Properties of the phage-shock-protein (Psp) regulatory complex that govern signal transduction and induction of the Psp response in *Escherichia coli*

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The phage-shock-protein (Psp) response maintains the proton-motive force (pmf) under extracytoplasmic stress conditions that impair the inner membrane (IM) in bacterial cells. In *Escherichia coli* transcription of the *pspABCDE* and *pspG* genes requires activation of ρ^54^-RNA polymerase by the enhancer-binding protein PspF. A regulatory network comprising PspF–A–C–B–ArcB controls *psp* expression. One key regulatory point is the negative control of PspF imposed by its binding to PspA. It has been proposed that under stress conditions, the IM-bound sensors PspB and PspC receive and transduce the signal(s) to PspA via protein–protein interactions, resulting in the release of the PspA–PspF inhibitory complex and the consequent induction of *psp*. In this work we demonstrate that PspB self-associates and interacts with PspC via putative IM regions. We present evidence suggesting that PspC has two topologies and that conserved residue G48 and the putative leucine zipper motif are determinants required for PspA interaction and signal transduction upon stress. We also establish that PspC directly interacts with the effector PspG, and show that PspG self-associates. These results are discussed in the context of formation and function of the Psp regulatory complex.

INTRODUCTION

The phage-shock-protein (Psp) response maintains the proton-motive force (pmf) under extracytoplasmic stress conditions (e.g. upon secretin pIV production) that impair the integrity of the inner membrane (IM) and dissipate the pmf (reviewed by Darwin, 2005; see also Jovanovic et al., 2006). This adaptation to stress has been shown to be important for growth and virulence of some enterobacterial pathogens (reviewed by Darwin, 2005, 2007; Rowley et al., 2006). Transcription of the *psp* genes is controlled by ρ^54^-RNA polymerase and activated by the bacterial enhancer-binding protein PspF (reviewed by Model et al., 1997; Darwin, 2005; Wigneshweraraj et al., 2008). In *Escherichia coli* the PspF regulon consists of the *pspABCDE* operon, the adjacent *pspF* gene and the *pspG* gene (reviewed by Model et al., 1997; Darwin, 2005). In many other enterobacteria, the *pspF* and *pspABCDE* genes are highly conserved (Huvet et al., 2009). Under non-stress growth conditions, PspA inhibits the ATPase activity of PspF and negatively controls expression of the *psp* genes (Elderkin et al., 2002, 2005; Joly et al., 2009). It is proposed that under many *psp*-inducing stress conditions, the IM-bound sensors PspB and PspC act as positive regulators that receive and transduce the stress signal(s) to PspA via protein–protein interactions (Adams et al., 2003; Maxson & Darwin, 2006; Jovanovic et al., 2006, 2009). This ultimately results in the release of the PspA–PspF inhibitory complex, leading to *psp* induction and greatly elevated expression of the Psp proteins.

Recently, we provided evidence that during microaerobic growth the ArcAB system contributes to a Psp signal-transduction pathway in a PspBC-dependent manner and proposed that at least two signals could be recognized by PspBC for full pIV-dependent *psp* induction – caused by the mislocalization of pIV in the IM (Jovanovic et al., 2009). These data suggested that a double check-point for PspBC-specific *psp* induction could exist, potentially explaining the requirement for two sensor proteins. Importantly, in the absence of stress, PspBC overexpression has been shown to strongly induce *psp* (Weiner et al., 1991; Maxson & Darwin, 2006), implying that under these conditions, PspA inhibits the ATPase activity of PspF and negatively controls expression of the *psp* genes (Elderkin et al., 2002, 2005; Joly et al., 2009). It is proposed that under many *psp*-inducing stress conditions, the IM-bound sensors PspB and PspC act as positive regulators that receive and transduce the stress signal(s) to PspA via protein–protein interactions (Adams et al., 2003; Maxson & Darwin, 2006; Jovanovic et al., 2006, 2009). This ultimately results in the release of the PspA–PspF inhibitory complex, leading to *psp* induction and greatly elevated expression of the Psp proteins.

Abbreviations: Δψ, electron potential; Ara, L-arabinose; BACTH, bacterial two-hybrid; β-Gal, β-galactosidase; IM, inner membrane; LeuZ, leucine zipper motif; LeuZm, leucine zipper mutant; pIV, protein IV; pmf, proton-motive force; Psp, phage-shock protein; Sol, soluble fraction; MU, Miller units; TM, transmembrane.

Supplementary material is available with the online version of this paper.
conditions PspBC may bypass upstream signalling pathways and directly release the PspA–PspF inhibitory complex. However, the mechanistic basis of this effect is unknown. In light of these findings a Psp[1–A–C–B]–ArcB regulatory complex (under non-stress conditions) was proposed (Jovanovic et al., 2009), but importantly only pairwise interactions between PspA–PspF, PspA–PspC, PspB–PspC and PspB–ArcB have been identified (Elderkin et al., 2002; Adams et al., 2003; Elderkin et al., 2005; Maxson & Darwin, 2006; Jovanovic et al., 2009).

PspG is an IM effector protein not required for psp induction (Lloyd et al., 2004), but it may co-localize with PspA (Engl et al., 2009). Interestingly, it was observed that overexpression of PspG upregulates genes involved in microaerobic and anaerobic respiration (most of them ArcAB-regulated) (Jovanovic et al., 2006). The role of PspG and its potential association with the Psp regulatory complex has yet to be established.

In this work we dissect further the Psp regulatory complex in terms of protein–protein interactions and define the functional determinants in PspB and PspC that are required for pIV-dependent psp induction.

METHODS

Bacterial strains. The bacterial strains used in this study are listed in Supplementary Table S1, available with the online version of this paper. Strains were constructed by transduction using the P1<sub>mr</sub> bacteriophage (Miller, 1992).

