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OPEN Natural variations in the biofilmassociated protein BsIA from the genus Bacillus

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BsIA is a protein secreted by Bacillus subtilis which forms a hydrophobic film that coats the biofilm surface and renders it water-repellent. We have characterised three orthologues of BsIA from Bacillus amyloliquefaciens, Bacillus licheniformis and Bacillus pumilus as well as a paralogue from B. subtilis called YweA. We find that the three orthologous proteins can substitute for BsIA in B. subtilis and confer a degree of protection, whereas YweA cannot. The degree to which the proteins functionally substitute for native BsIA correlates with their in vitro biophysical properties. Our results demonstrate the use of naturally-evolved variants to provide a framework for teasing out the molecular basis of interfacial selfassembly.

Many bacterial species live in multi-cellular communities known as biofilms^{1, 2}. The Gram-positive bacterium Bacillus subtilis forms biofilms with a characteristic wrinkled morphology and a surface that is highly hydrophobic. A critical component in generating the hydrophobicity of the biofilm is the amphiphilic protein BslA³. The structure of BslA contains a hydrophobic, surface-exposed 'cap' that sits atop a hydrophilic immunoglobulin-like domain, and is reminiscent of the fungal hydrophobins³⁻⁶. We recently characterised the mechanism by which BslA remains both stable and monomeric in aqueous environments and how its adsorbs to an interface⁷. We showed that the hydrophobic cap of BsIA, in contrast to the hydrophobins, is structurally plastic; it undergoes an environmentally-responsive conformational change upon partitioning from an aqueous to a hydrophobic environment. Once at an interface, BslA self-assembles into an ordered rectangular 2D lattice that macroscopically forms an elastic protein film^{3, 7, 8}. Like the hydrophobins, BslA has been recognized as a 'natural' surface active agent that could find a range of applications in the food and personal care industries e.g. as surface modifier, coating agent, emulsifier, and/or foam stabiliser9-13.

Although the gross structural changes that lead to the surface-stabilising function of BsIA are understood, we wished to tease out the mechanism by which BslA interacts with a hydrophilic-hydrophobic interface, i.e. what triggers the structural change between the conformational forms? Moreover, what interactions are critical for the formation of the regular 2D lattice? To address these questions, we have studied several BsIA orthologues that are present in the genome of Bacillus species closely related to B. subtilis. In addition, we have investigated a paralogue of BsIA called YweA. Using genetic modification and biophysical techniques, we characterize the behaviour of these variants both *in vivo* and *in vitro*. We find that key serine residues in the cap regions of the variants act as 'switches' that drive conformational re-arrangement at an interface. We propose a classification system for BsIA and its variants based upon the behaviour of the protein films under compression. This categorization allows us to tease out the interface-protein and protein-protein interactions and suggest amino acids important for the formation of stable, highly organised 2D films.

Results

Protein Alignment & Choice of BslA Variants. As previously reported by Kobayashi & Iwano, a number of Bacillus species possess well-conserved genes that encode for homologous proteins that fall into two groups: BslA-like and YweA-like⁸. Figure 1 shows a phylogenetic tree generated via maximum likelihood analysis of the

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Figure 1. Maximum likelihood phylogeny of BslA and YweA protein sequences from *Bacillus* species. Abbreviations used are as follows: B_teq, Bacillus tequilensis; B_sub, Bacillus subtilis; B_pum, Bacillus pumilis; B_amy, Bacillus amyloliquefaciens; B_lic, Bacillus licheniformis; B_son, Bacillus sonorensis; B_moj, Bacillus mojavensis; B_vall, Bacillus vallismortis; B_atro, Bacillus atrophaeus; B_siam, Bacillus siamensis. The tree was rooted using the midpoint method. The bootstrap value shown in red indicates the support for each branch (from 0 to 1). YweA sequences for *Bacillus siamensis* and *Bacillus pumilus* could not be found using BlastP. Sequences highlighted in blue were used in this study.

two paralogues YweA and BslA from various *Bacillus* species. The paralogue YweA shares 67% sequence similarity with BslA and is distinguished from BslA by its lack of an N-terminal domain after the signal sequence in addition to the absence of 10 amino acids at the C-terminus that contain a conserved 'CxC' motif (Fig. 2A). In this work we have chosen to study the BslA orthologues produced by three species, *Bacillus amyloliquefaciens* (Ba_BslA), *Bacillus licheniformis* (Bl_BslA), and *Bacillus pumilus* (Bp_BslA), as well as the BslA paralogue YweA from *B. subtilis*. These particular *Bacillus* species were chosen as representatives of each distinct branch of the maximum likelihood tree (Fig. 1). The amino acid sequences of these proteins share a high degree of similiarity to *B. subtilis* BslA (Bs_BslA), while still possessing variations that may influence their behaviour both *in vivo* and *in vitro* (Fig. 2A; see Fig. S1 for the amino acid alignments for all species in the maximum likelihood tree).

In vivo characterization of BsIA variants. First, we tested whether the orthologous genes, and therefore proteins, could substitute for Bs_BslA in vivo in B. subtilis. This was achieved using heterologous expression of complemented orthologous bslA genes (encoding for Ba_BslA, Bl_BslA, and Bp_BslA) in a mutant of B. subtilis possessing a bslA gene deletion ($bslA^-$). Resultant biofilm formation was assessed by four criteria: complex colony morphology, pellicle formation, colony surface hydrophobicity, and sporulation ability. Wild-type B. subtilis strain NCIB3610 produces characteristic biofilm and pellicle morphologies with a pronounced surface hydrophobicity, reflected in the high contact angle a water droplet makes with respect to the biofilm surface (Fig. 2D,E left column). The biofilms heterologously expressing Ba_BslA showed the greatest similarity to wild-type or Bs_BslA complemented biofilms, with a characteristic wrinkled morphology and pronounced surface hydrophobicity. The B. subtilis biofilms substituted with Bl_BslA were also highly hydrophobic and wrinkled, however, the wrinkles in the central 'disc' of the colony were less pronounced and the extent of wrinkling in the pellicle was reduced. For Bp_BslA, the colony surface showed marked hydrophobicity, however colony growth appeared diminished relative to NCIB3610 and the bslA⁻ mutant genetically complemented with Bs BslA. Moreover, like the bslA⁻ mutant genetically complemented with Bl_BslA, the wrinkles in the central 'disc' of the colony were less pronounced. Finally, attempts to genetically complement the bslA⁻ mutant with the yweA coding region did not reinstate either the characteristic wrinkled morphology or surface hydrophobicity of biofilms, consistent with YweA having only a minor role in the biofilm architecture (Fig. 3 and ref. 8).

