

1
2
3
4
5
6
7
8
9
10
11
12
13
14
15
16
17
18
19
20
21
22
23
24
25
26
27
28
29
30
31
32
33
34
35
36
37
38
39

Supplementary Information

Concurrent Atomic Force Spectroscopy

Carolina Pimenta-Lopes[#], Carmen Suay-Corredera*, Diana Velázquez-Carreras,
David Sánchez-Ortiz, Jorge Alegre-Cebollada*

**Centro Nacional de Investigaciones Cardiovasculares Carlos III (CNIC), 28029
Madrid, Spain**

*equal contribution

[#]current address: Department of Physiological Sciences, University of Barcelona, Spain

Correspondence to Jorge Alegre-Cebollada: jalegre@cnic.es

1 **Supplementary Table 1. Number of unfolding events per AFM experiment analyzed in this**
 2 **study, following TFP or OFP strategies.**

3

(C3) ₈ , TFP	(C3) ₈ , OFP with (C3-L) ₄	(C3-L) ₄ , TFP	(C3-L) ₄ , OFP with (C3) ₈	(C3-L) ₄ , OFP with (C3-SUMO1) ₄	(C3-SUMO1) ₄ , TFP	(C3-SUMO1) ₄ , OFP with (C3-L) ₄
117	117	14	29	55	69	54
84	71	35	52	42	7	71
140	182	17	81	82	119	16
141	225	42	136	14	148	92
65	30	31	13	6	149	52
224		38		42	166	57
96				67	47	176
191				47	37	29
52				43		41
75				65		38
149				178		230
				66		84
				81		70
				85		33

4

5

6

1 **Supplementary Table 2. Monte Carlo simulations considering different mechanical**
 2 **unfolding parameters.** The table reports the value of RSD of the distribution of ΔmF_u , at
 3 different values of r_0 and Δx . The remaining simulation parameters were the same as in Figure
 4 1B in the main text.

	<i>Calibration</i>	$r_0 = 0.01 \text{ s}^{-1}$	$r_0 = 0.015 \text{ s}^{-1}$	$r_0 = 0.01 \text{ s}^{-1}$	$r_0 = 0.015 \text{ s}^{-1}$
	<i>Uncertainty</i>	$\Delta x = 0.2 \text{ nm}$	$\Delta x = 0.2 \text{ nm}$	$\Delta x = 0.25 \text{ nm}$	$\Delta x = 0.25 \text{ nm}$
	(%)				
<i>Concurrent</i>	3.6	0.033	0.034	0.025	0.028
	18	0.035	0.037	0.035	0.036
<i>Traditional</i>	3.6	0.053	0.054	0.051	0.045
	18	0.238	0.232	0.227	0.231

7
 8
 9
 10

1 **Supplementary Text 1. Sequences of the proteins used in this report.**
2 We highlight the sequence of the domains in different colors: C3, red; protein L, bold type;
3 SUMO1, green). Linkers and extra amino acids are shown in regular black type.

4
5 (C3)₈
6 MRGSHHHHHHGSPVLITRPLEDQLVMVGQRVEFECEVSEEGAQVKWLKDGVELTREE
7 TFKYRFKKGQRHHLIINEAMLEDAGHYALCTSGGQALAEIVQEKRSPVLITRPLEDQ
8 LVMVGQRVEFECEVSEEGAQVKWLKDGVELTREETFKYRFKKGQRHHLIINEAMLE
9 DAGHYALCTSGGQALAEIVQEKRSPVLITRPLEDQLVMVGQRVEFECEVSEEGAQVK
10 WLKDGVELTREETFKYRFKKGQRHHLIINEAMLEDAGHYALCTSGGQALAEIVQEKR
11 RSPVLITRPLEDQLVMVGQRVEFECEVSEEGAQVKWLKDGVELTREETFKYRFKKGQ
12 RHHLIINEAMLEDAGHYALCTSGGQALAEIVQEKRSPVLITRPLEDQLVMVGQRVEFE
13 CEVSEEGAQVKWLKDGVELTREETFKYRFKKGQRHHLIINEAMLEDAGHYALCTSG
14 GQALAEIVQEKRSPVLITRPLEDQLVMVGQRVEFECEVSEEGAQVKWLKDGVELTRE
15 ETFKYRFKKGQRHHLIINEAMLEDAGHYALCTSGGQALAEIVQEKRSPVLITRPLED
16 QLVMVGQRVEFECEVSEEGAQVKWLKDGVELTREETFKYRFKKGQRHHLIINEAML
17 EDAGHYALCTSGGQALAEIVQEKRSPVLITRPLEDQLVMVGQRVEFECEVSEEGAQV
18 KWLKDGVELTREETFKYRFKKGQRHHLIINEAMLEDAGHYALCTSGGQALAEIVQE
19 KRSCC