Media and growth conditions. All strains were routinely grown under microaerobic conditions in Luria–Bertani (LB) broth or on LB agar plates at 37 °C (Miller, 1992). For microaerobic growth, overnight cultures of cells were diluted 100-fold (OD<sub>600</sub> ~0.025) and shaken at 100 r.p.m. For the bacterial two-hybrid (BACTH) assays, strains were grown in LB at 30 °C. For single-molecule fluorescence imaging of PspG–GFP, cells were grown in minimal medium as described previously (Engl et al., 2009). Induction of the pBAD ara promoter was achieved with either 0.001 % or 0.02 % (final 1 : 5000) 3-aminopyridine (ara). Antibiotics were used at the following concentrations: ampicillin, 100 μg ml<sup>−1</sup>; kanamycin, 25 or 50 μg ml<sup>−1</sup>, as indicated; chloramphenicol, 30 μg ml<sup>−1</sup>; and tetracycline, 10 μg ml<sup>−1</sup>.

DNA manipulations. Plasmids used in this study are listed in Supplementary Table S1. All constructs were verified by DNA sequencing, and protein production was verified using Western blotting. By site-directed mutagenesis (QuikChange, Agilent Technologies) using the appropriate primers and either pAJM2 or pGJ57. Using either pAJM8, pAJM13 or pAJM11 as a template, we constructed pAJM6–8, pAJM11–13, pGJ54, pGJ55 and pGJ56. The BACTH fusion template, we constructed pAJM6–8, pAJM11–13, pGJ54, pGJ55 and pGJ56. The BACTH fusion

In vivo bacterial BACTH system. The Cya-based BACTH assay was used to study in vivo protein–protein interactions (Karimova et al., 1998, 2005). Protein fusions (see Table 1 and Fig. 1) were assayed in BTH101 cells as described previously (Jovanovic et al., 2009). Interactions were quantified by measuring β-galactosidase activity in liquid cultures (see below). Chromosomal LacZ expression threefold above the negative control (vector alone) value was scored as a positive interaction signal.

To test protein–protein interactions in the absence of newly produced proteins that may provide or facilitate binding interactions by a bridging effect, we constructed a derivative of the BTH101 strain carrying the ΔpspF mutation – thereby preventing expression of the native Psp proteins (see Supplementary Table S2). The wild-type pspF gene was replaced by ΔpspF:: Kan, yielding MVA99 (see Supplementary Table S1). The Kan cassette from ΔpspF:: Kan was eliminated and the marker-less pspF99 variant (strain MVA100) constructed using plasmid pCP20 and the method described by Cherepanov & Wackernagel (1995) (see Supplementary Table S1). The strain was verified by PCR.

β-Galactosidase (β-Gal) assays. Activity from a chromosomal P<sub>lacZ</sub>–Gal transcriptional fusion was assayed to gauge the level of psp expression, while activity of the native chromosomal lacZ was measured in the BACTH assays. Overnight cultures grown at 37 °C (or 30 °C for the BACTH assays) were diluted 100-fold and grown under the same conditions until mid-exponential phase. Cultures were induced (with Ara or 0.5 mM IPTG for BACTH, as indicated) for 1 h and then assayed for β-Gal activity (Miller, 1992). For all β-Gal assays, mean values from six samples taken from technical duplicates of three independently grown cultures of each strain were used to calculate activity. The data shown in the figures are the mean values with ± standard error bars.

Bacterial cell fractionation. Bacterial cell were harvested at mid-exponential phase and resuspended in a mix of 30 μl 4 % SDS and 30 μl Laemml buffer (Sigma). Samples were normalized according to cell growth measured as OD<sub>600</sub> separated by SDS-PAGE and transferred onto PVDF membrane using a semidry transblot system (Bio-Rad). Western blotting was performed as described by Elderkin et al. (2002) using antibodies to PspA (1 : 10000) (Jones et al., 2003), PspB (1 : 5000), PspC (1 : 5000), PspG (1 : 1000) (Jovanovic et al., 2006) or pIV (1 : 10000). Proteins were detected using the ECL plus Western Blotting Detection Kit (GE Healthcare). Images were captured in a FujiFilm Intelligent Dark Box by an image analyser with a charge-coupled-device camera (LAS-3000). Densitometry analysis was performed with MultiGauge 3.0 software (FujiFilm USA) and quantification (results expressed in arbitrary units) was performed using the AIDA software.

Confocal fluorescence microscopy to assess membrane electron potential (ΔΨ). The ΔΨ component of pmf was measured using the JC-1 dye method (Becker et al., 2005) as described previously (Jovanovic et al., 2006; Engl et al., 2009). The green to red fluorescence emission ratio (530/590 nm) was calculated from 100 individual cells taken from technical duplicates of three independently grown cultures of each strain. The threshold to distinguish pmf differences was chosen based on the ΔΨ (equivalent to pmf under given experimental conditions) of the WT strain under non-stress growth conditions.

Single-molecule fluorescence imaging of PspG in vivo. Single molecules in fluorescent PspG–GFP complex (MG1655 ΔpspF/GspG– GFP) were imaged in vivo (Engl et al., 2009). PspG–GFP expressing cells
were mounted on a cover glass and images taken at 80 ms per frame. The fluorescent PspG–GFP complexes were analysed using the ImageJ software (Abramoff et al., 2004), with GFP from MG1655/pDSW209 cell lysates as a reference.

**Protein purification.** *E. coli* Top10 or Top10ΔpspA cells transformed with either pRD047 (encoding PspB and non-tagged PspC) or pRD047His (encoding PspB and His-tagged PspC) were grown in LB at 37 °C until they reached OD$_{600}$ ~0.5, at which point the culture was shifted to 30 °C. Overexpression was induced by addition of 0.02 % Ara and the cultures grown for a further 3 h. Cells were harvested at 8000 r.p.m. for 10 min at 4 °C; the pellet was resuspended in 20 ml buffer A (50 mM sodium phosphate pH 7, 300 mM NaCl, 5 %, v/v, glycerol) supplemented with Complete Protease Inhibitor (Roche) and lysed by sonication. The supernatant was loaded onto a 5 ml HiTrap metal-chelating column (GE Healthcare) pre-charged with Ni$_2^+$ and the protein eluted with buffer A supplemented with 1 M imidazole. Purified proteins were detected by Western blotting (see above).