The ability to sporulate is a sign of a mature biofilm matrix environment¹⁴. In *B. subtilis*, the *sspB* gene encodes for a small acid-soluble spore protein and expression is used as an indication of biofilm maturity. We utilized flow cytometry in conjunction with a P_{*sspB*}·yfp transcriptional reporter fusion to assess the degree of sporulation within each of the heterologously complemented biofilms. All heterologously expressed BsIA orthologues were found to confer high levels of sporulation relative to a negative control, albeit not to the degree exhibited by the wild-type strain (Table 1). Taken together, these data indicate that the orthologous proteins Ba_BsIA, Bl_BsIA, and Bp_BsIA can at least partially fulfil the role of Bs_BsIA within the *B. subtilis* biofilm, largely reinstating surface hydrophobicity and gross morphological characteristics. It is notable, however, that Bl_BsIA and, especially, Bp_BsIA do exhibit some differences in biofilm morphology. YweA, in contrast, does not recover wild-type biofilm morphology or hydrophobicity (Figs 2 and 3). Nonetheless, YweA does appear to contribute to the surface hydrophobicity, exacerbating the impact of the *bsIA* deletion on biofilm formation (Fig. 3).

Biophysical assessment of protein behaviour *in vitro*. Having studied the BsIA variants *in vivo*, we next characterised their behaviour in isolation using *in vitro* biophysical techniques. BsIA variant proteins were expressed using standard techniques (see Experimental Procedures). The resulting mature regions of the purified



Figure 2. Assessing biofilm formation of heterologously expressed bslA variants in B. subtilis. (A) Amino acid alignment of BsIA variants found in other Bacillus species. Abbreviations used are as follows: B sub, Bacillus subtilis; B_pum, Bacillus pumilis; B_amy, Bacillus amyloliquefaciens; B_lic, Bacillus licheniformis. Underlined and bolded amino acids signify the signal sequence, black represents 100% sequence identity. Blue amino acids represent the hydrophobic cap regions, where dark blue are conserved amino acids and light blue are conservative substitutions. The purple highlighted amino acids are glycine which are conserved across all species and always follow the amino acids comprising the caps. Aspartic acid is shown in red and serine residues within caps 1 and 2 are shown in green. Note that YweA is differentiated from Bs BslA and the BslA orthologues by the fact it lacks both the N-terminal region following the signalling sequence and the C-terminal domain. The * symbols indicate the cap regions containing serine residues. Biofilms and pellicles were formed from the wild type strain (NCIB3610), a strain possessing a B. subtilis bslA gene deletion mutant (bslA⁻), strains that heterologously express the bslA gene of each species studied (B. subtilis, B_sub_bslA; B. amyloliquefaciens, B_sub_amy; B. licheniformis, B_lich_bslA; B. pumilus, B_amy_bslA) complemented into the bslA⁻ mutant strain, and a strain expressing the yweA gene complemented into the bslA⁻ mutant. Biofilm phenotypes were characterized by assessing (B) complex colony morphology; (C) pellicle formation; (D,E) colony surface hydrophobicity (see Table S3 for details on strains).

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proteins were first analyzed using size exclusion chromatography and SDS-PAGE (Fig. 4). As demonstrated in previous work³, purified Bs_BslA is primarily composed of monomers and dimers, with a small sub-population of higher order oligomeric species (Fig. 4A,F). The orthologous BslA proteins showed similar monomer/dimer fractions as Bs-BslA, but higher order oligomers were absent. YweA, in contrast, is almost entirely monomeric (Fig. 4E).

Kinetics of interfacial self-assembly of BslA variants. After protein purification, we assessed the interfacial activity of the BslA variants using pendant drop tensiometry. The dynamics of interfacial protein adsorption can often be characterized by three kinetic regimes: Regime I is a 'lag time' where there is no apparent change in the interfacial tension, Regime II is when a sufficient proportion of protein adsorbs to the interface to produce a decrease in interfacial tension, and Regime III is reached when the interfacial tension plateaus to a roughly constant final value¹⁵. Importantly, it should be stressed that once an elastic film has formed there is no longer an interfacial tension between the vapour and liquid phases and the concept of an 'interfacial tension' no longer applies. A good indication of when a film has formed is an increase in the error in the fit of the Young-Laplace equation to the drop shape¹⁶. Bromley *et al.* showed that the only sensible data that can be extracted from interfacial tension measurements of Bs_BslA was the time at which the interfacial tension transitions between Regime I and Regime II (the 'Regime I time'), since at any time after this transition an elastic film is present on the droplet



Figure 3. Characterization of biofilm phenotypes of *yweA* mutants. Biofilm phenotypes were characterized by assessing (**A**) complex colony morphology; (**B**) pellicle formation; (**C**,**D**) colony surface hydrophobicity for *B. subtilis* wild-type strain 3610, a *bslA* mutant, a *yweA* mutant, and a double mutant *bslA yweA* (see Table S3). Row C shows a representative sessile drop resting atop a biofilm formed from one of the strains used above. Row D are the average contact angles; ND means that a contact angle could not be determined from the images.

