PspF-binding domain PspA₁₋₁₄₄ and the PspA-F complex: New insights into the coiled–coil-dependent regulation of AAA+ proteins

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Summary

Phage shock protein A (PspA) belongs to the highy conserved PspA/IM30 family and is a key component of the stress inducible Psp system in Escherichia coli. One of its central roles is the regulatory interaction with the transcriptional activator of this system, the σ^{54} enhancer-binding protein PspF, a member of the AAA+ protein family. The PspA/F regulatory system has been intensively studied and serves as a paradigm for AAA+ enzyme regulation by trans-acting factors. However, the molecular mechanism of how exactly PspA controls the activity of PspF and hence σ^{54} -dependent expression of the *psp* genes is still unclear. To approach this question, we identified the minimal PspF-interacting domain of PspA, solved its structure, determined its affinity to PspF and the dissociation kinetics, identified residues that are potentially important for PspF regulation and analyzed effects of their mutation on PspF in vivo and in vitro. Our data indicate that several characteristics of AAA+ regulation in the PspA·F complex resemble those of the AAA+ unfoldase ClpB, with both proteins being

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regulated by a structurally highly conserved coiledcoil domain. The convergent evolution of both regulatory domains points to a general mechanism to control AAA+ activity for divergent physiologic tasks via coiled-coil domains.

Introduction

Phage shock protein A (PspA), identified in filamentous phage infected cells of Escherichia coli 25 years ago, (Brissette et al., 1990) is the archetype of the conserved PspA/IM30 family that encompasses members in bacteria (Jordan et al., 2006), archaea (Bidle et al., 2008) and plant chloroplasts (Kroll et al., 2001). As part of a stress inducible system (the Psp system, for reviews see Darwin, 2005, Joly et al., 2010, Model et al., 1997, Yamaguchi and Darwin, 2012), PspA was found to interact with two small membrane proteins and putative sensors, PspB and PspC (Adams et al., 2003), as well as with PspF (Dworkin et al., 2000), the transcriptional activator of the system, resulting in a negative feedback-loop. Because of the propensity of PspA to associate with membranes (Brissette et al., 1990) and to oligomerize (Hankamer et al., 2004; Standar et al., 2008), it has been proposed that the Psp system is a membrane stress responsive system. Recent studies strengthen this interpretation, showing that PspA switches interaction partners from PspF to PspBC during overproduction of membrane-weakening, pore-forming secretins (Mehta et al., 2013; Yamaguchi et al., 2013; Flores-Kim and Darwin, 2015). Still, overproduction of several other proteins that do not affect membrane stability also highly induces psp (Horstman and Darwin, 2012), suggesting that multiple signals might exist and are integrated in psp regulation (Engl et al., 2011). Clearly, a thorough analysis of the regulatory interaction of PspA and PspF, the last, unifying switch in psp induction, is imperative for understanding of the system.

PspF is an extensively studied member of the bacterial enhancer-binding proteins (bEBP), specialized AAA+proteins (Neuwald *et al.*, 1999) needed for alternative sigma factor (σ^{54})-dependent transcription. Briefly, hexameric PspF binds σ^{54} and, via ATP hydrolysis, promotes the open complex formation of the DNA-bound RNA polymerase (Bose et al., 2008). The structure of PspF was solved a decade ago (Rappas et al., 2005), and numerous studies have probed the function of PspF in transcriptional activation (Chaney et al., 2001; Cannon et al., 2004; Rappas et al., 2005; 2006; Joly et al., 2006; Bose et al., 2008; Burrows et al., 2010; Zhang et al., 2013; Sharma et al., 2014). The interaction with its regulator PspA (which in trans takes the part of the usual cis-regulatory domains of other bEBPs) is however poorly understood. It is known that PspA and PspF can form a complex (Jolv et al., 2009). which leads to inhibition of ATPase activity (Elderkin et al., 2002) and down-regulation of psp expression in vivo (Dworkin et al., 2000). It is thought that the PspA·F complex likely consists of 6 PspA per 6 PspF (Elderkin et al., 2002; 2005; Joly et al., 2009; Zhang et al., 2013). An exposed loop on PspF around Trp56 has been identified as a PspA-binding determinant (Elderkin et al., 2005; Zhang et al., 2013), and the PspF-binding region (FBR) in PspA has been localized in the fragment PspA1-186 (Elderkin et al., 2005; Joly et al., 2009), with a recent study suggesting that residues 25-40 form an amphipathic helix important for PspF inhibition (Jovanovic et al., 2014). Still, the lack of any PspA/IM30 family structure precludes a detailed understanding of the proteobacterial PspA·F complex as well as other PspA-like proteins on a molecular level.

In vitro work with full-length or fragmented PspA has been challenging, as the stability of activity strongly relied on the addition of detergents (Elderkin et al., 2002). We overcame those issues by identifying the PspF regulatory core domain in PspA that lacks membrane-interacting or oligomerization properties and is fully functional in the absence of detergent over longer periods of time with respect to PspF-binding and inhibition in vivo and in vitro. We present the crystal structure of this domain at 1.8 Å resolution, identify the PspF-binding surface and characterize regulatory influences of single amino acid exchanges on the activity of PspF in vivo and in vitro. The PspA·F complex has a striking resemblance to the AAA+ unfoldase ClpB (Lee et al., 2003) and its regulatory middle domain, showing that regulation of AAA+ proteins via coiled-coil domains evolved convergently for AAA+ proteins of diverse functions.

Results

Identifcation of $PspA_{1-144}$, the PspF-inhibiting domain of PspA

The σ^{54} activator PspF is an important example for a AAA+ family protein that does not contain intrinsic regulatory domains, but instead is regulated by another protein, PspA, in *trans*. On a molecular and structural level, it is still largely unresolved how this regulation is

achieved, and we therefore intended to obtain more information about the involved structures and interactions. PspA is difficult to study as it can not only interact with PspF, but also with membrane components such as PspC or lipids (Brissette et al., 1990; Adams et al., 2003), and it can self-associate to form large superstructures (Hankamer et al., 2004; Standar et al., 2008). To circumvent issues arising from those characteristics that are unrelated to the PspA-PspF interaction, we sought to identify the minimal PspF-interacting domain of PspA. A previous fragmentation approach that was based on a helical domain prediction (HD1-4, Fig. 1A; Elderkin et al., 2005, Joly et al., 2009) already indicated that PspFbinding determinants are located in the not oligomerizing fragment PspA₁₋₁₈₆. However, PspA₁₋₁₈₆ was less effective than full-length PspA (PspA1-222) in PspF-ATPase inhibition (Joly et al., 2009), still partially associated with the membrane and was purified by a protocol that employs detergent to preserve solubility (Elderkin et al., 2005; Jovanovic et al., 2014), indicating that PspA1-186 still contains determinants unrelated to PspF-binding. It is thus important to recognize possible functional domains in PspA. In previous studies, PspA was predicted to comprise a large number of more or less likely coiled-coils that are organized in four helical domains (Joly et al., 2009). When we performed a coiled-coil prediction using COILS, the outcome was much less complex. COILS predicted three large coiled-coil regions (CC1-3) that did not correspond to the helical domains that were the basis of earlier fragmentation approaches (Fig. 1A). Importantly, CC2 ended at amino acid 144, which is 42 residues earlier than the end of the previous PspA₁₋₁₈₆ construct. As it is known that regions within the first 67 residues and after residue 110 are important for PspF regulation (Joly et al., 2009), it seemed possible that CC1 and CC2 are responsible for PspF regulation, and that the 42 amino acid extension of CC2 in PspA₁₋₁₈₆ somehow supports the membrane interaction of this construct (Jovanovic et al., 2014).

We thus generated PspA1-144, which comprises the native N-terminal region (NTR) and only the two predicted coiled-coil domains CC1 and CC2 (orange, Fig. 1A). PspA₁₋₁₄₄ and full-length PspA turned out to essentially indistinguishably inhibit PspF-dependent psp expression in a $\Delta pspA$ reporter strain over the course of 15 hours, suggesting that both interact similarly with PspF (Fig. 1B, left). Neither protein had a negative effect on growth (Fig. 1B, right). However, the two proteins differed in their subcellular localization. In agreement with previous analyses, overproduced full-length PspA localized to both the membrane and cytoplasmic fractions (Yamaguchi et al., 2010). In contrast, PspA1-144 was exclusively soluble, indicating that the membrane-interacting trait of PspA had been removed (Fig. 1C). Consequently, we were now able to purify his-tagged PspA₁₋₁₄₄ from the cytoplasmic frac-



Fig. 1. Identification of the minimal PspF-inhibitory domain of PspA, PspA₁₋₁₄₄.