**Bioinformatic methods.** Multiple sequence alignment was performed using Multalin software version 5.4.1 (http://bioinfo.genotoul.fr/multalin/) (Corpet, 1988). Protein domain analysis was performed using Pfam version 21.0 (http://www.sanger.ac.uk/pfam) (Finn et al., 2006). Transmembrane (TM) helices prediction, localization and topologies were performed using TMHMM (http://www.cbs.dtu.dk/services/TMHMM/, V 2.0) (Kahsay et al., 2005), HMMTOP (http://www.enzim.hu/hmmtop/, V 2.0) (Tusnády & Simon, 2001), TopPred2 (http://mobyle.pasteur.fr/cgi-bin/portal.py?form=toppred) (von Heijne, 1992) or TOPCONS (http://topcons.net/) (Bernsel et al., 2009).

**RESULTS AND DISCUSSION**

**PspG associates with Psp proteins known to form regulatory complexes**

Previous studies suggested the presence of Psp regulatory complexes (Adams et al., 2003; Maxson & Darwin, 2006; Jovanovic et al., 2009). To determine whether PspG is able to interact with any of the proteins that constitute these regulatory complexes we performed pairwise BACTH interaction assays with PspA, PspB, PspC, ArcB and PspG (Table 1 and data not shown). The topologies of PspB, PspC and PspG (Fig. 1a) (reviewed by Darwin, 2005; was loaded onto a 5 ml HiTrap metal-chelating column (GE Protease Inhibitor (Roche) and lysed by sonication. The supernatant was shifted to 30 °C. Overexpression was induced by addition of 0.02 % Ara and the cultures grown for a further 3 h. Cells were harvested at 8000 r.p.m. for 10 min at 4 °C; the pellet was resuspended in 20 ml buffer A (50 mM sodium phosphate pH 7, 300 mM NaCl, 5 %, v/v, glycerol) supplemented with Complete Protease Inhibitor (Roche) and lysed by sonication. The supernatant was loaded onto a 5 ml HiTrap metal-chelating column (GE Healthcare) pre-charged with Ni$_2^+$ and the protein eluted with buffer A supplemented with 1 M imidazole. Purified proteins were detected by Western blotting (see above).

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et al. (2005), Jones et al. (2003), Kleerebezem et al. (1996) and Engl et al. (2009), respectively. Per, periplasm; IM, inner membrane; Cyt, cytoplasm; N, N-terminus of the protein; C, C-terminus of the protein. (b) Representation of the predicted conserved domain organization of PspC: the cytoplasmic extrusion domain (Cyt, residues 1–39), the transmembrane portion (TM, residues 40–64), the periplasmic extrusion domain (Per, residues 65–119) and the conserved PspC domain (residues 1–68, light grey). The putative leucine zipper motif (LeuZ; residues 77–98) is as indicated. The numbering refers to E. coli PspC.

Results from the BACTH assay also indicated that PspG, when expressed from the high-copy-number plasmid pUT18, interacts with ArcB (data not shown). However, when PspG was expressed from the low-copy-number plasmid pKNT25 no interaction with ArcB was detected (Table 1). As we were unable to detect an interaction between PspG and ArcB with the low-copy-number plasmid expressing PspG, we infer that the observed interaction either may be non-specific or may reflect the action of PspG as an effector when overproduced (Jovanovic et al., 2006) – the later possibility has yet to be explored.

PspC does not stably self-associate

Peak fractions from the PspBC\textsubscript{6His} purification were analysed by SDS-PAGE (Fig. 2a) and found to contain a number of PspC cross-reacting species (Supplementary Figs S1Ai, S2C, Dii, S3Bi and C) – one which migrates as a monomer and one as a putative dimer. It has been proposed that changes in pmf that occur upon extracytoplasmic stress of PspC acts as a psp inducer, causing a drop in pmf – see below). We specifically selected PspC given that the pairwise interaction data suggest that PspC interacts with PspA, PspB and PspG (Table 1). We reasoned that PspC would, by means of protein–protein interactions, specifically pull down Psp proteins involved in the regulatory complex during protein purification (via the His-tag on PspC). As a control, we performed the chromatography assay with non-tagged PspC to demonstrate that any Psp proteins identified in the eluted fractions are present due to a specific interaction with PspC\textsubscript{6His}. When we analysed the PspC\textsubscript{6His} elution fractions we found that PspA, PspB and PspG all co-purified with PspC\textsubscript{6His} (Fig. 2a–d) – co-purification of PspA,B,C suggests that a Psp[A–C–B] regulatory complex exists, to which PspG may associate. PspG also co-purifies with PspC\textsubscript{6His} when expressed in cells lacking PspA (Fig. 2e), further indicating that PspG interacts directly with PspC.

Potentially, co-expression of PspBC leads to an ‘active’ form of Psp(B)C that can specifically interact with PspA and possibly PspG. Such complexes may have significant implications in understanding the signalling cascade and activation of the Psp response as well as the effector function of PspG. Importantly, a reduced psp operon, \textit{pspACG}, found in \textit{Aeromonas} species (M. P. H. Stumpf, personal communication) further suggests that functional interactions occur between PspA, PspC and PspG.

Given the ability of PspG to self-associate (as suggested in the BACTH assay; Table 1) and since the oligomeric state of PspG could be important for its effector function, we analysed PspG–GFP complexes in live MG1655\textit{paspG} cells using single-molecule fluorescence imaging and objective-type total internal reflection fluorescence (TIRF) (see Methods, Engl et al., 2009 and e.g. Haggie & Verkman, 2008). The results illustrate that PspG–GFP can form complexes as large as pentamers/hexamers but the majority are dimers/trimers (Fig. 3).