	% Sporulation positive ^a	
Strain	0 µM IPTG	$25\mu\mathrm{M}\mathrm{IPTG}$
3610 PsspB-yfp	26.6	24.8 ± 1.63
bslA PsspB-yfp	5.9 ± 0.6	4.9 ± 0.8
bslA P _{IPTG} -bslA _{B.sub} PsspB-yfp	3.2 ± 0.2	14.1 ± 2.3
bslA P _{IPTG} -bslA _{B.lic} PsspB-yfp	5.6 ± 0.3	15.6 ± 1.2
bslA P _{IPTG} -bslA _{B.amy} PsspB-yfp	5.6 ± 0.4	13.0 ± 3.7
bslA P _{IPTG} -bslA _{B.pum} PsspB-yfp	5.2 ± 0.2	15.6 ± 1.2

Table 1. Sporulation frequency assessed by flow cytometry. *a*The percentage sporulation was calculated using fluorescence form the *PsspB-ypf* fusion as a proxy after 48 hours incubated at 30 °C. NCIB3610 was used as the non-fluorescent control. For the full genotypes of the strains see Table S3. For each condition at least two independent samples were analysed. Expression of the variant gene was achieved using 25 μ M IPTG. The value presented is the mean and the error is the standard deviation.

surface⁷. Regime I times were determined in one of two ways: (1) the transition time between regimes I and II when the fit error was still low ($<0.4 \,\mu$ m) or (2) when the fit error increased to a threshold value ($>0.75 \,\mu$ m). Criterion (1) or (2) was chosen by whichever occurred first. We found that the Regime I times of the orthologues Ba_BslA and Bp_BslA were within error of Bs_BslA (Fig. 5A). In contrast, the Regime I time of Bl_BslA was nearly twice as long as those of the other orthologues while the Regime I time for YweA was faster than that of Bs_BslA by ~25\%.

Relaxation dynamics of elastic films. The formation of an elastic film at a hydrophobic/hydrophilic interface is demonstrated by the observation of wrinkles following compression of the droplet (achieved by withdrawing a small volume of fluid following equilibration). One way of characterizing these films is by monitoring the formation and subsequent disappearance of these wrinkles over time, reflecting relaxation of the interfacial protein film (Fig. 5B). We have assumed that wrinkle relaxation is due to the loss of protein into the subphase, although rearrangement of the protein at the interface cannot be ruled out. For these experiments an oil/water interface was used as the small difference in density between the two phases results in rounder droplets, allowing for better imaging of the wrinkles that form. As found previously⁷, wrinkles formed under compression in a Bs_BslA film do not relax over the timescale of the experiment. In contrast, Ba_BslA and Bl_BslA exhibit very



Figure 4. Size exclusion chromatography and SDS-PAGE analysis of Monomer-Dimer Formation. The *bslA* variant genes were cloned, overexpressed as a GST fusion and then purified. (**A**) Using size exclusion chromatography Bs_BslA was found to contain two primary peaks corresponding to monomer and dimer species, although there are higher-order additional peaks present. (**B**) Ba_BslA, (**C**) Bl_BslA, and (**D**) Bp_BslA all eluted in primarily two peaks. (**E**) YweA, in contrast, eluted in effectively one monomer peak. (**F**) SDS-PAGE analysis of recombinant protein without β -mercaptoethanol and (**G**) with β -mercaptoethanol.

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slow relaxation, and Bp_BslA relaxes within a minute. Strikingly, relaxation of YweA is extremely rapid, occurring immediately upon compression.

Film organisation of BslA variants. Bs_BslA forms highly ordered 2D rectangular lattices at an air/water interface⁷. We investigated the structure of the films formed by BslA variants using transmission electron microscopy (TEM). Protein lattices were observed for all variants, but exhibited varying degrees of order. The films formed by Ba_BslA showed domains of ordered protein, but these were not widespread (Fig. 5C). As shown by the Fast Fourier Transform (FFT), weak peaks can be observed, indicating some level of ordering (Fig. 5C). The Bl_BslA films exhibited larger domains of ordered protein, and the FFT shows clear peaks (Fig. 5D). In contrast, Bp_BslA showed very weak ordering with only very small patches of organized structure, similar to those previously observed for the BslA L77K mutant³ (Fig. 5E). Thus amongst BslA films are highly ordered and wrinkles formed under compression do not relax, Ba_BslA and Bl_BslA show some degree of ordering and display slow wrinkle relaxation dynamics, whilst Bp_BslA films are not well ordered and wrinkles formed upon compression relax rapidly.

The paralogue YweA, however, does not conform to the trend observed for BslA orthologues. Indeed, YweA formed clearly defined and well ordered films similar to those formed by Bs_BslA, despite its rapid relaxation (Fig. 5B,F), calling into question the relationship between the microscopic organization of the protein within a film and film robustness.

CD spectroscopy of solution and interfacial states. Finally, circular dichroism (CD) spectroscopy was used to study the conformation of the BslA variants in aqueous solution and at an oil/water interface (Fig. 5G). Qualitatively, the spectra of the different BslA variants in aqueous solution are similar, with only Bp_BslA showing the loss of a feature between 210 and 218 nm (Fig. 5G). When bound to refractive indexed matched emulsions (RIMEs; Fig. 5H) all of the proteins underwent a structural transition consistent with an increase in β -sheet structure, reflecting rearrangement of the hydrophobic cap upon insertion into the oil phase.