20
21 (C3-L)₄
22 MRGSHHHHHHGSPVLITRPLEDQLVMVGQRVEFECEVSEEGAQVKWLKDGVELTREE
23 TFKYRFKKGQRHHLIINEAMLEDAGHYALCTSGGQALAEIVQEKRSMEEVTIKANL
24 IFANGSTQTAEFKGTFEKATSEAYAYADTLKKNGEWTVADVADKGYTLNIKFRSP
25 VLITRPLEDQLVMVGQRVEFECEVSEEGAQVKWLKDGVELTREETFKYRFKKGQRH
26 HLIINEAMLEDAGHYALCTSGGQALAEIVQEKRSMEEVTIKANLIFANGSTQTAEFK
27 GTFEKATSEAYAYADTLKKNGEWTVADVADKGYTLNIKFRSPVLITRPLEDQLVMV
28 GQRVEFECEVSEEGAQVKWLKDGVELTREETFKYRFKKGQRHHLIINEAMLEDAGH
29 YALCTSGGQALAEIVQEKRSMEEVTIKANLIFANGSTQTAEFKGTFEKATSEAYAY
30 ADTLKKNGEWTVADVADKGYTLNIKFRSPVLITRPLEDQLVMVGQRVEFECEVSEEG
31 AQVKWLKDGVELTREETFKYRFKKGQRHHLIINEAMLEDAGHYALCTSGGQALAEI
32 VQEKRSMEEVTIKANLIFANGSTQTAEFKGTFEKATSEAYAYADTLKKNGEWTV
33 DVADKGYTLNIKFRSCC

34 (C3-SUMO1)₄
35 MRGSHHHHHHGSPVLITRPLEDQLVMVGQRVEFECEVSEEGAQVKWLKDGVELTREE
36 TFKYRFKKGQRHHLIINEAMLEDAGHYALCTSGGQALAEIVQEKRSMSDQEAKPST
37 EDLGDKKEGEYIKLVIGQDSSEIHFKVKMTTHLKKLKESYCQRQGVPMNSLRFLFEG
38 QRIADNHTPKELGMEEDVIEVYQEQTGGRSPVLITRPLEDQLVMVGQRVEFECEVSEE
39 GAQVKWLKDGVELTREETFKYRFKKGQRHHLIINEAMLEDAGHYALCTSGGQALAE
40 LIVQEKRSMSDQEAKPSTEDLGDKKEGEYIKLVIGQDSSEIHFKVKMTTHLKKLKESY
41 CQRQGVPMNSLRFLFEGQRIADNHTPKELGMEEDVIEVYQEQTGGRSPVLITRPLEDQ
42 LVMVGQRVEFECEVSEEGAQVKWLKDGVELTREETFKYRFKKGQRHHLIINEAMLE
43 DAGHYALCTSGGQALAEIVQEKRSMSDQEAKPSTEDLGDKKEGEYIKLVIGQDSSEI
44 HFKVKMTTHLKKLKESYCQRQGVPMNSLRFLFEGQRIADNHTPKELGMEEDVIEVYQ
45 EQTGGRSPVLITRPLEDQLVMVGQRVEFECEVSEEGAQVKWLKDGVELTREETFKYRF
46 KKGQRHHLIINEAMLEDAGHYALCTSGGQALAEIVQEKRSMSDQEAKPSTEDLGDK
47 KEGEYIKLVIGQDSSEIHFKVKMTTHLKKLKESYCQRQGVPMNSLRFLFEGQRIADNH
48 TPKELGMEEDVIEVYQEQTGGRSCC
49

1 **Supplementary Text 2. Estimation of uncertainty in cantilever calibration by the thermal**
2 **fluctuations method.**

3 The thermal fluctuations method estimates the spring constant of the cantilever (k_{sc}) from the
4 deflection signal of a laser beam focused on the cantilever, measured by a split photodetector
5 (Supplementary Figure 1). This method considers the cantilever as a harmonic oscillator and
6 applies the equipartition theorem to obtain Equation S1¹:

7
8
$$k_{sc} = \frac{k_b \cdot T}{S^2 \cdot \langle z^2 \rangle} \quad \text{Equation S1}$$

9
10 In Equation S1, k_b is the Boltzmann constant, T is the absolute temperature, S (deflection
11 sensitivity, in nm/V) is the slope of the change in voltage detected by the photodetector for
12 displacements of the surface while in contact with the cantilever, and $\langle z^2 \rangle$ is the mean squared
13 displacement of the free cantilever in units of V^2 . Hence, cantilever calibration by the thermal
14 fluctuations method requires estimation of S and $\langle z^2 \rangle$. Forces are then calculated from the
15 deflection signal (A-B, in units of V) according to:

16
17
$$F = (A - B) \cdot S \cdot k_{sc} \quad \text{Equation S2}$$

18
19 Combining Equations S1 and S2, we obtain

20
21
$$F = (A - B) \cdot \frac{k_b \cdot T}{S \cdot \langle z^2 \rangle} \quad \text{Equation S3}$$

22
23 We measured experimental distributions of S and $\langle z^2 \rangle$ for a single cantilever and found that the
24 RSD of S was 3.5% while the RSD of $\langle z^2 \rangle$ was 0.8% (Supplementary Figure 4). Hence, we
25 conclude that inaccuracies in the determination of deflection sensitivity, which are 4-5 times
26 larger than variations in $\langle z^2 \rangle$, are the main driver of interexperimental variation in spring
27 constants of AFM cantilevers estimated using the thermal fluctuations method, as proposed
28 before³.

29
30 Considering error propagation, we propose that a reasonable value for the minimum calibration
31 uncertainty in force is given by $\sqrt{0.035^2 + 0.008^2} = 3.6\%$. It is interesting to note that the
32 uncertainty in the determination of k_{sc} is higher since it depends on the square of S :
33 $\sqrt{(2 \cdot 0.035)^2 + 0.008^2} = 7.1\%$. This effect can be observed in the distributions in
34 Supplementary Figure 4.

35
36 To obtain errors (ε) in force for simulated AFM experiments, we drew random values from a
37 normal distribution centered in 100 and whose RSD corresponds to the % uncertainty being
38 considered. Then:

39
40
$$\varepsilon = \frac{\text{random value}}{100} \quad \text{Equation S4}$$

41
42 This error is considered when calculating the probability of unfolding (Equation 3 in the main
43 text).