In order to provide evidence that Psp proteins may indeed form a complex and to determine whether PspG is truly associated with PspC, we performed pull-down assays using PspC\textsubscript{6His} co-expressed with WT PspB (since overexpression of PspC acts as a psp inducer, causing a drop in pmf – see below). We specifically selected PspC given that the pairwise interaction data suggest that PspC interacts with PspA, PspB and PspG (Table 1). We reasoned that PspC would, by means of protein–protein interactions, specifically pull down Psp proteins involved in the regulatory complex during protein purification (via the His-tag on PspC). As a control, we performed the chromatography assay with non-tagged PspC to demonstrate that any Psp proteins identified in the eluted fractions are present due to a specific interaction with PspC\textsubscript{6His}. When we analysed the PspC\textsubscript{6His} elution fractions we found that PspA, PspB and PspG all co-purified with PspC\textsubscript{6His} (Fig. 2a–d) – co-purification of PspA,B,C suggests that a Psp[A–C–B] regulatory complex exists, to which PspG may associate. PspG also co-purifies with PspC\textsubscript{6His} when expressed in cells lacking PspA (Fig. 2e), further indicating that PspG interacts directly with PspC.

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may cause the stable self-association of PspC – reported by Adams et al. (2003) as an SDS-PAGE-stable dimeric form of PspC. Although we showed that co-expression of PspBC does not change pmf (see below), and BACTH analysis failed to detect pairwise PspC–PspC interactions in either Yersinia enterocolitica (Maxson & Darwin, 2006) or E. coli (Table 1), we cannot discount the possibility that either (i) the conformation of the fusion proteins is unfavourable for detecting PspC self-association or (ii) PspC may stably associate with an unknown protein to form the apparent ‘dimer’ species. Alternatively, PspC self-association may only occur upon pIV stress or when PspC is highly overproduced, since the apparent dimer was only observed when PspC was overproduced in a ΔpspC strain (Supplementary Fig. S3Bi and C). To investigate this further, and to address whether formation of a putative PspC dimer can be observed under physiological conditions, we analysed the expression of chromosomally encoded PspC in WT cells and cells lacking the negative regulator PspA (ΔpspA) in the presence and absence of pIV (Fig. 4). Fully consistent with its role in psp induction, pIV elevated Psp expression in WT cells (observed as an increase in monomeric PspC protein levels). Further, in cells lacking PspA, monomeric PspC expression remained high regardless of the presence of pIV (indicative of deregulated Psp expression). Notably, the intensity of the slower-migrating anti-PspC-reactive complex (the putative PspC dimer) remained unchanged under conditions where PspC was expressed at different levels (Fig. 4), suggesting that under physiological conditions

Fig. 2. Co-purification of PspA, PspB and PspG with PspC6His. PspC6His co-expressed with PspB (pRD047His) was purified using metal-affinity chromatography (see Methods). As a control non-tagged PspC co-expressed with PspB (pRD047) was also purified. Peak fractions from both purifications (corresponding to fractions 2–4), were visualized using Western blotting and (a) anti-PspC, (b) anti-PspB, (c) anti-PspA or (d) anti-PspG antibodies as indicated. In (a) the positions of monomeric PspC (PspC; arrow) and an additional slower-migrating anti-PspC-reactive (double asterisk) species are as indicated. PspB (b), PspA (and non-specific band, asterisk) (c) and PspG (d) co-purified with PspC6His fractions (2–4) but not with PspC (non-tagged). (e) PspC6His co-expressed with PspB was also purified from a ΔpspA strain and the peak fractions were visualized using Western blotting and anti-PspG antibodies. The positions of molecular mass marker proteins (kDa) are indicated.

Fig. 3. PspG–GFP forms higher oligomers in vivo. The number of molecules in fluorescent PspG–GFP complexes was estimated using ImageJ software. The intensity of a single pixel in a fluorescent PspG–GFP (pGJ7) cluster expressed in E. coli MG1655 ΔpspG cells (MVA40) was measured and compared to GFP fluorescence in E. coli MG1655 cell lysates harbouring pDSW209 (GFP alone). (a) The mean fluorescence intensity of 50 PspG–GFP complexes within living cells was on average at least three times higher than that of GFP spots, suggesting that PspG can form at least a dimer/trimer in vivo. (b) The frequency distribution among the 50 complexes analysed illustrates that PspG–GFP self-assembles into a single major distinct oligomeric class.
PspC does not form a stable complex with either itself or PspA (since an identical anti-PspC-reactive complex, the putative PspC dimer, is observed upon overexpression of PspC in a ΔpspA strain). The lower level of PspC expression in cells encountering pIV stress versus cells lacking negative regulation (Fig. 4) is fully consistent with our findings that even under pIV stress conditions, PspA is still imposing apparent negative regulation (Engl et al., 2009; Jovanovic et al., 2009).

These results show that the apparent dimer species recognized by the PspC antibodies does not follow PspC expression levels, and therefore should not be considered as a stable PspC dimer. Taking these findings together with the BACTH assays, we conclude that under the physiological conditions tested PspC does not stably self-associate.

