Functional roles of the pre-sensor I insertion sequence in an AAA+ bacterial enhancer binding protein

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Summary

Molecular machines belonging to the AAA+ superfamily of ATPases use NTP hydrolysis to remodel their versatile substrates. The presence of an insertion sequence defines the major phylogenetic presensor I insertion (pre-Sli) AAA+ superclade. In the bacterial σ^{54} -dependent enhancer binding protein phage shock protein F (PspF) the pre-Sli loop adopts different conformations depending on the nucleotidebound state. Single amino acid substitutions within the dynamic pre-Sli loop of PspF drastically change the ATP hydrolysis parameters, indicating a structural link to the distant hydrolysis site. We used a sitespecific protein-DNA proximity assay to measure the contribution of the pre-Sli loop in σ^{54} -dependent transcription and demonstrate that the pre-Sli loop is a major structural feature mediating nucleotide statedependent differential engagement with $E\sigma^{54}$. We suggest that much, if not all, of the action of the pre-Sli loop is mediated through the L1 loop and relies on a conserved molecular switch, identified in a crystal structure of one pre-Sli variant and in accordance with the high covariance between some pre-Sli residues and distinct residues outside the pre-Sli sequence.

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Introduction

The large AAA+ (ATPases associated with various cellular activities) protein family are responsible for a wide range of ATP-dependent molecular transformations that involve the disassembly and remodelling of protein and nucleic acid complexes (Thomsen and Berger, 2008). Characteristically, all AAA+ members have the ability to self-assemble into higher-order oligomeric structures, usually as hexamers arranged in a ring (Hanson and Whiteheart, 2005). They contain the highly conserved Walker A and Walker B motifs, involved in ATP binding and ATP hydrolysis, the sensor I sequence and the second region of homology that comprises at least one intersubunit catalytic R-finger (Zhang et al., 2002). Sequence insertions prior to the sensor I motif and the preceding α -helix are regarded as a synapomorphic trait that defines the phylogenetic presensor I insertion (pre-SIi) β-hairpin superclade within the AAA+ superfamily (lyer et al., 2004). AAA+ members belonging to this superclade include HslU, ClpX, Lon, MCM, RuvB, the Ltag helicase and bacterial enhancer binding proteins (bEBPs; Schumacher et al., 2006). Although these sequence insertions (typically 15-25 amino acids in length) are not highly conserved among different AAA+ members, they are generally located in, or at the edge of the inner pore defined by the common hexameric AAA+ ring assembly (lyer et al., 2004).

In a wide range of bacteria specialized AAA+ proteins (bEBPs), function to activate highly regulated adaptive responses mediated by the enhancer-dependent σ^{54} containing RNA polymerase ($E\sigma^{54}$) (Studholme and Dixon, 2003; Wigneshweraraj et al., 2008). Upon binding and hydrolysis of ATP, bEBPs remodel the initial closed promoter complexes formed by $E\sigma^{54}$ to create transcriptionally proficient open complexes (Popham et al., 1989). The bEBPs therefore effectively couple ATP hydrolysis to open complex formation where the DNA template strand is loaded within the active site of RNAP (Sasse-Dwight and Gralla, 1988; Morett and Buck, 1989; Popham et al., 1989). The major binding target for bEBPs is the σ^{54} factor (Bordes et al., 2003). Biochemical and structural analyses of bEBPs PspF, ZraR and NtrC1 (Lee et al., 2003; Rappas et al., 2005; Sallai and Tucker, 2005) have demonstrated that a direct interaction with σ^{54} occurs via the bEBP-specific GAFTGA motif, located within a structural

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insertion in Helix 3, termed the L1 loop (Bordes *et al.*, 2003; Chen *et al.*, 2007). The L1 loop is flanked by the pre-Sli [formerly referred to as the L2 loop, *Escherichia coli* residues 131–139 in PspF (Rappas *et al.*, 2005)], and co-ordinated action between these two loop structures has previously been suggested (Schumacher *et al.*, 2007). Allosteric co-ordination of nucleotide-dependent dynamics between the heterogeneously occupied subunits in the hexameric assembly are important for transcription activation, suggesting that only a subset of subunits directly act on $E\sigma^{54}$ (Joly *et al.*, 2006; Schumacher *et al.*, 2008).

Nucleotide analogues are commonly used to study structural transitions associated with ATP-dependent reactions (Wittinghofer, 1997). The ATP ground state analogue ADP-BeF causes a pronounced rising of the L1 and pre-Sli loops in the bEBP NtrC1, seen using small and wide angle X-ray scattering (SAXS/WAXS) analysis (Chen et al., 2007), a result consistent with the local changes observed in the ATP-bound crystal structure of PspF₁₋₂₇₅ (Rappas *et al.*, 2006). Further, σ^{54} can form stable complexes with NtrC1 and PspF in the presence of ADP-BeF (Chen et al., 2007). Interestingly in SAXS/ WAXS analysis, the ATP analogues AMP PNP and ATP_yS did not cause detectable movements in these loop structures, although AMP.PNP was observed to stabilize complexes formed between σ^{54} and the bEBPs NtrC1 or PspF. Taken together, these results suggest that ATP binding causes the L1 and pre-Sli loops to rise, thereby supporting an initial unstable interaction between the bEBP and σ^{54} (Chen *et al.*, 2007). Experimentally, stable complex formation between the bEBP and (E) σ^{54} occurs in the presence of the non-hydrolysable ATP hydrolysis transition state analogue ADP-AIF, which is thought to reflect the conformation of the bEBP– $E\sigma^{54}$ complex near to, or at the point of, an energy coupling step required for reorganization of the $E\sigma^{54}$ closed complex (Chaney *et al.*, 2001; Rappas et al., 2005). ADP-AIF presumably 'freezes' the local motions at the ATPase catalytic site, which are linked to other dynamic excursions including the presentation of the L1 and pre-Sli loops to the closed complex.

Here we report a detailed analysis of the function of the flexible pre-Sli loop of PspF₁₋₂₇₅. Single amino acid substitutions of the pre-Sli residues (131–139) to alanine invariably impacted on the ATPase kinetics, indicating a strong structural coupling between the ATP hydrolysis site and the distant pre-Sli loop. Differences in the functionalities of the pre-Sli variants enabled three classes of variants to be identified: (i) L1 loop facing; responsible for co-ordinating the position of the L1 loop; (ii) the pre-Sli loop tip; essential for all σ^{54} interaction and remodelling activities; and (iii) a more variable N-terminal side, which likely serves as a structural scaffold and contains residues important for ATPase activity. Our results establish that specific regions of the pre-Sli loop are important for

engaging and remodelling the $E\sigma^{54}$ closed complex in response to different stages of the ATP hydrolysis cycle. We suggest that much, if not all, of the action of the pre-Sli loop is mediated through the L1 loop and relies on a conserved molecular switch, identified in a crystal structure of one pre-Sli variant and in accordance with the high covariance between some pre-Sli residues and distinct residues outside the pre-Sli sequence.