Effects of BI_BsIA cap mutations on the properties of *in vitro* **protein films and** *in vivo* **bio-film morphology.** Pendant drop tensiometry showed that it takes BI_BsIA approximately twice as long as the other orthologues to become adsorbed at an interface (Fig. 5A). We have previously interpreted this slower-than-diffusion adsorption time in relation to the energy barrier between solution and interfacial conformations of BsIA⁷, implying that there is a larger energy barrier that must be overcome for BI_BsIA adsorption. Comparison of the amino acid sequence of Bs_BsIA with that of BI_BsIA reveals two amino acid differences in the hydrophobic cap region (Fig. 2A). Specifically, the first difference resides in 'Cap 1' (residues L77, S78, and L79 of Bs_BsIA) in which serine-78 in Bs_BsIA is replaced by an alanine (A76 in BI_BsIA). The second difference is in 'Cap 3' (residues L153, S154, and I155 of Bs_BsIA) in which S154 is replaced by a glycine (G152 in BI_BsIA). In contrast, these two serine residues are conserved in Ba_BsIA and Bp_BsIA, which have similar adsorption







wavelength (nm)

kinetics to the Bs_BslA. To investigate the role of these residues in the kinetics of adsorption and the properties of the resultant film, we produced two single amino acid mutants of Bl_BslA (A76S and G152S) and the double mutant A76S/G152S in which the cap region is identical to that of Bs_BslA. If these cap regions, and their re-orientation at an interface, determine the kinetics of adsorption, one would predict that the Regime I time of Bl_BslA(A76S/G152S) would coincide with those of Bs_BslA and the other orthologues. This is in fact what is observed as shown in Fig. 6B. In contrast, both single mutants result in Regime I times that are shorter than those



Figure 6. *In vitro* and *in vivo* effects of the Bl_BslA cap mutations. (A) Wrinkle relaxation measurements for Bl_BslA (black squares), Bl_BslA(A76S) (red circles), Bl_BslA(G152S) (blue triangles), Bl_BslA(A76S G152S) (upside down pink triangles). Error bars represent standard deviation in relaxation times for at least N = 10 separate wrinkles across the drop. (B) Regime I times of Bs_BslA (repeated from Fig. 3 for clarity) and the Bl_BslA single and double mutants. (C) Heterologous expression of Bl_BslA variants shows no deleterious effects on biofilm morphology or hydrophobicity relative to both wild-type Bs_BslA (as determined using the parental NCIB3610 strain (3610)) and parental Bl_BslA constructs.

of Bs_BslA and Bl_BslA(A76S/G152S) but very similar to that of YweA. Interestingly, the 'Cap 3' region of YweA possesses the same $S \rightarrow G$ substitution as Bl_BslA (Fig. 2A). The fact that the single mutants are more 'YweA-like' in terms of adsorption indicates that residues 76 and 152 of Bl_BslA help shape the energetic landscape of cap re-organization at an interface. To further explore this hypothesis, we also studied the relaxation of the films formed by these mutant proteins (Fig. 6A). Notably, wrinkles formed under compression of a Bl_BslA(A76S/G152S) droplet did not relax over the time scale of the experiment, unlike Bl_BslA which relaxes slowly, lending further support to a role for these cap residues in determining the stability and mechanical properties of the film. Consistent with this idea, the single cap mutants more closely recapitulate YweA-like behaviour, with films formed by these proteins exhibiting more rapid relaxation than both Bl_BslA and Bl_BslA(A76S/G152S) films. Indeed, films formed by Bl_BslA(A76S), the amino acid composition of which most closely resembles that of YweA, demonstrates the fastest relaxation rate of the cap mutants investigated. Intriguingly, the relaxation of wrinkles formed by this mutant appears to plateau to a non-zero value over the course of the experiment, indicating that wrinkles persist. We conclude from these results that the serine/glycine residue found in Cap 3 of BslA variants plays a prominent role in both the energetics of adsorption and the robustness of the resultant film.

Finally, we tested how the Bl_BslA cap mutations affect the structure and properties of the *B. subtilis* biofilm. When the genes encoding the Bl_BslA cap variants are heterologously expressed, the resultant biofilm morphology and hydrophobicity are indistinguishable from both wild-type Bs_BslA (as determined using the parental NCIB3610 strain) and parental Bl_BslA constructs (Fig. 6C). Thus, while the *in vitro* films produced from these mutants show distinctive physical characteristics, we do not observe any differences in *in vivo* phenotype.

Discussion

While the structural metamorphosis of the hydrophobic cap has been identified as the mechanism that allows BslA to be stable in both aqueous and hydrophobic environments, many questions remain regarding how BslA self-assembles into 2D ordered lattices and what the key amino acids are that contribute to network formation at interfaces. We have previously demonstrated that single amino acid mutations can have marked consequences for both the kinetics and energetics of adsorption along with the spatial ordering of the protein at the interface^{7, 17}. Instead of using a brute-force site-directed mutagenesis approach to further illuminate the biophysical properties of BslA at an interface, we have investigated the BslA variants found in three other *Bacillus* species, along with the *B. subtilis* BslA paralogue YweA. The aim of this strategy is to lead to more efficient targeting of important amino acids for future systematic mutagenesis studies of Bs_BslA. Such knowledge could be exploited to tailor both the kinetics of interfacial adsorption as well as the mechanical properties of the protein film for the application of BslA in multiphase formulations.

We have confirmed that YweA plays no obvious role in overall biofilm formation and morphology *in vivo*, as was demonstrated in previous studies⁸. However, the phenotype of the double deletion mutant *bslA yweA*

indicates that YweA is indeed present within the biofilm and makes a small but noticeable contribution to surface hydrophobicity⁸ (Fig. 3). The major difference between YweA and the other BslA variants is the lack of both the N- and C-terminal domains. Since YweA cannot recover the characteristics of the wild-type biofilm phenotype, it is possible that the lack of one or both of these domains is functionally important for biofilm formation, in either mediating interactions between other BslA proteins, or interactions with other protein or polysaccharide components of the biofilm matrix.