1 **Supplementary Text 3. A model for the propagation of force calibration errors to ΔmF_u .**

2 We consider that each value of measured unfolding force ($F_u^{measured}$) is affected by an error δ
3 coming from uncertain force calibration:

4
5
$$F_u^{measured} = F_u^{real} + \delta \quad \text{Equation S5}$$

6

7 Hence, the value of mF_u that results from averaging unfolding data from m experiments, each
8 one with a certain number of events n , can be estimated as:

9
10
$$mF_u^{measured} = \frac{\sum_{i=1}^{n_1}(F_{u,i}^{real} + \delta_{1,i}) + \dots + \sum_{i=1}^{n_m}(F_{u,i}^{real} + \delta_{m,i})}{n_1 + \dots + n_m} \quad \text{Equation S6}$$

11

12 We can consider that for every experiment j , $\sum_{i=1}^{n_j} F_{u,i}^{real} = n_j \cdot mF_{u,j}^{real}$, where n_j is the number
13 of events in experiment j and $mF_{u,j}^{real}$ is the mean unfolding force that would have been
14 measured in experiment j if there was no error in calibration. Similarly, considering that
15 $\sum_{i=1}^{n_j} \delta_{j,i} = n_j \cdot \bar{\delta}_j$, where $\bar{\delta}_j$ is the average error in force in experiment j , and $n_{events} = n_1 +$
16 $\dots + n_m$, we obtain:

17
18
$$mF_u^{measured} = mF_u^{real} + \frac{\sum_{j=1}^m n_j \cdot \bar{\delta}_j}{n_{events}} \quad \text{Equation S7}$$

19 Equation 1 in the main text is derived from Equation S7 considering comparison between two
20 proteins under the assumption that $\bar{\delta}_j$ is the same for both proteins when these are measured
21 concurrently in the same experiment.

22

1 **Supplementary Text 4. Interpretation of relative improvement in accuracy by concurrent**
2 **atomic force spectroscopy.**

3 Figure 1E shows that the degree of improvement in accuracy in ΔmF_u by concurrent relative to
4 traditional AFM increases with the number of events per experiment and remains insensitive to
5 the total number of experiments. Here, we provide a qualitative explanation for this observation.
6

7 The RSD of the distribution of ΔmF_u derives from two independent factors: the calibration
8 uncertainty (Figure 1F) and the limited number of unfolding events defining the distribution of
9 unfolding forces (Supplementary Figure 5B). In the case of balanced datasets obtained in
10 concurrent atomic force spectroscopy, the contribution of calibration uncertainty is zero as
11 predicted from Equation 1.
12

13 What are the consequences of increasing the number of events per experiment in the accuracy in
14 ΔmF_u achieved by atomic force spectroscopy measurements?
15

- 16 - In the traditional strategy, increasing the number of events per experiment leads to
17 better definition of the distribution of unfolding forces, but has no impact in the error
18 associated to calibration uncertainty. In fact, at high number of events, the RSD of the
19 distribution of mF_u is fairly insensitive to further increases in the number of events
20 since it is dominated by errors associated to calibration uncertainty (Supplementary
21 Figure 5B).
22
- 23 - In concurrent modes, the effect of better definition of distributions of unfolding forces is
24 the same as in traditional AFM. However, since in concurrent AFM the contribution of
25 calibration uncertainty to the RSD of the distribution of ΔmF_u is null (Equation 1), the
26 overall relative improvement in accuracy is higher than in traditional AFM.
27

28 Increasing the number of experiments minimizes the impact of calibration uncertainty and also
29 leads to an increase in the total number of events, both of which contribute to make the RSD of
30 the distribution of ΔmF_u smaller in traditional experiments (Figure 1C,D). In concurrent AFM,
31 since there is no error associated to calibration uncertainty, RSD decreases only as a
32 consequence of higher number of events. However, as explained above, the impact of higher
33 number of events on the RSD of ΔmF_u distributions is more prominent for concurrent
34 experiments. We suggest that this differential impact results in a fairly constant relative
35 performance of concurrent vs. traditional strategies regardless of the number of experiments
36 considered (Figure 1E).
37
38

1 **Supplementary Text 5. Validity of the linear approximation in the procedure used to**
2 **obtain distributions of unfolding forces.**

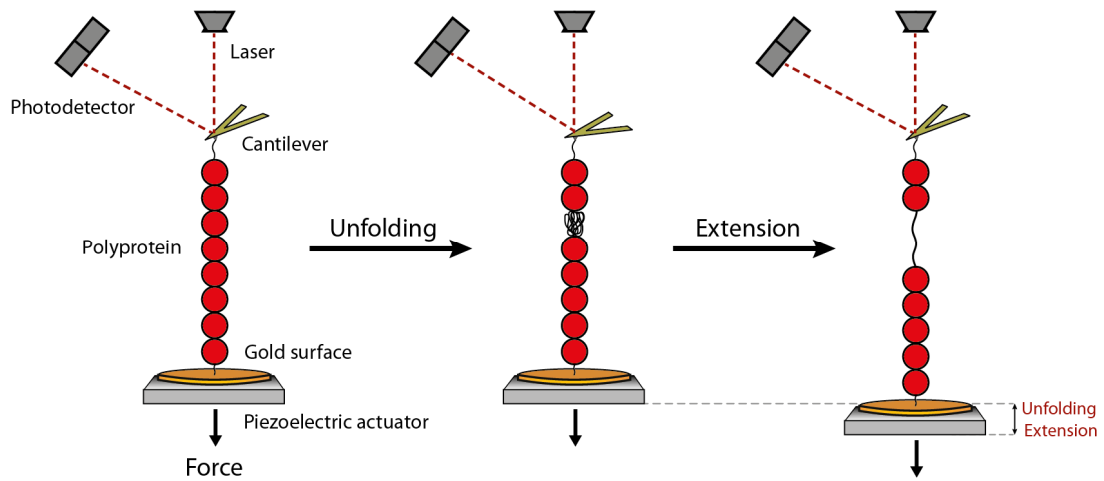
3 In our Monte Carlo simulations, we approximate the instantaneous probability of unfolding to
4 the linear regime (Equation 3 in the main text), which is valid at low values of $n \cdot r \cdot \Delta t^4$.