A. Bottom: COILS prediction for PspA. Top: Domain structure of PspA as derived from the COILS prediction, PspA₁₋₁₄₄ (*orange*) highlighted. For comparison, the proposed helical domains HD1-4 of PspA as suggested by Elderkin *et al.* (2005) are indicated earlier. B. Comparison of full-length PspA and PspA₁₋₁₄₄ in *psp* regulation. Both have strong and indiscernible inhibitory effects. Left: LacZ-activity assay in a MC3 $\Delta pspA$ reporter strain showing the inhibitory effect of PspA₁₋₁₄₄ (*orange*) and full-length PspA (*black*) relative to an empty vector control (*dashed line*) when overproduced. Right: Growth curves for the cultures used for LacZ-activity assays. C. Western blot with antibodies against the His-tag showing that PspA₁₋₁₄₄ localizes to the soluble fraction after ultracentrifugation, while full-length PspA is found in both the membrane and cytoplasmic fraction. Signals of the control proteins DnaK (CP) and YidC (M) in the samples are given to show that the fractionation was successful. CD, cell debris; M, membrane; CP, cytoplasmic fraction. D. Sedimentation equilibrium of PspA₁₋₁₄₄ during analytic ultracentrifugation indicates that PspA₁₋₁₄₄ is purely monomeric *in vitro*. The curve fit corresponds to a mass of 17.6 kDa (monomeric PspA₁₋₁₄₄ ≈ 17.55 kDa).

tion without the need of detergents for keeping the protein in solution. Subsequent equilibrium sedimentation experiments (analytic ultracentrifugation, AUC) showed that purified $PspA_{1-144}$ was exclusively monomeric (Fig. 1D), indicating that it also lacks the determinants for selfassociation. Thus, PspF interaction is mediated by a domain that comprises the N-terminal 2/3rds of PspA, whereas self-association as well as detectable membrane interaction require the truncated C-terminal regions, possibly including CC3.

Crystal structure of PspA₁₋₁₄₄ *reveals similarity to the M-domain of the* AAA+ *protein* ClpB

Having the stable, monomeric and soluble PspFinteracting PspA domain at our hands, we were able to crystallize this protein and solved its structure to a resoforms an extended monomeric structure in which the two coiled-coil regions CC1 and CC2 (see Fig. 1A) form an intramolecular anti-parallel coiled-coil (Pro25-Arg142) linked by a tip region (Ala75-Leu91). A short NTR (Ile3-Val11) covers a hydrophobic patch on CC1 (Fig. 2A, Fig. S1A-C), connected to CC1 by a flexible linker (residues Asn12-Glu23), with Asp24 acting as an *N*-cap to the CC1 helix. Three highly flexible residues in this linker (Lys20, Ala21, Glu23) could not be resolved. In atomistic molecular dynamics simulations that included these residues, the NTR remained attached to the coiled coil while the flexibility of the linker was confirmed (Fig. S1D,E). Conservation of residues contributing to intramolecular coiled-coil stabilization (Fig. 2B, green) and tip formation (red) within proteins of the PspA/IM30 family indicates

lution of 1.8 Å (Fig. 2A). This is the first member of the

conserved PspA/IM30 family to be crystallized. PspA₁₋₁₄₄

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Fig. 2. The crystal structure of PspA₁₋₁₄₄ shows the conserved architecture of PspA/IM30 family proteins. Conserved amino acids: green, redstructurally important amino acids; yellow- others.

A. Crystal structure of PspA₁₋₁₄₄ shown in *cartoon* representation, conserved residues as *sticks*. The short loop of amino acids 23–24 was modelled as no electron density for these residues was observed.

B. Overall conservation histogram and detailed excerpts for well-studied members of the PspA/IM30-family. For orientation, CC-domains of PspA₁₋₁₄₄ (as in Fig. 1A), N-terminal region (NTR) and tip domain (tip) are indicated earlier.

that the *E. coli* PspA₁₋₁₄₄ structure presented here reflects the general architecture of this family, including the wellstudied VIPP1 from chloroplasts and LiaH from *Bacillus subtilis*.

A search for structurally related proteins using the program DALI (Holm *et al.*, 2008) yielded a number of coiled-coil containing protein domains (as expected for such a common structural element) of varying lengths and little to no functional similarity. Interestingly, PspA₁₋₁₄₄ showed significant structural similarity (*Z*-score 6.6, top 10% of all hits) to the coiled-coil M-domain of the ClpB family of AAA+ unfoldases/disaggregases, although they have alternative permutations of the secondary structure elements, as CC1 of PspA is represented by two helical domains in ClpB (Fig. 3B and C), which is known to decrease the *Z*-score of similar folds (Holm and Rosenstrom, 2010). This domain (ClpB-MD), which is found as an insertion within the AAA+ domain and adoms

the peripheral surface of the hexameric ring, regulates the disaggregase activity of ClpB by modulating its ATPase activity (Schirmer *et al.*, 2004; Oguchi *et al.*, 2012). The highly similarly sized PspA₁₋₁₄₄ and ClpB-MD superimpose with an r.m.s.d. (root-mean-square deviation) of 1.1 Å for backbone atoms (Fig. 3A; see Fig. S2 for stereo images).

Variants of PspA1-144 reveal a FBR on CC1

We investigated residues of PspA₁₋₁₄₄ that are responsible for PspF binding and regulation by characterizing the phenotypes of amino acid substitutions in CC1 positions that are conserved in γ -proteobacteria (Fig. S3A), suggestive of a possible interaction interface with the likewise conserved PspF. When produced in a P_{pspA}-lacZ reporter strain with a $\Delta pspA$ background, several of these PspA₁₋ 144 variants resulted in less *psp* repression in comparison to the wild-type fragment (including R30A and E37A,



Fig. 3. PspA₁₋₁₄₄ (orange) is structurally highly similar to the M-domain of ClpB (ClpB-MD).

A. Superimposition of the backbones of coiled-coiled moieties of both proteins using Swiss-PdbViewer (Guex and Peitsch, 1997). For stereo representations see Fig. S3.

B. Side by side comparison of PspA and ClpB-MD structure as *cartoon* representation, schematically depicted in (C). CC1 of PspA is divided into two helices (H3/4 and H1) in ClpB-MD, but secondary structure and backbone directions are conserved (*arrows* indicating N- to C-terminal direction).



Fig. 4. Effects of single amino acid exchanges E37A (red) and R59E (blue) in the putative PspF-interacting region of PspA.

A. Variants that change PspF inhibition in vivo cluster on one side of PspA (cf. Fig. S3).

B. LacZ-activity assays (MC3 $\Delta pspA$) showing changes in PspF-inhibitory effect of PspA₁₋₁₄₄ variants relative to WT-PspA₁₋₁₄₄ (*orange*) and compared to the unregulated empty vector control (empty). PspA₁₋₁₄₄-E37A is a weaker, PspA₁₋₁₄₄-R59E is a stronger inhibitor of *psp*. Long-term induction was needed to dilute the high LacZ concentration of the unregulated $\Delta pspA$ reporter strain.

C. Subcellular localization of both PspA₁₋₁₄₄ variants. Both fragments are present in the cytoplasm, but PspA₁₋₁₄₄-E37A is also found in inclusion bodies and the membrane fraction. For localization of the wild-type fragment and used abbreviations see Fig. 1C.

D. LacZ-activity assays in a wild-type *pspA* background with constant expression of *pspF* (MC3 $\Delta pspF$ *pUL-pspF-strep*) showing that PspA₁₋₁₄₄-E37A induces *psp* above the level of the empty vector control (empty). PspA₁₋₁₄₄-R59E leads to slightly stronger repression than the wild-type. Effects are already detectable after 3 h of induction (left), and get more pronounced after long-term induction (right).