**PspB and PspC act together to directly release negative regulation of psp**

It is formally possible that the PspBC–PspA interactions that occur under non-stress conditions contribute to a steady-state level of a Psp[F–B]–ArcB regulatory complex. Alternatively, since the absence of PspBC does not release the negative regulation, and so cannot be required for formation of the inhibitory complex (Jovanovic et al., 2009; Gueguen et al., 2009), Psp[F–A] and Psp[C–B]–ArcB subcomplexes may communicate only in the presence of stress. Nevertheless, in both cases the stress-induced activation of psp may follow conformational changes in PspB and PspC that lead to release of negative regulation of PspA, presumably through altered protein–protein interactions. Interestingly, overexpression of PspC or PspBC, but not PspB alone, induces psp (Weiner et al., 1991; Maxson & Darwin, 2006) (also see Fig. 5a and Supplementary Fig. S1A). Since overexpression of PspC decreases pmf in ΔpspF cells (i.e. where the Psp response is absent) (Fig. 5b) it may, similarly to pIV (Jovanovic et al., 2006, 2009) or PulD (Guilvout et al., 2006), act as an inducing stress for psp (Fig. 5a). In contrast, overexpression of PspBC strongly induces psp (Fig. 5a), but does not reduce pmf (Fig. 5b). Apparently, PspB (in the context of the PspBC complex) appears to prevent PspC from changing pmf, consistent with the observation that the growth rate in exponential phase is normal in ΔpspC cells overexpressing PspBC but severely impaired (10-fold) in ΔpspC cells overexpressing PspC alone (data not shown). Since PspB counteracts the ‘toxic’ effect of PspC, the PspBC proteins may be acting as an antitoxin–toxin pair as suggested by Brown & Shaw (2003). A parallel study (Gueguen et al., 2009) showed that in Y. enterocolitica PspB stabilizes expression of WT PspC, a result differing from our E. coli data (Supplementary Fig. S1A).

Notably, overexpression of PspBC results in higher psp induction than does overexpression of PspC alone (Fig. 5a) (consistent with Weiner et al., 1991 and Maxson & Darwin, 2006) – at a level comparable to uncontrolled psp expression in ΔpspA cells (Fig. 5c). In line with this observation, co-expression of PspBC in the presence of pIV did not result in increased psp induction (Fig. 5c and Supplementary Fig. S1B). Since PspBC overexpression does not simply induce psp by decreasing pmf (i.e. the overproduced proteins do not in themselves generate a ‘stress’ signal) we suggest that overproduced PspBC resembles the ‘stressed on-state’ of the Psp response.

The effect of overproduced PspBC could directly alter the state of PspA, and hence negative regulation, thereby bypassing upstream signalling systems (e.g. the ArcAB system). To test this directly, we used a leucine zipper motif (LeuZ) mutant (LeuZm) of PspB (PspB[LeuZm]) that maintains the ability to interact with PspC but is no longer able to interact with ArcB and is therefore inactive in upstream signalling (Jovanovic et al., 2009). When co-expressed with PspC, this mutant form of PspB should support psp induction independent of any upstream signalling event. Significantly, overexpression of PspB[LeuZm]C resulted in a similar level of psp induction as overexpressed WT PspBC (Fig. 5c). In line with this, overexpression of either PspBC or PspB[LeuZm]C strongly induces psp in the absence of ArcB (ΔarcB strain) (Fig. 5c). These data argue that a PspBC

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**Fig. 4.** Oligomerization state of PspC. Western blots (using anti-PspC) illustrating PspC expression from the chromosome in either WT (MG1655) or ΔpspA (MG1655ΔpspA) cells (in the absence or presence of pIV). The positions of species that specifically cross-react with anti-PspC are highlighted as monomeric PspC (13.5 kDa; arrow) and a putative dimer (double asterisk). The positions of the marker proteins (kDa) are indicated. Below: the relative expression levels of monomeric PspC (labelled as PspC monomer) and the putative PspC dimer (double asterisk) were quantified within each strain tested, and the results expressed as a percentage of the induced corresponding protein band (+pIV; lanes 2 and 4, 100%). ‘Proteins’ refers to the loading control. Importantly, these results demonstrate that in the absence of PspA (lanes 3 and 4), the relative expression levels of monomeric PspC are clearly highly elevated compared to WT in the presence of pIV, whereas the putative dimer expression level remains relatively unchanged, suggesting that this band corresponds to an unspecific anti-PspC cross-reacting species.
complex is sufficient to induce the Psp response, by relieving PspA-imposed inhibition, in the absence of ArcB.

**PspC residues 40–119 are required for psp induction**

We next addressed which determinants were important for PspBC–PspA signalling. Recall that PspA–PspC and PspB–PspC, but not PspB–PspA, were shown to directly interact (see above), suggesting that PspC may have a central role in transducing the stress signal to PspA. We first truncated PspC based on its predicted domain structure and topology (Fig. 1) to form three PspC fragments: PspC<sub>1–68</sub> (cytoplasmic and TM domain), PspC<sub>40–68</sub> (TM domain) and PspC<sub>40–119</sub> (TM-periplasmic domain). In the absence of PspC (ΔpspC), induction of the psp response upon pIV stress no longer occurs (reviewed by Darwin, 2005; see also Jovanovic et al., 2009). We initially introduced into ΔpspC cells either full-length PspC (residues 1–119) or one of the three fragments to test their activity with regard to their ability to support psp induction upon pIV stress. Each protein was expressed at a level similar to the basal level that was insufficient to induce psp in the absence of stress, and was scored for supporting pIV-dependent induction of psp (Fig. 6a). Only full-length PspC complemented the ΔpspC mutation for psp induction by pIV. Notably, similar levels of pIV expression were detected in all the samples tested (Supplementary Fig. S2A), and expression of wild-type PspC was at least 20-fold lower than that following pIV induction of chromosomal psp (Supplementary Fig. S2B). These results establish that full-length PspC, but not its fragments, supports signal transduction to PspA in pIV-dependent induction of psp.