Results

Crystal structures of nucleotide-bound forms of $PspF_{1-275}$, derived from soaking apo-PspF₁₋₂₇₅ crystals with ADP, ATP and AMP PNP revealed striking and distinct movements of the pre-Sli loop (Rappas et al., 2006). These structures suggest a structural coupling between the nucleotide binding site and the pre-Sli loop (Fig. 1A), which probably reflects nucleotide-dependent movements of the pre-Sli loop in solution. SAXS/WAXS data on the PspF homologue NtrC1 also provide evidence for different conformations of these loops in different nucleotide states (Chen et al., 2007). Figure 1B shows the pre-Sli loop consensus sequence obtained from 289 annotated Pfam (00158) bEBPs sequences. Alignment of these sequences (Fig. 1B) suggests that conservation of the pre-Sli loop (residues 131-139 in PspF) varies, with the RVGG motif (residues 131-134 in PspF, particularly the first glycine present in 99% of the available sequences) being the most conserved segment. To study the function of the pre-Sli loop in PspF, single amino acid substitutions to alanine were made across residues 131-139. For experimental simplicity, we used a structurally characterized form of PspF that is deleted for its DNA binding domain (AAA+ domain residues 1–275; $PspF_{1-275}$), yet capable of activating transcription from σ^{54} -dependent promoters both in vivo and in vitro from solution (Bordes et al., 2003). Notably, the sequence alignment (Fig. 1B) also indicated that the proline at position 137 in PspF (present in 21% of the available sequences) is replaced by a threonine residue in 14% of bEBPs. In order to ascertain the specific effect of the threonine at position 137, we also constructed a P137T variant.

With the exception of G134A, all clones directed the synthesis of soluble variant forms of $PspF_{1-275}$. We infer that residue G134, which is located at the tip of the pre-Sli loop (see Fig. 1A), may be required for the structural stability of $PspF_{1-275}$ because expression was below our detection limit.

Transcription activation activities of some of the pre-Sli variants are compromised

The ability of the pre-Sli variants to support transcription activation was assessed using full-length transcription assays on the super-coiled *Sinorhizobium meliloti nifH*



Fig. 1. Location and sequence of the pre-Sli loop in $PspF_{1-275}$.

A. Crystal structure of the PspF₁₋₂₇₅ hexamer with the location of the L1 and pre-Sli loops, Helix 3 and Helix 4 highlighted (PDB 2BJW). The residues that comprise the PspF pre-Sli are indicated.

B. Left panel: The consensus pre-Sli insertion sequence obtained using the 289 annotated Pfam (00158) bEBP sequences. Alignment of these sequences suggests that conservation of the pre-Sli loop varies, with the RVGG motif being the most conserved sequence. Right panel: The pre-Sli sequences of structurally characterized bEBPs PspF₁₋₂₇₅, NtrC1, ZraR and DctD.

promoter. As shown in Fig. 2A, only two residues (Q136 and P137, lanes 7-9) in the pre-Sli loop were tolerant to alanine (or threonine) substitution, and supported near wild-type (WT, lane 2) levels of full-length transcription. These two residues are located farthest away from the L1 loop (see Fig. 1A) and do not appear to undergo any conformational or rotational changes in the presence of different nucleotides (Rappas et al., 2006). As these residues are not highly conserved among bEBPs (Fig. 1B), we suggest that Q136 and P137 may have little effect on the remodelling activities of PspF₁₋₂₇₅ in particular the positioning or stabilization of the σ^{54} interacting L1 loop. We note that the S135A variant (lane 6) shows significantly reduced activity (compared with lane 2) as do the R131A (lane 3) and L138A (lane 10) variants, but the others are essentially unable to activate transcription.

Using abortive transcription assays (to measure heparin-resistant open complex formation; Wigneshwe-

raraj et al., 2005) with two linear nifH promoter probes, a fully double-stranded (duplex) probe (Fig. 2B) and a preopened probe (Fig. 2C; mismatched between positions -10 and -1, thereby mimicking the DNA conformation within the $E\sigma^{54}$ open complex) (Wedel and Kustu, 1995; Cannon et al., 1999), we addressed whether pre-opening the DNA could rescue the transcription activation activity of any of the pre-Sli loop variants. Similar to the full-length transcription assays, only the pre-Sli variants Q136A and P137A/T were able to support open complex formation (Fig. 2B and C, lanes 7-9) at levels comparable to PspF₁₋₂₇₅WT (lane 2), even in the presence of pre-opened DNA (Fig. 2C). We also note that the activity of the PspF₁₋₂₇₅S135A variant still remained significantly lower than PspF₁₋₂₇₅WT (Fig. 2B, lane 6). The remainder of the pre-Sli loop variants demonstrated a significantly reduced ability to support $E\sigma^{54}$ open complex formation. As preopening the DNA had little effect on the transcription acti-



Fig. 2. Specific pre-Sli variants demonstrate defects in transcription activation. A. Some pre-Sli variants fail to support $E\sigma^{54}$ transcription from the super-coiled (SC) *Sinorhizobium meliloti nifH* promoter (10 min activation time). The full-length (FL) transcript is as indicated. The Q136A and P137A/T variants demonstrate significant levels of transcription activity. B. Top: Schematic and nucleotide sequence of the *S. meliloti nifH* duplex promoter probe with the consensus promoter elements GG (positions –26 and –25) and GC (positions –14 and –13) underlined. Bottom: Abortive transcription gel showing the pre-Sli variants ability to support $E\sigma^{54}$ open complex formation on the linear duplex probe. The abortive transcript UpGGG is indicated. The relative numbers of complexes formed on the duplex probe in the presence of the pre-Sli variants are indicated in the graph below the gel. C. As in B but using the pre-opened promoter probe. The lowercase letters in bold type-face indicate the non-complementary residues in the pre-opened promoter probe.

vation activities of the majority of pre-Sli variants, we suggest that these variants have defects in steps that precede DNA melting. Clearly the pre-Sli loop is important for transcription activation with some residues making greater contributions than others.

The transcription activation defective pre-SIi variants fail to stably interact with $\sigma^{\rm 54}$

To assess the contribution of the pre-Sli loop in supporting σ^{54} binding and hence the $E\sigma^{54}$ transcription activation

Fig. 3. The pre-Sli variants defective for open complex formation fail to interact with $\sigma^{\text{54}}.$

A. Top: Schematic and nucleotide sequence of the S. meliloti nifH mismatch promoter probe, the lowercase letters in bold type-face indicate non-complementary residues in the mismatch promoter probe. Bottom: SDS-PAGE gel showing the cross-linking profiles of σ^{54} -DNA complexes formed on the mismatch promoter probe in the presence of 4 mM dATP and either PspF₁₋₂₇₅WT or variants. The migration positions of the cross-linked σ^{54} –DNA and PspF₁₋₂₇₅–DNA species are indicated. Native-PAGE gel illustrating supershift complexes (σ^{54} ss–DNA) are formed in the presence of PspF₁₋₂₇₅WT (lane 2) and the S135A (lane 6), Q136A (lane 7) and P137A/T (lanes 8 and 9) variants. The migration positions of the supershift $(\sigma^{54}$ ss–DNA) and binary σ^{54} –DNA (σ^{54} –DNA) complexes, free DNA and percentage DNA bound in each complex is indicated. B. SDS-PAGE gel as in A but in the presence of core RNAP. The migration positions of the cross-linked β/β' –DNA and σ^{54} –DNA species are indicated. Cross-linked β/β' -DNA species are only observed with the transcription competent PspF₁₋₂₇₅ variants (WT, Q136A and P137A/T).

