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Designed anchoring geometries determine lifetimes of biotin–streptavidin bonds under constant load and enable ultra-stable coupling†

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The small molecule biotin and the homotetrameric protein streptavidin (SA) form a stable and robust complex that plays a pivotal role in many biotechnological and medical applications. In particular, the SA–biotin linkage is frequently used in single-molecule force spectroscopy (SMFS) experiments. Recent data suggest that SA–biotin bonds show strong directional dependence and a broad range of multi-exponential lifetimes under load. Here, we investigate engineered SA variants with different valencies and a unique tethering point under constant forces using a magnetic tweezers assay. We observed orders-of-magnitude differences in the lifetimes under force, which we attribute to the distinct force-loading geometries in the different SA variants. Lifetimes showed exponential dependencies on force, with extrapolated lifetimes at zero force that are similar for the different SA variants and agree with parameters determined from constant-speed dynamic SMFS experiments. We identified an especially long-lived tethering geometry that will facilitate ultra-stable SMFS experiments.

Introduction

The non-covalent, high-affinity binding of the small molecule biotin to streptavidin (SA) is ubiquitously used in a variety of biological, chemical, biophysical and pharmaceutical applications.^{2–4} Biotin can readily be covalently attached to nucleic acids,^{6–8} proteins,^{9,10} or linker molecules.¹¹ SA is stable over a wide range of conditions and easy to handle.² Owing to the specificity of the binding, as well as the robustness of the complex, the interaction has in particular become a popular tool in the context of single-molecule force spectroscopy (SMFS) assays.^{12–15} It serves as a molecular handle to anchor molecules of interest and apply forces and torques to them.^{7,16–23} The long lifetime of the SA–biotin complex under

external forces has enabled constant-force SMFS experiments lasting for hours and even up to weeks in magnetic tweezers (MT).^{24,25} Despite its widespread use, SA's tetra-valency poses a problem, in particular in SMFS applications, since it is *a priori* ambiguous which of the four subunits biotin binds to. This ambiguity results in four different force-loading geometries for a given attachment of the SA tetramer (Fig. 1).²⁶ Furthermore, if SA is non-specifically attached – as is the case in many commercially available SA-coated magnetic beads – a variety of attachment points combined with tetra-valency results in an even larger range of possible force-loading geometries.^{25,27}

Atomic force microscopy (AFM)-based constant-speed SMFS experiments have recently shown that the force needed to unbind biotin from the SA binding pocket is strongly dependent on the force-loading direction:^{28,29} tethering SA by a

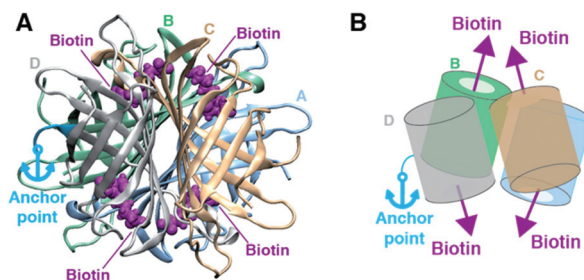


Fig. 1 SA's tetra-valency results in different force-loading geometries. (A) Crystal structure of the SA tetramer (PDB-ID: 6M9B,¹ rendered using VMD⁵) with the four subunits shown in different colors. Four bound biotin molecules are shown in purple. The light blue anchor marks the attachment point (C-terminus of subunit D). (B) Schematic representation of the tetramer structure. The colored barrels represent the four subunits. Arrows indicate the initial force-loading directions in the SMFS experiments: the light blue anchor marks the C-terminus of subunit D used for site-specific immobilization. Purple arrows indicate the four possible directions of pulling biotin out of the different binding pockets. Under constant load, the complex will rotate and rearrange in such a way that the sum of forces acting on it equals zero. Depending on which subunit biotin is bound to, the orientation of the complex will be different resulting in different force propagation pathways.

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single defined residue and pulling biotin out of one of the binding pockets results in different force-loading geometries, depending on which SA subunit the biotin has bound to. For some of the pulling directions, the SA subunit is deformed such that the energy barrier of the binding is decreased, causing lower biotin unbinding forces.²⁹ However, the influence of the tethering geometry of SA on the lifetime of the SA–biotin interaction under constant forces is currently unknown.

