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Rapid and Robust Polyprotein Production Facilitates Single Molecule Mechanical Characterization of BamA POTRA Domains

Toni Hoffmann‡†, Katarzyna M. Tych‡†, Thomas Crosskey†, Robert Schiffrin†, David J. Brockwell†‡ and Lorna Dougan†‡

†School of Physics and Astronomy, University of Leeds, Leeds, LS2 9JT, ‡Astbury Centre for Structural and Molecular Biology, University of Leeds, Leeds, LS2 9JT, §School of Molecular and Cellular Biology, University of Leeds, Leeds, LS2 9JT

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ABSTRACT: Single molecule force spectroscopy by atomic force microscopy exploits the use of multimeric protein constructs, namely polyproteins, to decrease the impact of non-specific interactions, to improve data accumulation and to allow the accommodation of benchmarking reference domains within the construct. However, methods to generate such constructs are either time- and labour-expensive or lack control over the length or the domain sequence of the obtained construct. Here, we describe an approach that addresses both of these shortcomings, that uses Gibson Assembly (GA) to rapidly generate a defined recombinant polyprotein using novel linker sequences. To demonstrate the feasibility of this approach we used GA to make a polyprotein comprised of alternating domains of I27 and Tm Csp, (I27- Tm Csp), and showed the mechanical fingerprint, mechanical strength and pulling speed dependence are the same as the same polyprotein constructed using the classical approach. After this benchmarking we exploited this approach to facilitate the mechanical characterization of POTRA domain 2 of BamA from E. coli (EcPOTRA2), by assembling the polyprotein (I27-EcPOTRA2)–I27GA. We show that, as predicted from the α+β topology, EcPOTRA2 domains are mechanically robust over a wide range of pulling speeds. Furthermore, we identify a clear correlation between mechanical robustness and brittleness for a range of other α+β proteins that contain the structural feature of proximal terminal β-strands in parallel geometry. We thus demonstrate that the GA approach is a powerful tool as it circumvents the usual time- and labour-expensive polyprotein production process, and allows for rapid production of new constructs for single molecule studies. As shown for EcPOTRA2, this approach allows the exploration of the mechanical properties of a greater number novel proteins and their variants. This improves our understanding of the relationship between structure and mechanical strength, increasing our ability to design proteins with tailored mechanical properties.

Single-molecule force spectroscopy using the atomic force microscope (AFM) is an important technique to study the mechanical stability and energy landscape of single proteins by mechanically perturbing their structure. Using this approach, protein molecules are immobilized on a surface, and the AFM probe is repeatedly brought into contact with the surface. Upon immobilization to the tip of the AFM cantilever, through a non-specific interaction, the tethered protein is extended via retraction of the tip at a constant velocity or by varying the tip-substrate separation to apply a constant force. As both methods increase the distance between the surface and the cantilever tip, a mechanical force is exerted, and the protein is stretched leading to successive domain unfolding which finally results in a fully unfolded protein. While single protein domains can be used to polyprotein chains (which comprise repeats of identical or alternating protein domains) are typically employed. These polyproteins provide clear mechanical fingerprints in a single molecule force spectroscopy experiments, such as the saw-tooth pattern seen in force-extension traces collected in a constant-velocity experiment. The use of polyproteins is important as it reduces the frequency of non-specific interactions between the AFM probe and the surface relative to bona fide protein unfolding events and it increases the number of data points collected for a given approach and retraction cycle. Polyproteins can either consist of identical protein domains, termed homopolyproteins or of alternating domains of different proteins, termed chimeric- or heteropolyproteins.

A number of different techniques have been employed to engineer such polyproteins. A classical approach is based on the assembly of DNA cassettes that together encode the full-length polyprotein. This method uses specific restriction sites between DNA fragments that encode respective protein domains. Sequential enzymatic digestions and ligations generate the full length polyprotein DNA in a step-wise manner. This method allows for the precise control of the number of domains in the polyprotein and the order of protein domains within a chimeric polyprotein. This method has consequently been employed to make a wide range of different homo- and chimeric polyprotein constructs, making it a versatile method for polyprotein production. However, in this approach the substitution of single DNA fragments occurs sequentially, making the process both laborious and time-expensive. Variants of this method such as ligating multiple copies of a single ‘cassette’ encoding the non-palindromic restriction site (AvaI) at its termini or the iterative ligation of cassettes that en-
code BamHI and BglII or NheI and SpeI restriction sites 5′ and 3′ to the gene are quicker but yield less defined products or cannot be used to construct more complex polyproteins. An alternative method is based on the chemical coupling of identical protein monomers or dimers. Here, proteins can be linked through either disulfide bridge formation between cysteine pairs at designed locations or by maleimide coupling of sulhydryl groups within the protein. This chemical coupling allows for a faster one-step construction of polyproteins. It also has the advantage of allowing for the study of different pulling directions by changing the position of linkage groups on the surface of the protein. However, this method precludes the generation of more complex protein scaffolds and generates an ensemble of low- to high-order multimers which may have an impact on the subsequent analysis of the experimental data.