Importantly, the biophysical characterization of YweA interfacial activity and self-assembly provides new insights into the relationship between the microscopic organization of the protein at the interface and resulting macroscopic properties of the surface layer. YweA interfacial films showed extremely rapid relaxation under applied compression (Fig. 5B), yet YweA films imaged by TEM showed the greatest order of all the variants (Fig. 5F). In previous studies, the cap mutant Bs_BslA(L77K) also formed films that relaxed within very short time scales. However, this variant demonstrated significant 2D disorder^{3,7}, and we suggested that the single mutation in the cap disrupted the ability of Bs_BslA(L77K) to form a space-spanning network by altering the orientation of the protein at an interface¹⁷. The behaviour of YweA runs counter to this hypothesis, indicating that film order does not necessarily correlate with film robustness. It can be concluded, then, that YweA has the ability to form interfacial protein-protein lateral interactions but that these are weak, particularly compared to those formed by Bs_BslA.

As Bl_BslA differed markedly in its adsorption kinetics relative to the other BslA variants investigated here, this orthologue formed the focus of further analyses. The amino acids of Bl_BslA differ from Bs_BslA and the other variants by two non-conservative substitutions in Cap 1 ($S \rightarrow A$) and Cap 3 ($S \rightarrow G$). Examination of the interfacial conformation of BslA reveals the two serines to be immediately adjacent to one another, both orientated towards the interior of protein (Fig. S1), and are likely to form a sidechain-sidechain hydrogen bond. In contrast, when the protein is in aqueous solution and the cap region is disordered, these serines are oriented outward towards the aqueous environment. The switch from an outward to an inward orientation would facilitate the conformational switch of the cap region upon contact with a hydrophobic interface, with hydrogen bonding between the residues further stabilising the interfacial conformation.

The Bl_BslA cap is more hydrophobic than Bs_BslA and the other orthologues - the polar serines are replaced by nonpolar amino acids (A76 and G152). Counter-intuitively, the higher hydrophobicity of the cap region overall does not result in faster adsorption (Fig. 5A). The resolution of this apparent paradox resides in the fact that the cap is a flexible, plastic structure. In an aqueous environment, the more hydrophobic residues of the Bl_BslA cap will most likely be oriented inwards, away from the aqueous phase, whereas in Bs_BslA the equivalent hydrophilic side chains are oriented outwards. We propose that the driving force for cap rearrangement in Bs_BslA, is reorientation of these serines away from the hydrophobic interface and that this mechanism is lost in Bl_BslA, resulting in slower adsorption. Thus, we hypothesize that it is the hydrophilic residues within the cap that act as a 'sensor' for hydrophobic interfaces. Interaction with an apolar interface induces a reorientation of the hydrophilic amino acid(s), which in turn triggers the larger scale reorganization of the cap such that the hydrophobic residues become surface-exposed and subsequently restructure to form a three-stranded β -sheet.

Interestingly, the single cap mutants of Bl_BslA show shorter adsorption times, very similar to the Regime I time of YweA (Figs 5A and 6B). YweA has a similar amino acid cap content as the Bl_BslA(A76S) mutant, with a serine in Cap 1 and a glycine in Cap 3. For Bl_BslA(G152S), there is a hydrophobic residue in Cap 1 and a hydrophilic residue in Cap 3. We hypothesize that this combination of a hydrophilic residue in one cap strand, and a hydrophobic residue in the other, may cause a tension within the aqueous conformation of the cap as a whole or, alternatively, facilitate a rapid interconversion between states. This tension or interconversion then primes the cap to more easily undergo the conformational change when it encounters an interface, effectively lowering the energy barrier to re-structuring. Moreover, when in the interfacial conformation, the remaining serine will be unfavourably partitioned into the hydrophobic protein core, but without a second serine present with which to form a hydrogen bond. This may explain why the proteins are displaced from the interface under compression.

Taking the result of this work together, we propose a categorization of BsIA and its variants based upon their wrinkle relaxation behaviour:

- Surfactant-like: The surfactant-like proteins comprise YweA, Bp_BslA and the Bs_BslA mutants L77K and L79K³. Whilst YweA differs from the other three BslA variants in its ability to form ordered lattices at an interface, wrinkles formed under compression of all four proteins relax within very short time scales. This implies that the interfacial lateral protein-protein interactions either cannot form correctly due to incorrect orientation at the interface (in the case of L77K and L79K¹⁷) or are weak due to substitution of critical amino acids.
- **Transiently wrinkling film formers**: The proteins that belong to this category are Ba_BslA, Bl_BslA, Bl_BslA(A76S), and Bl_BslA(G152S). In these cases, wrinkles persist for longer time scales than those formed by the surfactant-like proteins, but do eventually relax over time. Such behaviour suggests relatively weak lateral interactions between proteins at the interface, but stronger than those exhibited by the surfactant-like proteins.
- Rigid film formers: Bs_BslA and Bl_BslA(A76S + G152S) are members of this group. The stresses imposed
 via our methods are insufficient to dislodge protein from the interface, and moreover the entire droplet shape
 distorts under compression. This implies that there exist not only strong protein-interface interactions but
 well established protein-protein interactions within the film.

Categorizing the BslA variants in this manner allows us to identify links between the *in vivo* morphology of *Bacillus* biofilms (Figs 3 and 4) and the interactions the proteins experience at an interface. Thus the extent to which each BslA variant rescues the wild type *B. subtilis* biofilm may be ranked and mapped to the classification system outlined above (Table 2). Bs_BslA produces rigid, elastic films *in vitro* and creates the highly wrinkled

Rank	Protein	Film category
1	Bs_BslA	Rigid film
2	Ba_BslA	Transient wrinkling film
3	Bl_BslA	Transient wrinkling film
4	Bp_BslA	Surfactant-like
5	YweA	Surfactant-like

Table 2. Ranking of biofilm phenotype vs in vitro film classification.