activities (of the L1 loop) we used a site-specific protein-DNA cross-linking method to study protein-DNA relationship changes for σ^{54} –DNA, β/β' –DNA and PspF_{1–275}–DNA interactions. The photo-reactive DNA templates were constructed by strategically placing the cross-linking reagent p-azidophenacyl bromide (APAB; Sigma), between positions -1 and +1 (-1/+1) in the context of the duplex, pre-opened and mismatch (at positions -12 and -11, mimicking the DNA conformation of the closed complex) promoter probes. The -1/+1 site was chosen based on previous observations (Burrows et al., 2009) that: (i) only $\sigma^{\rm 54}\text{--}{\rm DNA}$ interactions are detected within the closed complex; (ii) in the ADP–AIF (intermediate) complex σ^{54} – DNA and PspF₁₋₂₇₅-DNA cross-linked species are detected; and (iii) only within the open complex are β/β' -DNA interactions observed corresponding to the loading of DNA within the active site of RNAP (Burrows et al., 2008; 2009).

Initially, we analysed a binary σ^{54} -promoter complex formed on the mismatch promoter probe in the presence of PspF₁₋₂₇₅ (WT and pre-Sli variants) and dATP. Previously we have shown that $PspF_{1-275}WT$ and dATP can remodel this binary σ^{54} -DNA complex to a 'supershift' complex (Fig. 3A labelled σ^{54} ss–DNA; Cannon *et al.*, 2000; 2001). Using this assay, we can establish whether the defects observed in open complex formation by the pre-Sli variants (Fig. 2) are due to an inability to efficiently interact with σ^{54} . In line with the transcription activation data (Fig. 2) only the Q136A, P137A/T and (to a lesser extent) S135A variants supported σ^{54} supershift complex formation (Fig. 3A, lanes 6-9); however, the remainder of the pre-Sli variants failed to support σ^{54} ss–DNA complex formation (Fig. 3A, lanes 3-5 and 10-11). When these reactions were analysed by cross-linking we observed that where supershift complexes were present, the profile of the cross-linked $\sigma^{\rm 54}\text{-}{\rm DNA}$ species was altered from a single band (Fig. 3A, SDS-PAGE; lane 1 arrowed) to a



double band (Fig. 3A, lanes 2 and 7–9, lane 2 arrowed), suggesting that the organization of σ^{54} is changed in the σ^{54} -DNA complex. However, no change in the σ^{54} -DNA profile was observed in reactions containing S135A (Fig. 3A, lane 6), where a weak σ^{54} ss–DNA complex was observed, or the other pre-Sli variants (131–133 and 138–139; lanes 3–6 and 10–11) where no σ^{54} ss–DNA complexes were detected. We also note a weak cross-linked PspF₁₋₂₇₅–DNA species was detected in the supershift reactions, particularly with the P137T variant (Fig. 3A, SDS-PAGE; lanes 2 and 7–9, lane 9 arrowed).

In the presence of core RNAP, the Q136A and P137A/T variants (Fig. 3B, lanes 7–9) supported a change in the σ^{54} –DNA interactions (from a single band; lane 1 arrowed,

Table 1.	ATP hydrolysis	parameters of	f the PspF ₁₋₂₇₅	pre-Sli proteins.
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$PspF_{1-275}$ variant	$V_{\rm max}$ (min ⁻¹)	Standard error	<i>K</i> _m (μM)	Standard error
Wild type	30.57	3.46	178	61
R131A	< 0.1	ND	ND	ND
V132A	0.41	0.04	10	5
G133A	2.72	0.75	91	50
S135A	< 0.1	ND	ND	ND
Q136A	3.98	0.39	8	2
P137A	26.72	4.29	300	130
P137T	47.49	4.30	140	61
L138A	2.50	0.66	160	60
Q139A	< 0.1	ND	ND	ND

ND, not determined.

to a double band, lane 7 arrowed) and formation of β/β' -DNA cross-links (lane 9 arrowed) comparable to PspF₁₋₂₇₅WT. Together these data confirm that the transcription activation defect associated with some pre-Sli variants is due to their inability to productively interact with σ^{54} in the σ^{54} -DNA complex or in the E σ^{54} closed complex in the presence of dATP. Hence, defects in the pre-Sli variants occur at the level of the σ^{54} isomerization event, prior to the loading of DNA within the active site of RNAP.

Pre-Sli variants impact on the ATP hydrolysis active site

Eσ⁵⁴ open complex formation requires the coupling of energy derived from bEBP-dependent ATP hydrolysis to remodelling of the Eσ⁵⁴ closed complex. Effective energy coupling relies on co-ordinated ATP binding and hydrolysis between different subunits of the hexameric PspF₁₋₂₇₅ AAA+ ring assembly (Joly *et al.*, 2007; Schumacher *et al.*, 2008). Failure of the pre-Sli variants to activate transcription could therefore be a consequence of a failure to bind or hydrolyse ATP and/or to efficiently couple hydrolysis to restructuring of the Eσ⁵⁴ closed complex.

To ascertain whether this was the case we measured the ATP binding activities of the pre-Sli variants using non-equilibrium UV cross-linking of $[\alpha^{-32}P]$ -ATP (Schumacher et al., 2004; 2007). Results (Fig. S1) demonstrate that the PspF₁₋₂₇₅ pre-Sli variants bind ATP with similar affinity as PspF1-275WT, indicating that the pre-Sli loop does not contribute to the nucleotide binding activities of PspF. In contrast, the pre-Sli variants appear to have a striking effect on the steady-state ATP hydrolysis rates (Table 1). We note however, there is no clear correlation between low ATP hydrolysis rates and defects in transcription activation activity, because the Q136A and T137A variants are similarly active in the transcription assays despite a more than 80% reduction in the ATPase activity of Q136A. The decrease in V_{max} observed for all the pre-Sli variants, except P137, indicates a finely tuned coupling between the structure of the pre-Sli loop and the distant ATP active site. These observations functionally distinguish the pre-Sli loop from the adjacent L1 loop, where alanine substitutions do not negatively impact on PspF₁₋₂₇₅ ATPase activity (Rappas *et al.*, 2005).

