isolation, however, can unravel their intrinsic properties in molecular detail and allow these to be compared with the known properties of intact sarcomeres and measurable enzymatic properties. We therefore used atomic force microscope (AFM)-based single-molecule force spectroscopy, molecular dynamics simulations, and enzymatics to investigate the molecular details of mechanical TK activation.

Results

Sequential Unfolding of TK at Low Forces. We expressed a TK construct A168-M2, encompassing the human kinase domain flanked by its naturally surrounding Ig/Fn domains [Fig. 1B and C; see details in [supporting information \(SI\) SI Text](#) (section 1), [Fig. S1](#), and [Table S1](#)]. The TK construct was attached to an

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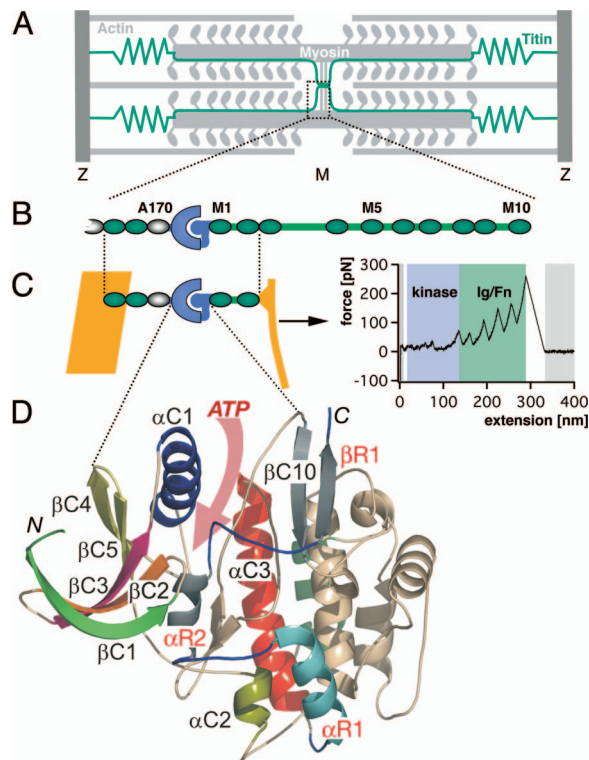


Fig. 1. Sarcomeric location and structure of the investigated TK protein construct. (A) Schematic diagram of the sarcomere showing the transverse Z- and M-bands and actin and myosin filaments, linked by the elastic titin filament. M-bands cross-link myosin filaments by a complex of titin (green), obscurin, and myomesin (15). (B) Domain structure of M-band titin showing the array of structural Ig (green), Fn3 domains (white), and unique sequences (green lines) surrounding titin kinase domain (TK) (blue). (C) The titin construct A168-M2 contains the kinase domain surrounded by Fn3 and Ig domains. ATP binding requires relief of the C-terminal autoinhibitory tail (blue) from the active site, which can be achieved by external force. In mechanical single-molecule experiments, A168-M2 is pulled off the gold support (yellow line) by a cantilever, resulting in a unique force spectrum when the protein is stretched and domains sequentially unfold. Analysis of the unfolding force spectrum (see *SI Text*) identifies the peaks shaded in blue as kinase unfolding peaks; the five unfolding peaks shaded in green correspond to sequential Ig and Fn domain unfolding. (D) Kinase domain structure, with the ATP binding site highlighted by the pink arrow and individual secondary structure elements color-coded. Numbering is from N to C terminus, where C1 to C10 refer to catalytic core structures, and R1 to R3 (in red) refer to the regulatory tail (5). The N and C termini are marked.