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and hydrophobic biofilm phenotype associated with *B. subtilis* pellicles and colonies. Ba_BslA and Bl_BslA share similar film characteristics *in vitro* and in both cases the biofilm morphology is similar to the *B. subtilis* wild-type phenotype with some small, but discernible differences. Bp_BslA falls within the surfactant-like class and is the least effective of the orthologues at recovering the wild-type *B. subtilis* biofilm morphology. Finally, the paralogue YweA is the most surfactant-like of the proteins investigated and clearly cannot recover any features of the biofilm phenotype. Relating the biofilm morphology to the classes of film formed *in vitro* shows a clear correlation between film robustness and biofilm structure.

This classification provides useful insights into which residues are important for protein-protein interactions within the film. A large proportion of outward-facing, non-cap residues are conserved across all four variants. However, amino acids that are conserved between Bl_BslA and Bs_BslA but not between these two proteins and Ba_BslA and Bp_BslA are likely to affect the strength of the interaction between the proteins in the film. This suggests roles for S49, T105, Q107, N111 and E128 in the formation of the optimal lateral interactions that give rise to stable films (Fig. 7). Specifically, comparing the amino acid sequences of each orthologue reveals that Bp_BslA possesses non-conservative substitutions in four out of these five positions, Ba_BslA has one non-conservative substitution, and Bl_BslA is identical to Bs_BslA. This correlates well with our observations of interfacial protein lattice structure (Fig. 5C-F). The subtle difference in lattice structure between Bl_BslA and Bs_BslA, despite the conservation of these putative lateral interaction sites, we hypothesize originates from the key serine residues (A76S, G152S) identified in this work. These residues may influence how the protein is tilted at an interface, which may in turn influence the extent and strength of lateral interactions between adjacent proteins. We have previously modeled how changes in the cap region influence average protein tilt¹⁷.

Further *in vivo* investigation will be required to determine whether the biophysical differences measured as part of this study have evolved specifically to generate a fitness benefit in the distinct environmental habitat or niche occupied by each species of bacteria.

Experimental Procedures

Bacterial strains and growth conditions. Escherichia coli and Bacillus subtilis strains used and constructed in this study are detailed in Table S1. All strains were routinely grown in Lysogeny broth (LB; 10 g NaCl, 5 g yeast extract and 10 g tryptone per litre) or on LB solidified with 1.5% (w/v) select agar (Invitrogen) at 37 °C. *B. subtilis* biofilms were grown in MSgg medium (5 mM potassium phosphate and 100 mM MOPS at pH 7.0 supplemented with 2 mM MgCl₂, 700 μ M CaCl₂, 50 μ M MnCl₂, 50 μ M FeCl₃, 1 μ M ZnCl₂, 2 μ M thiamine, 0.5% (v/v) glycerol, and 0.5% (v/v) glutamate). When required, antibiotics were used at the following concentrations: 100 μ g ml⁻¹ ampicillin, 100 μ g ml⁻¹ spectinomycin, 25 μ g ml⁻¹ kanamycin, 1 μ g ml⁻¹ erythromycin and 25 μ g ml⁻¹ lincomycin. Ectopic gene expression was induced with 25 μ M isopropyl β -D-1 thiogalactopyranoside (IPTG). *E. coli* strain MC1061 [*F*¹acIQ lacZM15 Tn10 (tet)] was used for the routine construction and maintenance of plasmids. *B. subtilis* 168 derivatives were generated by transformation of competent cells with plasmids using standard protocols¹⁸. SPP1 phage transductions, for the introduction of DNA into *B. subtilis* strain NCIB3610, were performed as previously described¹⁹.

Plasmid construction. The primers and plasmids used in this study are presented in Tables S2 and S3 respectively. Single amino acid substitutions were generated by PCR site-directed mutagenesis using KOD Hot Start DNA Polymerase (Novagen) and the appropriate primer pairs from Table S2, with reaction conditions calculated according to the Stratagene manual for QuikChange Site-Directed Mutagenesis. The resulting mutant plasmids were digested with DpnI and transformed into competent *E. coli* MC1061 cells. All mutations were verified by DNA sequencing. For expression of BslA orthologues in *B. subtilis* under control of the IPTG-inducible Phyper-spank promoter, the *bslA* gene was amplified from genomic DNA isolated from *Bacillus licheniformis* DSM13; *Bacillus amyloliquefaciens* FZB42; and *Bacillus pumilus* SAFR-032 using the primer pairs NSW812/NSW813; NSW829/NSW830; and NSW819/NSW820 respectively. Resulting PCR products were digested with HindIII and SphI, and ligated into the *B. subtilis* shuttle vector, pDR111 (also digested HindIII/SphI), to enable subsequent gene integration into the non-essential *amyE* locus of the *B. subtilis* chromosome (final pDR111-derived plasmids are listed in Table S3). Expression of *yweA* was achieved in a similar manner. The *yweA* gene was amplified from *B. subtilis* NCIB3610 genomic DNA using primers NSW810 and NSW811, digested HindIII/SphI and ligated into HindIII/SphI-digested pDR111.