Figure 1. Gibson assembly cloning. The Gibson assembly of a polyprotein contains 21 nucleotides guided by the following principles: (i) the linkers should have similar physico-chemical properties at the protein level but sufficiently distinct at the DNA level to allow specific and ordered assembly and (ii) the chosen amino acids should ensure stearic freedom and minimize any potentially interfering interactions with neighbouring protein domains. Figure 2A shows the sequence of each linker based on a symmetrical design with a central glycine (G) and a combination of uncharged, polar (S, serine; T, threonine) and non-polar residues (A, alanine; I, isoleucine; L, leucine; V, valine). By shuffling these residues while keeping the location of the central glycine constant, six unique linker peptides emerged possessing a global palindromic-like symmetry while maintaining the same amino acid composition and length (Figure 2A). At the DNA level, the specificity of the complementary region at cassette termini was further increased by switching between the two most frequently used codons for each amino-acid (Figure 2B). Gibson assembly was carried out as described in the Methods section. Briefly, DNA cassettes that encode individual protein domains (Figure 1, where X designates the previously mechanically characterized I27 domain and Y the protein under study) as well as the linearized backbone containing the first I27 domain (Table 1, Supporting Information) was obtained by standard PCR methods and joined to form a circular DNA molecule by Gibson assembly cloning. Full length assembled constructs were identified by colony PCR and their successful assembly confirmed by DNA sequencing.

RESULTS AND DISCUSSION

Polyprotein construction by Gibson assembly. Gibson assembly cloning depends on the presence of compatible overhanging ends in the DNA ‘cassettes’. As polyproteins used in force spectroscopy experiments consist of protein domains joined by unstructured peptide linkers, these regions can be used to encode complementary overlap regions between Gibson-assembly-compatible cassettes allowing for the ordered assembly of DNA molecules. The recommended length for Gibson-assembly-compatible overhangs is 20 nucleotides. Thus, we designed linker peptides made of seven amino acids encoded by 21 nucleotides guided by the following principles: (i) the linkers should have similar physico-chemical properties at the protein level but sufficiently distinct at the DNA level to allow specific and ordered assembly and (ii) the chosen amino acids should ensure stearic freedom and minimize any potentially interfering interactions with neighbouring protein domains. Figure 2A shows the sequence of each linker based on a symmetrical design with a central glycine (G) and a combination of uncharged, polar (S, serine; T, threonine) and non-polar residues (A, alanine; I, isoleucine; L, leucine; V, valine). By shuffling these residues while keeping the location of the central glycine constant, six unique linker peptides emerged possessing a global palindromic-like symmetry while maintaining the same amino acid composition and length (Figure 2A). At the DNA level, the specificity of the complementary region at cassette termini was further increased by switching between the two most frequently used codons for each amino-acid (Figure 2B). Gibson assembly was carried out as described in the Methods section. Briefly, DNA cassettes that encode individual protein domains (Figure 1, where X designates the previously mechanically characterized I27 domain and Y the protein under study) as well as the linearized backbone containing the first I27 domain (Table 1, Supporting Information) was obtained by standard PCR methods and joined to form a circular DNA molecule by Gibson assembly cloning. Full length assembled constructs were identified by colony PCR and their successful assembly confirmed by DNA sequencing.
of cardiac titin (I27) and cold shock protein B from the hyperthermophilic organism Thermatoga maritima (TmCsp, (I27-TmCsp)_3-I27<sup>GA</sup>) was constructed, expressed and purified. Its mechanical unfolding properties were then characterised by AFM (Figure S1) and compared to those of a homologous polyprotein constructed using the cassette approach ((I27-TmCsp)_3-I27<sup>GA</sup>). Full details of the sequences of both constructs can be found in the Supporting Information. The interpeak distance (x<sub>p2p</sub>), defined as the distance from one unfolding peak to the same force value on the following curve, and the peak unfolding force (F<sub>p1</sub>) was then measured for each unfolding event and frequency histograms constructed for both x<sub>p2p</sub> and F<sub>p1</sub>. Excellent agreement can be found for x<sub>p2p</sub> and F<sub>p1</sub> values measured for the constructs made using each method (Supplementary Information and Figure S1). We followed the same procedure to obtain force-extension traces at three other pulling velocities: 100, 200 (Figure S1) and 2000 nms<sup>−1</sup> (Figure S2 and Table S2). At each pulling velocity we completed three experiments to measure F<sub>p1</sub> for each type of unfolding event and constructed three histograms of F<sub>p1</sub>. We found that the pulling speed dependence of F<sub>p1</sub> for TmCSP and I27 were the same for both constructs, (I27-TmCsp)_3-I27<sup>GA</sup> and (I27-TmCsp)_3-I27 (Figure S2, Table S2).