AFM cantilever [see *SI Text* (sections 4 and 5) and *Fig. S2*] and stretched with nanometer accuracy. The resulting force, recorded with piconewton precision (*Fig. 1D*), showed a characteristic saw-tooth appearance as TK was gradually stretched and unfolded, mimicking the mechanical stress in muscle. (In a very simple comparison, the slowest experimental pulling speed per folded protein length amounts to $300 \text{ nm/s}/25 \text{ nm} = 12 \text{ s}$ and is close to physiological rates. A rabbit sarcomere of $2 \mu\text{m}$ length can contract with $6 \mu\text{m/s}$, yielding a contraction rate of 3 s .) Typically, a series of five initial low-force peaks below 50 pN was followed by up to five distinct saw-tooth-shaped high-force peaks that correlated exactly with the number and contour lengths of the flanking Ig/Fn-domains (18, 19) [*Fig. 1*; and see *SI Text* (sections 6–8) and *Figs. S3–S5*, and *Table S2*]. Therefore, the low-force peaks, occurring before Ig/Fn unfolding, derive from unfolding events within the kinase domain (see *Fig. S2b* for a schematic). These low-force unfolding events are strictly ordered, although their height is similar. In contrast to the independently unfolding Ig/Fn domains, their fixed sequence is

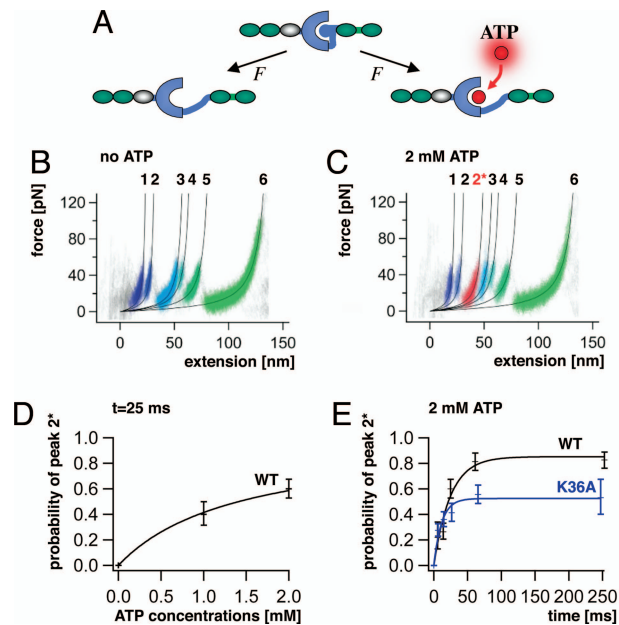


Fig. 2. Unfolding profile of TK and kinetics of mechanically induced ATP binding. (A) External force can open the ATP binding site of TK by unfolding of the autoinhibitory domain (blue ball). (B) Superimposed traces of 66 single-molecule unfolding events in TK show a fixed sequence of local unfolding events, numbered 1–5. (C) Mechanically induced ATP binding leads to a distinctly altered force profile with the appearance of an extra force peak, 2^* , absent in unfolding events in the absence of ATP (44 traces). (D) ATP binding probability (i.e., the occurrence of peak 2^*) depends on ATP concentrations, giving access to virtual direct kinetics though the system is not in equilibrium (ATP peaks detected at pulling rates of 720 nm/s , or after 25 ms). (E) Probability of ATP binding depends on pulling rates, decreasing at faster speeds because of reduced opening time of the ATP binding site. ATP binding is strongly reduced by mutation of lysine-36 to alanine (K36A).

not determined by mechanical stability but rather by topology. The forces required even for complete unfolding of the kinase do not exceed 50 pN at 23°C , or 30 pN at 37°C [see *SI Text* (section 9) and *Fig. S6*] and at pulling speeds of $1 \mu\text{m/s}$. Such low forces were also predicted from force probe simulations (11). The fact that the mechanically more stable Ig/Fn domains always unfold after the kinase domain shows that the force acts on all domains in series, and that the protein construct is therefore completely stretched in the beginning of a retraction cycle.

