α-catenin switches between a slip and a cooperative catch bond with F-actin to regulate cell junction fluidity

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α-catenin is a crucial protein at cell junctions that provides connection between the actin cytoskeleton and the cell membrane. At adherens junctions (AJs), α-catenin forms heterodimers with β-catenin that are believed to resist force on F-actin. Outside AJs, α-catenin forms homodimers that directly connect the cell membrane to the actin cytoskeleton, but their mechanosensitive properties are inherently unknown.

Surprisingly, by using ultra-fast laser tweezers we found that a single α-β-catenin heterodimer does not resist force but instead slips along F-actin in the direction of force. Conversely, the action of 5 to 10 α-β-catenin heterodimers together with unidirectional force applied to F-actin engaged a molecular switch in α-catenin, which unfolded and strongly bound F-actin as a cooperative catch bond. Similarly, an α-catenin homodimer formed an asymmetric catch bond with F-actin triggered by protein unfolding under force. Our data reveal that α-catenin clustering together with
intracellular tension engage a fluid-to-solid phase transition at the membrane-cytoskeleton interface.

In any living cell, an array of mechanotransducer proteins responds to mechanical cues to trigger complex molecular signaling, driving cell morphology and gene expression profiles\(^1\). Among these proteins, \(\alpha\)-catenin has been reported to act as a mechanosensor that regulates adherens junctions (AJ) in response to mechanical cues from neighboring cells in a tissue\(^2\). It is widely accepted that \(\alpha\)-catenin in AJ binds \(\beta\)-catenin, which, in turn, connects to the cytoplasmatic portion of E-cadherin to constitute the cadherin-catenin complex (CCC). Since \(\alpha\)-catenin also binds F-actin, \(\alpha\)-catenin has been indicated as a major candidate for providing a link between the CCC and the actin cytoskeleton. This molecular link is required to provide mechanical stability to AJ. According to the most accepted model of \(\alpha\)-catenin mechanosensitivity, tension on the actin cytoskeleton is transmitted to bound \(\alpha\)-catenin inducing a conformational change that opens up (i.e. unfolds) its central M-domain and increases the bond strength\(^2\). \(\alpha\)-catenin unfolding is predicted to expose cryptic binding sites for vinculin and other actin-binding proteins, which further reinforce the connection to the actin cytoskeleton. However, the role of \(\alpha\)-catenin as the mechanical link between the CCC and F-actin has been challenged by the finding that the binding of mammalian \(\alpha\)-catenin to F-actin and \(\beta\)-catenin is mutually exclusive\(^3\).

Using a *Danio Rerio* \(\alpha\)-catenin\(^4\), Buckley et al. recently demonstrated that the bond between CCC and F-actin is reinforced by increasing force (i.e. it is a catch bond)\(^5\). However, it remains to be determined whether mammalian \(\alpha\)-\(\beta\)-catenin heterodimers show similar catch-bond properties. Moreover, it is still unclear whether a single \(\alpha\)-\(\beta\)-catenin heterodimer can bind to F-actin and bear force on it. CCCs have been recently shown to be organized in
clusters in cells\textsuperscript{6}. This finding raises the intriguing question whether CCC clustering and the interaction of F-actin with multiple $\alpha$-catenins might be important in defining the mechanosensitivity of AJ.

Recent reports revealed the importance of $\alpha$-catenin outside AJ as well, where it regulates F-actin organization\textsuperscript{7} and promotes cell adhesion and migration by providing a direct link between the cell membrane and the actin cytoskeleton\textsuperscript{8}. While $\alpha$-catenin forms $\alpha$-$\beta$-catenin heterodimers in AJ, outside AJ $\alpha$-catenin binds itself and is prevalently found as homodimers. Despite its biological relevance, no data is currently available on the force-dependence of the interaction between $\alpha$-catenin homodimers and F-actin.

Finally, it is still unclear whether $\alpha$-catenin unfolds during its interaction with F-actin. Using a truncated $\alpha$-catenin that lacked the actin and $\beta$-catenin binding domains, Yao et al. showed that forces of about 5 pN trigger reversible unfolding of the $\alpha$-catenin M-domain and allow vinculin binding\textsuperscript{9}. However, there is no direct evidence that forces applied to $\alpha$-$\beta$-catenin heterodimers or $\alpha$-catenin homodimers through F-actin are sufficient to induce $\alpha$-catenin unfolding.