Here, we employ engineered variants of SA with different defined valencies and a unique tethering point to restrict and control the number of possible force-loading geometries for SMFS measurements. We use AFM imaging to verify the valencies by showing that only the competent subunits can bind biotin. Furthermore, we employ isothermal titration calorimetry (ITC) to directly measure the binding enthalpies of the different SA variants. With an MT assay we assess the stability of the SA–biotin interaction under different levels of constant load and demonstrate large differences in the lifetime depending on the attachment geometry. The different stabilities give rise to multi-exponential lifetime distributions for multivalent constructs. We observe an exponential decrease of lifetimes with increasing force, with parameters for the force dependencies fully consistent with findings from constant-speed SMFS experiments. By using one well-defined attachment and a monovalent SA construct, a single extremely stable population is achieved. We expect our results to be highly relevant for force spectroscopy, and, in general, to improve assays where the SA–biotin bond is under load, *e.g.* through fluid flow or rinsing steps.

Results and discussion

To systematically investigate the stability of the SA–biotin complex under constant mechanical load, we prepared tetra-, tri-, and monovalent variants of SA. These comprise four, three, and one functional subunit(s), while the remaining subunits are incapable of biotin binding (4SA, 3SA, and 1SA; Fig. 2A) due to three mutations located around the binding pocket (N23A, S27D, S45A).³⁰ In addition, a variant consisting of four non-functional subunits (0SA) was prepared. All variants possess a single cysteine residue at the C-terminus of their subunit D, allowing for site-specific immobilization.^{19,26,29} For 3SA and 0SA, subunit D is non-functional, whereas for 1SA and 4SA, it is functional (Fig. 2A; for details on protein engineering see ESI Materials and Methods†).

AFM imaging reveals binding stoichiometry

To verify the valency of the different variants, we incubated them with biotinylated 250 bp double-stranded DNA constructs and directly visualized the resulting SA–biotinylated DNA complexes by AFM imaging (Fig. 2B and ESI Fig. S1–S4†). An excess of biotinylated DNA over SA (approximately twenty-fold for 4SA and 3SA, and four-fold for 1SA and 0SA) was used to ensure that SA molecules with DNA strands bound to all functional subunits could be observed. Indeed, a maximum of

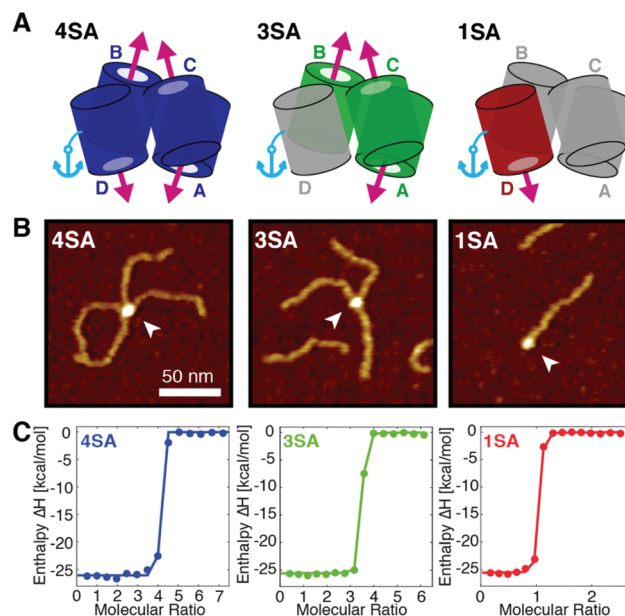


Fig. 2 Probing SA variants with different valencies by AFM and ITC. (A) Schematic structure of SA constructs with different valencies. 4SA (left), 3SA (middle), and 1SA (right) have four, three and one functional subunit(s) (colored), respectively. The remaining subunit(s) (gray) are incapable of binding biotin. All constructs have a single C-terminal cysteine at their subunit D –nonfunctional for 3SA, functional for 1SA and 4SA– for site-specific immobilization (light blue line). The light blue anchors mark the anchoring site of SA for the SMFS experiments, while the purple arrows indicate the possible directions in which biotin can be pulled out of the binding pockets. (B) AFM images of 4SA (left), 3SA (middle), and 1SA (right) with the maximal number (four, three, and one, respectively) of biotinylated DNA strands bound. Arrows mark the SA molecules. Height range of color scale is 2 nm. (C) Isothermal titration calorimetry data of free biotin binding to SA of different valencies. Colored dots are the measured heat release per mole upon adding biotin to SA plotted against the molecular ratio (biotin per SA) in the measurement cell. Lines are fits to the data (taking the discreteness of the measurement into account). For details of the fits see ESI.†

four, three, and one bound biotinylated DNA strand(s) was observed for 4SA, 3SA and 1SA, respectively, confirming the expected valencies (ESI Fig. S5†). In the case of 0SA, no SA–biotinylated DNA complexes were observed.