Mechanically Activated ATP Binding Detected by AFM. Mechanical activation of TK must at an early stage uncover its ATP binding site while leaving the active site intact. Subsequent ATP binding alters the stability of the enzyme (*Fig. 2A*) and should therefore give rise to ATP-dependent changes in the TK unfolding profile (20). In the absence of ATP, five energy barriers separated by 9.1 , 28.6 , 7.3 , 18.0 , and 57.9 nm in contour length are observed (*Fig. 2B*). In the presence of Mg^{2+} -ATP at physiological concentrations (2 mM), a certain fraction of the traces shows an additional well pronounced peak (peak 2^*) at 51.6 nm (*Fig. 2C*). This peak denotes an early interaction with ATP during the sequential unfolding of the kinase and thus demonstrates the initial opening of the active site.

The probability of observing the ATP-dependent peak should depend on the likelihood for ATP binding during the time span between the opening of the binding site and the moment when the ATP barrier (peak 2^*) is probed and, therefore, on the ATP concentration (*Fig. 2D*). More interestingly, because this time span is controlled by the pulling speed, we gain direct experimental access to the ATP binding kinetics (see *SI Text*, section 10). This

Table 1. Kinetic parameters for ATP of autoinhibited titin kinase (WT and K36A) and the constitutively activated WT-kin3 measured by AFM or in solution

Construct	K_{on} , 1/M (AFM)	K_{off} , 1/s (AFM)	K_d , μ M (AFM)	K_M , μ M (solution)	Autophosphorylation
WT	$1.8 \pm 0.3 \times 10^4$	6 ± 3	347	No activity	–
K36A	$2.3 \pm 0.5 \times 10^4$	41 ± 11	1,810	No activity	–
WT-kin3				270	+

experiment can be seen as a mechanical pump–probe experiment: first the binding pocket of the TK is “pumped” open, and after a certain time ATP binding is probed. Variation of this time window provides the kinetic constants. The calibration of the time axis was estimated by the ratio of the MD-determined extension, during which the binding pocket is open but not deformed, and by the pulling speed. This mechanical pump–probe experiment showed saturation after ≈ 100 ms (Fig. 2E). Toward higher pulling rates, the probability of ATP binding decreased strongly and approached zero (Fig. 2E). This time dependence demonstrates that ATP binding is mechanically induced and perfectly agrees with the absence of catalytic activity of autoinhibited TK in solution (Table 1). Furthermore, this experiment allows estimates of the apparent on and off rates and resulting dissociation constant, which compare with affinity values observed for titin (Table 1) and other kinases in solution (21).

Following a suggestion from our MD simulations, we mutated lysine-36 to alanine (K36A), a highly conserved residue equivalent to lysine-72 interacting with the α/β phosphates of ATP in cAMP-dependent protein kinase (22, 23). This mutation abolishes kinase activity in TK (5). Now the ATP affinity of TK was dramatically reduced, with a >6 -fold increase of k_{off} and a concomitant increase of K_d to millimolar values (Fig. 2D, *SI Text*, section 10 and Table 1). These results localize mechanically induced ATP binding to the canonical site in TK and confirm that the conserved lysine residue, known to play a crucial part in ATP binding of homologous protein kinases, is also a key residue in the TK binding pocket.

Molecular Mechanism of TK Activation by Force. We used force–probe MD simulations (24, 25) to characterize the force-induced unfolding of TK at the atomic level and to correlate the structural states with the energy barriers observed by the single-molecule force spectroscopy experiments. Force–probe molecular dynamics simulations (24, 25) used the TK x-ray structure [Protein Data Bank entry 1TKI (5)] as the starting structure, with the autoinhibitory tail partly removed [see *SI Text* (sections 11–16) and *Tables S3 and S4*]. Two sets of simulations (five each) were carried out for this truncated TK: one set with an empty binding pocket, and one set with an ATP molecule and magnesium ions inserted into the (closed) binding pocket. As a control, the autoinhibited complete TK was also subjected to force–probe MD simulations (see *SI Text*, section 17). As in the experiment, the two force profiles obtained from the simulations of the truncated TK (Fig. 3B *Top* and *Middle*) are largely similar. A notable exception is the more pronounced force peak seen in the presence of ATP (see Fig. 3B *Inset*) at the position of the measured force peak 2*. To allow direct comparison of the unfolding pathways between experiment and simulation, we transformed the force extension traces of Fig. 2 into barrier position histograms (14) and derived the same from our simulations (see *SI Text*, section 18). The two histograms agree well both in the presence and absence of ATP [Fig. 4 (dashed lines) and Fig. S7], allowing the conclusion that the main unfolding events are correctly described by the simulations.