E. Co-elution of strep-tagged PspF with his-tagged PspA₁₋₁₄₄-E37A from the reporter strain after 3 h shows that both proteins interact *in vivo* while the variant induces *psp*. Wild-type PspA₁₋₁₄₄ is shown as comparison. Western blots of Ni-IMAC using equal amounts of the flow-through (FT), last wash fraction (W) and the seven elution fractions (E1–E7), using antibodies against the Strep- and His-tag, respectively.

Fig. S3B). Additionally, amino acid substitutions at the very C-terminal end of this conserved region caused more effective repression than the wild-type (R59E, Fig. S3C). Two results of this preliminary screen were especially noteworthy: First, all single amino acid exchange variants retained PspF inhibition to a certain degree since all measured *psp* levels were markedly below the level of the uninhibited control indicating that these single amino acid exchanges did not fully abolish the functional interaction.

Second, the surface-exposed exchanges with pronounced effects on *psp* regulation clustered on one side of PspA, which suggested to us that we might have identified the FBR on PspA (Fig. 4A and Fig. S3D).

To address this important aspect further, we extensively characterized two $PspA_{1-144}$ derivatives in more detail: $PspA_{1-144}$ -E37A, the strongest loss-of-repression variant of a surface residue, exhibiting a ~20-fold loss in repression; and $PspA_{1-144}$ -R59E, the only derivative causing

enhanced repression relative to the wild-type fragment (0.3-fold *psp* level, Fig. 4B). Both variants were present in the cytoplasm at a concentration roughly similar to the wild-type fragment (Fig. 4C, compared with Fig. 1C, all localizations were done in parallel). If anything, the amount of PspA₁₋₁₄₄-R59E seemed to be slightly lower, indicating that the enhanced repression due to this mutation is not an effect of higher protein concentration. We found PspA₁₋₁₄₄-E37A to also form inclusion bodies that resulted in a pronounced accumulation in the cell debris and membrane fractions. As will be shown later, the protein in the soluble fraction nevertheless is monomeric and functional.

We undertook a detailed in vivo analysis of the effects of the two regulatory variants on PspF using a wild-type pspA reporter strain. Our previous experiments were performed in a $\Delta pspA$ reporter strain with a strong uninhibited psp promoter activity (up to 25 000 Miller Units shortly after inoculation from an overnight culture, Fig. 1B). This made prolonged induction times necessary to dilute previously accumulated LacZ, not leaving the option to examine differential effects of variants on PspF directly after induction. We therefore tested effects of our variants in the wild-type pspA genetic background, which has a balanced psp expression that is maintained by the autoregulatory feedback-loop of PspA-dependent PspF inhibition. To ensure that our repression-phenotypes were not caused by varying PspF levels in our reporter strains, endogenous pspF was deleted, and strep-tagged PspF was maintained at a constant level by expressing its gene constitutively from a low-copy plasmid. Western blots confirmed that PspF levels were indeed comparable between the strains at all times (Fig. S4A).

As expected in this pre-regulated Psp system, changes in psp levels because of the expression of PspA1-144 variants in the PspA wild-type background were already visible after 3 h (Fig. 4D, left) and became more pronounced after 18h (Fig. 4D, right). PspA₁₋₁₄₄-R59E caused a slight increase in PspF repression relative to WT-PspA₁₋₁₄₄ at 3 h with significantly enhanced repression after 18 h, consistent with the data obtained in the $\Delta pspA$ strain (Fig. 4B). PspA₁₋₁₄₄-E37A induced *psp* above wild-type level at both time points, turning from a less-effective inhibitor in the $\Delta pspA$ background to an activator in the wild-type pspAbackground. We performed co-elution experiments of the soluble fraction to investigate whether this activating effect was still due to direct interaction with PspF, or whether it was a secondary effect because of the partial inclusion body formation of that variant. Both PspA₁₋₁₄₄ and PspA₁₋ 144-E37A co-elute PspF-strep with indiscernible strength, indicating that PspA1-144-E37A directly interacts with PspF in the cytoplasm of E. coli while simultaneously inducing the Psp system (Fig. 4E). These co-elution experiments were performed using the same induction time and strength at which we observed the activating effect. These results are in full agreement with the regulatory data obtained with the $\Delta pspA$ strain, where PspA₁₋₁₄₄-E37A significantly inhibited the PspF-dependent promoter activity, which can most easily be explained by an interaction of PspA₁₋₁₄₄-E37A with PspF in vivo (Fig. 4B). We ensured that this effect was not due to the elevated PspF levels by showing that the effect of PspA1-144-E37A was even stronger in the wild-type psp background, i.e. a strain without deletion and *trans*-complementation of the *pspF* locus (Fig. S4B). The activating effect of the E37A mutation in a wild-type psp background was unexpected, as the intrinsic repression by endogenous PspA should override expression of a loss-of-function variant as PspA₁₋₁₄₄-E37A. It is therefore highly remarkable that a variant of PspA₁₋₁₄₄ is dominant over the wild-type PspF inhibition mechanism without being able to regulate PspF at least as effectively as intrinsic PspA. We also found evidence that the activating effect of the E37A mutation was not confined to the PspA1-144 fragment, but that it also existed in the context of full-length PspA: In a wild-type psp background, the E37A mutation led to a 6.7-fold induced psp level relative to the empty vector control (Fig. S4C). It has to be mentioned that this up-regulation, although strong and significant, cannot fully be attributed to the E37A mutation alone, as un-mutated full-length PspA already slightly induced psp after long-term production (2.8-fold), which was a largely PspBC-dependent effect in this experimental setup and thus not a direct effect of the PspA-PspF interaction per se (Fig. S4D). We could further confirm that the E37A variant of full-length PspA still bound PspF comparable with the wild-type full-length protein (Fig. S4E). While the E37A mutation therefore seems to have a comparable physiologic effect in the full-length and truncated fragment, this experiment once more shows the advantage of working with PspA₁₋₁₄₄, as it is not prone to secondary effects on PspA·F interaction that can result from oligomerization with the pool of intrinsic PspA as well as from membrane and/or PspBC interaction.

Six PspA monomers bind one hexamer of PspF

We further characterized the PspA–PspF interaction *in vitro*. Size exclusion chromatography experiments using purified PspA₁₋₁₄₄ and PspF₁₋₂₆₅ (a stably folding variant lacking the C-terminal DNA-binding domain) clearly showed complex formation of PspA₁₋₁₄₄ and PspF₁₋₂₆₅ (Fig. S5A). Using AUC, we investigated the dynamics of PspA₁₋₁₄₄·PspF₁₋₂₆₅ complex formation (for simplicity, called PspA₁₋₁₄₄ to PspF₁₋₂₆₅ demonstrated a micromolar dissociation constant ($K_D \approx 1 \mu M$, Fig. 5A). The most strongly *psp*-inducing FBR-variant PspA₁₋₁₄₄-E37A bound PspF₁₋₂₆₅ with slightly reduced affinity ($K_D \approx 6 \mu M$), whereas the



Fig. 5. PspA and PspF form a highly stable complex with a basal ATPase activity in vitro. (A/B) In analytic ultracentrifugation experiments (AUC), PspA1-144, PspA1-144-E37A or PspA1-144-R59E were titrated in variable concentrations to PspF1-265 and the change of the sedimentation coefficient (Svedberg) of the fast-sedimenting PspA-F species as a function of PspA concentration was determined. A. Addition of PspA to 10 µM PspF₁₋₂₆₅, a concentration at which PspF readily self-oligomerizes. The K_D was determined for both WT and variants, determination of complex stoichiometry (1 to 1 binding of WT-PspA) is shown as dashed lines that cross at around 10 µM PspA. B. Stabilizing effect of PspA and its variants at low concentrations of PspF. At 1 µM PspF₁₋₂₆₅, where PspF is mostly non-hexameric, addition of very low amounts of PspA1-144 already lead to formation of a fast-sedimenting PspA F species. Variants show either slightly less

(PspA₁₋₁₄₄-R59E) or nearly no (PspA₁₋₁₄₄-E37A) stabilization at sub-stoichiometric concentrations. C. PspF₁₋₂₆₅ exhibits K_p-dependent inhibition of its ATPase activity by PspA variants under non-saturating conditions.