PspC and its fragments (tested separately or co-expressed with PspB) were strongly overexpressed in ΔpspC cells at a level that was sufficient for PspC or PspBC (see Fig. 5a) to induce psp. Under these conditions the PspC<sub>40–119</sub> fragment induced psp expression even in the absence of co-expressed PspB (Fig. 6b). We do note that expression levels of the PspC fragments varied greatly (Supplementary Fig. S2C); however, these differences do not account for the inability of some fragments to induce psp. For example, the PspC<sub>1–68</sub> and PspC<sub>40–119</sub> fragments are expressed similarly in the absence of PspB (Supplementary Fig. S2i and Di), yet only the PspC<sub>40–119</sub> fragment induces psp (Fig. 6b). Further, the fragments did not appear to mislocalize (Supplementary Fig. S2D). Notably, expression of the PspC<sub>40–68</sub> and PspC<sub>40–119</sub> fragments seemed to be stabilized in the presence of co-overexpressed PspB, potentially via interactions between the TM domains of these two proteins (see below). As an additional control, we determined Δψ in ΔpspF cells (where psp-inducing agents, such as pIV, decrease pmf in the absence of the Psp response) over-expressing the PspC fragments, including PspC<sub>40–119</sub> (in either the absence or presence of PspB), and found that pmf was not decreased (Fig. 6c), establishing that PspC<sub>40–119</sub> does not act as a typical psp-inducing stress agent. Importantly, overexpression of the PspC<sub>1–68</sub> fragment in the absence of co-expressed PspB decreased pmf (Fig. 6c) and impaired cell growth in ΔpspC cells similarly to full-length PspC (data not shown), but was unable to induce psp. Therefore, even when pmf was reduced, the PspC<sub>1–68</sub> fragment was unable to complement a ΔpspC mutation and support psp induction. These data suggest that the PspC<sub>1–68</sub> sequence lacking the putative periplasmic domain is important for the ‘toxic’ effect of PspC (i.e. a drop in pmf, see above). The PspC putative periplasmic domain (residues 69–119, present in the PspC<sub>40–119</sub> fragment that induces psp but not in the PspC<sub>1–68</sub>
or PspC 40–68 fragments, which do not induce psP) can function independently of upstream signalling and so may be directly involved in the release of the PspA–PspF inhibitory complex and hence functions positively in psP induction.

The integrity of the putative leucine zipper motif of PspC is required for psP induction

The fragmentation approach demonstrated that the putative periplasmic region of PspC may be involved in psP induction and hence presumably might receive the inducing signal or contact PspA in order to release the negative regulation that PspA imposes on PspF.

Initially we analysed the ability of each of the PspC fragments to interact with PspB and PspA. Significantly, we observed that all the PspC fragments were capable of interacting with PspB or PspBLeuZm, demonstrating that this interaction (and also PspB self-association) does not require the PspB LeuZ (Table 1 and data not shown). These data also suggest that the determinants in PspC required for the interaction with PspB lie within the putative TM domain (residues 40–68) (Table 1 and Supplementary Fig. S2C). Importantly, and in contrast to WT PspC, two of the three PspC fragments failed to interact with PspA; only the C-terminal PspC 40–119 Cya fusions interacted with PspA (Table 1 and Supplementary Table S2). This is somewhat surprising given the proposed topology of PspC, where the C-terminus is facing the periplasm (Kleerebezem et al., 1996), suggesting that in the absence of the predicted cytoplasmic domain, the PspC 40–119 fragment may adopt another topology.

The sequence assigned to the PspC periplasmic domain contains a putative LeuZ (Kleerebezem et al., 1996) (Fig. 1b). When we examined the activities of LeuZ mutants (LeuZm) in the context of full-length PspC (PspCLeuZm) and the PspC 40–119 fragment (PspC 40–119 ) (see Supplementary Table S1) we found that both proteins failed to interact with PspA (Table 1 and Supplementary Table S2) but maintained an ability to interact with PspB (Table 1). Given that the LeuZ of PspC appears important for PspA interactions, we reasoned that PspC proteins that contain a disrupted LeuZ would no longer be active for psP induction (assuming that they no longer relieve the inhibition imposed by PspA that is mediated by protein–protein interactions). Since full-length PspC is strictly required for psP induction upon pIV stress (Fig. 6a), and to avoid affecting pmf (Fig. 6c), the signal transduction complementation assays were performed in ΔpspF cells co-expressing...
PspB and full-length PspC (Fig. 7a). PspBC\textsubscript{leuZm} when expressed at a low level in the presence of pIV was unable to fully induce the psp response (Fig. 7a; compared to WT PspBC). The expression level of pIV under all conditions assayed was similar (Supplementary Fig. S3A), indicating that differences that arise in psp induction are unlikely to be due to variations in the level of the primary stress signal. The LeuZ of PspC appears to be important for signalling and/or the outcome of signalling to PspA.

When co-expressed with PspB, the PspC LeuZm in the context of either WT PspC (PspC\textsubscript{LeuZm}) or the PspC\textsubscript{40–119} fragment (PspC\textsubscript{LeuZm}{40–119}) was either able (to some extent) or unable, respectively, to induce psp (Fig. 7b). Significantly, the expression level and localization of PspC\textsubscript{LeuZm} were similar to those of WT PspC (Supplementary Fig. S3B and C), further suggesting that a loss of productive interactions with PspA and the subsequent reduction in psp induction is not simply a consequence of protein misfolding or mislocalization.

Recent parallel studies (Gueguen \textit{et al.}, 2009) demonstrated that in \textit{Y. enterocolitica} the periplasmic region and LeuZ of PspC were important in psp induction. Gueguen \textit{et al.} (2009) proposed a model in which the periplasmic region of PspC (more specifically the LeuZ) receives the stress signal rather than directly interacting with PspA. In addition, they showed that complementation of a chromosomal pspC deletion with PspC variants harbouring cytoplasmic domain truncations led to constitutive psp expression. These outcomes resemble the activities of the PspC\textsubscript{40–119} fragment reported here. Gueguen \textit{et al.} (2009) proposed that the cytoplasmic domains of PspC and PspB are involved in the negative regulation of psp in the absence of stress, and then interact with the Psp\textsubscript{[A–F]} inhibitory complex upon stress to release PspF negative regulation. However, Gueguen \textit{et al.} (2009) found that the PspC cytoplasmic domain truncations still required the putative periplasmic domain to exhibit constitutive psp expression. Our results suggest that the PspC putative periplasmic domain may function in a direct interaction with PspA for release of negative regulation rather than just in receiving the inducing signal.