Thermodynamic parameters determined by isothermal titration calorimetry

Next, we performed isothermal titration calorimetry (ITC) measurements to determine the thermodynamic parameters of biotin-binding to the different SA constructs in the absence of force (Fig. 2C). In principle, ITC allows determination of the stoichiometry, the affinity, and the binding enthalpy. To ensure good comparability across measurements, we used the same biotin stock solution with an estimated 5% uncertainty in absolute concentration for all measurements. The uncertainty in the concentrations of the SA stocks was estimated to be 10% (see ESI Materials and Methods†). Fits to the ITC data give values for the binding stoichiometries of 1.0 ± 0.2 for 1SA, 3.3 ± 0.5 for 3SA, and 3.9 ± 0.6 for 4SA (Fig. 2C), in excellent

agreement with the results from AFM imaging (Fig. 2B and ESI Fig. S5†). The largest contributions to the measurement errors result from the uncertainties in concentration. The uncertainties of the values increase with the number of available binding sites, because a given uncertainty in protein concentration has a larger impact on the uncertainty with increasing stoichiometry. Due to limitations of our instrument and the very high affinity of biotin to SA, the binding constant could not be obtained directly and we can only determine that the affinity is higher than 1 nM. We obtained binding enthalpies per binding site of $-(25.0 \pm 1.3)$ kcal mol⁻¹ for 1SA, $-(25.6 \pm 1.4)$ kcal mol⁻¹ for 3SA and $-(26.1 \pm 1.3)$ kcal mol⁻¹ for 4SA (ESI Fig. S6†). These results agree well with enthalpies measured in previous studies.^{26,31,32} Within experimental errors, the binding enthalpies per binding site for all SA variants are the same, suggesting that in the absence of force all subunits are equivalent with regard to biotin binding and that no effects of binding geometries or binding cooperativity come into play.

Single-molecule MT measurements determine lifetimes under force

To directly measure the lifetimes of the SA–biotin interactions under constant force and to investigate the influence of different force-loading geometries, we performed MT measurements using the different SA variants (Fig. 3). In MT, the molecular construct of interest is tethered between the bottom surface of a flow cell and a superparamagnetic bead (Fig. 3A). By applying a magnetic field, generated by permanent magnets, a constant force is exerted on the bead and thereby on the tether.^{33,34} We track the 3D position of the bead and the extension of the tether can be determined with nanometer resolution. Importantly, with our MT setup we can track approximately 100 beads in parallel, enabling us to obtain good statistics in a short amount of time.²⁵ In addition, MT provide excellent force and drift stability, facilitating long measurements,²⁵ which are critical due to the high stability of the SA–biotin bonds.

For the MT measurements, the small protein domain ddFLN4 (fourth F-actin cross-linker filamin rod domain of *Dictyostelium discoideum*³⁵) was biotinylated and covalently coupled to the bottom surface of a flow cell by an elastin-like polypeptide (ELP) linker.³⁶ The different SA variants (4SA, 3SA, or 1SA) were site-specifically and covalently immobilized on magnetic beads *via* polyethylene glycol (PEG) linkers, by reacting the C-terminal cysteine of subunit D with a thiol-reactive maleimide group on the PEG linker (Fig. 3A). The SA-functionalized beads were introduced into the flow cell and one of the functional subunits of the respective SA construct bound to the biotinylated ddFLN4, thereby tethering the magnetic bead to the surface. Upon force application, the molecular linkers are stretched and ddFLN4 unfolds in a characteristic two-step manner^{25,37} (Fig. 3B). We use the distinct two-step unfolding pattern as fingerprint to identify specific, single-tethered beads, *i.e.* beads that are bound to the surface *via* a single SA–biotin interaction.

MT measurements at 65 pN reveal different lifetime populations

In a first set of measurements, beads were subjected to a constant force of 65 pN and the time until bead rupture was recorded. The rupture events are attributed to the unbinding of biotin from SA, as this is the only non-covalent bond within the tether connecting the beads with the surface and as the ddFLN4 protein fingerprint allows us to limit the analysis to correctly tethered beads.

Measurements of 1SA, the monovalent variant, exhibited a survival time distribution that is well described by a single-exponential fit (Fig. 3C, red) with a lifetime of $\tau_1 = 7.2$ h \pm 0.2 h (2.61×10^4 s \pm 680 s; see ESI Materials and Methods for details of the fits†). The fitted lifetime is in good agreement with the 6.7 h reported recently for a smaller data set.²⁵ The single-exponential lifetime suggests the presence of a single population, consistent with the expectation that for 1SA only subunit D (attached to the bead *via* its C-terminus) is capable of binding biotin. All 1SA-functionalized beads are thus tethered in the same geometry, resulting in one well-defined force-loading direction.