**Investigating the mechanical strength of EcPOTRA2.** Having benchmarked polyproteins constructed by Gibson assembly against a homologous, previously characterized polyprotein, we employed this method to facilitate the rapid construction of a novel polyprotein that comprised I27 and a mechanically uncharacterized domain – the second POTRA domain of BamA from the E.coli BAM complex (EcPOTRA2). The function of BAM (β-barrel assembly machinery) is to fold and insert β-barrel outer membrane proteins (OMPs) into the outer membrane of Gram-negative bacteria, with homologous machinery present in mitochondria and chloroplasts (Figure 3). BamA, which carries out OMP insertion by an as yet unknown mechanism is essential for cell viability and comprises a β-barrel domain and five N-terminal, tandemly-arrayed POTRA (polypeptide transport associated) domains that extend into the periplasm.

**Reduction in time for polyprotein engineering.** Compared to standard cloning techniques, Gibson assembly cloning reduces the time to obtain polyprotein samples that can be mechanically tested on an AFM. The process starts with the PCR-based amplification and purification of Gibson cloning compatible fragments followed by the Gibson assembly reaction itself. On the same day, competent E. coli cells are transformed and spread onto selective plates. On day two, positive clones are selected by colony PCR, and liquid overnight cultures are inoculated, allowing identification of successful constructs after only three days. To date, this method has been used to construct seven polyproteins-encoding plasmids. Of these plasmids, 13 % (min = 4 %, max = 35%) contained a fully-assembled construct. By contrast to the rapidity of Gibson assembly, the classical construction of polyprotein takes significantly longer. Firstly, a PCR-amplified DNA fragment is subcloned into a bacterial expression vector that contains a polyprotein construct (e.g. (X<sub>3</sub>)), using restriction digestion of a unique pair of sites to replace the second protein domain encoding fragment in (X<sub>3</sub>), for example. This results in a X-Y-(X<sub>3</sub>) construct that needs to be selected by analytical digest / colony PCR and confirmed by sequencing. This will take a minimum of six days. Once confirmed, this construct is used for subsequent rounds of cloning to replace further DNA fragments (e.g. the fourth and the sixth domain) totaling in about 18 days of experimental work before a full construct ((X-Y)<sub>3</sub>-X) is confirmed. In reality, the subsequent repetitions of subcloning and selection increase the likelihood of failures and further delays.

**Single-molecule AFM experiments on (I27-TmCSP) polyprotein prepared by Gibson assembly.** To validate this approach, a polyprotein made by Gibson assembly comprising of alternating domains of the 27<sup>th</sup> immunoglobulin-like domain

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**Figure 2.** Peptide linker design. A) Gibson assembly-compatible linker peptides were designed comprising a balanced, symmetrical combination of seven polar and non-polar uncharged residues. B) Higher specificity between adjacent cassettes was achieved by varying codon usage by switching between the two most frequently used codons for each amino-acid in E. coli.

**Figure 3.** (A) Schematic of the BAM (β-barrel assembly machinery) complex in E. coli. The central component of the machinery is the outer membrane β-barrel protein BamA. The N-terminal periplasmic component of BamA consists of five POTRA domains (P1-P5). A ribbon representation of the tertiary structure for EcPOTRA2 is shown inset. In the AFM
protein unfolding experiments EcPOTRA2 is extended from the amino- and carboxy-terminal ends (N and C).