\textbf{Results}

\textbf{A single $\alpha$-catenin homodimer unfolds to form a catch bond with F-actin}

Here, we used ultrafast force-clamp spectroscopy, a technique with sub-millisecond and sub-nanometer resolution based on laser tweezers\textsuperscript{10–13}, to dissect the mechanosensitivity of single mammalian $\alpha$-catenin homodimers and $\alpha$-$\beta$-catenin heterodimers. We first analyzed single purified recombinant $\alpha$-catenin homodimers (named as $\alpha$-catenin below). A single actin filament was suspended between two optically trapped beads and brought in close proximity to a single $\alpha$-catenin that was attached to a micron-sized glass bead on the
coverslip surface (Fig. 1a, Methods section). A constant force was applied to the bead-actin-bead complex (herein named “dumbbell”) through a double force-clamp system. The applied force caused the dumbbell to move through the assay solution at a constant velocity and the force was alternated in direction to maintain the dumbbell within few hundred nanometer excursion (Fig 1b, “1. Unbound”). When α-catenin bound to the actin filament and became loaded with the applied force, the dumbbell motion stopped within ~30 µs (Fig 1b, “2. Bound”). The interactions between α-catenin and the actin filament were detected through the change in the dumbbell velocity and their lifetimes and position assessed under forces in the range of ±17 pN (see Methods).

The lifetime of the interactions visibly increased with force (Fig. 1c and Fig. 1d-f). Plot of the event lifetime versus force showed that the bond between α-catenin and F-actin is reinforced by increasing force and responds asymmetrically with respect to the force direction (Fig. 1c). Event lifetime at 2.3 pN (7.5 ± 0.8 ms) is slowed down about 19-fold at 8.1 pN (145 ± 16 ms), reduces to 9.5 ± 2.6 ms at 11.5 pN and shows a second maximum at 13.8 pN (61 ± 11 ms). A similar effect was observed for force applied in the opposite direction, although the bond lifetime was about 7-fold shorter in the 0-8 pN range and with maximum lifetime at slightly different force values. Event lifetime at -4.3 pN (1.25 ± 0.07 ms) is slowed down about 16-fold at -7.7 pN (20.5 ± 2.9 ms), reduces to 9.0 ± 1.8 ms at -8.6 pN, and shows a second maximum at -10.3 pN (42 ± 6 ms). Overall, these results show that a single α-catenin forms an asymmetric catch bond with F-actin that resists forces below 15 pN. The presence of two peaks in the interaction lifetime suggests the emergence of two conformational changes of α-catenin induced by force. Accordingly, we could fit each peak in the lifetime distribution with a two-state catch-bond model in which force triggers switching between a weak-binding and a strong-binding state (green and blue curves in Fig. 1c; see also methods and supplementary figure 1). The distance parameters for the transitions between the two
states indicate that large conformational changes (> 10 nm) occur during these transitions (supplementary table 1).

Analysis of α-catenin binding position at different forces further supports this view. Observation of position records showed that interactions occurred at single locations for forces below ~3-4 pN (Fig. 1d and supplementary figure 2a), whereas above ~5 pN most interactions showed "steps" (red arrowheads in Fig. 1e,f and supplementary figure 2b,c).

This behavior raises the question whether the observed steps were a consequence of (i) α-catenin detaching and rapidly reattaching to the actin filament or (ii) α-catenin undergoing large conformational changes (i.e. unfolding) under force\(^9\), as suggested by the analysis of the interaction lifetime. The former hypothesis would imply that steps occurred only in the direction of the force, following the periodicity of the actin filament. However, analysis of step size distribution at about +5 pN showed a main peak in the force direction centered around 12.1 ± 5.8 nm and the presence of a step in the opposite direction of similar amplitude (-11.2 ± 5.1 nm) (Fig. 1h). Likewise, step size distribution at -5 pN showed steps of similar amplitude in both directions (-11.4 ± 6.4 nm and 10.3 ± 4.2 nm) (Fig. 1g). Analysis of individual records around 5 pN force display α-catenin jumping back and forth between two position levels separated by about 10-20 nm (supplementary figure 2d). This indicates that, around 5 pN, α-catenin is near the equilibrium between the folded and the unfolded state and switches between the two states because of thermal noise. Our results are in very good agreement with those from Yao et al., who measured a reversible unfolding step on the α-catenin M-domain occurring around the same force and with similar amplitude\(^9\). At higher force (±11 pN), the negative step is largely reduced (Fig. 1j) or disappears (Fig. 1i and supplementary figure 3), confirming the idea that force drives the unfolding of the protein in the force direction and that the reverse folding step is inhibited at large forces. Moreover, larger irreversible steps of about 35 and 71 nm become apparent in the position records.
Fig. 1: A single α-catenin homodimer unfolds and resists force on F-actin. 