3SA is complementary to the 1SA variant, in the sense that all but the attached subunit are functional, so that three different pulling geometries are possible for 3SA. The lifetime measurements reveal much shorter overall lifetimes for 3SA compared to 1SA (Fig. 3C, compare red and green data points). In addition, the data are not well described by a single exponential, suggesting that the different possible pulling geometries for 3SA give rise to different lifetimes. The 3SA data are well described by a fit with the sum of three exponentials and we find two relatively short and one longer lifetime (fitted lifetimes are 98 s \pm 12 s, 365 s \pm 30 s, and 3100 s \pm 230 s). A simpler model that combines the two shorter lifetimes into one exponential, $f(t) = f_2/3 [1 \exp(-t/\tau_2) + 2 \exp(-t/\tau_3)]$, fits the 3SA data almost equally well (Fig. 3C, green). From this fit we obtain the two distinct lifetimes as $\tau_2 = 3600$ s \pm 350 s and $\tau_3 = 199$ s \pm 10 s, where the weighting factors of the fit formula are chosen such that two thirds of the interaction of biotin and 3SA exhibit the short lifetime and one third exhibits the long one. We hypothesize that the longer-lived population corresponds to binding to one subunit, while the other two subunits exhibit lifetimes under mechanical tension that are similar.

Using steered molecular dynamics simulations, Sedlak *et al.*²⁹ have shown that for pulling biotin out of subunit A or C of SA tethered by the C-terminus of subunit D, the molecular linker adjacent to biotin gets pushed against a flexible peptide loop, significantly lowering the mechanical stability of the binding pocket. For pulling biotin out of subunit B, the same effect occurs, yet markedly less pronounced. Therefore, we assign the longer lifetime for the 3SA construct to biotin unbinding from subunit B. The shorter-lived population that comprises approximately two thirds of all unbinding events is consequently assigned to the sum of unbinding events from subunits A and C. Remarkably, the lifetime of this shortest-lived population at 65 pN is 130-fold lower than the one