Next, we investigated which molecular interactions determine the observed force peaks. For the ATP peak 2*, two strong interactions

are seen, a salt bridge from lysine-36 to the α -phosphate group of ATP, and a contact between methionine-34 and the adenine moiety of ATP (Fig. 3C). Both interactions break irreversibly upon β C3– β C4 rupture, giving rise to the significantly larger force peak of 270 ± 39 pN in the simulations with bound ATP as compared with 188 ± 13 pN without ATP (Fig. 3B *Inset* and *Movie S1*). Notably, in the AFM experiment, the contour length of 51.6 nm for the ATP peak position (Fig. 4, peak 2*) also points to a residue close to lysine-36. An additional peak is seen at 18 nm for the simulation with ATP present (plus sign in Fig. 3B *Top*). Here, a force-induced deformation of the N-terminal domain triggers the transient rupture and reformation of the methionine-34–ATP and lysine-36–ATP interactions.

Closer structural analysis of our simulations suggests the following sequence of events (colors in Fig. 3A and B, and Fig. 4). Peak 1 (Fig. 4) is caused by unfolding of the 23-residue linker at the N terminus of TK, which is not present in the simulations (see *SI Text* for details). At peak 2, the autoinhibitory tail is unfolded and removed, rendering the ATP binding site accessible (region shaded in gray in Fig. 3B *Bottom*). Subsequently, N-terminal β -sheets β C1– β C2 and β C2– β C3 rupture (regions B and C). For these events, no force peak is seen in the experiment, because it would fall into the lag time after force peak 2. Peak 2* described above is dominated by interactions of ATP with the binding pocket. The truncated construct necessarily lacks part of the autoinhibitory tail stabilizing the adjacent C-terminal α -helix α R1 in the full-length TK. Accordingly, α R1 unfolds first in the truncated kinase (Fig. 3B *Top* and *Middle*, region A) but after β R1 and α R2 in the autoinhibited kinase (Fig. 3B *Bottom*). Hence, and in agreement with the complete TK unfolding simulations (Fig. 3B *Bottom*), peaks 3 and 4 are assigned to unfolding of α C1 and β C4– β C5, respectively (regions D and E). Finally, peak 5 arises from the combined effect of α C2 and α C8 rupture (Fig. 3B *Bottom*). At peak 6, the complete TK is unfolded and stretched. Taking the diameter of the folded TK into account (5.5 nm), the contour length increment to peak 1 (121 nm) corresponds to $(5.5 + 121 \pm 2) \text{ nm} / 0.365 \text{ nm} = 346 \pm 5$ residues, in agreement with the 344 aa of TK including its N-terminal linker (see *SI Text*, section 7).

Autophosphorylation of TK. Our simulations show that the open ATP binding site does not relieve autoinhibition of the catalytic base aspartate-127 by tyrosine-170. However, our model of the ATP-bound state of TK suggests that this semi-opened state might autophosphorylate, in agreement with previous predictions of the open apo-enzyme (11). We tested this notion by assaying recombinant TK with its ATP binding site released (TK-kin3, mimicking the mechanically induced open state after peak 2), and found that the release of ATP binding not only activates kinase activity toward substrates like telethonin, but also allows tyrosine autophosphorylation (see *SI Text*, sections 1 and 2, for experimental details). As shown in Fig. 5, although low levels of phosphotyrosine are detected by the 4G10 antibody before incubation with ATP, tyrosine phosphorylation is strongly stimulated by ATP, with a preference toward Mn^{2+} , similar to other enzymes (26, 27).