To determine whether the reduced ATP hydrolysis rates observed in the pre-Sli variants are due to defects in hexamer formation, a prerequisite for maximal hydrolysis rates (Schumacher et al., 2004), we performed gelfiltration experiments in the absence and presence of ADP. PspF₁₋₂₇₅WT normally exists in equilibrium between a dimer and a hexamer at low protein concentrations (below 10 µM), but in the presence of either ATP or ADP the equilibrium shifts to the hexameric species (Joly et al., 2006). The pre-Sli variants eluted predominantly as dimers (Fig. 4A) in the absence of nucleotide and hexamers in the presence of ADP (Fig. 4B), although we note that the apparent dimer/hexamer ratios differed from PspF₁₋₂₇₅WT, especially the V132A, S135A, Q136A and P137T variants. In the case of the S135A and Q136A variants a significant shift towards the hexameric species was observed, whereas the V132A and P137T variants demonstrate a significant defect in their ability to form hexamers, which may account for the reduced ATPase activity of PspF₁₋₂₇₅V132A. However, we note that PspF₁₋₂₇₅P137T exhibits PspF₁₋₂₇₅WT activities in terms of transcription activation and $\sigma^{\rm 54}$ binding. As none of the pre-Sli variants failed to form the apparent hexameric species in the presence of ADP, we suggest that the lack of ATPase activity for the majority of pre-Sli variants is not simply due to the inability to form a hexamer or bind nucleotide.

The pre-SIi loop stabilizes distinct interactions $\text{PspF}_{\text{1-275}}$ makes with σ^{54}

To ascertain the contribution the pre-Sli loop makes to $PspF_{1-275}$ energy coupling activities, nucleotidedependent interactions between $PspF_{1-275}$, σ^{54} and/or $E\sigma^{54}$ promoter complexes were studied using the protein–DNA cross-linking assay and different ATP analogues: (i) the transition state analogue ADP–AIF; (ii) the ground state



Fig. 4. Gel filtration profiles of $PspF_{1-275}$ (WT and variants) in the absence and presence of ADP. Nucleotide-dependent apparent dimer/hexamer equilibrium of $PspF_{1-275}$ proteins as obtained by analytical HPLC gel-filtration chromatography using a Biosep 3000 column (Phenomenex), in the absence (A) and presence (B) of ADP. Chromatographs are overlaid and offset but not normalized. The $PspF_{1-275}$ variants are as indicated.

analogues AMP–AIF and ADP–BeF; and (iii) the slowly hydrolysable ATP analogue ATP γ S.

The ability of the PspF₁₋₂₇₅ pre-Sli variants to support stable (trapped) complex formation with σ^{54} and/or $E\sigma^{54}$ was first assayed in the presence of the ATP transition state analogue, ADP–AIF (Fig. 5). As illustrated in Fig. 5A (Native-PAGE), only in the presence of the pre-Sli variants S135A, Q136A and P137A/T were ADP–AIF-dependent trapped complexes formed at a similar level to PspF₁₋₂₇₅WT. When these reactions were analysed by photo-cross-linking (Fig. 5A; SDS-PAGE), we observed a σ^{54} -dependent PspF₁₋₂₇₅–DNA cross-link (PspF₁₋₂₇₅–DNA; Fig. 5A, lanes 6–9 and Burrows *et al.*, 2009). However, we also observed a cross-linked PspF₁₋₂₇₅–DNA species in reactions containing the V132A (lane 4 arrowed) and L138A variants (lane 10). These data demonstrate that in the presence of ADP–AIF, the effect of the V132A and

L138A substitutions are less apparent than in the transcription assays (Fig. 2), although the stability of ADP-AIF-dependent complexes are greatly reduced (as judged by Native-PAGE analysis; Fig. 5A). Similar to the reactions containing PspF₁₋₂₇₅WT, the V132A (lane 4), S135A (lane 6), Q136A (lane 7), P137A/T (lanes 8-9) and L138A (lane 10) variants caused a change in the σ^{54} –DNA profile (Fig. 5A; from a single lane 1 arrowed, to no band lane 6 arrowed). We also observed very weak PspF₁₋₂₇₅-DNA interactions with the R131A (lane 3), G133A (lane 5) and Q139A (lane 11) variants, although no alteration in the σ^{54} –DNA profile was evident. Interestingly, when core RNAP was added to these reactions (Fig. 5B) the very weak PspF₁₋₂₇₅-DNA interactions with the R131A, G133A and Q139A variants were no longer evident (compare Fig. 5A and B, lanes 3, 5 and 11 respectively); however, we do note a change in the σ^{54} –DNA profile (from a double



Fig. 5. The activities of the V132A and L138A variants are rescued by ADP-AIF. A. SDS-PAGE gel showing the cross-linking profiles of σ^{54} –DNA complexes formed on the mismatch promoter probe in the presence of ADP-AIF. The migration positions of the cross-linked σ^{54} –DNA and PspF₁₋₂₇₅–DNA species are indicated. A cross-linked PspF₁₋₂₇₅-DNA species is observed in reactions containing the V132A (lane 4), S135A (lane 6), Q136A (lane 7), P137A/T (lanes 8-9) and L138A (lane 10) variants. Native-PAGE gel illustrating that stable ADP-AIF trapped complexes are only observed in the presence of PspF1-275 WT (lane 2) and the S135A (lane 6), Q136A (lane 7) and P137A/T (lanes 8-9) variants. The migration positions of the $\sigma^{\rm 54}\text{-}\text{DNA-PspF}_{\rm 1-275}\text{:}\text{ADP-AIF}$ (trapped) and binary σ^{54} –DNA (σ^{54} –DNA) complexes, free DNA and percentage DNA bound in each complex are as indicated. B. SDS-PAGE gel as in A but on the duplex promoter probe in the presence of core RNAP. The migration positions of the cross-linked β/β' –DNA, σ^{54} –DNA and

PspF1-275-DNA species are indicated.

band to a single band) in the reaction containing the R131A variant (lane 3 arrowed), similar to that observed for PspF₁₋₂₇₅WT (lane 2 arrowed) and the V132A, S135A, Q136A, P137A/T and L138A variants (lanes 4 and 6-10).

In the presence of the ATP ground state analogues AMP–AIF (Joly *et al.*, 2008) and ADP–BeF (Burrows *et al.*, 2009), similar to the ADP–AIF reactions, only the S135A, Q136A and P137A/T variants supported stable trapped complex formation (Fig. S2A, Native-PAGE). Importantly, only these variants (S135A, Q136A and P137A/T) supported PspF₁₋₂₇₅–DNA interactions (espe-

cially in the presence of core RNAP) and an alteration in the σ^{54} –DNA cross-linking profile (from a single band, lane 1 arrowed, to no band, lane 6 arrowed). For the remainder of the pre-Sli variants, no cross-linked PspF₁₋₂₇₅–DNA species was detected and neither was a change in the σ^{54} –DNA profile. In the presence of the slowly hydrolysable ATP analogue, ATP γ S, we observed a PspF₁₋₂₇₅–DNA cross-linked species and an altered σ^{54} –DNA profile with the S135A, Q136A and P137A/T variants (Fig. S3). The S135A variant displays a significantly reduced level of PspF₁₋₂₇₅–DNA cross-linked species, which we attribute to the lower ATPase activity of S135A.

Taken together these data indicate that residues V132 and L138 are organized differently with the ATP hydrolysable (dATP and ATP γ S) and ATP ground state (AMP–AIF and ADP–BeF) analogues, than with the ATP transition state analogue (ADP–AIF).