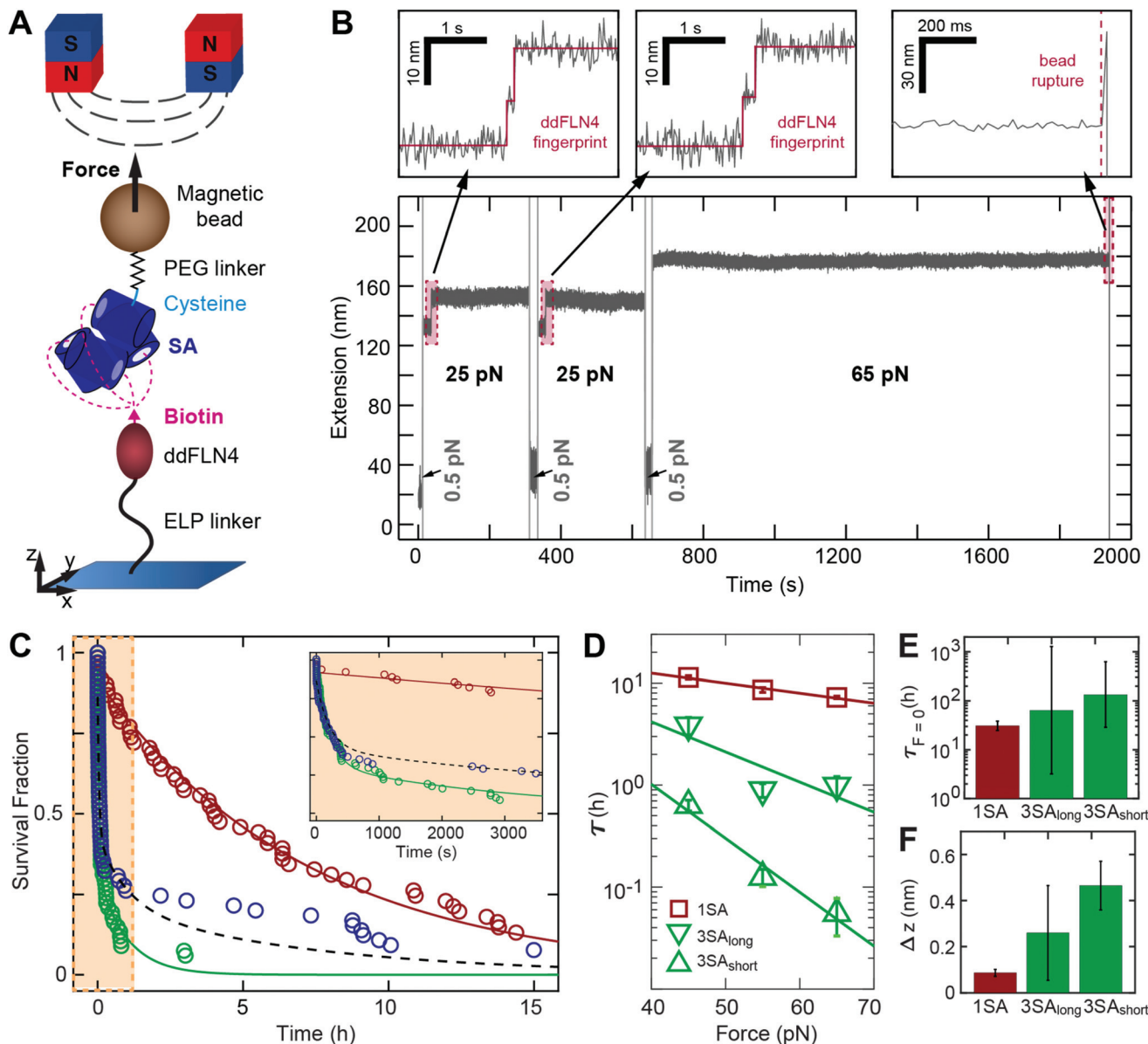


Fig. 3 Lifetimes of SA–biotin interactions under constant force probed in MT. (A) Schematic of MT experiments (not to scale). SA (4SA, 3SA, or 1SA) is site-specifically and covalently immobilized on magnetic beads *via* the single C-terminal cysteine at its subunit D using a PEG linker with a thiol-reactive maleimide group. Biotinylated ddFLN4 is covalently immobilized on the bottom slide of the MT flowcell *via* an ELP linker. Binding of the biotin to one of the functional subunits of the respective SA construct tethers the beads by a single SA–biotin bond. Force is exerted on the magnetic beads by permanent magnets positioned above the flowcell. (B) Time trace of the tether extension during an MT measurement. At the beginning of the measurement, beads are subjected to two 5 min intervals at 25 pN, to observe unfolding of ddFLN4 in a characteristic two-step pattern (left and middle zoom-in), which serves as fingerprint to identify specific, single-tethered beads. Short low-force intervals (0.5 pN) allow for ddFLN4 refolding. Tethers are then subjected to a constant force of 65 pN and the time until bead rupture due to unbinding of biotin from SA is monitored (right zoom-in). (C) Survival fractions at 65 pN as a function of time for 1SA (red), 3SA (green), and 4SA (blue). 1SA data were fit with a single exponential (red line) with a mean lifetime of 2.6×10^4 s. 3SA data were fit with a two-exponential model (ESI Materials and Methods[†]) with a short lifetime of 199 s and a long lifetime of 3.6×10^3 s (green line). 4SA data are well described by the predicted response from the combination of 1SA and 3SA lifetimes that takes into account the binding site stoichiometry (black dashed line, not a fit). The inset shows a zoom on the first hour of the data. (D) Lifetimes for 1SA (τ_1) and for the long- and short-lived 3SA interactions (τ_2 and τ_3) as a function of applied force. Data points are from fits of the survival fractions in panel C and in ESI Fig. S8. [†] Error bars are from a bootstrap analysis. Solid lines are fits of the Bell model. (E) Lifetimes at zero force τ_0 from fits of the Bell model in panel D. (F) Distances to the transition state Δz from fits to the Bell model. Error bars in (E) and (F) correspond to 95% confidence intervals from the fits.

observed for the 1SA construct and even the long lifetime observed for 3SA is still approximately one order of magnitude shorter than the lifetime observed for 1SA.

For 4SA, we observe a rapid initial decay of bonds, but also very long-lived tethers (Fig. 3C, blue). Since for 4SA all force-loading geometries realized in 3SA and 1SA are possible, we expect a combination of the short-lived populations observed for 3SA and the long-lived population observed for 1SA constructs. Based on this assumption, we co-plotted a prediction for the 4SA survival fraction over time as given by $f(t) = f_3/4 [\exp(-t/\tau_1) + \exp(-t/\tau_2) + 2 \exp(-t/\tau_3)]$ using the lifetimes obtained from fitting the 1SA and 3SA data (Fig. 3C, black dashed line). The prediction using the fitted lifetimes from the 1SA and 3SA data closely matches the experimentally determined lifetimes for the 4SA variant, confirming the validity of our lifetime model and suggesting essentially random binding to the different subunits.