**a.** Schematic of ultrafast force-clamp spectroscopy applied to a single α-catenin homodimer. An α-catenin molecule is attached to a glass bead stuck on the coverslip. An actin filament is suspended between two beads trapped in optical tweezers. Black arrows are the force clamped on the right ($F_1$) and left bead ($F_2$), red arrow is the net force ($F = F_1 - F_2$) on the dumbbell. $F$ is alternated in direction to maintain the dumbbell within a limited excursion. 

**b.** Left: illustration showing the position of the right bead when: (1) α-catenin is unbound and the dumbbell oscillates; (2) α-catenin binds to the actin filament. Right: example trace showing displacement and force ($F \sim 6$ pN) during the corresponding phases of dumbbell oscillation and α-catenin attachment, under positive and negative loads. 

**c.** Load-dependent lifetime of the interaction between a single α-catenin homodimer and F-actin. Green and blue lines are, respectively, fit of the low-force and high-force peak with the two-state catch-bond model (see methods). $n = 23234$. Error bars, s.e.m. 

**d-f.** Example traces under force of about 3 pN, 5 pN, and 10 pN, respectively. Red arrowheads indicate steps occurring during the interaction. 

**g-j.** Distribution of step size at various forces. Gaussian fits of the main peaks are also shown with their center and s.d. $g$, $F = -5$ pN, $s_1 = -11.3\pm6.4$ nm, $s_2 = 10.3\pm4.3$ nm $h$, $F = +5$ pN, $s_1 = 12.1\pm5.8$ nm, $s_2 = -11.2\pm5.1$ nm $i$, $F = -11$ pN, $s_1 = -15\pm12$ nm $j$, $F = +11$ pN, $s_1 = 14\pm12$ nm, $s_2 = 35\pm24$ nm, $s_3 = 71\pm15$ nm. Error, s.d.
(Fig. 1f and supplementary figure 2c) and step distributions (Fig. 1j) in the direction of the force. These larger steps are compatible either with additional irreversible unfolding steps as observed by Yao et al. at about 12 pN⁹, or from rapid unbinding-rebinding of α-catenin from the actin filament, although the presence of the above mentioned two peaks in the interaction lifetime supports the former hypothesis.

**A single α-β-catenin heterodimer forms a slip bond with F-actin**

We then analyzed the interaction between a single α-β-catenin heterodimer and an actin filament (Fig. 2a). Plot of the event lifetime versus force showed that the bond between a single α-β-catenin heterodimer and F-actin does not bear force (Fig. 2b). The bond lifetime decreased with force, occurring in the millisecond time scale at forces around 5 pN and rapidly dropping below 1 ms for F > 7pN. We could detect such rapid interactions thanks to the unprecedented time resolution of our technique¹⁰. The lifetime vs force plot was well fitted by the Bell slip-bond model, in which the lifetime decreases exponentially with force (green lines in the log plot of Fig. 2b). Position records show that interactions of the α-β-catenin heterodimer also included “steps” (supplementary figure 4). However, the step size distribution was very different than in the case of the α-catenin homodimer, displaying (i) negligible number of negative steps and (ii) strong 5.5 nm periodicity, which is characteristic of the actin monomer separation¹⁴,¹⁵ (Fig. 2c,d and Supplementary figure 5). The actin filament is made of two protofilaments composed by 13 monomers per helix turn, giving a periodicity of 13 x 5.5 nm = 71.5 nm. The two protofilaments are separated by a stagger of 2.75 nm, giving an apparent periodicity of 6 x 5.5 nm + 2.75 nm = 35.75 nm (Fig. 2e). Figure 2f shows the peak position, obtained from the step size distribution in Fig. 2d, against the peak number, as defined in Fig. 2d. Linear regression analysis showed that the points
Fig. 2: a single α-β-catenin heterodimer forms a slip bond with F-actin that does not bear force.  

**a**, Schematic of ultrafast force-clamp spectroscopy applied to a single α-β-catenin heterodimer.  

**b**, Load-dependent lifetime of the interaction between a single α-β-catenin heterodimer and F-actin. Green lines are fits of lifetimes $\tau$ with the Bell-bond equation $\tau = \tau_0 \exp \left( \frac{-d_{\alpha\beta}}{k_B T} \right)$ (see Methods). Fitting parameters are $d_{\alpha\beta} = 0.88 \pm 0.05$ nm, $d_{\alpha\beta}^- = 1.22 \pm 0.08$ nm, $\tau_0^+ = 6.1 \pm 0.8$ ms, $\tau_0^- = 16 \pm 4$ ms. $n = 360486$. Error bars, s.e.m.  

**c**-**d**, Distribution of step size versus force for negative (c) and positive force (d).  

**e**, Illustration showing the actin filament structure and highlighting the two protofilaments (red and green), the helix and apparent periodicity, the distance between consecutive monomers on the same protofilament, and the stagger between monomers on two consecutive protofilaments.  