5 Considering that the mF_u obtained in our simulations is around 100 pN, we can estimate the
6 maximum number of events at the midpoint of the unfolding distribution that still satisfy $n \cdot r \cdot$

7 $\Delta t < 0.05$, which according to Equation 3, is 385 (considering $r_0 = 0.01 \text{ s}^{-1}$ and $\Delta x = 0.2 \text{ nm}$).

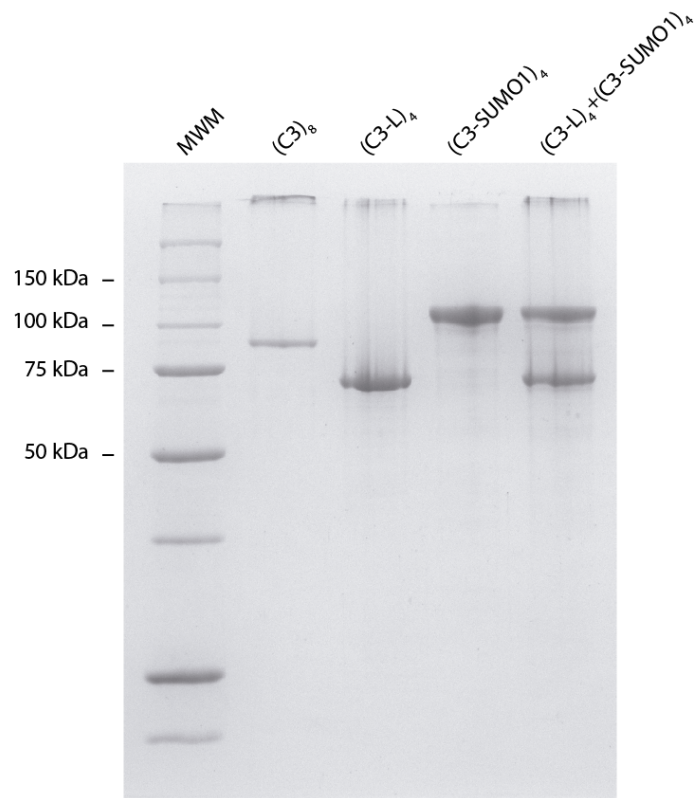
8 Hence, we always kept the number of simulated unfolding events below 2×385 .

9



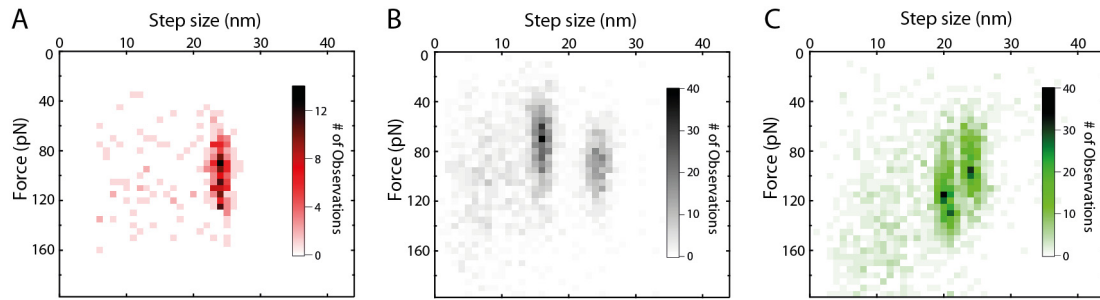
Supplementary Figure 1. Schematic representation of a single-molecule force-clamp AFM experiment. A single polyprotein is tethered between an AFM cantilever and a gold-coated surface. *Left:* The piezoelectric actuator moves away from the cantilever, which results in a pulling force applied to the polyprotein. The magnitude of the pulling force is calculated from the difference in voltage between the two regions of a split photodetector that is reached by a laser beam reflecting on the cantilever. *Middle:* When a domain unfolds, the force relaxes momentarily, changing the laser deflection. *Right:* To recover the programmed force set point in force-clamp experiments, the piezoelectric actuator is displaced, stretching the polyprotein and producing an unfolding step in experimental recordings (Figure 1A).