Lifetimes depend exponentially on applied load

To determine the force dependencies of the lifetimes, we performed a set of experiments at lower forces on the 1SA and 3SA constructs. The lifetime data at 45 and 55 pN are well described by the single and double exponential models used at 65 pN for 1SA and 3SA, respectively (ESI Fig. S8†). We found that all observed lifetimes (τ_1 – τ_3) systematically increase with decreasing force (Fig. 3D). For example, τ_1 for 1SA increases to $11.2 \text{ h} \pm 0.4 \text{ h}$ at 45 pN. The force dependencies of the lifetimes are well described by the Bell model³⁸ with an exponential dependence on the force: $\tau(F) = \tau_0 \exp(-F \cdot \Delta z / k_B T)$ where Δz is the distance to the transition state, τ_0 the lifetime in the absence of force, and $k_B T$ the Boltzmann constant times the absolute temperature (Fig. 3D). From fits of the Bell model, we find similar lifetimes in the absence of force for τ_1 , τ_2 , and τ_3 , in the range of $\tau_0 \sim 50 \text{ h}$ (Fig. 3E). In contrast, the fitted distances to the transition state Δz are significantly different for the three lifetimes. Δz is smallest for 1SA and increases for the long-lived and again for the short-lived 3SA population (Fig. 3F). The observed convergence of lifetimes, within error, at zero force would be expected, since in the absence of force the force-loading direction should be irrelevant. The orders-of-magnitude differences between lifetimes under force for the different force-loading directions is accounted for in the Bell model by the different distances to the transition state, which correspond to the slopes of $\log(\text{lifetime})$ vs. applied force (Fig. 3D). With the caveat that extrapolation over orders-of-magnitude to zero force is necessarily somewhat imprecise, we find off-rates at zero force $k_{0,\text{off}} = \tau_0^{-1}$ of 2 to $8 \times 10^{-6} \text{ s}^{-1}$ from our constant force measurements in the MT, well within the range of results from dynamic AFM force spectroscopy measurements that reported values of 10^{-5} s^{-1} to 10^{-7} s^{-1} for the 1SA construct from fits of the Bell–Evans model to data at defined loading rates.^{28,29,39} The observed off-rates at zero force from force spectroscopy are also in reasonable agreement with the value of $6.1 \times 10^{-5} \text{ s}^{-1}$ determined in bulk from a radiolabeled biotin assay.³⁰ Further, we find reasonable agreement between $\Delta z_{1\text{SA}} = 0.09 \text{ nm} \pm 0.03 \text{ nm}$ from fits of the Bell

model to constant force MT data and the values in the range of 0.13 to 0.23 nm from fits of the Bell–Evans model to constant retraction speed AFM data.^{28,29} The small value of $\Delta z_{1\text{SA}}$ corresponds to a highly cooperative unbinding transition and ensures high stability even under load. In contrast, Δz for the force-loading directions probed with the 3SA construct are larger, which can likely be attributed to the molecular mechanism observed in constant-speed force spectroscopy experiments combined with all-atom steered molecular dynamics simulations: for certain pulling directions, the SA binding pocket is deformed before biotin leaves the pocket and, consequently, the unbinding pathway is altered, resulting in lower unbinding forces for measurements at constant retraction velocities,²⁹ and in shorter lifetimes for constant force experiments.

More importantly, from an application perspective, the force-loading geometry that yields the longest lifetime corresponds to pulling biotin out of the binding pocket of the subunit that is C-terminally tethered. The lifetime for this geometry is, at the forces probed here, almost two orders-of-magnitude larger than for the other possible geometries. Thus, it is highly beneficial to utilize this geometry in applications for which high yield of tethers with high force stability is desirable. Importantly, this can straightforwardly be realized employing the 1SA variant used in our experiments.

Finally, we note that the lifetimes obtained for the site-specifically attached 4SA used here were, both for the longest- and for the shortest-lived population, appreciably higher than the respective lifetimes measured for commercially available SA-coated beads (Dynabeads M-270 Streptavidin, Invitrogen/Thermo Fisher)²⁵ or beads randomly coupled *via* lysine residues.¹⁴ This difference may be explained considering that the SA–biotin complex can withstand higher forces when loaded with force from the C-terminus as compared to pulling from the N-terminus, as it has recently been demonstrated for 1SA in AFM SMFS.^{28,40} The attachment of commercially available beads is likely not site-specific, resulting in a variety of pulling geometries, whereas in the custom SA constructs, force was specifically applied from the C-terminus, ensuring highest stability.

Conclusions

To conclude, we show that the lifetimes of the SA–biotin interaction subjected to constant mechanical load strongly depend on the force-loading geometry and exponentially decrease with increasing force. Different geometries arise from binding of biotin to one of the four binding pockets of SA and result in lifetimes under force that differ by orders-of-magnitude, despite identical thermodynamic stabilities for binding to the different subunits and similar extrapolated off-rates at zero force. Our results illustrate that it is, in general, not possible to infer the mechanical stability of a receptor–ligand complex from its affinity and binding enthalpy.