Evolutionary covariance of the E81, E97 and R131 residues support a conserved switch between the pre-Sli and L1 loops in bEBPs

Our structural and functional data suggest that (i) pre-Sli residues facing the L1 loop are probably required to position the L1 loop for engagement with σ^{54} at distinct steps during the ATP hydrolysis cycle and (ii) some pre-Sli residues, despite having no obvious contact features outside the pre-SIi loop, impact on the ATP hydrolysis and σ^{54} engagement activities of PspF. As the pre-Sli loops of bEBPs demonstrate a degree of sequence conservation (Fig. 1B), this sequence is well suited for predictive covariance analysis as an independent means to identify interacting co-evolved residues. As bEBPs function in a conserved manner, it seems reasonable to suggest that evolutionary variations are susceptible to selective pressures to maintain functionally linked residues. To explore covariation we first aligned the 289 seeded nonredundant sequences of the AAA+ domains of bEBPs (Pfam 00158) using the ClustalW program. Using three established covariance algorithms (Fodor and Aldrich, 2004) [observed minus expected squares, statistical coupling analysis and McLachlan-based substitution correlation (Olmea et al., 1999)] we determined the covariance of residues within the conserved pre-Sli sequence and the entire AAA+ domain of bEBPs (Fig. S4). We also determined the covariance between PspF residues 80-98 (which includes the L1 loop sequence) and the PspF pre-Sli loop sequence (Fig. 6A). As shown in Fig. 6A PspF residue E97 (arrowed) exhibits strong covariance with pre-Sli residue R131 and residue E81 (arrowed) exhibits covariance with the pre-Sli residues R131, G134 and L138, supporting the existence of a functionally conserved mechanism by which bEBPs relate the nucleotidebound state to pre-Sli loop dynamics.

Conformational changes are apparent in the crystal structure of $PspF_{1-275}R131A$

The severely reduced ATP hydrolysis rates observed for the majority of the pre-Sli variants suggest the existence of an important structural link coupling the pre-Sli loop to the distant ATP hydrolysis site. We reasoned that mutations in the pre-Sli loop would result in local, and potentially global, conformational changes that could reveal mechanistic details of this structural link. Based on our covariance analysis we determined the crystal structure of apo-PspF₁₋₂₇₅R131A in order to determine the effect of the R131A mutation on the conformation of the pre-Sli loop compared with apo-PspF₁₋₂₇₅WT (Fig. 6B and Fig. S5A-C). Clear local differences are evident in the conformation of the pre-Sli loop, as well as a significant rotation in the side-chain of residue E81 of the adjacent L1 loop (Fig. 6B). These observations strongly support the view that residue R131 of the pre-Sli loop communicates with residue E81 of the L1 loop in a nucleotide-dependent manner that appears essential for PspF₁₋₂₇₅ activity. Disruption of the R131-E81 link (by introduction of the R131A mutation) causes the pre-Sli to rotate, so that the alanine side-chain is now pointing towards the other stem of the pre-Sli loop (away from the L1 loop). More globally, the alanine substitution at residue R131 results in conformational changes in Helix 4 (most notably residue E125) and, somewhat surprisingly, the orientation of residue R168 a putative R finger (as shown by the root mean square deviation plot between PspF₁₋₂₇₅R131A and PspF₁₋₂₇₅WT; Fig. S5D). Although these differences are modest, they are the most pronounced changes evident over the entire length of the PspF₁₋₂₇₅ structure outside the pre-Sli and L1 loops.

Discussion

We have systematically examined the activities of the pre-Sli sequence in PspF₁₋₂₇₅, characterized by crystallographic studies as a well-defined loop structure that changes its organization and orientation in the presence of different nucleotides (Rappas *et al.*, 2006). In PspF₁₋₂₇₅ and the related bEBPs NtrC1 and ZraR, a direct interaction between the pre-Sli loop and the L1 loop is evident (Lee *et al.*, 2003; Sallai and Tucker, 2005). The importance of specific pre-Sli residues in PspF₁₋₂₇₅ functionality now establishes that particular regions of the pre-Sli loop are important for engaging and remodelling the E σ^{54} closed complex in response to the ATP hydrolysis cycle.

PspF₁₋₂₇₅ pre-Sli integrity is required for high ATP hydrolysis rates

The majority of the pre-Sli variants showed no marked defects in self-association, although we do note that the V132A and P137T variants exhibited lower levels of hexamer formation (which may account for the low ATPase activity of V132A) and the S135A and Q136A variants showed a propensity for the hexameric state (even though the ATPase activity of S135A is severely reduced). Taken together these results suggest that the pre-Sli loop does not contribute greatly to the energetically favourable interactions (at the protomer–protomer interface) needed for hexamer formation. Clearly, alanine substitutions in the pre-Sli loop impact on the catalytic site

в



Fig. 6. A conserved switch between the pre-Sli and L1 loops exists within bEBPs. A. The covariance between the PspF pre-Sli sequence (RVGGSQPLQ; colour-coded as shown) and PspF residues 80–98 was calculated and depicted graphically. The strong covariance of residue E81 with pre-Sli residues R131, G134 and L138 (arrowed) and residue E97 and pre-Sli residue R131 (arrowed) is indicated by high covariance scores.

B. The crystal structures of the apo-PspF₁₋₂₇₅R131A (vellow) and apo-PspF₁₋₂₇₅WT (blue; PDB 2BJW) demonstrate the effect of the R131A mutation on the pre-Sli loop conformation. The two structures were aligned on the main chain atoms of residues 35-42. The positions of residues E81 (L1 loop), E125 (Helix 4), R168 (putative R-finger), R131 and V132 (pre-Sli) and the R131A mutation (in the context of the PspF₁₋₂₇₅R131A structure) are indicated. Structural features relevant to bEBPs such as the L1 and pre-Sli loops, Helix 3 and Helix 4 are labelled. Clear local differences between the apo-PspF₁₋₂₇₅WT and apo-PspF₁₋₂₇₅R131A structures are apparent in the pre-Sli loop conformation, as well as a significant rotation of residue E81 (L1 loop) resulting in the disruption of the E81-R131 link.



architecture, reflected by the decrease in ATP hydrolysis rates. Differences in K_M, notably for the V132A and S135A variants probably reflect differences in hydrolysis rates rather than in ATP binding due to the reported high offrates (Rombel *et al.*, 1999; Schumacher *et al.*, 2004). The similar apparent nucleotide binding affinities of all the pre-Sli variants tested (Fig. S1) are consistent with the spatial separation of residues involved in nucleotide binding and hydrolysis (Rappas *et al.*, 2005). Apart from the P137A/T variants, all the pre-Sli variants showed drastically reduced ATP hydrolysis rates, which is notable because the pre-Sli is a flexible surface-exposed loop that is spatially distant from the ATP hydrolysis site, suggesting that linked dynamic modes underpin the nucleotide-dependent functions of PspF (Rappas *et al.*, 2006).