**f**, Plot of the position of peaks obtained from the step histogram in (d) against peak number. Red squares correspond to monomers on the same protofilament. Black squares correspond to monomers on the adjacent protofilament. Red and black dashed lines are the least-squares fittings of the red and black points, giving slopes of $5.56 \pm 0.01$ and $5.54 \pm 0.04$ nm/monomer respectively, and a stagger between protofilaments of $2.56 \pm 0.31$ nm. Errors, s.e.m.
nm. The line slopes clearly corresponded to the distance between contiguous actin monomers, while the distance between the two lines corresponded to the protofilament stagger.

These results strongly indicate that the observed steps are due to a single \( \alpha \)-catenin that rapidly unbinds from an actin monomer, slips along the actin filament in the direction of the force, and rebinds to a neighbor actin monomer. Moreover, no apparent unfolding was detected. In fact, all peaks corresponded to the actin monomer separation while a negligible number of negative steps was detected. Inspection of records at any force did not show \( \alpha \)-catenin jumping back and forth between two position levels as in the case of the \( \alpha \)-catenin homodimer. Overall, these results indicate that a single \( \alpha \)-\( \beta \)-catenin heterodimer forms a slip bond with F-actin that does not resist force. Force applied by the actin filament to the \( \alpha \)-\( \beta \)-catenin complex causes a rapid slip to a neighbor actin monomer rather than inducing \( \alpha \)-catenin unfolding and strong binding to actin.

**Multiple \( \alpha \)-\( \beta \)-catenin heterodimers form a cooperative catch bond with F-actin**

Since single mammalian \( \alpha \)-\( \beta \)-catenin complexes cannot bear force on actin, we questioned whether the cooperative action of multiple complexes might resist the physiological forces that occur at AJ. We, thus, increased the \( \alpha \)-\( \beta \)-catenin heterodimer concentration 10-fold to have multiple proteins interacting simultaneously with the actin filament (Fig. 3a). Similarly to what we observed using \( \alpha \)-catenin homodimers, the bond between \( \alpha \)-catenin and F-actin was reinforced by increasing force and responded asymmetrically with respect to the force direction, with lifetime peaks centered about 7.2 pN and 11.4 pN (Fig. 3b). Moreover, an additional peak was detected at the highest force that we could probe (~17 pN). The asymmetric response to force was more pronounced here, resulting in a catch bond for force...
applied in one direction and a slip bond in the opposite direction (Fig. 3b and supplementary figure 6).

**Fig. 3:** Multiple α-β-catenin heterodimers restore force-bearing capability on F-actin. 

- **a:** Schematic of ultrafast force-clamp spectroscopy applied to multiple α-β-catenin heterodimers. 
- **b:** Load-dependent lifetime of the interaction between multiple α-β-catenin heterodimers and F-actin. \( n = 55171 \). Error bars, s.e.m. 
- **c-f:** Distribution of step size at various forces. Gaussian fits of the main peaks are also shown with their center and s.d. 

(c) \( F = -5 \) pN, \( s_1 = -11.7 \pm 2.9 \) nm, \( s_1 = 10.2 \pm 1.8 \) nm, 

(d) \( F = +5 \) pN, \( s_1 = 12.8 \pm 3.8 \) nm, \( s_1 = -12.1 \pm 4.0 \) nm, 

(e) \( F = -15 \) pN, \( s_1 = -11.9 \pm 2.9 \) nm 

(f) \( F = +15 \) pN, \( s_1 = 11.6 \pm 3.1 \) nm, \( s_1 = -10.5 \pm 2.1 \) nm. Error, s.d.

Position records were also similar to the α-catenin homodimers, mainly composed of single events at forces below ~3 pN and multiple “steps” above ~5 pN (supplementary figure 7). Step size distribution at about -5 pN showed a main peak centered around -11.7 ± 2.9 nm in the force direction and the presence of a step in the opposite direction of similar amplitude (10.2 ± 1.8 nm) (Fig. 3c). Similarly, step size distribution at about +5 pN showed a main
peak centered around 12.8 ± 3.8 nm and the presence of a negative step of similar amplitude (-12.1 ± 4.0 nm) (Fig. 3d). Analysis of individual records at force greater than 5 pN often displayed α-catenin jumping back and forth between two position levels separated by 10-20 nm (supplementary figure 7). The negative step rapidly disappeared at increasing negative forces in which the bond slips, but it was still present at the highest positive forces in which the bond lifetime is maximum (Fig. 3f and extended figure 8b).