Plasmid pNW1420, for over-expression of YweA₃₁₋₁₅₅ in *E. coli*, was generated as follows: *yweA*₃₁₋₁₅₅ was amplified from *B. subtilis* NCIB3610 genomic DNA using primers NSW1853 and NSW1854 and ligated into the expression vector pET15bTEV using the NdeI and XhoI restriction sites. The resulting plasmid encodes the N-terminally His₆-tagged fusion protein, His₆-YweA₃₁₋₁₅₅, which contains the recognition site of Tobacco Etch Virus (TEV) protease between His₆ and YweA₃₁₋₁₅₅ coding regions, allowing removal of the His₆ tag by digestion



Figure 7. Possible amino acids that mediate interfacial protein-protein interactions. 3D structure of BsIA highlighting the hydrophobic residues involved in interfacial adsorption (orange) and the residues identified as putatively involved in protein-protein interaction S49, T105, Q107, N111 and E128 (green = S, light green = T, cyan = Q, purple = N, red = E).

with TEV. Plasmids for production of BslA orthologues from *B. amyloliquefaciens*, *B. licheniformis*, and *B. pumilus* encoded truncated proteins corresponding to *B. subtilis* $BslA_{42-181}$ (determined by sequence alignment; see Fig. 2A), on which previous analyses were based^{3, 8}, and were generated as follows. For over-expression of *B. amyloliquefaciens* $BslA_{42-181}$ (referred to herein as Ba_BslA), the appropriate DNA fragment was amplified from *B. amyloliquefaciens* FZB42 genomic DNA using primers NSW2011 and NSW2012, and ligated into the GST expression vector pGEX-6-1P via BamHI and XhoI sites. The resulting plasmid, pNW1422, encodes the N-terminally tagged fusion protein GST-Ba_BslA, from which the GST tag can be removed by TEV protease digestion Plasmid pNW1423 for over-expression of *B. licheniformis* $BslA_{40-179}$ (Bl_BslA) and pNW1424 for over-expression of *B. pumilus* $BslA_{37-177}$ (Bp_BslA), were generated in an identical manner using primer pairs NSW2013/NSW2014 and *B. licheniformis* DSM13 genomic DNA, and NSW2015/NSW2016 and *B. pumilus* SAFR-032 genomic DNA respectively. All resulting constructs were sequence-verified.

Biofilm analysis. Biofilm analysis was performed as previously described, with minor modifications¹⁹. For complex colony formation, $10 \,\mu$ L of the appropriate *Bacillus* culture, grown to mid-exponential phase in LB, was spotted onto MSgg medium solidified with 1.5% (w/v) agar and incubated at 30 °C for 48 hours. For pellicle analysis, the LB starter culture was diluted 1:100 (v/v) in MSgg medium and incubated at 25 °C for 72 hours. Both colony and pellicle biofilms were imaged using a Leica MZ16 stereoscope (Leica Microsystems).

Colony hydrophobicity contact angle measurements. Biofilms were grown on MSgg agar plates (as described above). A section through the mature biofilm was extracted using a scalpel and placed on a microscope slide. A 5μ l droplet of sterile double-distilled water was placed on the colony using the ThetaLite Optical Tensiometer with OneAttension software. The drop was allowed to equilibrate for 5 minutes prior to imaging and contact angle measurement.

Sporulation assays and FACS analysis. Strains harbouring the promoter *PsspB* fused with *yfp* coding sequence were used to assess sporulation levels in mature colonies by flow cytometry²⁰. The experiments used bio-films grown for 2 days at either 30 °C on MSgg agar without IPTG or supplemented with 25 μ M IPTG. After incubation, biofilms were collected, washed and fixed with 4% paraformaldehyde (PFA), as previously described¹⁴. Fluorescence was analysed at the single-cell level using a BD FACSCalibur (BD Biosciences)²¹.

Protein Purification. BslA proteins were purified as previously described⁷, with minor changes as follows. E. coli BL21 (DE3) pLysS cells were transformed with the appropriate plasmids for over-expression of GST-BsIA fusion proteins (described above; Table S3). For protein production, the transformed cells were grown in autoinduction medium²² supplemented with ampicillin (100 μ g mL⁻¹) at 37 °C and 200 rpm until an OD₆₀₀ of 0.9 was reached, at which point protein expression was induced by reducing the temperature to 18 °C for further incubation overnight. Cells were collected by centrifugation at $4000 \times g$ for 45 min and stored at -80 °C until further use. For purification, bacterial pellets were thawed and resuspended in purification buffer (50 mM HEPES pH 7.5, 300 mM NaCl (for YweA), or 50 mM HEPES pH 7.5, 250 mM NaCl (for all other purifications)) supplemented with Complete EDTA-free Proteinase Inhibitors (Roche) before lysis using an Emulsiflex cell disruptor (Avestin). Unlysed cells and cell debris were removed by centrifugation at $27000 \times g$ for 20 min, and the cleared lysates incubated with Glutathione Sepharose 4B resin (GE Healthcare; for BslA) or Ni-nitrilotriacetic acid (Ni-NTA) agarose (Qiagen; for YweA) at a ratio of 750 µL resin per 1 L bacterial culture with gentle rotation for 4 h at 4 °C. To isolate the bound fusion proteins, the mixture of lysate plus beads was passed through a single-use 25-mL gravity flow column (Bio-Rad), and the collected beads washed twice with 20 mL of the appropriate purification buffer. Untagged BslA variants were generated by resuspending the washed beads in 25 mL purification buffer supplemented with 1 mM DTT and 0.5 mg TEV protease prior to gentle rotation overnight at 4 °C. To generate untagged YweA from the His₆-YweA fusion protein, the overnight mixture was additionally supplemented with 250 mM imidazole. Following overnight cleavage, mixtures were again passed through gravity flow columns and the flow-through collected. To the flow-through, $750\,\mu\text{L}$ fresh Glutathione Sepharose plus $250\,\mu\text{L}$ Ni-NTA agarose (for BslA variants), or 100 µL Ni-NTA agarose (for YweA), was added and the solution incubated with gentle rotation overnight at 4 °C to remove the TEV protease and any unbound GST or His₆. The mixtures were passed through the gravity flow columns for a final time, and purified proteins collected in the flow-through. The purified proteins were then concentrated (with simultaneous exchanges into 25 mM phosphate buffer pH 7.0 where appropriate) and further purified by size exclusion chromatography (SEC) as previously described⁷.