The crystal structure of apo-PspF₁₋₂₇₅R131A illustrates that destabilization of the pre-Sli loop–L1 loop interaction results in a significant conformational change in the pre-Sli loop that propagates to the C-terminal moiety of Helix 4 (Fig. 6B). This change appears to weaken the intrasubunit E125–R168 interaction and the orientation of the putative R-finger residue R168. Previously we showed that the R168A variant forms nucleotide-independent hexamers that are ATPase defective (Schumacher *et al.*, 2004). These prior observations suggested that residue R168 inhibits hexamer formation in apo-PspF₁₋₂₇₅WT while being required for intersubunit catalysis in the ATP-

bound state (Schumacher *et al.*, 2004). Consistent with these findings, we propose that the structural conformation of the pre-Sli loop of apo-PspF₁₋₂₇₅WT, via Helix 4 residue E125, supports a conformation of R168 that inhibits self-association.

Pre-Sli residues support different steps during transcription activation

Using different ATP analogues three broad classes of activities associated with the pre-Sli loop were identified. Class I contained variant forms of the pre-Sli loop that had activities that were indistinguishable from PspF₁₋₂₇₅WT, indicating that these pre-Sli residues (Q136 and P137) are largely unimportant for nucleotide-dependent engagement of the $E\sigma^{\scriptscriptstyle 54}$ closed complex. We also assign S135 to class I because it behaves similarly to PspF₁₋₂₇₅WT in the nucleotide-dependent interaction studies and the significantly reduced transcription activation and $\sigma^{\rm 54}$ isomerization activities could be attributable to its low ATPase activity. These residues are located farthest away from the L1 loop (Fig. 1A) and are therefore unlikely to interact with the L1 loop. Class II variants (G133, Q139 and to some extent R131) essentially failed in all the nucleotidedependent interaction assays tested. The severe functional defects observed for PspF1-275G133A are not surprising given its high sequence conservation and high $\phi-\psi$ angle indicating that backbone flexibility at the tip of the pre-Sli loop appears indispensable for bEBP activity. Crystal structures of PspF₁₋₂₇₅ illustrate that nucleotide binding causes the pre-Sli loop to twist around its longitudinal axis (Rappas et al., 2006). Diminishing the glycine-associated flexibility (in the G133A variant) strongly suggests that the potential for 'structural twist' is critically important for PspF functionality. Class II variants maintain the nucleotide binding and self-association activities but cannot engage the $E\sigma^{54}$ closed complex, presumably because the contact features for binding σ^{54} (i.e. the L1 loop) are inappropriately presented. Class III variants (represented by V132A and L138A) supported PspF₁₋₂₇₅-DNA cross-links in the presence of the ATP transition state analogue (ADP-AIF) but not with the ATP ground state analogues (AMP-AIF and ADP-BeF) or the slowly hydrolysable ATP analogue (ATP_γS), suggesting that the pre-Sli loop plays at least two distinct roles in productive $E\sigma^{54}$ –PspF interactions that are dependent on the nucleotide state: weak interactions that exist in the nucleotide ground state and strong interactions in the ATP transition state.

The high probability of pair-wise co-evolution of the L1 residues E81 and E97 (in PspF) with the pre-Sli residue R131 (in PspF) suggests that functionally important contacts between the pre-Sli and L1 loops exist which may provide a mechanistic rational for how different nucleotide

states could orientate the L1 loop for E σ^{54} engagement. R131 would function as switch which pivots between E81 in the ATP ground state and E97 in the ATP transition state. Disruption of this switch may account for the failure of PspF₁₋₂₇₅R131A to stably interact with σ^{54} , as demonstrated by the lack of σ^{54} -dependent PspF₁₋₂₇₅–DNA crosslinks with the R131A variant in the presence of the metal fluoride analogues (Fig. 5 and Figs S2 and 3, lane 3). The covariance of residues E81 (L1 loop) and R131, G134 and Q136 (pre-Sli) indicates that additional residues are required for maintaining a pre-Sli loop conformation suitable for a productive E81–R131 interaction to occur (Fig. 6A). Overall, the functionally important R131-E81/ E97 switch (studied in PspF) appears highly conserved within the bEBP family.

The pre-Sli loop does not appear to contact promoter DNA

The DNA-bound model of the electron microscopy structure of $E\sigma^{54}$ -PspF₁₋₂₇₅:ADP–AIF indicates that the pre-SIi loop lies proximal to both σ^{54} and promoter DNA (Bose et al., 2008). One important observation from the nucleotide-dependent studies is that in the presence of different DNA templates (Fig. 5 and Figs S2 and 3) no large scale changes in the amounts of complexes formed (in Native-PAGE analysis compared with PspF₁₋₂₇₅WT) were observed, nor was the stability of these complexes increased or reversed, inferring that the pre-Sli loop does not directly contact promoter DNA and/or σ^{54} . In contrast, interactions between $\sigma^{\scriptscriptstyle 54}$ and the L1 loop variant PspF₁₋₂₇₅T86S can be recovered when promoter DNA sequences downstream of the -12 position are missing (Dago et al., 2007). We propose that much of the action of the pre-Sli loop is manifest through the L1 loop interacting with σ^{54} and that the pre-Sli loop in PspF and other related bEBPs works mainly on the L1 loop rather than its substrate.

Comparison with the pre-Sli sequence of other AAA+ proteins

The similarity in the location of the synapomorphic pre-Sli feature, which is found in or at the edge of the inner pore of the hexameric AAA+ ring assembly (lyer *et al.*, 2004), raises an important question regarding a common role for the pre-Sli in other AAA+ proteins. In RuvB, which contacts RuvA via the pre-Sli β -hairpin, mutations in the pre-Sli result in decreased RuvB ATPase activity (Han *et al.*, 2001; Yamada *et al.*, 2002), while in E1 helicase of Papilloma virus, the pre-Sli β -hairpin interacts with single-stranded DNA (Enemark and Joshua-Tor, 2006). In these cases the pre-Sli contacts auxiliary proteins or substrates directly, for which we found no evidence in PspF. In MCM, a pre-Sli mutation modestly increased the K_d for DNA binding, but

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significantly reduced ATPase and abolished helicase activity (McGeoch *et al.*, 2005), suggesting tight structural coupling between the pre-Sli and the hydrolysis site, similar to our observations with PspF. Distinct functional roles that depend on the nucleotide-bound state have, as far as we know, not been shown for other AAA+ pre-Sli sequences. However, structural studies of the E1 helicase (Enemark and Joshua-Tor, 2006) and the E1-related Ltag helicase (Gai *et al.*, 2004) suggest that different nucleotide states result in distinct orientations of the pre-Sli. More recently, nucleotide state-dependent dynamics for the pre-Sli have also been proposed for MCM (Barry *et al.*, 2009) and ClpX (Martin *et al.*, 2008a,b).

Clearly, the divergence in pre-Sli sequences among members of the pre-Sli β -hairpin superclade of AAA+ proteins and their distinct direct or indirect involvements in substrate remodelling do not allow the simple assignment of common roles to this structural feature. Rather, the strategic placement of the pre-Sli near or in the pore of the AAA+ ring assembly, and the potential structural coupling to the ATP hydrolysis event may explain the marked evolutionary prevalence of the pre-Sli.