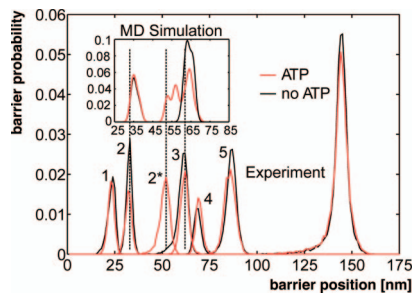


Fig. 4. Contour length histograms obtained from single-molecule force spectroscopy experiments (transformation with QM-WLC and $P = 0.8$ nm) and from MD simulations (*Inset*). The folded kinase construct has a length of 25 nm. The peak positions with (red) and without (black) ATP are similar in both histograms (dashed lines), except for one additional peak in the presence of ATP (red peak at ≈ 51.6 nm). The experimentally determined contour length increments are, in the absence of ATP, 9.1, 28.6, 7.3, 18.0, 57.9 nm; and, in the presence of ATP, 9.1, 19.4, 10.1, 7.5, 16.4, 58.3 nm—with an estimated error of $\pm 2\%$. The position of the initial peak (24 nm) reflects the mean length of the TK construct with completely folded domains.

the homologous nematode kinases, which retain catalytic activity in their inhibited form (13), we show that TK is completely inactive in its autoinhibited form. Mechanical switching of its ATP-binding site thus confers a significant signal between active and inactive kinase, as expected for a signal that modulates energy-costly processes like protein breakdown and transcriptional activity (12).

Our surprising observation that a protein kinase can be activated by local protein unfolding induced by mechanical force may find analogies in the small GTPase Rab8, whose activation by the nucleotide exchange factor MSS4 also involves local protein unfolding (35). The mechanoenzymatic sensor found in titin kinase may therefore be paradigmatic also for other members of the family of cytoskeletal autoregulated protein kinases, a branch of the calcium-calmodulin-regulated enzymes of the human kinome (36), containing myosin light-chain kinase and obscurin kinases. These enzymes share with titin the N- and C-terminal cytoskeletal association (37) or specific residues involved in autoinhibition (38) and may thus bear features of mechanical modulation. Furthermore, other autoregulated cytoskeletal signaling domains, like GDP-GTP exchange factor

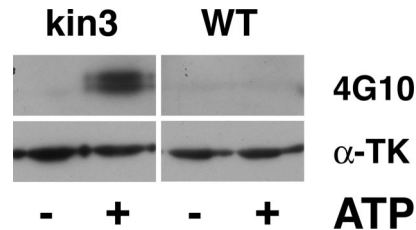


Fig. 5. Autophosphorylation of TK on tyrosine. Incubation of the highly purified TK-kin3 enzyme in the absence (–) and presence (+) of ATP and Mn^{2+} ions leads to tyrosine phosphorylation detected by Western blot, using the phosphotyrosine antibody 4G10. The autoinhibited kinase construct A168-M2 (WT) shows no appreciable phosphotyrosine incorporation under any tested condition. Lower blot: loading control, detection with anti-titin kinase antibody (α -TK).

domains, may be similarly activated. Our single-molecule approach will therefore be useful for investigating the mechanochemistry of many cellular systems that may share similar mechanosensitive regulation mechanisms.

Materials and Methods

Titin kinase expression was carried out in sf9 insect cells by using a recombinant baculovirus system essentially as described in ref. 5. Purification and enzymatic assays were performed essentially as described in refs. 5 and 12 (for details, see *SI Text*). Atomic force microscopy using a custom-built instrument, and analysis of the data were carried out essentially as described (14, 19); for details see the SI. Force-probe molecular dynamics simulations (24, 25) used the TK x-ray structure [Protein Data Bank entry 1TKI (5)] as the starting structure, with the autoinhibitory tail partly removed. Two sets of simulations (five each) were carried out for this truncated TK, one set with an empty binding pocket, and one with an ATP molecule and magnesium ions inserted into the (closed) binding pocket. As a control, the autoinhibited complete TK was also subjected to force-probe MD simulations. For further details, see *SI Text*.

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