D. Variants change the basal ATPase activity (grey) of the PspAF complex. This effect does not correspond to their differences in K₀.

super-inhibitory variant PspA1-144-R59E showed binding comparable to the WT (K_D $\approx 0.7~\mu M$). Furthermore, the stoichiometry of binding could be shown to be 1:1 in all cases. As PspF is mostly hexameric at these concentrations (10 µM, Fig. S5B), this ratio corresponds to a complex of six $PspA_{1-144}$ with six $PspF_{1-265}$, which is in full agreement with earlier studies (Joly et al., 2009; Lenn et al., 2011; Mehta et al., 2013; Jovanovic et al., 2014). As PspA₁₋₁₄₄ is a monomer, these results suggest that one hexamer of PspF possesses six distinct binding sites for monomeric PspA, analogous to the hexameric AAA+ core of ClpB that can interact with the six coiled-coil regulator domains independently.

One PspA monomer is sufficient to stabilize one hexamer of PspF

While examining PspF hexamerization using AUC, we observed that PspF₁₋₂₆₅ does not oligomerize spontaneously when present in low concentration (1 μ M, Fig. S5B), which is relevant as PspF is thought to exist in low concentration, i.e. just ~130 copies per cell (Jovanovic et al., 1997, PspA: ca. 1500 copies, Valgepea et al., 2013). We therefore wondered whether PspA would also interact with monomeric PspF, or whether PspF needed to hexamerize first. Published gel filtration data using PspF variants suggest that a PspA·F complex is formed independently of the starting oligomeric state of PspF, yet in these experiments both proteins were pre-incubated at high concentrations and PspA was added in excess (Joly et al., 2009).

To our surprise, addition of very low concentrations of PspA (0.08 µM) to 1µM PspF resulted in the emergence of one fast-sedimenting species of PspA·F (9.4 S, Fig. 5B). At these concentrations, PspA only contributed less than 2 % of the overall UV absorption in the sample cell. The fast-sedimenting species therefore comprised around 0.46 μ M PspF in the sample cell (~ 46% of total UV absorption, Fig. S5C). Taking the ratio of applied PspA to shifted PspF into account (0.08:0.46 µM or 1:5.8), these data show that one monomer of PspA₁₋₁₄₄ is able to bind and stabilize one hexamer of PspF1-265 over the course of the AUC measurement. No intermediary complexes could be observed. In contrast, PspA1-144-E37A was found to stabilize PspF hexamers only above equimolar concentrations (Fig. 5B, Fig. S5C), while PspA₁₋₁₄₄-R59E showed slightly impaired ability to form complexes at low concentrations. These results indicate that sub-stoichiometric PspA stabilizes PspF hexamers and also supports the idea that we have identified a PspA interface important for binding. We would like to note that, while certainly interesting for PspA-PspF interaction studies, PspAdependent stabilization cannot be a basic prerequisite for PspF hexamerization *in vivo*, as PspF is active in a $\Delta pspA$ strain (Fig. 1B). Therefore, most likely the C-terminal DNA-binding region of PspF, which had to be truncated for effective purification in our experiments, stabilizes PspF hexamers in vivo, as suggested before (Schumacher et al., 2004). The PspA-mediated sub-stoichiometric stabilization of PspF hexamers may thus play only a very minor physiological role, possibly by contributing to the

long-term stability of active PspF. Typically, AAA+ proteins are also strongly bound and stabilized by nucleotides. PspF is unusual in this respect as it binds ATP or ADP only weakly. (Joly *et al.*, 2006) Although PspF hexamers are stabilized by nucleotides at protein concentrations above 9 μ M (Joly *et al.*, 2006), AUC measurements with near-physiologic 1 μ M PspF₁₋₂₆₅ indicated no stabilizing effect, even at ADP concentrations of 0.5 mM (ca. fivefold K_D, Fig. S6D, cf. Joly *et al.*, 2006).

Fully saturated PspA·F complex has a basal ATPase activity that can be increased or decreased by PspA variants

The central function of the PspA·F complex, regulation of PspF-dependent transcriptional activation, is tightly linked to the ATPase activity of PspF (Elderkin et al., 2005). Therefore, we examined the influence of WT-PspA₁₋₁₄₄ and its FBR-variants PspA1-144-E37A and PspA1-144-R59E on the ATPase activity of PspF in vitro. As predicted, PspA₁₋₁₄₄ strongly reduced PspF-ATPase activity (Fig. 5C). However, even the saturated PspA·F complex showed a significant residual ATPase activity of ~18% (Fig. 5D), in accordance with the previous finding that inhibition of PspF-ATPase remains incomplete in the presence of either fragments or full-length PspA (Elderkin et al., 2002; 2005; Joly et al., 2009). As this basal ATPase activity might have been due to an association-dissociation equilibrium of the complex (leading to brief periods of uninhibited PspF following PspA dissociation), we measured the increase of PspF activity after jump dilution, which is dependent on PspA dissociation (Fig. S7A). From these data we determined the half-life for PspA·F to be 43 min $(k_{off} = 2.69 \cdot 10^{-4} \text{ s}^{-1})$, which strongly argues against fast exchange rates of PspA protomers of PspA·F complexes. In ATPase assays with PspA in sub-saturating concentrations, we found that the strength of ATPase inhibition was dependent on the K_D of the PspA-PspF interaction (Fig. 5C). PspA₁₋₁₄₄ and PspA₁₋₁₄₄-R59E with virtually the same K_D inhibited PspF stronger than PspA₁₋₁₄₄-E37A, which has a sixfold higher K_D). However, in assays with PspA-saturated complexes, this K_D-dependence disappeared: In the presence of saturating PspA, the residual ATPase activity of PspF was 30% lower with PspA1-144-E37A and 30% higher with PspA₁₋₁₄₄-R59E as compared to WT-PspA₁₋₁₄₄ (Fig. 5C). Thus, wild-type or mutated variants can have distinct effects on PspF ATPase that do not necessarily correlate with effects on transcriptional activity of PspF: The exchange E37A leads to a lower ATPase activity in vitro, but higher psp induction in vivo, and vice versa for R59E. As we will discuss in detail later, this disconnection of ATPase and transcriptional activity is counterintuitive, but in agreement with previously reported effects of mutations in the PspF- σ^{54} system.

Atomistic simulations indicate a possible PspA·F complex structure

We tested PspA·F complex formation and stability in a series of atomistic molecular dynamics simulations based on the assumption that six PspA monomers should contact PspF with their FBR and that one of the interaction sites on PspF should be at the W56-loop (Elderkin et al., 2002; Zhang et al., 2013). In one upright orientation in which the binding region contacted both the W56-loop of one PspF protomer and an ATP binding helix (sensor-II domain) on the adjacent PspF, PspA remained bound to PspF during the time course of the simulation (138 ns, four independent simulations) (Fig. S6A, complex depicted in Fig. S6B). In control simulations (in which the binding epitopes on PspA pointed away from PspF), PspA failed to maintain a stable interaction (Fig. S6A). While these simulations are by no means exhaustive, they demonstrate the possibility that PspA engages two PspF protomers, which might explain the stabilization of PspF hexamers by sub-stoichiometric PspA.

Discussion

In this study, we examined the regulatory mechanism of PspA·F-dependent psp induction in vivo and in vitro. We identified the PspF regulating domain of PspA, PspA₁₋₁₄₄, a large coiled-coil formed by the helices CC1 and CC2 (Fig. 1A). This large fragment, comprising two-thirds of PspA, was as effective as the full-length protein in psp regulation (Fig. 1B). In previous studies, shorter fragments showed impaired PspF interaction (Elderkin et al., 2005; Joly et al., 2009), which clearly indicates that this domain is necessary and sufficient for PspF regulation under non-stress conditions. Because of its monomeric and stably soluble nature (Fig. 1C and D), PspA₁₋₁₄₄ allowed us to look at characteristics of the PspA·F complex that eluded observation so far, and it furthermore led us to the first crystal structure of the PspA/IM30 protein family. The conservation of residues that are important for this structure indicates that the PspA architecture must be highly conserved throughout the PspA/ IM30 family (Fig. 2B).