1
2
3



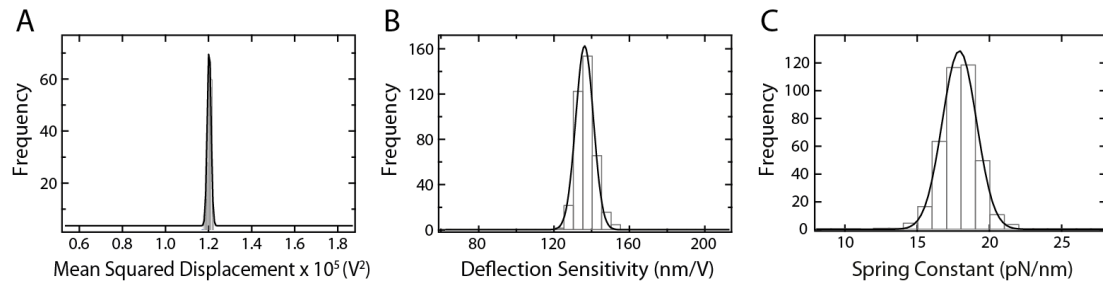
Supplementary Figure 2. 12% SDS-PAGE analysis of the purified proteins used in this report. We note that the C3-containing polyproteins have a tendency to show high-molecular weight aggregates that do not enter the resolving gel. We interpret this effect as an artifact of the electrophoresis, since equivalent aggregates do not appear in the void volume of the size-exclusion chromatography. MWM: Precision Plus Protein Unstained Standards (Bio-Rad). The last lane shows results from simultaneous purification of (C3-L)₄ and (C3-SUMO1)₄.

1
2



Supplementary Figure 3. Definition of fingerprinting unfolding lengths of C3, protein L and SUMO1. Polyproteins were pulled at a rate of 40 pN/s. All traces containing at least two events of the same step size were included in these analyses. The bidimensional histograms show the frequency of the size and force of all steps in the selected traces. In all cases, the main populations of unfolding lengths are clearly identified from a background of steps that correspond to non-specific interactions. For subsequent analysis, only traces showing the fingerprinting unfolding lengths were considered. **(A)** Results obtained for the mechanical unfolding of $(C3)_8$ ($n = 293$ steps), showing a single population of unfolding events at 24 ± 1 nm and around 90 pN. **(B)** Results obtained for the mechanical unfolding of $(C3-L)_4$ ($n = 2555$ steps). The two well-defined populations correspond to unfolding of L domains (step size 16 ± 1 nm at around 70 pN) and to unfolding of C3 (step size 24 ± 1 nm at around 90 pN). **(C)** Results obtained for the mechanical unfolding of $(C3-SUMO1)_4$ ($n = 1998$ steps). The two well-defined populations correspond to unfolding of SUMO1 (step size 20 ± 1 nm at around 115 pN) and C3 (step size 24 ± 1 nm at around 90 pN).

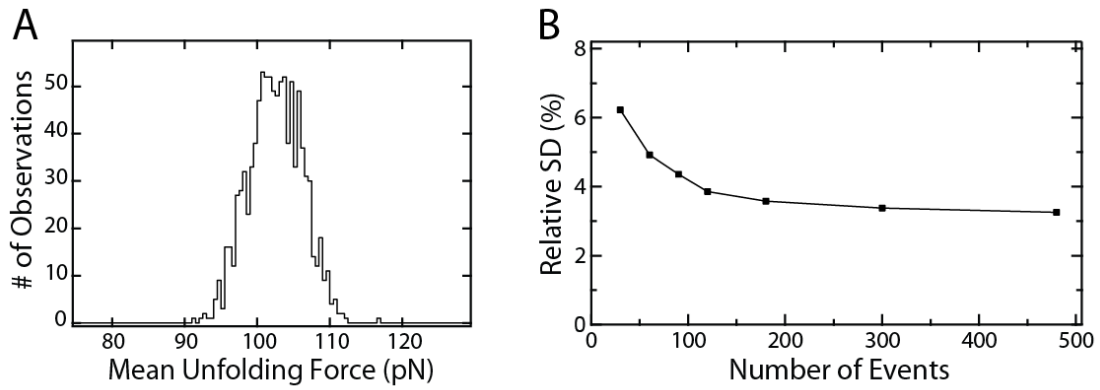
1
2



Supplementary Figure 4. Experimental estimation of calibration uncertainty from multiple rounds of calibration of the same cantilever using the thermal fluctuations method. (A) Distribution of 250 measurements of mean squared displacement ($\langle z^2 \rangle$) for a single cantilever ($1.2 \cdot 10^{-5} \pm 9.2 \cdot 10^{-8} \text{ V}^2$). (B) Distribution of 388 measurements of the deflection sensitivity (S) for a single cantilever ($136.7 \pm 4.9 \text{ nm/V}$). (C) Considering $\langle z^2 \rangle = 1.2 \cdot 10^{-5} \text{ V}^2$, we show the distribution of spring constants that arise from the values of deflection sensitivity in panel B ($17.9 \pm 1.3 \text{ pN/nm}$). Solid lines are Gaussian fits to the data. Mean \pm SD of the distributions are indicated. In the three panels, the relative range of the x axes with respect to the mean value is the same.