Figure 4. Investigating the mechanical properties of a POTRA domain. (A) Mechanical unfolding of the (I27-EcPOTRA2)3-I27 polyprotein constructed using Gibson assembly at 600 nm s⁻¹. The Worm Like Chain (WLC) model fit to the data is shown in grey (B) Schematic showing a single (I27- EcPOTRA2)-I27 molecule (EcPOTRA2 and I27, pink and yellow, respectively) attached to a gold surface (bottom) and the tip of an AFM cantilever (top). (C) The scatter plots of EcPOTRA2 (pink) and I27 (yellow) unfolding forces and inter-peak distances for 96 unfolding events are shown as the same symbol with colored regions to indicate their separation based on the force histogram. (D) Pulling speed dependence of the mechanical unfolding of (I27-EcPOTRA2)3-I27. The mechanical unfolding forces for EcPOTRA2 (pink squares) and I27 (yellow triangles) at 23°C. Each pair of data points at a given pulling speed show the average of the Gaussian mean values of the unfolding force for EcPOTRA2 and I27 from three experiments completed under the same conditions. The error bars indicate the standard deviation between the three experiments. Solid lines are the best fit to the data. The Monte Carlo fits to the experimental data are shown as dashed lines. (E) Schematic of the unfolding energy landscape for EcPOTRA2 with parameters obtained using the Monte Carlo fits to experimental results. These include the activation energy barrier height (ΔG*) and the distance from the native, folded state to the unfolding transition state (ΔxU).

POTRA domains are thought to perform three functions: (i) to act as a scaffold for complex formation; (ii) to interact with periplasmic chaperones and (iii) to form a binding site for OMP substrates. The mechanism(s) by which POTRA domains facilitate OMP folding and insertion is unclear but may involve a scaffolding role (by β-sheet augmentation between POTRA β-strands and the nascent OMP) or even a more active role by driving OMP β-hairpin formation by cycling between bent and extended conformations. Single molecule force unbinding studies are ideally suited to address this question, but a pre-requisite to such a study is the full characterisation of the mechanical properties of the POTRA domains themselves. In addition to its function where mechanical unfolding may play a role, EcPOTRA2 (residues 92-173 of BamA) was selected for study as these domains have an ideal topology to further our understanding of the relationship between protein structure, sequence and mechanical strength of topologically-simple proteins. Accordingly, these α+β domains are ~ 80 residues in length and comprise a three-stranded β-sheet packed against two helices (β1-α1-α2-β2-β3) whose N- and C-termini are located on two neighbouring parallel β-strands (Figure 3). From a mechanical unfolding perspective this is an attractive topology as these two strands might form a mechanical clamp region that conveys resistance to an applied force. Mechanically unfolding (I27-EcPOTRA2)3-I27 revealed a sawtooth-like unfolding force-
extension pattern corresponding to the sequential unfolding of single domains of either EcPOTRA2 or I27 (Figure 4A). While the spacing between each unfolding event appears similar (~22 nm), unfolding appears to occur at two distinct forces (~112 and 175 pN). A scatter plot (Figure 4C) that combines the data for $x_{p2p}$ and $F_U$ reveals two populations of events, one population at lower forces (highlighted by the pink squares) and a second population at higher forces (highlighted by the yellow squares). The latter distribution agrees well with the forces measured previously for I27 at this pulling speed (163 ± 15 pN). The shorter $x_{p2p}$ distribution (highlighted by the pink squares) reflects the slightly smaller size of the folded core of EcPOTRA2 relative to I27 (81 and 89 residues, respectively). These data thus demonstrate that EcPOTRA2 exhibits significant mechanical stability and unfolds in a single step.

**Unfolding energy landscape of EcPOTRA2**

Force-extension traces at three other pulling velocities (100, 280, and 2000 nm s$^{-1}$) were obtained to access the pulling speed dependence of EcPOTRA2. The $F_U$ histograms at each speed and the statistics for each experiment can be found in the Supporting Information (Figure S3 and Table S3). Figure 4D shows that $F_U$ for EcPOTRA2 (pink squares) and I27 (yellow circles) both increase as a function of pulling speed. Parameters which describe the underlying unfolding energy landscapes of the proteins, namely $\Delta G_U^*$, the height of the activation energy barrier and $\Delta x_{U}$, the distance between the folded state and the transition barrier on the mechanical unfolding pathway, were then obtained from this dependence using a Monte Carlo method. For EcPOTRA2 we obtained $\Delta x_{U} = 0.35 ± 0.01$ nm and $k_U = 0.0200 ± 0.0020$ s$^{-1}$ (dashed line, Figure 4E). Assuming a value of the pre-factor $A = 10^6$ s$^{-1}$ this gives an activation energy barrier height, $\Delta G_U^* = 44$ kJ mol$^{-1}$ for EcPOTRA2. Comparing this with I27, we obtained $\Delta x_{U} = 0.32 ± 0.04$ nm, $k_U = 0.0013 ± 0.0001$ s$^{-1}$, and $\Delta G_U^* = 45$ kJ mol$^{-1}$, in agreement with previous studies.