Full-length PspA is partially membrane localized and oligomeric whereas PspA₁₋₁₄₄ is soluble and monomeric. This is fully compatible with evidence in the literature that point toward a cooperative role of the N- and C-terminal parts of PspA in oligomerization and functions at the membrane. Previous studies have indicated that either deletion of an N-terminal helix (Jovanovic *et al.*, 2014) or point mutations in a part or the C-terminus (Yamaguchi *et al.*, 2010) lead to a change in membrane-dependent functions rely on having both N- and C-termini present, it

is not surprising that we find an exclusively cytoplasmic location for PspA₁₋₁₄₄, lacking its C-terminus. Importantly, the short N-terminal amphipathic helix (residues 2-19) that has previously been implied to interact with membranes (Jovanovic et al., 2014), is laterally docked against CC1 and CC2 in our structure (Fig. S1D). Our structure may thus represent a membrane-detached state with the NTR back-folded to the coiled-coil in some kind of 'parking' position, ready to be released for membrane interactions upon some signal. However, this helix alone is clearly not sufficient to mediate a stable membrane interaction of monomeric PspA₁₋₁₄₄. The PspF-regulatory domain (CC1 and CC2, residues 23 to 144) is likely not involved in membrane binding, since there is no evidence from the crystal structure for a direct membrane interaction: No amphipathic patches (with exception of those in the folded coiled-coils) are present in this part of the protein, and there are no discernible regions of positively charged surface amino acids, either of which would presumably be important for a direct membrane interaction.

The monomeric state of PspA₁₋₁₄₄ likely results from lack of CC3 and the far C-terminal region. This view is in full agreement with previous analyses that suggested a key role of C-terminal regions in PspA for oligomerization and effector function, as a PspA₁₋₁₈₆ fragment was already strongly affected in both respects (Joly et al., 2009; Jovanovic et al., 2014). We note that CC3 contains a consensus-motif for short parallel trimeric coiled-coils (amino acids 174-179, see Kammerer et al., 2005), which might indicate that PspA forms trimers when initializing oligomerization. It is however uncertain at this point how exactly oligomerization is achieved and whether or not parts of PspA₁₋₁₄₄ have to interact with C-terminal regions directly. The lack of membrane interaction and oligomerization made PspA₁₋₁₄₄ suitable for crystallization and PspA·F complex studies. It must be kept in mind that the PspA₁₋₁₄₄ construct and the analyzed derivatives thereof are useful to selectively address the regulatory PspF interaction without interfering aspects of membrane interaction, such as recruitment of PspA by PspBC or the effector function. However, we would like to stress that the PspA–PspF interaction certainly has to be seen in the context of the full-length protein and the Psp system it is part of to fully understand the physiologic role of proteins of the PspA/IM30 family. Furthermore, the now established coiled-coil structure of PspA1-144 has to be kept in mind when interpreting effects of PspA fragments that have helices partially removed from within the coiled-coil, which certainly can result in non-physiologic secondary effects because of structural destabilization of the protein and exposition of amphipathic helices that lack their coiled-coil counterpart.

Using $PspA_{1-144}$ for mechanistic studies, we were able to characterize the $PspA \cdot F$ complex as a highly stable

complex with a 6:6 stoichiometry and a basal ATPase activity *in vitro*. If the stability of the PspA₁₋₁₄₄·PspF₁₋₂₆₅ complex *in vitro* (half-life ~ 43 min, Fig. S5E) resembles that *in vivo*, then the high stability of the complex constrains the mechanism of regulation of *psp in vivo* and an entirely dissociation-based mechanism that relies on indirect activation of *psp* via sequestering free PspA to the membrane does not suffice for a rapid *psp* response. It may thus be required that PspA has to be more actively 'peeled off' PspF during induction either by conformational changes in PspA (e.g. mediated by the C-terminus) and/or by conformational changes upon interactions with PspBC or the membrane (Yamaguchi *et al.*, 2013).

Our results support the possibility that PspF activity can in principle be altered without a requirement for PspA·F complex dissociation. An in vivo screen for PspF-inhibition variants of PspA₁₋₁₄₄ allowed us to identify a surface patch on PspA that is likely responsible for the regulatory PspF interaction, stretching along the length of the coiled-coil on the side of the protein opposite to the back-folding N-terminus. Detailed characterization of two PspFregulation variants of PspA1-144 show that mutations in this region can alter the activity of PspA-regulated PspF, likely while bound to PspF (a modulatory interaction), leading to a change in psp levels in vivo as well as in the basal ATPase activity of PspF in vitro: The up-regulating effect of PspA1-144-E37A is a dominant phenotype in a wild-type background while this fragment clearly binds PspF (Fig. 3D and E). Therefore, the bound PspA₁₋₁₄₄-E37A represses PspF less effectively than the wild-type protein, which may mimic a post-binding level of PspF regulation in the case of our PspA fragment. Similar post-binding effects on the ATPase activity can be observed in vitro, where the basal activity of the saturated PspA·F complex seems to be modulated by variants relative to the wildtype (Fig. 4D). Interestingly, the mutation E37A, which led to an induction of psp in vivo relative to the wild-type fragment (Fig. 3B, also visible in the full-length PspA, Fig. S4C), exhibited a lower ATPase level in the saturated complex in vitro, and the R59E mutation that led to a higher basal ATPase level in vitro repressed psp stronger than wild-type PspA₁₋₁₄₄ in vivo (Fig. 5C).

With the current model of PspF-dependent *psp* regulation in mind, these observations are counterintuitive. However, our results are not unprecedented, as mutations in either PspF or σ^{54} have similar effects. Although wildtype PspA–PspF- σ^{54} exhibits a linear correlation between ATPase activity and transcriptional activation, the mutated system does not: Zhang *et al.* (2013) produced PspF variants essentially without ATPase activity yet hyperactive transcriptionally (e.g., variant G58C). Additionally, enhancer-bypass mutations of σ^{54} can fully alleviate the requirement for a bEBP (Syed and Gralla, 1997; Chaney and Buck, 1999). Hence, the energy dependence of transcriptional activation in the natural EBP- σ^{54} system is likely used for tight transcriptional control (see e.g.Sharma et al., 2014). Thus, even though it was unexpected that PspA seems to change the coupling of ATPase and transcriptional activity in PspF while bound to it, the observed effects are similar to those of described variants of PspF and σ^{54} . How could the effect of PspA variants be explained mechanistically? With the emerging pattern for AAA+ function being that controlled, sequential ATPase activity of subunits is highly important (see e.g., Carroni et al., 2014, Glynn et al., 2009, Sysoeva et al., 2013). The PspA variant R59E might lead to a slight desynchronization of PspF subunits, resulting in a higher raw ATPase activity, but decrease in physiologic efficacy of the bEBP, while variant E37A acts vice versa. A modulatory role of PspA would be in agreement with previous reports, which indicated that PspA can interact with PspF in all its conformational states during transcriptional activation: a PspA·F complex still binds to σ^{54} , and PspA interacts stably with ADP-AIF_x trapped PspF $\cdot \sigma^{54}$ complexes (Joly *et al.*, 2009) that are thought to mimic the transcription activating state of PspF (Chaney et al., 2001; Burrows et al., 2010).