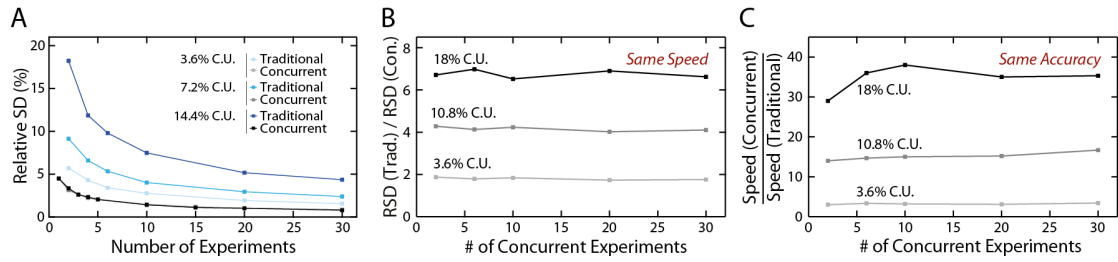
1

2



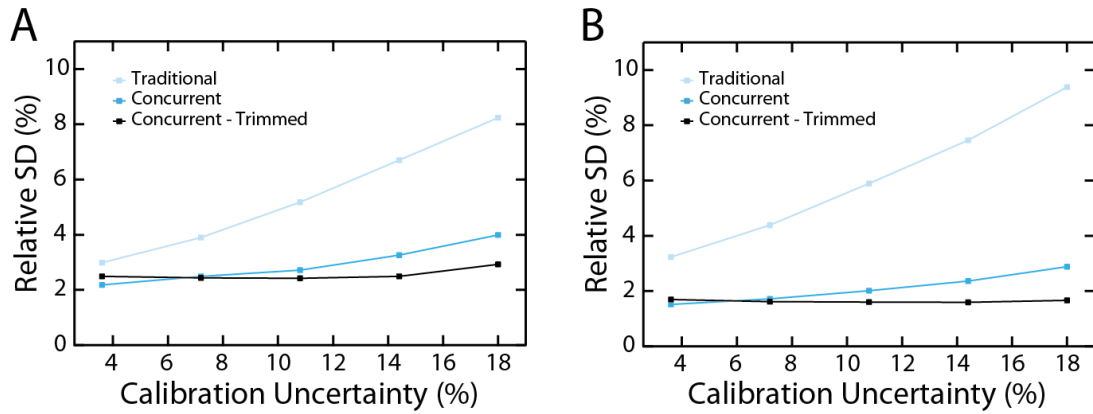
Supplementary Figure 5. Distribution of mean unfolding force for an individual protein. **(A)** Distribution of mF_u corresponding to one traditional experiment (200 events), as estimated from Monte Carlo simulations. **(B)** Dependence of the RSD of the distribution of mF_u with the number of unfolding events, obtained from simulations of a single protein measured following traditional atomic force spectroscopy. We considered a 3.6% calibration uncertainty. At low number of events, the distribution of mF_u is not well defined and the RSD is high. At higher number of events, the distribution of mF_u is well sampled, and the major contributor to RSD is the calibration uncertainty.

1
2
3



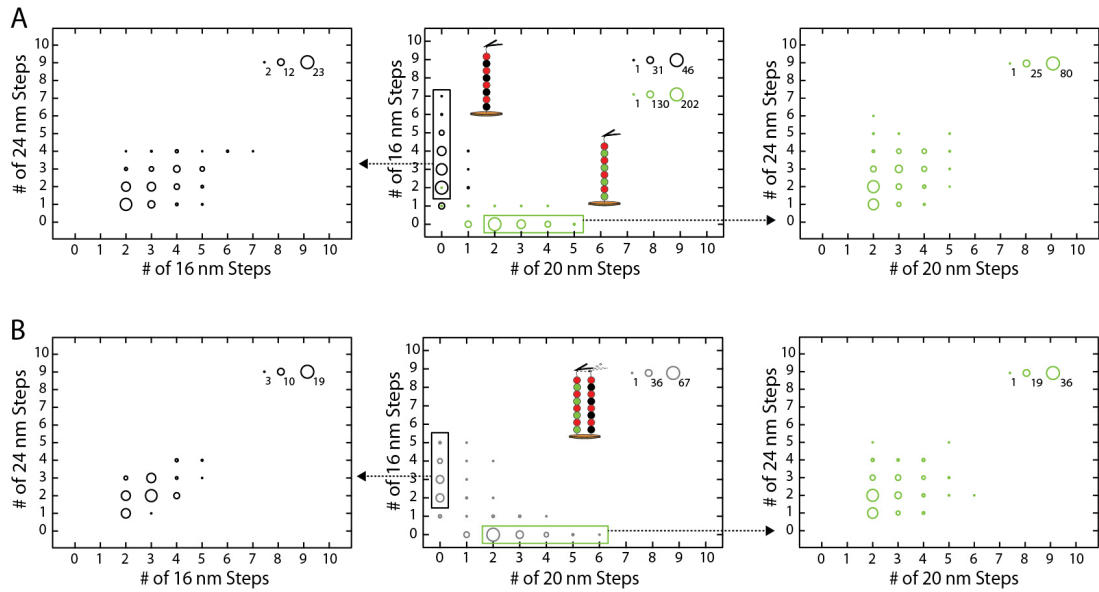
Supplementary Figure 6. Increased speed of data acquisition and accuracy in ΔmF_u by concurrent atomic force spectroscopy. (A) RSD of the distribution of ΔmF_u estimated from Monte Carlo simulations at different calibration uncertainties (C.U.) and number of experiments. Note that in the case of concurrent measurements, the RSD values overlap for the three calibration uncertainties. (B) Relative increase in accuracy in ΔmF_u achieved by concurrent with respect to traditional strategies (at equivalent speeds of data acquisition), at different calibration uncertainties and number of experiments. (C) Relative increase in throughput achieved by concurrent with respect to traditional strategies (at equivalent accuracy), at different calibration uncertainties and number of experiments. Speed(Concurrent)/Speed(Traditional) was calculated as the ratio between the number of traditional and concurrent experiments that are needed to achieve the same RSD of the distribution of ΔmF_u . All simulations considered 100 unfolding events per protein and experiment.