While the height of the activation energy barrier to unfolding is very similar for EcPOTRA2 and I27, the distance to the unfolding transition state is slightly larger for EcPOTRA2.

**Figure 5.** Mechanical stability of α + β proteins. (A) α + β proteins studied using single molecule force spectroscopy using the AFM, and for which a pulling speed dependence of the mechanical stability has been obtained. Proteins are shown in ribbon representations with β-strands as blue arrows and α-helices as green ribbons. The proteins are extended from their amino- and carboxy-terminal ends (N and C). The relevant PDB accession code is shown for each protein. The proteins are ordered by their unfolding force $F_U$ at 600 nm s$^{-1}$, with ubiquitin having the largest $F_U$ and RNaseA the smallest $F_U$. (B) Histograms show the percentage of α-helical (green) and β-sheet (blue) content in each protein, calculated from the PDB structure. The proteins are again ordered by their unfolding force $F_U$ at 600 nm s$^{-1}$. (C) The unfolding force $F_U$ at a pulling speed of 600 nm s$^{-1}$ and the unfolding distance $\Delta x_{U}$ are shown for each of the α + β proteins which have been experimentally unfolded at least at two different speeds. Where required, the expected unfolding force at 600 nm s$^{-1}$ was interpolated. The data can be described by a bootstrapped, non-linear fit following a power law with $\Delta x_{U} = 45.0 ± 10.5 /F_U$ ($R^2 = 0.86 ± 0.01$). Examination of the structure of the 12 α + β proteins reveals a sub-set of
proteins (blue circles) which possess a similar structural feature - directly-connected, parallel alignment of the N- and C-terminal β-strands. α + β proteins which lack this structural feature are shown as dark blue squares.

**Figure 6.** The number of inter-residue contacts correlates with Δx_U and k_U. (A) Contact maps for 6 α + β proteins, protein G, protein L, SUMO2, ubiquitin, EcPOTRA2 and SUMO1. The protein name is followed by its PDB accession code. (B) The number of inter-residue contacts correlates with Δx_U (upper) and with k_U (lower) for the 6 α + β proteins