Such differences between thermal and forced dissociation of molecular complexes are plausible considering the high-dimensional binding energy landscape. Unbinding pathways under mechanical load can be very different from each other and also very different from the thermally preferred ones, as it is also observed *e.g.* for the force-induced melting of double-stranded DNA in shear- or zipper-geometry.⁴¹ For proteins, similar behavior of monovalent SA has recently been employed by Erlich *et al.* to create a force hierarchy of receptor–ligand complexes.⁴² Also, the mechanically most stable receptor–ligand complex measured to date⁴³ has just ordinary thermodynamic binding characteristics.

Our work provides a clear route to improving the yield of force spectroscopy experiments and in general of assays where SA–biotin is used for attachment and experiences mechanical loads, *e.g.* due to fluid flow or magnetic actuation. For measurements utilizing the SA–biotin interaction as a handle, and in particular for constant force SMFS measurements, it is highly beneficial to implement a specific SA tethering geometry that yields a single population with high lifetime to enable long measurement durations even at high forces. The tethering geometry that we identified as the one yielding the longest lifetimes can be easily realized in experiments by employing the 1SA variant presented in this study. Thus, our results give a straightforward approach for highly specific and stable experiments that employ the SA–biotin linkage.

Conflicts of interest

There are no conflicts to declare.

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Notes and references

- S. Basu, A. Finke, L. Vera, M. Wang and V. Olieric, *Acta Crystallogr., Sect. D: Struct. Biol.*, 2019, **75**, 262–271.
- O. H. Laitinen, V. P. Hytonen, H. R. Nordlund and M. S. Kulomaa, *Cell. Mol. Life Sci.*, 2006, **63**, 2992–3017.
- C. M. Dundas, D. Demonte and S. Park, *Appl. Microbiol. Biotechnol.*, 2013, **97**, 9343–9353.
- M. Wilchek and E. A. Bayer, *Anal. Biochem.*, 1988, **171**, 1–32.
- W. Humphrey, A. Dalke and K. Schulten, *J. Mol. Graph.*, 1996, **14**, 33–38.
- V. V. Didenko, *Anal. Biochem.*, 1993, **213**, 75–78.
- F. Kriegel, W. Vanderlinden, T. Nicolaus, A. Kardinal and J. Lipfert, *Methods Mol. Biol.*, 2018, **1814**, 75–98.
- I. D. Vilfan, J. Lipfert, D. A. Koster, S. G. Lemay and N. H. Dekker, in *Handbook of Single-Molecule Biophysics*, ed. P. Hinterdorfer and A. Oijen, Springer US, New York, NY, 2009, pp. 371–395, DOI: 10.1007/978-0-387-76497-9_13.
- E. A. Bayer and M. Wilchek, *Methods Enzymol.*, 1990, **184**, 138–160.
- M. G. Cull and P. J. Schatz, *Methods Enzymol.*, 2000, **326**, 430–440.
- S. M. Cannizzaro, R. F. Padera, R. Langer, R. A. Rogers, F. E. Black, M. C. Davies, S. J. Tendler and K. M. Shakesheff, *Biotechnol. Bioeng.*, 1998, **58**, 529–535.
- V. T. Moy, E. L. Florin and H. E. Gaub, *Science*, 1994, **266**, 257–259.
- R. Merkel, P. Nassoy, A. Leung, K. Ritchie and E. Evans, *Nature*, 1999, **397**, 50–53.
- C. Danilowicz, D. Greenfield and M. Prentiss, *Anal. Chem.*, 2005, **77**, 3023–3028.
- F. Rico, A. Russek, L. Gonzalez, H. Grubmuller and S. Scheuring, *Proc. Natl. Acad. Sci. U. S. A.*, 2019, **116**, 6594–6601.
- J. Lipfert, J. W. Kerssemakers, T. Jager and N. H. Dekker, *Nat. Methods*, 2010, **7**, 977–980.
- J. Lipfert, M. M. van Oene, M. Lee, F. Pedaci and N. H. Dekker, *Chem. Rev.*, 2015, **115**, 1449–1474.
- J. A. Rivas-Pardo, E. C. Eckels, I. Popa, P. Kosuri, W. A. Linke and J. M. Fernández, *Cell Rep.*, 2016, **14**, 1339–1347.
- F. Baumann, M. S. Bauer, L. F. Milles, A. Alexandrovich, H. E. Gaub and D. A. Pippig, *Nat. Nanotechnol.*, 2016, **11**, 89–94.
- W. Ott, M. A. Jobst, C. Schoeler, H. E. Gaub and M. A. Nash, *J. Struct. Biol.*, 2017, **197**, 3–12.
- R. Walder, M. A. LeBlanc, W. J. Van Patten, D. T. Edwards, J. A. Greenberg, A. Adhikari, S. R. Okoniewski, R. M. A. Sullan, D. Rabuka, M. C. Sousa and T. T. Perkins, *J. Am. Chem. Soc.*, 2017, **139**, 9867–9875.
- Z. Ganim and M. Rief, *Proc. Natl. Acad. Sci. U. S. A.*, 2017, **114**, 11052–11056.
- M. Krieg, G. Fläschner, D. Alsteens, B. M. Gaub, W. H. Roos, G. J. L. Wuite, H. E. Gaub, C. Gerber, Y. F. Dufrêne and D. J. Müller, *Nat. Rev. Phys.*, 2019, **1**, 41–57.
- H. Chen, G. Yuan, R. S. Winardhi, M. Yao, I. Popa, J. M. Fernandez and J. Yan, *J. Am. Chem. Soc.*, 2015, **137**, 3540–3546.
- A. Löf, P. U. Walker, S. M. Sedlak, S. Gruber, T. Obser, M. A. Brehm, M. Benoit and J. Lipfert, *Proc. Natl. Acad. Sci. U. S. A.*, 2019, **116**, 18798–18807.
- S. M. Sedlak, M. S. Bauer, C. Kluger, L. C. Schendel, L. F. Milles, D. A. Pippig and H. E. Gaub, *PLoS One*, 2017, **12**, e0188722.