Experimental procedures

Proteins and DNA probes

Escherichia coli core RNAP was purchased from Epicentre technology (Cambio). Klebsiella pneumoniae o54 was purified as described (Wigneshweraraj et al., 2003). Plasmid pPB1 encoding Escherichia coli PspF residues 1-275 (PspF1-275 with an N-terminal His6 tag in pET28b+) was mutagenized (Quickchange Mutagenesis Kit, Stratagene) resulting in plasmids pPB1(R131A), pPB1(V132A), pPB1(G133A), pPB1(G134A), pPB1(S135A), pPB1(Q136A), pPB1(P137A), pPB1(P137T), pPB1(L138A) and pPB1(Q139A). Proteins were overproduced as described (Wigneshweraraj et al., 2003). Proteins were purified using a gravity flow method. Briefly Ni-NTA resin (Qiagen) was equilibrated with buffer A_{NI} (20 mM sodium phosphate pH 7.0, 500 mM NaCl, 5% (v/v) glycerol) prior to applying the cell supernatant. The resin was then washed with buffer $A_{\mbox{\tiny NI}},$ followed by wash buffer 1 (buffer A_{NI} plus 40 mM imidazole) and wash buffer 2 (buffer A_{NI} plus 80 mM imidazole). The protein was eluted using buffer B_{NI} (buffer A_{NI} plus 1 M imidazole). The S. meliloti nifH phosphorothioated (Operon) DNA probes were derivatized with p-azidophenacyl bromide as described (Burrows et al., 2004). The modified promoter strands were ³²P labelled and annealed to the complementary strand as described (Wigneshweraraj et al., 2003; Burrows et al., 2008).

In vitro full-length or abortive transcription assays

Full-length or abortive transcription assays were performed in STA buffer (25 mM Tris-acetate pH 8.0, 8 mM Mg-acetate, 10 mM KCl, 3.5% w/v PEG 6000) in a 10 μ l reaction containing 100 nM E σ^{54} (reconstituted at a 1:4 ratio of E: σ^{54}), 4 mM dATP

and 20 nM promoter DNA probe. The mix was incubated at 37°C for 5 min and the reaction started by addition of 5 μ M PspF₁₋₂₇₅WT or variants and incubated for a further 10 min at 37°C. Full-length transcription was initiated by adding an elongation mix containing 100 μ g ml⁻¹ heparin, 1 mM ATP, CTP, GTP, 0.05 mM UTP and 3 μ Ci (α -³²P]-UTP and incubated at 37°C for 20 min. The full-length transcription products were analysed on a 6% denaturing gel. Synthesis of the abortive transcript (5' UpGGG) was initiated by adding an abortive mix containing 100 μ g ml⁻¹ heparin, 0.5 mM 5' UpG and 4 μ Ci (α -³²P]-GTP and incubated at 37°C for 20 min. The solution of 5 min containing 100 μ g ml⁻¹ heparin, 0.5 mM 5' UpG and 4 μ Ci (α -³²P]-GTP and incubated at 37°C for 20 min. The abortive transcription products were analysed on a 20% denaturing gel. Full-length and abortive transcription gels were visualized and quantified using a Fuji FLA-5000 PhosphorImager.

ATP binding and hydrolysis

ATP binding assays of PspF₁₋₂₇₅ proteins by UV cross-linking were performed as described (Schumacher et al., 2004). ATPase reactions were carried out as described (Schumacher et al., 2008). ATPase assays were performed in reaction buffer (35 mM Tris-acetate pH 8.0, 70 mM K-acetate, 5 mM Mg-acetate, 19 mM ammonium acetate, 0.7 mM DTT) containing 15 mM MgCl₂ and incubated at 23°C, using 0.06 µCi $[\alpha^{-32}P]ATP$ or $[\alpha^{-32}P]$ -dATP as nucleotide tracers. Prior to ATP titration experiments, protein concentration titrations were carried out at 0.1 mM ATP to verify maximal turnover rates were achieved. Reactions were guenched with 5 vols of 2 M formic acid. [α -³²P]-ADP was separated from [α -³²P]-ATP by thin-layer chromatography [on Polygram Cel 300 PE (polyethyleneimine)] and visualized and quantified using a Fuji FLA-5000 PhosphorImager. ATP titration experiments and all PspF₁₋₂₇₅ titrations were designed so that only 10–30% of the initial amount of ATP was hydrolysed thereby ensuring that any ADP produced had no measurable effect on ATP hydrolysis rates. It also ensures that ATP was not limiting in the assays and that non-linearity of turnover rates was minimal. V_{max} and $K_{\rm m}$ values as well as standard errors were determined at initial concentrations of 0.05-1 mM ATP using non-linear regression Graphit software (Erithacus Software Limited).

Gel filtration chromatography

Gel filtration chromatography was carried out with 50 μ M PspF₁₋₂₇₅ (WT and variants) at room temperature in running buffer (20 mM Tris pH 8.0, 50 mM NaCl, 15 mM MgCl₂, 0.02% (w/v) azide) at a 0.7 ml min⁻¹ flow rate using a BioCad Sprint HPLC system and a Bio-Sep-S 3000 column (Phenomenex). To test for nucleotide-dependent oligomerization the column was pre-equilibrated with five bed volumes of 0.5 mM ADP, prior to loading 10 μ M PspF₁₋₂₇₅ (WT or variants), supplemented with 0.5 mM ADP. The column was calibrated using molecular weight standards (Sigma): Blue Dextran (200 kDa), Thyroglobulin (669 kDa), Ferritin (440 kDa), Catalase (232 kDa), Aldolase (158 kDa), Albumin (67 kDa), Ovalbumin (43 kDa), Chymotrypsin (25 kDa) and Ribonuclease (13.5 kDa).

Photo-cross-linking assays

Cross-linking reactions were conducted at 37° C in STA buffer in a 10 µl reaction volume as described (Burrows *et al.*, 2008;

2009). Briefly, where indicated either 1 μ M σ^{54} or 200 nM E σ^{54} (reconstituted using a 1:2 ratio of $E:\sigma^{54}$) and 20 nM modified ³²P-labelled promoter DNA probe was incubated for 5 min at 37°C. Open complexes were formed by 4 mM dATP and 5μ M PspF₁₋₂₇₅ and the reaction incubated for 10 min at 37°C. Trapped complexes were formed in situ by adding 5 µM PspF₁₋₂₇₅, 5 mM NaF and either 1 mM ADP, 0.2 mM BeCl₃ for the ADP-BeF reactions; 1 mM ADP, 0.2 mM AICl₃ for the ADP-AIF reactions; and 1 mM AMP, 0.2 mM AICl₃ for the AMP-AIF reactions and incubated for 10 min at 37°C. To eliminate free core RNAP from binding the promoter probe, reactions contained 100 ng ml⁻¹ salmon sperm DNA. Reactions were UV irradiated at 365 nm for 30 s using a UV-Stratalinker 1800 (Stratagene). A 2 µl sample of the reaction was analysed by Native-PAGE (4.5%), run at 100 V for 55 min. The remainder of the cross-linking reaction was diluted using 5 µl 10 M Urea and 5 µl 2× SDS loading buffer (Sigma), heated at 95°C for 3 min and 10 µl loaded on a 7.5% SDS-PAGE gel run at 200 V for 50 min. The gels were visualized and guantified using a FLA-500 PhosphorImager. The cross-linked proteins were identified using antibodies as described (Burrows et al., 2008).