Discussion

Here, we provided direct evidence that a single mammalian α-catenin bound to β-catenin forms a slip bond with F-actin. The weak and rapid interactions that we observed could be detected thanks to the 10-100 μs time resolution of our setup, which is about two orders of magnitude better than conventional optical trapping assays. The limited time resolution is likely the reason why previous reports could not detect the slip bond between a single α-β-catenin heterodimer and F-actin\(^5\), whereas they could possibly detect the action of multiple molecules, which behave as a catch bond and produce much longer interactions than single molecules under force.

Our results indicate that α-catenin binds F-actin as an unconventional slip bond that cooperatively switches into a catch bond. A cooperative action of multiple α-catenin molecules is required because a mere summation of multiple slip bonds would result again in a slip bond. Further insight into this mechanism can be gained by analysis of the duty ratio \( r \), i.e. the fraction of time that a molecule spends attached to the actin filament: \( r = \frac{\tau_{on}}{\tau_{on} + \tau_{off}} \)^\(^16\). When the number of α-catenin molecules that can interact with the actin filament is above \( N_{min} = 1/r \), at least one molecule is bound to actin on average and bear force on it (see Supplementary Methods). The duty ratio of a single α-catenin homodimer rapidly switches from \( r < 0.2 \) for low forces (\( F < 5 \) pN) to \( r = 0.8-0.9 \) in the force range where
also the lifetime is maximized (5 < F < 15 pN, Fig. 4a). Therefore, α-catenin homodimers under force remain bound to actin most of the time and an average number of 1.1-1.2 molecules can sustain force on an actin filament for a prolonged time. Contrary, the duty ratio of a single α–β-catenin heterodimer is roughly constant to \( r \sim 0.1-0.2 \) independently of force (Fig. 4b), whereas the cooperative action of multiple molecules is required to increase the duty ratio to \( r \sim 0.7 \) under directional force (Fig. 4c). The \( r \sim 0.1-0.2 \) for a single α–β-catenin heterodimer indicates that about 5-10 molecules are required to maintain continuous contact with the actin filament. Strikingly, CCC in AJ have been found to cluster in nanodomains composed by 5-6 molecules\(^\text{17}\). Our data can, thus, provide an explanation of the biological relevance of CCC clustering\(^\text{6,17–19}\). In fact, below 5-10 molecules the probability of finding multiple molecules bound simultaneously to an actin filament is small. The engagement of the cooperative catch bond mechanism observed here is unlikely, and AJ would not be able to bear force on actin. Previous studies indicate that α-catenin binds cooperatively to F-actin by altering the actin filament conformation\(^\text{20}\), which might be the molecular mechanism at the base of the change in the load-dependence from single to multiple α–β-catenin heterodimers that we observed. Moreover, such asymmetric cooperative load-dependence might itself (i) cause unidirectional flow of CCC clusters in cell-cell junctions\(^\text{21}\) and (ii) drive cluster formation through increased diffusional trapping mediated by α-catenin cooperativity and tension applied by myosin motors on the actin cytoskeleton\(^\text{6}\).

Notably, our data show that the catch-bond behavior is accompanied by a large conformational change (unfolding) of the α-catenin molecule, both in a single α-catenin homodimer and in multiple α-β-catenin heterodimers. Unfolding is not observed in a single α-β-catenin heterodimer slip bond, indicating that α-catenin unfolding provides the molecular switch that converts the link to F-actin into a catch bond. The unique ultra-fast
force-clamp configuration used here allowed us to directly observe under which conditions folding/unfolding transitions of the α-catenin molecule actually occur when force is applied to the interacting actin filament, as previously hypothesized⁹.

Fig. 4: α-catenin clustering and intracellular tension regulate cell junction fluidity. a, Top: Illustration showing α-catenin in filopodia. α-catenin is present in the form of homodimers bound to PIP₃ on the plasma membrane. Middle: duty ratio of a single α-catenin homodimer. Forces above 5 pN induce α-catenin unfolding and large increase in duty ratio. Bottom: As a consequence, interaction between α-catenin and F-actin switches from a fluid to a solid response. b, Top: Illustration showing α-catenin in immature adherens junctions. α-catenin is present in the cadherin-catenin complexes at low density. Middle: duty ratio of a single α-β-catenin heterodimer is not affected by force. Bottom: Interaction between α-catenin and F-actin is fluid regardless of the tension on the actin cytoskeleton. c, Top: Illustration showing α-catenin in mature adherens junctions. α-catenin is present in the cadherin-catenin complexes which cluster in groups of 5-6. Middle: duty ratio of a multiple α-β-catenin heterodimers. Forces above 5 pN induce α-catenin unfolding and large increase in duty ratio. Bottom: As a consequence, interaction between α-catenin and F-actin switches from a fluid to a solid response.
Based on our results, we developed a simple chemo-mechanical model to relate the sliding velocity of an actin filament interacting with $\alpha$-catenin ($v$) to the force applied to it ($F$), as a function of the $\alpha$-catenin duty ratio ($r$): $F = \frac{\gamma}{1-r} v$, where $\gamma$ is the cytosol viscous drag (see supplementary methods). This relation tells us that when the duty ratio is significantly smaller than one and independent of force, as in the case of a single $\alpha$-$\beta$-catenin heterodimer (Fig. 4b), the $\alpha$-catenin-actin interaction creates additional viscous drag on the actin filament, but the system behaves as a fluid ($F \propto v$). However, if the duty ratio gets closer to one, as it occurs when force is applied to a single $\alpha$-catenin homodimer (Fig. 4a) or to multiple $\alpha$-$\beta$-catenin heterodimers (Fig. 4c), the velocity drops to zero and the system behaves as a solid.