- 27 G. Fonnum, N. Hofsløkken, E. Aksnes, L. Killas, P. Stenstad, R. Schmid, J. Bjørgum, T. Nilsen and A. Berge, *EU Patent EP1*, 2006, 693.
- 28 S. M. Sedlak, L. C. Schendel, M. C. R. Melo, D. A. Pippig, Z. Luthey-Schulten, H. E. Gaub and R. C. Bernardi, *Nano Lett.*, 2019, **19**, 3415–3421.
- 29 S. M. Sedlak, L. C. Schendel, H. E. Gaub and R. C. Bernardi, *Sci. Adv.*, 2020, **6**, eaay5999.
- 30 M. Howarth, D. J. Chinnapen, K. Gerrow, P. C. Dorrestein, M. R. Grandy, N. L. Kelleher, A. El-Husseini and A. Y. Ting, *Nat. Methods*, 2006, **3**, 267–273.
- 31 A. Chilkoti and P. S. Stayton, *J. Am. Chem. Soc.*, 1995, **117**, 10622–10628.
- 32 M. Sarter, D. Niether, B. W. Koenig, W. Lohstroh, M. Zamponi, N. H. Jalarvo, S. Wiegand, J. Fitter and A. M. Stadler, *J. Phys. Chem. B*, 2020, **124**, 324–335.
- 33 T. R. Strick, J. F. Allemand, D. Bensimon, A. Bensimon and V. Croquette, *Science*, 1996, **271**, 1835–1837.
- 34 J. Lipfert, X. Hao and N. H. Dekker, *Biophys. J.*, 2009, **96**, 5040–5049.
- 35 I. Schwaiger, A. Kardinal, M. Schleicher, A. A. Noegel and M. Rief, *Nat. Struct. Mol. Biol.*, 2004, **11**, 81–85.
- 36 W. Ott, M. A. Jobst, M. S. Bauer, E. Durner, L. F. Milles, M. A. Nash and H. E. Gaub, *ACS Nano*, 2017, **11**, 6346–6354.
- 37 M. S. Bauer, L. F. Milles, S. M. Sedlak and H. E. Gaub, bioRxiv, 2018.
- 38 G. I. Bell, *Science*, 1978, **200**, 618.
- 39 E. Evans and K. Ritchie, *Biophys. J.*, 1997, **72**, 1541–1555.
- 40 L. C. Schendel, S. M. Sedlak and H. E. Gaub, *Nanoscale*, 2020, **12**, 6803–6809.
- 41 S. K. Kufer, E. M. Puchner, H. Gumpf, T. Liedl and H. E. Gaub, *Science*, 2008, **319**, 594–596.
- 42 K. R. Erlich, S. M. Sedlak, M. A. Jobst, L. F. Milles and H. E. Gaub, *Nanoscale*, 2019, **11**, 407–411.
- 43 L. F. Milles, K. Schulten, H. E. Gaub and R. C. Bernardi, *Science*, 2018, **359**, 1527–1533.