**Mechanical stability of α + β proteins.** The data set from protein mechanical unfolding experiments has greatly expanded in the last decade, permitting understanding of the measured differences in mechanical strength for proteins. Previously we completed a review of all experimental data on single molecule characterization of proteins using the AFM and showed that proteins which are mechanically strong, and exhibit high unfolding forces, are typically “brittle”, with small Δx_U values. Conversely, proteins which exhibit low unfolding forces are typically “soft”, with large Δx_U values. While β-strand-rich proteins populate the strong and brittle end of the spectrum, α-helical-rich proteins are comparatively weak and soft. An interesting intermediate region is populated by proteins which are α-helical and β-strand-rich, like the EcPOTRA2 studied here. An updated review of the current literature shows experiments that measure the pulling speed dependence of the unfolding force, F_U (such as those shown in the Figure 4D) have been completed for 11 other α + β proteins (Figure 6A). These include proteins such as the extensively studied ubiquitin, protein G and protein L, as well as the more recently studied SUMO proteins. The 12 α + β proteins range in size from 56 amino acids (protein G) to 155 amino acids (RNase H) and have varied α-helical and β-sheet content (Figure 5B). There is no correlation between the amount of α-helix or β-sheet content and F_U. Comparing the experimentally measured F_U at 600 nm s⁻¹ and Δx_U for each of the 12 α + β proteins (Figure 5C) reveals a power law dependence (solid line, Figure 5C) of the form Δx_U = 45.0/F_U. The scaling law indicates that mechanically weaker α + β proteins have a larger value of Δx_U, indicative of a ‘softer’ protein structure which can be deformed by a greater amount before reaching the unfolding transition state and unfolding. The power law dependence measured for α + β proteins (Figure 5C) is in agreement with previous work on a broader range of protein structures, including all-β proteins and all-α proteins. Here, by focusing on a specific class of protein, all α + β, we can obtain a more accurate correlation between F_U and Δx_U, including an improved chi-squared value. Of the 12 α + β proteins studied to date, eight are mechanically strong (F_U > 100 pN at 600 nm s⁻¹) and relatively brittle (i.e. small Δx_U < 0.51 nm). While 4 α + β proteins are mechanically weak (F_U < 70 pN at 600 nm s⁻¹) and malleable (i.e. large Δx_U > 0.68 nm). It is interesting to consider whether the mechanically strong and brittle proteins share any common features, which the mechanically weak and malleable proteins lack. Inspection of the structure of the α+β proteins studied to date by AFM (Figure 6A) shows that six proteins possess directly-connected, parallel alignment of the N- and C-terminal β-strands (protein G, protein L, SUMO2, ubiquitin, EcPOTRA2 and SUMO1). This geometry has previously been hypothesized as the main reason for mechanical stability in β-strand containing proteins, as proteins with this geometry at the termini have been found to have higher unfolding forces than those which lack this feature. To further understand the mechanical properties of α + β proteins, we calculated the inter-residue contacts in the proteins which possess connected, parallel β-strands at their termini and constructed contact maps for each protein (Figure 6A). An inter-residue contact was assumed to be present if the shortest distance between side-chain atoms of different residues was < 5 Å, the same cut-off used in a similar study on SUMO proteins. For the six identified α + β proteins we find that the number of contacts increases from 92 contacts in protein G to 171 in the SUMO1 protein. Interestingly, we find a clear correlation between both Δx_U and k_U and the number of contacts (Figure 6B), where the Δx_U and k_U from each study has been obtained using a similar method as employed in the present study using EcPOTRA2. This correlation was not observed when all twelve of the α + β proteins which have been studied using single molecule force spectroscopy are included (Figure S4). As the number of contacts increases for the six identified α + β proteins, Δx_U increases (Figure 6B, upper) and k_U decreases (Figure 6B, lower), resulting in enhanced malleability and mechanical stability. Possessing a large number of inter-residue contacts may be an important feature for mechanical stability, providing the protein with increased options for contacts breaking and reforming else-
Previous studies on the unfolding processes of protein L, protein G, ubiquitin and SUMO1 and 2 suggest that these proteins unfold by a similar structural mechanism. The transition state to unfolding is thought to involve the breaking of contacts between two well-defined structural units. For example, for protein L one structural unit comprises β-strands I and II and the helix (light blue in protein L schematic Figure 7A), and the second unit involves the β-hairpin of strands III and IV (dark blue in protein L schematic Figure 7A). As the transition states to unfolding for several of the proteins described here are known to involve the breaking of contacts between two well-defined structural units, we divided each of the 6 α+β proteins identified into two structural units (shown as light blue and dark blue in Figure 7A). Contacts between these two structural units were defined as long-distance contacts. We examined the contact maps of side-chain contacts (nearest distance between atoms of two residues < 5 Å) using CMView software and found that there is a correlation between the number of long-range contacts and the measured Δxₜ (upper) and kₜ (lower) for 5 of the α+β proteins (Figure 7B). For example, protein G has only 26 contacts and a relatively small Δxₜ, while EcPOTRA2 has 50 contacts and a relatively large Δxₜ. The higher number of long-range contacts between mechanically important structural units may cause the higher mechanical resilience in EcPOTRA2, as these residues must first be broken apart, before the protein can be extended. The exception is SUMO 1, which has 42 contacts and a higher Δxₜ (Figure 7B, upper) than the other 5 α+β proteins. To examine this further, we calculated the number of inter-residue hydrogen bonds between N- and C-terminal β-strands in the 6 α+β proteins and found that it did not correlate with Δxₜ or kₜ (Figure S5). Therefore, the best correlation we have identified for the 6 α + β proteins is that between the total number of inter-residue contacts and Δxₜ (Figure 6B, upper) and kₜ (Figure 6B, lower).

This suggests that other features of the protein structure must be responsible for tailoring their mechanical properties. By fine-tuning our understanding of protein stability and malleability we obtain more robust predictive tools for understanding the mechanical properties of previously mechanically uncharacterized proteins. Such insight will be increasingly important for the rational design of novel, protein-based materials.