PspA-dependent modulation of PspF could be employed in two ways: Either to ensure a controlled lowlevel psp production under non-stress, or to serve as a second mode of stress-dependent induction. The activity of psp in the exponential growth phase (~1500 copies of PspA;Li et al., 2014, Valgepea et al., 2013) has been so far attributed to the presence of slight membrane stresses (Jovanovic et al., 2014), but our data allow another interpretation: The basal ATPase level of the saturated PspA·F complex indicates that PspA-dependent PspF activity might be kept at a controlled basal level by bound PspA. Our R59E variant indeed indicates that psp could be more repressed in vivo than it actually is by bound wild-type PspA, and the long half-life might prevent the complex from sensing varying PspA levels via PspA dissociation under non-stress conditions. A tightly controlled basal induction of psp might therefore be advantageous for the cell to keep the Psp system in check, and could explain why the PspA-dependent regulation of PspF does not interfere with PspF oligomerization or σ^{54} interaction (Joly et al., 2009). Alternatively, modulation of PspF could also add a second layer of *psp* induction that may be achieved by slight conformational changes upon stress signals, which would enable a rapid and fine-tuned stress response. While strong induction of psp undoubtedly relies on the dissociation of PspA, as clearly observed for secretin stress (Yamaguchi et al., 2013), it will be interesting to see how the PspA-PspF interaction changes under other stresses where *psp* is comparably slightly induced, e.g. salt stress (Weber et al., 2006), and if a modulatory action of bound PspA could play a role there. Hence, although we found strong evidences suggesting a modulatory regulation of PspF by PspA, it is unclear at this point how relevant these effects are under different conditions *in vivo*. Most importantly, with the current level of molecular understanding, we do not know whether the variants of PspA that we produced mimic actual states in the cycle of PspF regulation or not. We also cannot exclude that, although unlikely in our opinion, the tendency of PspA₁₋₁₄₄-E37A to form inclusion bodies might have indirect effects on the Psp system. We see however a strong interaction of the variant with PspF in the cytoplasm of *E. coli* as well as clear binding, a modulation of PspF-ATPase activity and no sign of aggregation or inactivity *in vitro*, suggesting that the inclusion body formation does not interfere with assessment of PspF inhibition in our experimental setup.

While being an insulated occurrence in bEPB regulation, we found that PspA·F system shares several structural and mechanistic features with ClpB. It is interesting to note that the typical regulation of bEBPs happens at the level of hexamerization: Regulatory domains act exclusively on the assembly state of the AAA+ATPase, allowing formation of an active oligomeric bEBP only in the presence of an inducing signal (Doucleff et al., 2005; De Carlo et al., 2006). PspF, however, is active per default (Jovanovic et al., 1996), and regulated by PspA in a post-assembly mechanism (Elderkin et al., 2002). While PspA·F therefore differs from canonical bEBP regulation, our studies reveal that a structurally similar regulator can be found in members of the ClpB family of AAA+ proteins, where the middle domain (ClpB-MD), a coiled-coil resembling PspA (Fig. 3), regulates the activity of the protein. From a strictly physiologic perspective, ClpB and PspA·F have completely different roles: One is a disaggregase, responsible for unfolding of misfolded proteins under stress conditions (Woo et al., 1992), the other is involved in the regulation of gene expression (Jovanovic et al., 1996). Also, ClpB's middle domain is fused to the AAA+ domain, while in PspA·F, PspA acts in trans. Their common feature is the hexameric AAA+ core that provides the driving force behind both protein functions. We found several similarities: PspA·F and ClpB have the same stoichiometry (6:6), and both show a basal ATPase activity (Seyffer et al., 2012, this study). Our simulations of the PspA·F complex also hints toward a possible interaction of PspA with two protomers of PspF (Fig. S6), which is a feature of ClpB, where the middle domain likely interacts with two neighboring subunits (Oguchi et al., 2012). Additionally, helix 3 of ClpB-MD and its counterpart on PspA, the N-terminal part of CC1, contain residues crucial for the regulation of the AAA+-domain (e.g. Y503A in ClpB,Oguchi et al., 2012; E37A in PspA, this study), indicating at least partial structural overlap of regions of AAA+ interaction. Nevertheless, this does not mean that both system are regulated in exactly the same way, as ATPase activity in both systems is harnessed for two different tasks: ClpB pulls aggregated peptides apart using repeated ATPase cycles (Lum et al., 2004), while PspF, as discussed earlier, adds an energy barrier to a process that does not need energy per se, allowing for highly specific induction. As a consequence, we note that the effects of amino acid variants on ATP hydrolysis are different and seem to be stronger in ClpB than in PspA·F (PspA E37A; ATPase activity 30 % down, psp levels ~20-fold up), but translate into comparably weaker effects on physiologic activity (e.g. ClpB K476C; ATPase ~15-fold up, disaggregase activity ~3-fold up, Oguchi et al., 2012). Future studies will shed more light on the molecular mechanism of PspF regulation, and the crystal structure of the PspA·F complex, followed by a thorough mutational study of the PspA-PspF interface, is certainly needed to reach the level of understanding that exists for middle domain-dependent ClpB regulation (see Haslberger et al., 2007, Mogk et al., 2003, Oguchi et al., 2012, Schirmer et al., 2004).

At last, it remains to be seen whether this general regulatory mechanism is used by other, less wellcharacterized AAA+ proteins as well, and how the mechanism has adapted to its respective task. Although different in certain aspects, the similarities of ClpB and PspA·F show that a coiled-coil domain, either covalently fused to the AAA+ domain or non-covalently bound, can act as a regulator of diverse AAA+ proteins. A tempting, but speculative guestion that remains is, why a member of the conserved PspA/IM30 family 'hijacked' a bEBP for its own regulation in the proteobacterial system: Did the conserved PspA/IM30 family coincidentally happen to have a structure that made bEBP-dependent regulation of its own production in proteobacteria possible, even though the structure evolved for a physiologically different reason, such as membrane stabilization? Or did the PspA/IM30 family per se (co-)evolve as regulators of the ubiquitous AAA+ domain, which in turn allowed PspA in proteobacteria to regulate its own production?

While we hope to have laid the foundations for a more detailed understanding of the PspA/IM30 protein family in general, and the intricate regulatory features of the proteobacterial PspA·F complex in particular, many features concerning the action of PspA/IM30 remain obscure. Twenty-five years after its first description, there is still much to discover in the field of PspA.

Experimental procedures

Strains and cultivation

For all protein purifications, *E. coli* strain BW25113 (Datsenko and Wanner, 2000) with indicated plasmids was used. The strain MC3 (Bergler *et al.*, 1994), a derivative of *E. coli* MC4100 (Casadaban, 1976) harboring a *pspA* promoter fused to *lacZ* integrated into the λ attachment site, was used to

investigate changes in WT *psp* level via LacZ-activity assays. Prior to first use, the arabinose-resistance of this strain was assured via plating on LB containing 1 % (w/v) arabinose. The same clone was used for all further transformations and transductions to ensure that observed differences in *psp* induction were not a result of different intracellular arabinose levels in the tested strains (Lindenstrauss *et al.*, 2010). Its derivative MC3 $\Delta pspA::kan$ was constructed via P1-phage transduction (Thomason *et al.*, 2007) of MC3 using the $\Delta pspA::kan$ containing JW1297 from the Keio collection (Baba *et al.*, 2006). MC3 $\Delta pspF::kan$ was obtained similarly, using JW1296. All cultures were grown while shaking at 37°C in LB medium (1% (w/v) tryptone, 1% (w/v) NaCl, 0.5% (w/v) yeast extract) if not otherwise specified and supplemented with ampicillin (100 µg ml⁻¹) where appropriate.

Genetic methods and plasmids

The fragment of *pspA* coding for PspA₁₋₁₄₄ was cloned using the respective primers (Table S1) and chromosomal DNA as template. The primers contained restriction sites (5'-Ncol; 3'-Xhol) for cloning into the *pBAD-pspA-H*₆ plasmid previously constructed (Standar et al., 2008), leading to a short leucineglutamate linker in front of the C-terminal hexahistidine tag. Site-directed mutagenesis based on the QuikChange protocol (Agilent, Waldbronn, Germany) was performed to exchange bases in pBAD-pspA-H₆. Primer pairs and the resulting change on amino acid level are given in Table S1 ('pspA-ex-'). To allow easier molecular access to the region coding for the PspF-binding patch, a BspHI restriction site was introduced by a silent mutagenesis at base pairs 92-98 of pspA (coding for amino acids Leu31 to Ile33) resulting in pBAD-pspA-H₆-BspHI. This plasmid was then used for several base exchanges that were coded in a primer overhang and cloned into the plasmid using either Ncol/BspHI or BspHI/Xhol respectively (Tab. S1, 'pspA-BspHI-'). For cloning of pspF₁₋₂₆₅ into pBW22 (Wilms et al., 2001), pspF₁₋₂₆₅ was amplified using chromosomal DNA as template. Primers contained restriction sites (5'-Ndel; 3'-BamHI). For construction of the pSC101based constitutive low-copy expression system pUL-P_{tat}, the Plac-promoter-containing Ndel/Xbal fragment of pCHAP418 (Possot et al., 1992) has been removed by excision, Klenow treatment and religation, and a 436 bp fragment containing the constitutive E. coli tatA promoter with an engineered Ndel site at the tatA start codon has been amplified using pABStatABC (Berthelmann and Brüser, 2004) as template, and ligated into the Pstl/HindIII sites of the vector. The coding region of *E. coli pspF* was then cloned into pUL-P_{tat} (Tab. S1) using Ndel/HindIII. All plasmid constructs were verified by DNA sequencing.