SDS-PAGE analysis. SDS-PAGE analysis of BslA variants was performed using 30 μ g purified samples of BslA diluted 4:1 (v/v) in 4X loading buffer (6.2 g SDS, 40 mL 0.5 M Tris pH 6.8, 6.4 mL 0.1 M EDTA, 32 mL glycerol, 1 mg Bromophenol blue) either with or without β -mercaptoethanol at 4% (v/v). Samples to which β -mercaptoethanol was added were boiled at 100 °C for 5 min prior to analysis, whilst samples without β -mercaptoethanol were not subjected to boiling. Proteins were run on a standard 14% polyacrylamide denaturing gel at 200 V for 60 min and visualised by staining with Coomassie Blue.

Mass spectrometry analysis. Purified proteins were identified by excision of protein bands from SDS-PAGE gels and LC-MS-MS following tryptic digest. Protein size estimation was performed using MS-TOF and peptides identified from the MASCOT database. All procedures were performed by FingerPrints Proteomics service, School of Life Sciences, Dundee.

Bioinformatics. BslA orthologues were identified using BLASTP^{23, 24} using the non-reduntant protein sequence database for each of the organisms of interest using the protein sequence of BslA from *B. subtilis* as the query. BslA orthologues were distinguished from YweA based on the presence of a C-terminal region not present in YweA⁸ (see Fig. 2A and Table S4). BslA and YweA protein sequences were aligned using Clustal Omega with the default settings²⁵. The signal sequences were predicted using the SignalP 4.1 server²⁶ and the aligned sequences were manually coloured to highlight the 'cap' regions and identity. A maximum likelihood tree was calculated from the Clustal Omega alignment using phylogeny.fr's standard workflow²⁷, where Gblocks²⁸ was used to eliminate divergent and poorly aligned columns, and a tree computed with PhyML²⁹ using a WAG substitution model³⁰ and 100 bootstrap replicates. The outputted tree was visualised using TreeDyn³¹.

Pendant drop tensiometry & and surface wrinkle relaxation quantification. Pendant drop experiments were performed on a Krüss EasyDrop tensiometer. Protein samples were diluted in buffer and immediately placed in a syringe with a needle diameter of 1.83 mm. Images of the pendant drop are captured by a CCD camera and Krüss software fits the Young-Laplace equation to the drop shape to determine the interfacial tension. For measuring the dynamic interfacial tension, samples were prepared by diluting each protein in phosphate buffer to a concentration of 0.03 mg ml⁻¹. Droplets were expelled in air and the interfacial tension was determined by fitting the droplet shape to the Young-Laplace equation. For the film relaxation experiments, a protein concentration of 0.2 mg ml⁻¹ was used. A droplet of the protein in aqueous solution was expelled into glyceryl trioctanoate oil and allowed to equilibrate at room temperature for 30 minutes. Images were acquired at 2 fps using a digital camera. Wrinkle relaxation experiments were performed following compression of the droplet, which was achieved by retracting 5 μ L of the protein solution. Compression induced the formation of wrinkles in the surface layer. The wrinkles were monitored over a 10 minute period. To analyze the relaxation of the wrinkles, a line profile was drawn across the wrinkles. The line profile was plotted using the greyscale values (from 0 to 255) of each pixel along this line using the ImageJ. At least 10 wrinkles were monitored over the time course of the experiment. To plot the relaxation rate, the greyscale value of the pixels was normalized and background corrected.

Transmission electron microscopy (TEM). Protein samples were deposited onto carbon-coated copper grids (Cu-grid) (TAAB Laboratories Equipment Ltd) and imaged using a Philips/FEI CM120 BioTwin transmission electron microscope. A 5 μ L droplet of protein (0.025 mg mL⁻¹) was pipetted onto a grid and allowed to equilibrate for 5 mins before being wicked with filter paper from the side. A 5 μ L droplet of 2% uranyl acetate was

then applied to the grid and similarly left for 5 min before being wicked from the side. The Fast Fourier Transform (FFT) of the images was performed using ImageJ software.

Circular dichroism analysis. Circular dichroism spectropolarimetry (CD) was performed using a Jasco J-810 spectropolarimeter. Samples were analysed at a concentration of 0.03 mg mL^1 in a 1 cm quartz cuvette. Measurements were performed with a scan rate of 50 nm sec⁻¹, a data pitch of 0.1 nm and a digital integration time of 1 sec. Twenty accumulations were measured and averaged to produce the final curve. Refractive index matched emulsions (RIMEs) were made by first preparing a 20% (v/v) decane emulsion with 0.2 mg ml⁻¹ of protein. The emulsion was mixed for 1 min using a rotor stator (IKA Ultra-Turrax T10) at 30,000 RPM. The emulsion was washed three times in order to remove any residual protein not adsorbed to an oil/water interface. Washes were performed by centrifuging at 1000 rpm for 20 sec, a portion of subphase was removed and replaced with buffer, then gently re-dispersed. Finally, subphase was removed and replaced with glycerol such that the final wt% of glycerol was 59% (w/v). The emulsion was then gently remixed on a rollerbank and then allowed to cream. The cream was placed in a 1 mm pathlength quartz cuvette for spectrum measurement. To prevent creaming during the experiment, the cuvette was briefly inverted between measurements to re-disperse the droplets.

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Author Contributions

R.J.M. designed research, performed experiments & data analysis, wrote & edited manuscript; M.S. performed experiments & data analysis, edited manuscript; R.M.C.G. performed experiments & data analysis, edited manuscript; A.S.F. provided experimental materials, performed experiments & data analysis; L.B. performed experiments & data analysis; C.E. performed experiments & data analysis; A.O. performed experiments; L.H. performed experiments; K.M.B. performed experiments & data analysis; T.S. provided experimental materials; S.A. performed experiments; N.S.W. designed research, wrote & edited manuscript; C.E.M. designed research, wrote & edited manuscript, C.E.M

Additional Information

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