Therefore, $\alpha$-catenin acts as a cooperative molecular switch that responds to tension to regulate the fluidity of its link to actin. This result is particularly notable in light of recent findings of the occurrence of fluid-to-solid transitions in cultured epithelial monolayers\textsuperscript{22} and in-vivo\textsuperscript{23}. Notably, disruption of E-cadherin clusters in living cells has been shown to increase cell-cell junction fluidity\textsuperscript{24}. We thus propose that, as a consequence of $\alpha$-catenin cooperative mechanosensitivity, AJ might switch from a fluid phase in the absence of CCC clustering to a solid phase when CCC become organized into clusters and tension builds up on the actin cytoskeleton.
Methods
Protein expression and purification

The pET28a vector containing the mouse αE-catenin was an offering from Dr. Noboru Ishiyama\textsuperscript{25}. The plasmid encoding the protein fused with a His-tag at the N-terminus was transformed in BL21 DE3 pLysS strains (Agilent). Recombinant αE-catenin was expressed via isopropyl 1-thio-β-d-galactopyranoside (IPTG, Sigma-Aldrich) induction. Cultures were grown to an OD of 0.7 and induced with 0.5 mM IPTG for 3 h at 37 °C. Protein was purified via FPLC (AKTA prime system, GE Healthcare) using HisTrap HP columns and eluted in 25 mM Tris, pH 7.5, 300 mM NaCl, 300 mM Imidazole. Size exclusion chromatography was performed in 25 mM Tris, pH 7.5, 150 mM NaCl, 1 mM DTT through with a HiLoad Superdex 200 (GE, Healthcare) as a further purification step. Monomeric and homodimeric fractions were pooled separately and purity was evaluated by SDS-PAGE gel electrophoresis. Protein concentration was measured using absorbance at 280 nm. Human β-catenin, GST tagged was bought from Sigma-Aldrich (SRP5172). G-actin and biotinylated actin were bought from Cytoskeleton, Inc. (AKL99, AB07-A). Actin filaments were obtained by incubating 0.4 μg·μl\textsuperscript{−1} biotinilated actin and 0.2 μg·μl\textsuperscript{−1} actin in 50 mM KCl, 2 mM MgCl\textsubscript{2} and 1 mM ATP in 10 mM Tris, pH 7.5 (actin polymerization buffer, Cytoskeleton, Inc. BSA02) for an hour. F-actin filaments were labeled overnight with 10 μM rhodamine phalloidin and 20 mM DL-Dithiothreitol.

Pull down and flow-cell assay

For the flow-cell assay, a 1% nitrocellulose-pentyl acetate solution was smeared on a coverslip and assembled to make a flow chamber as described previously\textsuperscript{26}. For reactions containing β-catenin, 80 μg/ml of protein was allowed to react with the surface for 5 minutes. αE-catenin was then incubated for 5 minutes to allow binding. Unbound protein was washed
away with a solution containing 20 mM MOPS pH 7.2, 50 mM KCl, 1 mM MgCl₂, 0.1 mM EGTA and 1 mg/ml bovine serum albumin, following incubation with a solution containing 100 nM rhodamine phalloidin F-Actin in 20 mM MOPS pH 7.2, 50 mM KCl, 1 mM MgCl₂, 0.1 mM EGTA, 1.2 µM glucose-oxidase, 0.2 µM catalase, 17 mM glucose and 20 mM DTT. Flow-cell assays were performed on an inverted fluorescence microscope (Nikon ECLIPSE TE300) equipped with a 532 nm laser (Coherent Sapphire) for rhodamine excitation (~3 mW on the sample). Images were acquired in total internal reflection configuration, through Nikon Plan Apo TIRF, 1.45 oil immersion objective on an EMCCD camera (Andor, iXon X3) with 82 nm pixel size, and 40x40 um² field of view. Integration time was 0.2 s, and EM gain 40.