Biochemical methods

Standard protein purification. Overnight cultures were diluted to an OD of 0.05. After induction with 0.1 % (w/v) arabinose (or rhamnose in case of pBW- $pspF_{1-265}$ - H_6) and further growth for 3 h, cells were centrifuged at $6000 \times g$ (4°C) and pellets stored at -18° C. Cells were suspended in 20 mM Tris/HCl, 100 mM NaCl and 20 mM imidazole (pH 8.0) and disrupted by two French Press passages at 138 MPa

and 4°C. Afterwards, cell debris was removed using lowspeed centrifugation at $6000 \times g$ (20 min, 4°C), and membranes and soluble fractions were further separated using ultracentrifugation (Beckman Optima L-80 XP, Beckman Coulter, Krefeld, Germany, 140 000 \times *q*, 1 h, 4°C). Protein was taken exclusively from the soluble fraction and purified via Ni-NTA agarose resin (Qiagen, Hilden, Germany) using standard Ni-affinity protocols. Samples were further purified by anion exchange (Resource Q, GE Healthcare, Freiburg, Germany, linear gradient from 0 to 1 M NaCl in 20 mM Tris/ HCl pH 8.0) resulting in pure protein as controlled via Coomassie Brilliant Blue stained SDS-PAGE gels and SEC. For long-term storage, the ionic strength was readjusted to 100 mM NaCl with HiTrap desalting columns (GE Healthcare, Freiburg, Germany) and protein was concentrated by ultrafiltration (Vivaspin 10 000 MWCO, Sartorius Stedim, Goettingen, Germany). Because of the high stability of PspA1-144 and PspF₁₋₂₆₅, proteins could be stored at -80°C in 20 mM Tris/ HCl and 100 mM NaCl without additives and behaved like freshly prepared samples in all assays (ATPase activity, AUC, SEC) after thawing. All protein concentrations were calculated via their extinction coefficient and absorption at 280 nm. To assess and compare the subcellular localization of PspA. PspA₁₋₁₄₄, and its variants, the pellet of low-speed (cell debris) and ultracentrifugation were resuspended in a buffer volume equal to the supernatant, and aliquots were analyzed via SDS-PAGE/Western blotting, using polyclonal anti-PspA antibodies (Standar et al., 2008), anti-DnaK, anti-YidC or monoclonal anti-His-tag antibodies (Qiagen, diluted 1:5000) and the corresponding secondary antibodies coupled to horse radish peroxidase for enhanced chemoluminescence (ECL) detection. All steps of the localization experiments shown in Fig. 1C and Fig. 4C were performed simultaneously. For co-elution experiments, Ni-affinity purifications from the cytoplasmic fraction of cell cultures were performed as described earlier. After Western blotting, his-tagged PspA (or its fragments and variants) and strep-tagged PspF were detected in the samples using monoclonal anti-StrepTag II (EMD Millipore, Billerica, USA) and His-probe (Santa Cruz Biotechnology, Dallas, USA) antibodies and detected using the respective goat secondary antibodies (IrDye 800CW and 680LT) and the Odyssey system (Li-Cor, Lincoln, USA).

Purification of selenomethionine labeled PspA₁₋₁₄₄ (Se-PspA). Se-PspA-producing cells were grown in minimal medium M9 (Sambrook and Russell, 2001) with 0.4 % (w/v) glucose as carbon source and 0.1 % (v/v) SL12 trace element solution (Overmann *et al.*, 1992). Incorporation of selenomethionine (Acros Organics, now Thermo Fisher Scientific, Geel, Belgium) was assisted by suppression of methionine biosynthesis (Van Duyne *et al.*, 1993) 15 min prior to induction. Purification of Se-PspA was performed similar to that of unlabeled protein, except that 5 mM 2-mercaptoethanol was added throughout all purification steps and in the storage buffer to keep Se-PspA in a reduced state.

Crystallization, data collection and structure determination of $PspA_{1-144}$. Purified $PspA_{1-144}$ in 10 mM Tris/HCI buffer (pH 8.0) and 50 mM NaCI at a final concentration of up to 9 mg ml⁻¹ was used for crystallization by the vapor diffusion hanging drop method. 1 μ l PspA₁₋₁₄₄ was mixed with an equal

volume of crystallization buffer containing 0.1 M HEPES/ NaOH (pH 7.5), 10 % (w/v) polyethyleneglycol 6000 and 5 % (v/v) 2-methyl-2,4-pentanediol and incubated at 15°C. Final high resolution diffracting crystals where obtained by macroseeding of initial PspA₁₋₁₄₄ crystals at a protein concentration of 2 mg ml⁻¹ using a refined crystallization buffer containing 0.1 M HEPES/NaOH (pH 7.2), 10% (w/v) polyethyleneglycol 6000 and 5% (v/v) 2-methyl-2,4-pentanediol. Se-PspA1-144 in 10 mM Tris/HCl buffer (pH 7.5), 50 mM NaCl and 5 mM 2-mercaptoethanol was crystallized in the same manner, but using native PspA₁₋₁₄₄ crystals for cross-seeding. Crystals of both native and Se-containing PspA₁₋₁₄₄ appeared after 3 days and reached their final size within 1 week. Prior to flash-freezing, the crystals were cryo-protected by addition of 15% (v/v) (R,R)-(-)-2,3-butanediol (Merck Millipore, Darmstadt, Germany) to the mother liquid. Collection of the native and the three MAD datasets was carried out under cryogenic conditions (100 K) at the BESSY synchrotron beamline 14.1 (Helmholtz Zentrum, Berlin, Germany) and processed with the XDS package (Kabsch, 2010) (see Table S2 for statistics). Phase determination was carried out by multiple wavelength anomalous dispersion (MAD) using the Se-PspA₁₋₁₄₄ datasets. The heavy atom substructure (four selenium sites) was determined and refined using SHELX (Sheldrick, 2010) and SHARP followed by density modification using SOLOMON within the autoSHARP pipeline (Vonrhein et al., 2007). The resulting electron density map was of sufficient quality to allow automated tracing and model building using the programs ARP/wARP (Langer et al., 2008) and BUCCA-NEER from the CCP4 suite (Winn et al., 2011). The generated model was subjected to further cycles of manual building and refinement employing the programs COOT (Emsley et al., 2010) and Refmac5 (Murshudov et al., 1997) using the highest resolution (1.8 Å) native dataset and Translation/ Libration/Screw (TLS) refinement for final refinement cycles. The final structure model of PspA covers residues Ile3 to Glu20 and Asp24 to Arg142, remaining residues were not visible in the electron density. Residues Asn12 to Leu31 show significantly higher conformational diversity than the rest of the structure as reflected by higher B-factors and poorly defined electron density in this region. It is assumed that the less ordered regions in the PspA structure are also the origin of the structural refinement converging at slightly higher R-factors (see Table S2). The structure was validated using Molprobity (Chen et al., 2010) and deposited at the Protein Data Bank (PDB) under the accession ID 4WHE.

LacZ-activity assays. Activity of *psp* reporters in different strains was assessed using the classic activity assay by Miller (Miller, 1972). Generally, all overnight cultures were diluted to an OD of 0.05 and allowed to grow for 3 more hours (induced with 0.1 % arabinose) before LacZ-activity was assessed. Where indicated ('18 h'), prolonged induction was used to discriminate between *psp* inhibiting and *psp*-inducing fragments of PspA. Only in those assays, media for overnight cultures already contained 0.1 % (w/v) arabinose to induce production of *psp* regulating fragments from pBAD vectors. All measurement were done in triplicate (error bars in figures correspond to one standard deviation). Where indicated, intrinsic controls [at least a *pBAD22* empty vector control (Guzman *et al.*, 1995) and a PspA₁₋₁₄₄ producing strain] were used for normalization to ensure comparability.