CONCLUSIONS

We have successfully demonstrated that Gibson assembly cloning can be used to rapidly produce chimeric polyproteins for use in AFM-based force spectroscopy. We designed new linker peptides (Figure 2) to allow for an ordered assembly of DNA fragments (Figure 1). These novel linker peptides did not affect the observed mechanical properties of TmCsp and I27 compared to a previously characterized polyprotein that only differed in its linker sequences (Figure S1 and 2). Most importantly, this cloning method is able to significantly reduce the time required to obtain protein samples. Being able to assemble polyproteins from a set of pre-made DNA building blocks will increase the design flexibility for the generation of future constructs. It is also feasible to combine constructs by subsequent assembly reactions. This would allow the generation of longer polyproteins (e.g. (X-Y)₉,I27 or (X-Y)₁₅(U-W)₁₅-I27) that would otherwise be too labour- and time-intensive to produce. It is our hope that the application of Gibson assembly cloning to the production of polyproteins will contribute to increasing the number of protein domains that will be mechanically studied in single-molecule experiments. Thus it will help to broaden our understanding of the forces that define the folding and topologies of amino acid chains. Moreover, this technique enables the generation of previously im-

Figure 7. Two distinct structural units are identified in each of the 6 α + β proteins, protein G, protein L, SUMO2, Ubiquitin, EcPOTRA2 and SUMO1. The units are shown in light blue and dark blue. (B) The number of long-range contacts between the structural units is obtained using CMView software, using contact maps of side-chain contacts nearest distance between atoms of two residues > 5 Å. The number of long-range contacts in protein G (26), protein L (31), SUMO2 (46), Ubiquitin (48), EcPOTRA2 (50) and SUMO1 (42). The number of long-range contacts is shown against Δxₜ (upper) and kₜ (lower).

Figure 6B, lower).
practical constructs, increasing the variety of possible experimental set-ups.

Using this method we generated a chimeric polyprotein containing the previously unstudied Ec POTRA2 domain (Figure 3). We show that EcPOTRA2 is mechanically robust, and that it is a suitable domain for force spectroscopy studies (Figure 4). By comparing the structure of EcPOTRA2 with other α + β proteins (Figure 5) studied using single molecule AFM, we find a similar structural feature, namely inter-connected, terminal, parallel geometry β-strands. For this subset of proteins we identify a clear correlation between their mechanical robustness and brittleness and the number of inter-residue contacts (Figure 6).

METHODS

Polyprotein construction by Gibson assembly. Specific primers were used to amplify six DNA cassettes by the polymerase chain reaction (PCR) that encode individual protein domains as well as the linearized backbone (a modified pET8c vector containing the N-terminal I27 domain (Table 1, Supporting Information). This step added the cassette-specific ends (which encode the amino-acid sequence of the linkers) to enable an ordered assembly reaction (Figure 1). PCR products were purified by gel extraction (Qiagen Gel Extraction Kit, UK). The six purified PCR-amplified DNA cassettes plus the vector backbone were joined to form a circular DNA molecule by Gibson assembly cloning following the manufacturer’s (NEB, MA, USA) instructions, using 100 ng of linearized vector backbone and a 5-fold molar excess of inserts. A summary of the method is shown in Figure 1. The reaction was incubated for 1 hour at 65 °C in a thermost-cycler (PTC-100, MJ Research Inc., MA, USA) and 1 μl of the reaction mix subsequently used to transform 25 μl competent E. coli cells. Plasmid DNA encoding full length assembled constructs was identified by colony PCR using T7 promoter (5′-TAATACGACTCACTATAGGG-3′) and T7 terminator (5′-GCTAGTTATGCTACGCGG-3′) primers and their sequences verified (Beckman Coulter Genomics, Takeley, UK).

Protein expression and purification. The polyprotein-encoding expression vector was transformed into E. coli BLR (DE3) pLysS (Novagen, Nottingham, UK). A 100 mL overnight starter culture (LB medium) was used to inoculate 10 × 1 litre LB medium supplemented with 25 μg mL⁻¹ chloramphenicol and 100 μg mL⁻¹ carbenicillin to obtain a starting optical density at 600 nm (OD₆₀₀) of 0.05. Culture incubation was carried out at 37 °C in an incubator shaker. At an OD₆₀₀ = 0.5 – 0.6 protein expression was induced with 1 mM (final concentration) isopropyl β-D-thiogalactopyranoside (IPTG). Three hours later the cells were harvested by centrifugation. The cell pellet was resuspended in lysis buffer (20 mM Tris, 300 mM NaCl, 0.025 % (v/v) sodium azide, 0.15 % (v/v) triton X-100, 5 mM imidazole, pH 8), and the cells were mechanically disrupted under high flow pressure (TS series Cell Disrupter, Constant Systems Ltd, Warwick, UK). The hexahistidine-tagged polyprotein was isolated from the cleared lysate using a histidine-binding Ni-NTA affinity chromatography resin (Ni sepharose HP, GE Healthcare, Sweden). The eluted protein was dialyzed into distilled deionized water and freeze-dried. The protein was then purified to homogeneity by size-exclusion chromatography. 5 mL of protein resuspended in 63 mM sodium phosphate, pH 7.4 was separated by application to a 320 mL HiLoad 26/60 Superdex 75 column (GE Healthcare, Sweden). Captured fractions containing purified polyprotein were pooled and dialyzed against distilled deionized water before being aliquoted and stored at -20 °C.