\textbf{Characterization of the PspC\textsubscript{69–119} fragment}

To further dissect the possibility that the putative PspC periplasmic domain (residues 69–119) directly interacts with PspA, we characterized the PspC\textsubscript{69–119} fragment (see Supplementary Table S1), which contains the sequence predicted to reside in the periplasm. PspC\textsubscript{69–119} expressed in the absence (Supplementary Fig. S2Ci) and presence (Supplementary Fig. S2Cii) of PspB was detected within the soluble cell fraction (Supplementary Fig. S2Dii). Overexpression of PspC\textsubscript{69–119} does not significantly change $\Delta$V (data not shown). Similarly to PspC\textsubscript{40–119}, overexpression of PspC\textsubscript{69–119} in the absence of stress strongly induced psp but only when co-expressed with PspB (Fig. 7b). In addition, induction of psp, by overexpressed

\begin{figure}[h]
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\includegraphics[width=\textwidth]{Fig7}
\caption{PspC determinants involved in signal transduction and induction of psp. (a) The PspC LeuZ and residue G48 are required for pIV-induced psp expression. Induction of the chromosomal $\phi(pspA\text{-}lacZ)$ fusion in a $\Delta$pspC strain (MVA13) expressing low-level PspBC (pAJM3) or PspBC mutants (PspC mutants: G48A, pGJ54; G74A, pGJ55; LeuZm, pGJ57) (using 0.001% Ara) in the absence or presence of pIV (pGJ4) (see Methods). As a control, pIV-dependent induction of psp in WT cells (MVA4) is presented. (b) High-level co-expression of PspBC\textsubscript{G48A}, PspBC\textsubscript{LeuZm}, PspBC\textsubscript{LeuZm}{40–119} (pGJ60) and PspBC\textsubscript{LeuZm}{69–119} (pGJ61) mutants failed to directly induce psp expression. Induction of the chromosomal $\phi(pspA\text{-}lacZ)$ fusion in a $\Delta$pspC strain (MVA13) by overexpression of PspBC or PspBC mutants (using 0.02% Ara). (c) The PspC periplasmic region may exist in two topologies: schematic illustrating the potential topologies of PspC (A, B or C) and PspC\textsubscript{40–119} (the periplasmic region containing the LeuZ; D).}
\end{figure}
PspC₄₀₋₁₁₉ or PspC₆₉₋₁₁₉ is independent of ArcB (data not shown). These data clearly indicate that PspB, in concert with the periplasmic domain of PspC, directs the release of negative regulation of psp expression. The Cya–PspC₄₀₋₁₁₉ fusion interacted with Cya–PspA (but not PspB–Cya) only when co-expressed with WT PspB (Table 1 and Supplementary Table S2). PspC₄₀₋₁₁₉ (see Supplementary Table S1 and Supplementary Fig. S2Cii), when co-overexpressed with PspB, was unable to induce psp (Fig. 7b) or interact with PspA (Table 1 and Supplementary Table S2). These results are consistent with the previously demonstrated PspA–PspB interaction observed only in the presence of PspC (Adams et al., 2003) and here we show that the PspC LeuZ is probably needed for this interaction. Taken together with results obtained with the PspC₆₉₋₁₁₉ fragment (see above), direct PspC₆₉₋₁₁₉ binding interactions with PspA seem to support induction of psp.

Our results show that the LeuZ located in the putative periplasmic domain of PspC is strictly required for PspA interactions and by clear inference psp induction – potentially explaining why PspC constitutive mutants still require the periplasmic domain for activation of psp expression (Gueguen et al., 2009). The results obtained with the PspC₄₀₋₁₁₉ and PspC₆₉₋₁₁₉ fragments in terms of psp induction and the BACTH assays, and assuming the periplasmic domain of PspC should interact with the cytoplasmic face of PspA, suggest that the periplasmic domain of PspC would favour one topology when pIV is expressed in order to directly transduce the stress signal to the Psp[A–F] regulatory complex. A phenocopy of the stress-induced ‘topology switch’ may be simply realized by mass-action effects: in the absence of stress PspC may exist in equilibrium between active and inactive conformations potentially reflecting two distinct topologies of PspC (and so controlling basal level expression of psp genes); overproduction of PspC would result in more of the activating ‘on-state’ PspC topology being present in the cell, resulting in psp induction.

PspC residue G₄₈ is a determinant in psp induction

The possibility that the periplasmic domain of PspC may exist in two states ('in' and 'out'), possibly subcellular localization in response to stress, prompted us to apply the positive-inside rule (von Heijne, 1992) to the PspC sequence using different topology prediction programs (see Methods and Supplementary Fig. S4). The results were as follows: TMHMM – N-terminus inside, C-terminus outside; HMMTOP – N-terminus outside, C-terminus inside; TopPred – N terminus outside, C-terminus inside; TopCONS – N-terminus inside, C-terminus inside with two potential TM domains and a small periplasmic loop (see Supplementary Fig. S4). These programs indicate that both topology scenarios (with respect to the periplasmic domain) are predicted for PspC, suggesting that the barrier between these two states is not high and that a topology switch potentially exists (Supplementary Fig. S4). In contrast, if we use the same programs with either PspB or PspG a single topology outcome is obtained, suggesting a relatively stable single topology state (data not shown). The localization of positive residues in extramembrane domains is important for the positive-inside rule application and depends on Δψ, so protein topology can be dynamic (Bogdanov et al., 2002, 2008; Zhang et al., 2003, 2005). Potentially relevant for the Psp system and the PspBC sensors is that the observed reversible topological organization within some E. coli membrane proteins (e.g. LacY, PheP, GabP) is governed by a change in membrane phospholipid composition (reviewed by Bogdanov et al., 2009; Dowhan & Bogdanov, 2009). Further, kinked hydrophobic TM helices (such as the ones obtained in the PspC topology predictions; Supplementary Fig. S4), particularly those containing glycine (G) residues, are proposed to bend more easily within the membrane than those with straight TM helices (Jiang et al., 2002). Notably, these glycine residues are also thought to mediate the protein–protein interactions in polytopic membrane proteins (Javadpour, et al., 1999). PhoA fused to the PspC region predicted to reside in the periplasm showed activity (Kleerebezem et al., 1996), but it has not been discounted that this region can also at times be localized in the cytoplasm (e.g. using a fusion to β-galactosidase).