ATPase assays. Measurements were performed in assay buffer containing 20 mM Tris/HCl, 100 mM NaCl and 10 mM MgCl₂ at pH 8.0 in triplicate. As PspA stabilizes PspF hexamers and the PspA-PspF interaction was found to be extremely stable, care was taken to ensure that PspF₁₋₂₆₅ was equilibrated at nearly the final concentration for 1 h prior to addition of PspA at the indicated final concentrations from concentrated stock solutions. Following another 15 min of preincubation at 30°C, the assay was started by the addition of 2 mM ATP (final concentration) and samples were taken at the indicated time points. The amount of accumulated phosphate released during ATP hydrolysis was subsequently measured using a colorimetric assay (Lanzetta et al., 1979), modified as described (Turgay et al., 1997). In all cases, negative controls without PspF were carried along to subtract (the generally very low) background effects. For jump dilution experiments, PspA1-144 (30 µM) and PspF1-265 (20 µM) were preincubated for 60 min and the assay was started by diluting the sample 1:40 into assay buffer already containing 2 mM ATP. Samples were taken at the indicated time points. Samples containing 0.5 µM PspF and 0.75 µM PspA were prepared as described earlier and equilibrated for 60 min before starting the measurement. Their ATP hydrolysis rate (triplicate) was determined to be 0.1387 μ M s⁻¹ and served as reference. The k_{off} was subsequently estimated numerically with the nls algorithm of $R^3(200\ 000\ iterations)$ using the reference hydrolysis rate and the data obtained for the jump diluted sample (Copeland et al., 2011). With dissociation being a first-order reaction, the half-life of the complex could

then be calculated as $t_{1/2} = \frac{0.693}{k_{off}}$ (Tummino and Copeland, 2008).

Size exclusion chromatography. SEC (ÄKTA Explorer, GE Healthcare) was used to investigate complex formation of PspA₁₋₁₄₄ and PspF₁₋₂₆₅. Proteins (PspF₁₋₂₆₅ at 130 μ M, PspA₁₋₁₄₄ at 210 μ M) were preincubated at indicated molar ratios for 10 min at 4°C, 1 μ M of ATP (Carl Roth, Karlsruhe, Germany) or AMPPNP (Sigma-Aldrich, Taufkirchen, Germany) and 10 mM MgCl₂ were added where indicated. A 100 μ L aliquot of the sample was then applied to a Superose 6 10/300 GL column (GE Healthcare) equilibrated with 20 mM Tris/HCI and 100 mM NaCl adjusted to pH 8.0 at a flow rate of 0.5 ml min⁻¹ and 4°C. Elution profiles were obtained measuring the absorption of the effluent at 230 nm.

AUC. All measurements were performed in 20 mM Tris, pH 8, 100 mM NaCl at 20°C using an Optima XL-A centrifuge (Beckman, Palo Alto, CA, USA), an An50Ti rotor, and double-sector cells. Depending on protein concentration and addition of cofactors, e.g. ADP, the distribution of the protein in the cell was monitored at 230, 260, 280 or 300 nm. Data were analyzed using the software SedFit (Schuck, 2000). Isolated PspA₁₋₁₄₄ was investigated at concentrations of 3, 10 and 30 μ M. Sedimentation velocity measurements were made at 40 000 r.p.m. for 4 h, sedimentation equilibrium was performed at 14 000 r.p.m. Sedimentation of PspF was measured at concentrations of 1–30 μ M at 40 000 r.p.m. (velocity run) and 14 000 or 5000 r.p.m. (sedimentation equilibrium). Complex formation

of PspF with PspA₁₋₁₄₄ was determined at initial concentrations of PspF of 1, 10 and 30 μ M, respectively by titration of PspA₁₋₁₄₄ to maximal 60 μ M. Sedimentation velocity of the complex was measured at 40 000 r.p.m., sedimentation equilibrium at 5000 r.p.m.

In silico methods

Coiled-coil prediction. Coiled-coil predictions for PspA (Uniprot accession: P0AFM6) were performed using the COILS algorithm (Lupas *et al.*, 1991) with a prediction frame of 21 amino acids for most accurate identification of coiled-coil ends (according to the COILS/PCOILS manual, see also Gruber *et al.*, 2006) and MTIDK matrix. The window size of 21 amino acids also increases the prediction accuracy relative to smaller window sizes, yet it can be assumed to be sufficiently small to not decrease the resolution of the prediction (Gruber *et al.*, 2006). Results were similar whether weighting was enabled or disabled and did not change significantly using an aligned input sequence (PCOILS algorithm).

Consensus sequences. Sequences of γ -proteobacterial members of PspA/IM30 were obtained from the Pfam database (Punta *et al.*, 2012). Clearly too short sequences lacking larger parts of the primary sequence were discarded, yielding 715 sequences. Redundancies were reduced to prevent overrepresentations and the consensus logo histogram was calculated using Jalview (Waterhouse *et al.*, 2009). The overall PspA/IM30 consensus was similarly prepared using the Pfam RP35 representative proteome sequences of the PspA/IM30 family (165 sequences from all phyla). In all consensus histograms, a score of 0 indicates no conservation at this position, 100 indicates full conservation in all proteins.

Structure comparison and modelling. A DALI (Holm et al., 2008) search was performed with the PspA structure (PDB-ID 4WHE). For subsequent detailed comparison of ClpB and PspA coiled-coil structures, backbones of the coiled-coil forming protein moieties (ClpB₃₉₉₋₅₁₃, PDB-ID 4HSE; PspA₂₇₋ 141, PDB-ID 4WHE) were superimposed in the Swiss-PdbViewer (Guex and Peitsch, 1997) using 'magic fit' and 'explore fragment alternative fit' algorithms, as primary sequence alignment failed because of the difference in domain order in PspA and ClpB (see Fig. 3C). The highest scoring fit had an r.m.s.d. of 1.10 Å for 284 overlapping backbone atoms with an overall score of 68. This fit was used for all subsequent comparisons and simulations. Missing loops of PspF (PDB-ID 2BJW) and PspA (PDB-ID 4WHE) were modelled with COOT (Emsley et al., 2010). For oligomeric complex prediction of PspA·F, the hexameric oligomer of PspF was constructed using GalaxyGemini (Lee et al., 2013) with subsequent energy minimization. ClpB was then fitted into the PspF hexamer with Pymol (Schrodinger, 2010) using only the AAA+-domain of ClpB (PDB-ID 4HSE) for superimposition. PspA was then fitted into the M-domain of the previously aligned ClpB to obtain a model for the fully saturated PspA·F complex.

Simulations. The atomistic molecular simulations of PspA·F complexes were performed with Gromacs (Hess *et al.*, 2008) (version 4.6) using the AMBER99SB-ILDN force field

756 H. Osadnik et al.

(Lindorff-Larsen *et al.*, 2010) with the TIP4P water model. Virtual sites were used to allow the usage of a larger integration time step (0.003 fs). The Particle Mesh Ewald method was used to calculate electrostatic interactions. A simulation box was fitted around the protein complex consisting of 6 PspA and 6 PspF, which allowed 2 nm distance from the periodic boundaries. The temperature in the simulations was coupled to an external heat bath of 303 K using the velocity rescale method (tau_t = 0.1 ps). The external pressure was coupled to 1 bar using the Berendsen barostat (compressibility = $4.5 \times 10^{-5} \text{ bar}^{-1}$, tau_p = 1.0 ps). Four independent simulations were performed for PspA·F complexes either based on ClpB-like M-domain orientation (18 ns of simulated time) or with various upright orientations of PspA at the PspF hexamer surface for a simulated time of 129–145 ns (138 ns on average).

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Author contributions

HO and TB designed the experiments, MS did the crystallization and MS, CP, and MTS solved the structure of PspA, EH and DM contributed localization data and part of the activity data, HL did AUC analyses, HJR and HG contributed MD simulations, HO performed all other experiments, HO, HL, MTS, and TB evaluated the data and HO, MTS, and TB wrote the article.

Conflict of interest

The authors declare that they have no conflict of interest.

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