For the pull-down assay, 3 µM F-actin was incubated with increasing concentrations of αE-catenin (0.05 µM-8 µM) in 20 mM MOPS pH 7.2, 50 mM KCl, 0.1 mM EGTA, 1 mM MgCl₂. Binding reactions were performed at room temperature for 20 min and spun at 13000 rpm for 2h at 4°C. The supernatant was removed and the pellet resuspended in the original volume of assay buffer. Laemmli Sample Buffer (Sigma-Aldrich) was added to the samples and pellet fractions were run on a 4-12% BIS-Tris gel (Thermo Fisher Scientific). The gel was stained with EZBlue reagent (Sigma-Aldrich) and scanned with a ChemiDoc™ MP Imaging system (Bio-Rad) and analyzed by densitometry with Image Lab™.

**Optical trapping experiments**

Preparation of neutravidin beads has been described in detail previously²⁷. Briefly, 0.5 % carboxylated microspheres (0.9 µm diameter, Sigma Aldrich) were incubated with 7.4 mg/ml EDC (N-(3-Dimethylaminopropyl)-N′-ethylcarbodiimide hydrochloride, Sigma-Aldrich) and 0.2mg/ml biotin-x-cadaverine (Molecular Probes, A1594) for 30 minutes at room temperature. Biotinilated beads were then incubated with 0.1 M glycine, 0.5 mg/ml neutravidin and 0.02 mg/ml streptavidin-alexa532 (Thermo Fisher Scientific).
Silica beads (1.2 µm, diameter, Bangslabs), dispersed in 1% nitrocellulose-pentyl acetate solution, were smeared on a coverslip and allowed to dry before the flow chamber was assembled\textsuperscript{27,26}. For reactions containing β-catenin, 80 µg/ml of protein was allowed to react with the surface for 5 minutes. αE-catenin (4 µg/ml for single molecule experiments or 40 µg/ml for multiple molecule experiments) was incubated for 5 min to form the α/β catenin complex, or in the absence of β-catenin to bind non-specifically the nitrocellulose surface. The flow chamber was then incubated with 1 mg/ml bovine serum albumin (Sigma-Aldrich) for 5 minutes to minimize non-specific interactions. A solution containing the neutravidin coated-beads and the biotinilated actin was loaded into the chamber. The final reaction was composed by 0.005 % neutravidin functionalized beads, 1 nM rhodamine phalloidin F-Actin in 20 mM MOPS pH 7.2, 50 mM KCl, 1 mM MgCl\textsubscript{2}, 0.1 mM EGTA, 1.2 µM glucose-oxidase, 0.2 µM catalase, 17 mM glucose and 20 mM DTT. Measurements were carried out after sealing the flow chamber with high vacuum grease. A single biotinilated actin filament was anchored through specific binding to the trapped neutravidin-coated microspheres and pre-tensioned to ∼3 pN. At low concentration of αE-catenin, approximately 1 in 4 silica beads showed interaction with the actin filament, providing evidence that the large majority of the beads contained at most a single catenin molecule. Control experiments to probe non-specific interactions were performed without αE-catenin. Few tens of beads were tested in different slides and no non-specific interactions were detected in these conditions.

**Ultrafast force-clamp spectroscopy setup.** Ultra-fast force-clamp spectroscopy and the experimental apparatus are described in detail elsewhere\textsuperscript{10,27}. In brief, dual optical tweezers and fluorescence microscopy were combined in a custom inverted microscope. The two laser traps were independently controlled by two acousto-optic deflectors (AODs). The forward scatter signal from the tweezers was filtered by two interferential filters before
reaching the two quadrant detector photodiodes (QDPs) for particle position measurements. The applied force was calculated from the displacement of the bead (x) and the trap stiffness (k). The trap stiffness was always calibrated prior experiments for the entire range of the trap positions using a power spectrum method\textsuperscript{28}. A high-magnification camera (CCD 2000X) monitored the coordinates of the pedestal bead and with the aid of a 3D piezo translator system nm-stabilization against thermal drifting and low-frequency noise was achieved\textsuperscript{29}. Additionally, a 532 nm laser (Coherent Sapphire) was used for the excitation of rhodamine-labelled F-actin. Both data acquisition and force-clamp feedback were controlled at 200-kHz sample rate through an FPGA board (NI-PCI-7830R) by a custom LabVIEW software.