1
2



Supplementary Figure 7. Application of Monte Carlo simulations to estimate RSD of the distribution of ΔmF_u associated to real datasets. (A) Monte-Carlo-estimated RSD of the distributions of ΔmF_u obtained from the atomic force spectroscopy experiments that compare the mF_u of C3 in the context of $(C3)_8$ and $(C3-L)_4$ (see Figure 2 in the main text). Simulations are fed with the actual number of unfolding events measured experimentally (Supplementary Table 1). Simulations were also run with trimmed concurrent datasets, by removing data to equal the number of events per protein within a concurrent experiment. **(B)** Monte-Carlo-estimated RSD of the distributions of ΔmF_u obtained from the atomic force spectroscopy experiments that compare the mF_u of C3 in the context of $(C3-L)_4$ and $(C3-SUMO1)_4$ (see Figure 3 in the main text). Simulations are fed with the actual number of unfolding events measured experimentally (Supplementary Table 1). Simulations were also run with trimmed concurrent datasets, by removing data to equal the number of events per protein within a concurrent experiment.

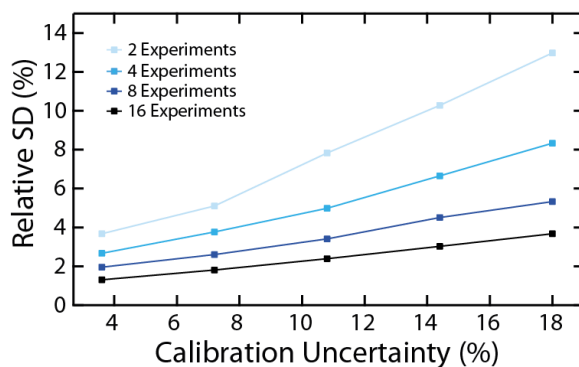
1
2



Supplementary Figure 8. Protein gating in dual-marker OFP. (A) Gating on data obtained in TFP experiments. **(B)** Gating on data obtained in OFP experiments. All the plots show the frequency of the traces that have different combinations of unfolding events, as indicated by the size of the dots. (C3-L)₄ and (C3-SUMO1)₄ unfolding traces were first classified according to the number of marker 16 and 20 nm steps (middle panels, same data as in Figure 3A,B). Sorted traces were further classified according to the number of 24 nm steps, which correspond to C3 unfolding events (left and right panels). Results show that highly similar distributions are obtained in traditional and orthogonal fingerprinting experiments, providing further support to the gating protocol to sort traces coming from mixtures of proteins. Number of traces in (A): 135 (left), 419 (right). Number of traces in (B): 102 (left), 174 (right).

1
2

1

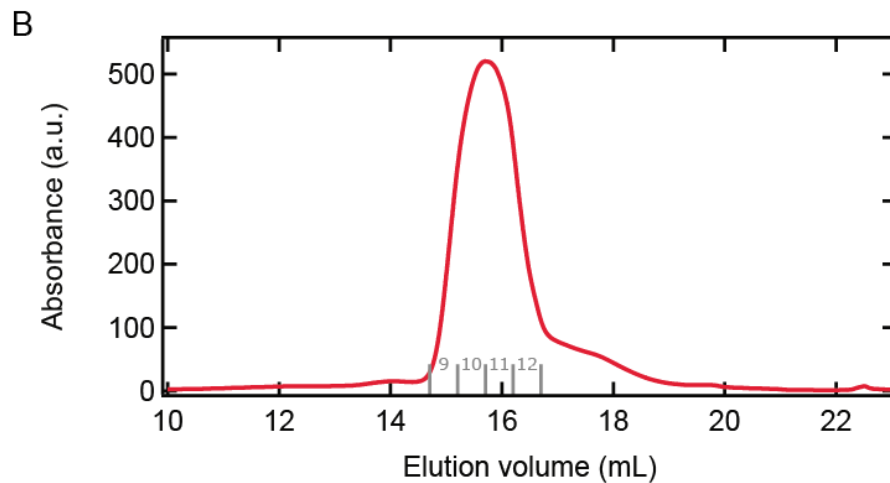
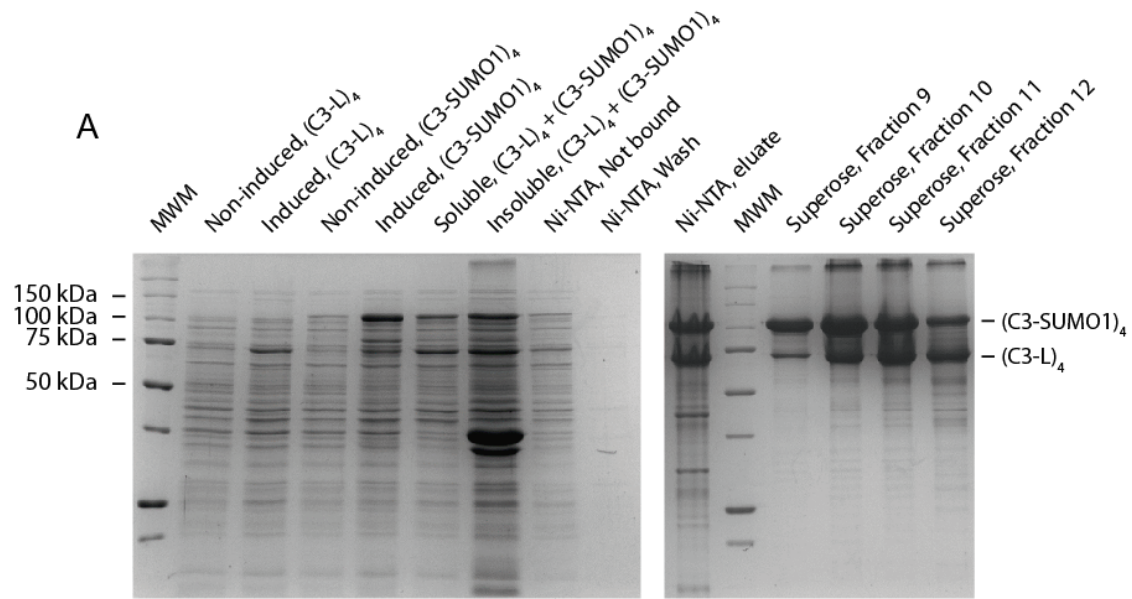