In light of the data concerning polytopic membrane proteins, we analysed the sequence of the putative TM helix (residues 40–68) of PspC and located a highly conserved glycine residue at position 48. When we mutated this residue to alanine (PspC₄₄₈ₐ₅ₐₚ₅ₐ) we found that, similar to PspC₄₄₈ₐ₅ₐₚ₅ₐ, PspC₄₄₈ₐ₅ₐₚ₅ₐ failed to induce the psp response (Fig. 7a, b and Supplementary Fig. S3B, C). In agreement with these observations, the BACTH assay illustrates that PspC₄₄₈ₐ₅ₐₚ₅ₐ is no longer able to interact with PspA (or, interestingly, PspG); however, its ability to interact with PspB remains unaffected (Table 1). As a control we constructed the PspC₄₄₈ₐₓₕ₆₅ₐ mutant, where residue G₇₄ is predicted to lie within the periplasmic domain and as such should not affect the topology of the TM domain. As predicted, this mutant retained WT PspC activities in terms of signal transduction and psp induction (Fig. 7a, b and Supplementary Fig. S3B, C) and Psp protein interactions (Table 1). These results suggest that the highly conserved residue G₄₈ may act as a determinant in regulating or establishing the activity of PspC in the stress signalling pathway. It is unlikely that G₄₈ directly contacts PspA, given that this residue is predicted to lie within the IM and PspA is a peripheral IM protein. Rather, G₄₈ might be required to establish the correct conformation and topology of PspC for an interaction with PspA.

Notably, the inability of PspC₄₄₈ₐ₅ₐₚ₅ₐ to interact with PspG (in contrast to the PspC₄₄₈ₐ₅ₐₚ₅ₐ variant) (Table 1) indicates that residue G₄₈ may direct a specific interaction with PspG, which could have implications for the function of PspG under stress. Consistent with this view, the PspC₄₀₋₆₈ fragment retains the ability to interact with PspG (Table 1),
further suggesting that PspC IM interactions (with PspG) are critical for PspG functionality.

Concluding remarks

The analyses presented here demonstrate that PspB and PspC interact within the IM and may well serve as a scaffold within the Psp regulatory complex to which ArcB, PspA and PspG bind. We show, for the first time, that both PspB and PspG have a clear tendency to self-associate – in contrast to PspC – and identify determinants within PspB and PspC required for regulatory complex formation and psp induction. Since the experiments were conducted in the presence of co-expressed PspBC (to prevent alterations in pmf, which could ultimately lead to psp induction), and within the signalling experiments the amount of PspBC production was not elevated above basal levels, we expect that the outcomes discussed here are physiologically relevant.

In particular, we have established that residues 40–119 of PspC, which include the putative LeuZ (in the periplasmic domain) and residue G48 (in the TM domain), are important determinants required for signal transduction during pIV-dependent psp induction and consequently appear to have important effects on whether PspC interacts with PspA (and PspG for G48). We suggest that PspC may function as a membrane protein with more than one topology, possibly dynamic protein topologies (Fig. 7c). PspC topologies may response to changes in the physical/chemical properties of the IM and $\Delta \psi$. Apparently the PspC_{40–119} fragment (which has no predicted cytoplasmic domain), which localizes in the IM, has the ability (according to the positive-inside rule) to position the periplasmic domain within the cytoplasm, and to induce psp in the absence of PspB (Fig. 7c). Moreover, the soluble PspC_{69–119} fragment (carrying only the periplasmic domain) in the presence of co-expressed PspB interacts with PspA and induces psp. Since neither interaction of PspB with PspC_{69–119} nor interaction of PspA with PspB was detected, PspA–PspC_{69–119}–PspB interactions may be highly co-operative, with very weak initial single component interactions. The interaction with PspA and induction of psp is dependent on the PspC periplasmic domain LeuZ. Considering that PspB–PspC interactions are both PspC LeuZ- and PspB LeuZ-independent (see Table 1) and that PspC_{40–119} but not PspC_{69–119} can interact with PspA in the absence of PspB (see Table 1 and Supplementary Table S2), we argue against PspC working through PspB. Additionally, we show that one ‘toxic’ effect of PspC (the drop in pmf) is counteracted by PspB. This finding indicates that PspC and PspB may be acting as a ‘toxin–antitoxin’ pair that may help to stabilize PspC in a topology or ‘active on-state’ favourable for interacting with PspA (Fig. 7c).

Finally, considering a potential mechanism of psp induction involving two distinct PspC topologies, we suggest that it is formally possible that another signal specifically recognized by PspC (or PspBC) may involve changes in the physical/chemical properties of the membrane, given that inhibitors of phospholipid biosynthesis (Bergler et al., 1994) or application of free fatty acids (Weiner & Model, 1994) strongly induce the psp response. In addition, transcription profiling revealed a link between PspA and use of glycerol 3-phosphate in the biosynthesis of phospholipids (Jovanovic et al., 2006). Further, PspA was shown to bind specific IM phospholipids, phosphatidylglycerol and phosphatidylserine to exhibit its effector function (Kobayashi et al., 2007). In agreement, outer-membrane secretins mislocalized within the IM or a block in protein translocation were shown to induce psp (reviewed by Model et al., 1997; Darwin, 2005). Negatively charged phospholipids were found to restore translocation of outer-membrane precursor proteins across the phosphatidylglycerol-depleted IM of E. coli cells (Kusters et al., 1991). Notably, archaea and acidophilic bacteria that have an inverted $\Delta \psi$ and different membrane content have a Psp system that contains only the PspA homologue and no recognizable PspBC proteins (Biddle et al., 2008; Vrancken et al., 2008), perhaps because PspBC function with a particular membrane potential and membrane lipid composition.

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