**Data analysis.** The analysis of the ultra-fast-force-clamp was carried out by a custom MATLAB algorithm which detects catenin-protein interactions by identifying changes in the derivative of the position signal i.e. dumbbell velocity which drops to zero upon binding. Limits were imposed to ensure that false events were kept at a maximum of 1% of the total event number\textsuperscript{10,27}. Step detection was performed as in Gardini et. with small modifications\textsuperscript{12}. Steps were detected from velocity changes due to proteins experiencing conformational changes or rapid detachment and reattachment to the filament. In this case, step length (>5 ms) and absolute amplitude (>5 nm) thresholds were set to limit detection of false steps due to thermal fluctuations, while step amplitude sign switch was allowed to account for both forward and backward steps. In order to distinguish if the steps were within an interaction or belonged to new interactions, maximum time interval (<10 ms) and step size (<75 nm) were set.
Slip-bond model

We fitted the lifetime vs force plot of a single α–β-catenin heterodimer with a Bell-bond model in which the lifetime $\tau$ decreases exponentially with force: $\tau = \tau_0^\pm \exp\left(-\frac{d_{\pm}}{k_BT}\right)$, where $k_B$ is the Boltzmann constant, $T$ the absolute temperature, $F$ is the absolute value of the force, $\tau_0^\pm$ and $d_{\pm}$ are the unloaded lifetime and the distance parameter under positive (+) and negative (−) force respectively. The force sign is arbitrarily chosen. Fitting parameters are reported in the caption of Fig. 2b.

Two-step catch-bond model

We divided the plot of the lifetime vs force for the α-catenin homodimer interaction with actin into a region of low force and high force to separate the lifetime peaks (Supplementary Fig. 1a). Assuming that within each force region α-catenin can transition between two structural states, we developed a two-step kinetic model to fit each lifetime peak. The model is based on two bound states, respectively weakly (W) and strongly (S) bound to actin, and one unbound state (U) (Supplementary figure 1b). α-catenin can switch between state W and S and vice-versa with transition rates $k_{WS}$ and $k_{SW}$, respectively. Detachment can occur from both states W and S with rates $k_{SU}$ and $k_{WU}$, respectively, with $k_{WU} \gg k_{SU}$. All rates depend exponentially on force $k_{ij} = k_{ij}^0 \exp\left(\frac{d_{ij}F}{k_BT}\right)$, with distance parameters $d_{WS}$, $d_{WU}$, $d_{SU} > 0$ and $d_{SW} < 0$ when $F > 0$. With $F < 0$ signs of distance parameters are changed. Therefore, if $k_{0SW} > k_{0WS}$, α-catenin spends most of its time in state W at zero force. As the force increases, $k_{WS}$ increases and $k_{SW}$ decreases, so that, above a threshold force, state S becomes more populated than state W. The kinetic model can be solved following Nolting$^{30}$ to get the measured lifetime $\tau(F)$, which depends on the detachment rates $k_{WU}$ and $k_{SU}$ from both.
bound states W and S and on the transition rates $k_{WS}$ and $k_{SW}$ between them. The probability density function of the bound state $([W]+[S] = 1-[U])$ is:

$$f(t) = -C_2 \lambda_1 \exp(-\lambda_1 t) - C_4 \lambda_2 \exp(-\lambda_2 t)$$

$$C_2 = \frac{k_{SW} + k_{WS} + k_{SU} - \lambda_3}{\lambda_1 - \lambda_2}$$

$$C_4 = -\frac{k_{SW} + k_{WS} + k_{SU} - \lambda_2}{\lambda_1 - \lambda_2}$$

$$\lambda_{1,2} = \frac{1}{2} \left( a \pm \sqrt{a^2 - 4b} \right)$$

$$a = k_{SW} + k_{WS} + k_{WU} + k_{SU}$$

$$b = k_{SW} k_{WU} + k_{WS} k_{SU} + k_{WU} k_{SU}$$

where $C_2$ and $C_4$ are calculated with initial conditions $[U]=[S]=0$, $[W]=1$. From the probability density function, we can calculate the average lifetime of the bound state:

$$\tau = \frac{k_{SW} + k_{WS} + k_{SU}}{k_{SW} k_{WU} + k_{WS} k_{SU} + k_{WU} k_{SU}}$$

that is used to fit experimental data (Fig. 1C and Supplementary Figure 6). Fit parameters are represented graphically in Supplementary Figure 1c and in Supplementary Table 1.
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Author contributions

M.C. designed the research and supervised the experiments, C.A. expressed and purified α-catenin constructs, C.A. set up and performed optical trapping experiments, M.S. analyzed data, M.S., L.G. and M.C. developed the step detection analysis, C.A. performed cosedimentation experiments, C.A. and L.G performed flow-cell experiments, M.C., M.S., C.A. wrote the paper, M.S. prepared the figures, M.C., M.S., C.A., L.G. and F.S.P. revised the paper.

Competing financial interests

The authors declare no competing financial interests.

Data availability

Data supporting the findings of this manuscript are available from the corresponding author upon reasonable request.