Supplementary Figure 9. Unbalanced datasets affect the accuracy of concurrent ΔmF_u determinations differentially according to the total number of experiments. Monte-Carlo-estimated RSD of the distributions of ΔmF_u taking into account increasing number of concurrent experiments. We have simulated unbalanced datasets with alternating 50/150 events for each protein in each experiment.

2

3

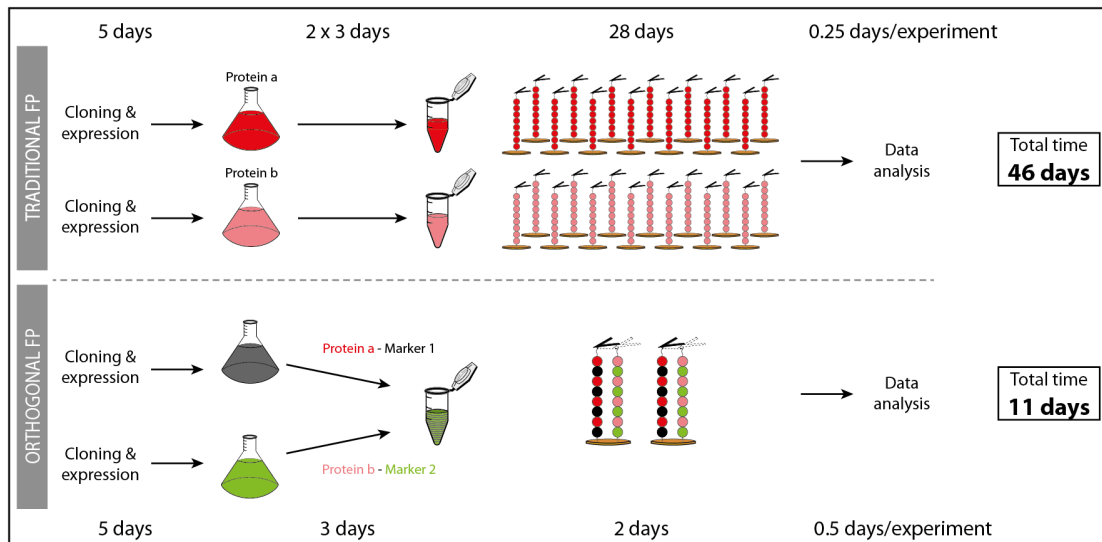
4



Supplementary Figure 10. Simultaneous purification of (C3-SUMO1)₄ and (C3-L)₄. (A) 12% SDS-PAGE to monitor the steps of expression and purification. *E. coli* cells containing the expression plasmid for (C3-SUMO1)₄ or (C3-L)₄ proteins are induced separately. The expression of the proteins is detected in the induced samples. Both cell cultures are lysed together, and the resulting soluble fraction is loaded in a Ni-NTA column. The fractions of highest protein concentration after elution with imidazole are subject to size-exclusion chromatography in an FPLC system using a Superose 6 Increase 10/300 GL column. Fractions 9-12 contain different proportions of a mixture of (C3-SUMO1)₄ and (C3-L)₄. OFP experiments can be set with these fractions directly. The preferred fraction gives similar number of unfolding events for both proteins. MWM: Precision Plus Protein Unstained Standards (Bio-Rad). (B) Chromatogram corresponding to the size exclusion chromatography shows that (C3-SUMO1)₄ and (C3-L)₄ are not resolved and co-elute in fractions 9-12.

1

2



Supplementary Figure 11. Total decrease in experimentation time by OFP-based concurrent atomic force spectroscopy. We take into account the full workflow of single-molecule AFM required to characterize differences in unfolding forces between protein a and protein b, including cloning, protein expression, protein purification, AFM measurements and data analysis. We have considered that OFP allows simultaneous purification of proteins and that data analysis of OPF datasets can be more time consuming due to the protein gating steps. Improvement in throughput of data acquisition is estimated from Monte Carlo simulations at 10.8% calibration uncertainty with 100 events per experiment and protein (see Figure 5 in the main text).

1 **Supplementary References**

- 2 1.Hutter, J. L. & Bechhoefer, J. Calibration of atomic-force microscope tips. *Review of Scientific*
3 *Instruments* **64**, 1868-1873. (1993).
- 4 2.Popa, I., Kosuri, P., Alegre-Cebollada, J., Garcia-Manyes, S. & Fernandez, J. M. Force
5 dependency of biochemical reactions measured by single-molecule force-clamp
6 spectroscopy. *Nature protocols* **8**, 1261-1276. (2013).
- 7 3.Ohler, B. Practical Advice on the Determination of Cantilever Spring Constants. Veeco
8 Instruments Incorporated, Technical Report., (2007).
- 9 4.Shi, J., Gafni, A. & Steel, D. Simulated data sets for single molecule kinetics: some limitations
10 and complications of data analysis. *European biophysics journal : EBJ* **35**, 633-645.
11 (2006).

12