Single-molecule force spectroscopy. Single molecule force spectroscopy experiments were completed for the chimeric polyproteins (I27-TmCsp)-I27GA and (I27-EcPOTRA2)-I27 using a method described previously[31]. Briefly, a custom-built AFM was mounted with silicon nitride cantilevers (MLCT, Veeco (Santa Barbara, CA)). The spring constant of the cantilever was found in buffer by applying the equipartition theorem[23] and was typically found to be 35 ± 3 pN nm⁻¹. A freeze-dried protein sample (0.1 mg) was reconstituted to 0.5 mg mL⁻¹ in sterile sodium phosphate buffer (63 mM, pH 7.4) and centrifuged for 5 mins at 14,500 x g (Espresso Personal Microcentrifuge, Thermo Scientific, UK). 40 μL of the protein solution was applied onto a coverslip with a freshly stripped gold surface that resulted in the immobilisation of polyproteins via covalent attachment between the sulphuryl groups of two cysteine residues at the C-terminus of the polyprotein and the gold surface. After incubation for 15 minutes, the surface was rinsed with buffer. Mechanical unfolding experiments were performed at pulling velocities of 100, 200, 600 and 2000 nms⁻¹ at room temperature (23 °C) over a distance of 400 nm. Three datasets, each containing at least 28 unfolding events for the domain under study (TmCsp or EcPOTRA2) were accumulated at each pulling velocity using a fresh sample and a new cantilever for each dataset.

Analysis of single-molecule force-extension data. Data from single-molecule force-extension experiments for the chimeric polyproteins (I27-TmCsp)-I27GA and (I27-EcPOTRA2)-I27 were filtered to only include traces that show the unfolding of a single polyprotein chain. This is characterized by displaying seven or fewer unfolding events. The spectra were subsequently analysed using in-house software written for IGOR Pro (Version 6, Wavemetrics, Lake Oswego, OR). Initially, a model-free analysis was carried out. Here, the peak unfolding force of each protein unfolding event, and the interpeak distance between unfolding events (xₚ₋ₚ), defined as the distance between one peak and the same force value on the following saw-tooth curve, were recorded. The unfolding forces and interpeak distances were binned and plotted in histograms, allowing Gaussian plots of the data to be obtained. To obtain an estimate of the number of amino acids that are released from a compact, globular native state to an extended unfolded state during each unfolding event, the data were analysed by fitting a worm-like-chain (WLC) model for polymer elasticity to the rising edge of each saw-tooth, as described previously[32]. The WLC model is given by:

\[
F(x) = \frac{k_B T}{p} \left( \frac{1}{4} - \frac{x^2}{L_c^2} - \frac{x}{4} + \frac{x}{L_c} \right)
\]

where F(x) is the force as a function of extension, x; k₋ is Boltzmann’s constant, p is the persistence length and L_c is the contour length.

Monte Carlo (MC) Simulations MC simulations were performed using a two-state model for unfolding, as described previously[33]. This technique is based on the assumption that
each protein unfolds via a two-state all-or-none process governed by a rate constant, $k_U$, and the distance from the native state to the transition state along the measured reaction coordinate, $\Delta x_U$. The simulations were used to generate histograms of unfolding forces for protein domains at a particular pulling velocity and compared to those generated experimentally. The pair of $k_U$ and $\Delta x_U$ values that provided the best global fit to the experimental data over all pulling velocities was obtained. The uncertainty in the experimental data, defined as the standard error in the straight line fit to the dependence of $F_U$ on the pulling speed, was used to quantify the uncertainty in $k_U$ and $\Delta x_U$. The range of $k_U$ and $\Delta x_U$ values that provided a fit to the data within the experimental uncertainty gave the value of the uncertainty for each parameter.

ASSOCIATED CONTENT

Supporting Information. Further details of primers used for the Gibson Assembly method as well as the single molecule protein unfolding experiments can be found online, including unfolding force histograms and tables with number of events for each experiment and a comparison of the pulling speed dependence of the I27 protein in the three different polyprotein constructs. This material is available free of charge via the Internet at http://pubs.acs.org.

AUTHOR INFORMATION

I These authors contributed equally to the work

Corresponding Author

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