Covariance analysis

Covariance analysis was performed as described (Fodor and Aldrich, 2004). Briefly, the 289 sequences in Pfam (00158) were first aligned using ClustalW. For a multiple alignment of N sequences, an NxN matrix (with dimensions k and I) was constructed for each pair of residues (i and j). Each entry in the matrix was then scored using the McLachlan substitution matrix (Olmea *et al.*, 1999), producing high scores for identities/conservative substitutions and low scores for non-conservative substitutions. The correlation between the two columns (*i* and *j*) was calculated using the

equation $r_{ij} = \frac{1}{N^2} x \frac{\sum_{kl} (S_{ikl} - \langle S_i \rangle) (S_{jkl} - \langle S_j \rangle)}{\sigma_i \sigma_j}$ where $\langle S \rangle$ is the

average and $\sigma_{\!\scriptscriptstyle i}$ is the standard deviation of all entries in the matrix.

Crystallization and structure determination

PspF₁₋₂₇₅R131A was expressed, purified, crystallized and flash frozen as described (Rappas et al., 2005). Diffraction data were collected (in-house) using a March 345 image plate mounted on a Rigaku RU-H3RHB rotating-anode X-ray generator (operated at 50 kV and 100 mA) fitted with Osmic confocal optics and a copper target (Cu Ka: λ = 1.542 Å). All the data sets were processed using MOSFLM (Leslie, 1992) and SCALA (Evans, 2006) using the CCP4 program suite (Collaborative Computational Project, 1994). Phases for PspF₁₋₂₇₅R131A crystals were obtained by molecular replacement using the PspF₁₋₂₇₅WT (PDB 2BJW) structure as a search model and the PHASER program (Storoni et al., 2004). Flexible regions such as the L1 (residues 79-93) and pre-Sli loops were first removed from the search model and built-in during rounds of refinement. The F₀-F_c and 2F₀-F_c maps showed some connected electron density for residues 79-81, 91-93 and 129-135, which were manually built-in using program O (Jones *et al.*, 1991). After several rounds of rebuilding, minimization and *B*-individual refinements, water molecules were added in the F_o - F_o map automatically (using CNS) and manually. A final cycle of TLS refinement was carried out using the REFMAC5 program (Murshudov, 1997) and the two TLS groups: the α/β subdomain (residues 7–177) and the α -helical domain (residues 180–255). The refinement statistics are summarized in Table S1. All figures were prepared using Pymol (DeLano, W.L. The PyMOL Molecular Graphics System 2002).

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References

- Barry, E.R., Lovett, J.E., Costa, A., Lea, S.M., and Bell, S.D. (2009) Intersubunit allosteric communication mediated by a conserved loop in the MCM helicase. *Proc Natl Acad Sci* USA 106: 1051–1056.
- Bordes, P., Wigneshweraraj, S.R., Schumacher, J., Zhang, X., Chaney, M., and Buck, M. (2003) The ATP hydrolyzing transcription activator phage shock protein F of *Escherichia coli*: identifying a surface that binds sigma 54. *Proc Natl Acad Sci USA* **100**: 2278–2283.
- Bose, D., Pape, T., Burrows, P.C., Rappas, M., Wigneshweraraj, S.R., Buck, M., and Zhang, X. (2008) Organization of an activator-bound RNA polymerase holoenzyme. *Mol Cell* **32:** 337–346.
- Burrows, P.C., Severinov, K., Buck, M., and Wigneshweraraj, S.R. (2004) Reorganisation of an RNA polymerasepromoter DNA complex for DNA melting. *EMBO J* 23: 4253–4263.
- Burrows, P.C., Wigneshweraraj, S.R., and Buck, M. (2008) Protein–DNA interactions that govern AAA+ activatordependent bacterial transcription initiation. *J Mol Biol* **375**: 43–58.
- Burrows, P.C., Joly, N., Cannon, W.P.C.B., Rappas, M., Zhang, X., Dawes, K., *et al.* (2009) Coupling sigma Factor Conformation to RNA Polymerase Reorganisation for DNA Melting. *J Mol Biol* **387**: 306–3019.
- Cannon, W., Gallegos, M.T., Casaz, P., and Buck, M. (1999) Amino-terminal sequences of sigmaN (sigma54) inhibit RNA polymerase isomerization. *Genes Dev* 13: 357–370.
- Cannon, W.V., Gallegos, M.T., and Buck, M. (2000) Isomerization of a binary sigma-promoter DNA complex by transcription activators. *Nat Struct Biol* **7**: 594–601.
- Cannon, W., Gallegos, M.T., and Buck, M. (2001) DNA melting within a binary sigma(54)-promoter DNA complex. *J Biol Chem* **276**: 386–394.
- Chaney, M., Grande, R., Wigneshweraraj, S.R., Cannon, W., Casaz, P., Gallegos, M.T., *et al.* (2001) Binding of tran-

scriptional activators to sigma 54 in the presence of the transition state analog ADP-aluminum fluoride: insights into activator mechanochemical action. *Genes Dev* **15:** 2282–2294.

- Chen, B., Doucleff, M., Wemmer, D.E., De Carlo, S., Huang, H.H., Nogales, E., *et al.* (2007) ATP ground- and transition states of bacterial enhancer binding AAA+ ATPases support complex formation with their target protein, sigma54. *Structure* **15:** 429–440.
- Collaborative Computational Project, Number 4. (1994) The CCP4 Suite: Programs for Protein Crystallography. *Acta Crystallog D* **50**: 760–763.
- Dago, A.E., Wigneshweraraj, S.R., Buck, M., and Morett, E. (2007) A role for the conserved GAFTGA motif of AAA+ transcription activators in sensing promoter DNA conformation. *J Biol Chem* **282**: 1087–1097.
- Enemark, E.J., and Joshua-Tor, L. (2006) Mechanism of DNA translocation in a replicative hexameric helicase. *Nature* **442:** 270–275.
- Evans, P. (2006) Scaling and assessment of data quality. *Acta Crystallogr D Biol Crystallogr* **62**: 72–82.
- Fodor, A.A., and Aldrich, R.W. (2004) Influence of conservation on calculations of amino acid covariance in multiple sequence alignments. *Proteins* **56**: 211–221.
- Gai, D., Zhao, R., Li, D., Finkielstein, C.V., and Chen, X.S. (2004) Mechanisms of conformational change for a replicative hexameric helicase of SV40 large tumor antigen. *Cell* **119:** 47–60.
- Han, Y.W., Iwasaki, H., Miyata, T., Mayanagi, K., Yamada, K., Morikawa, K., and Shinagawa, H. (2001) A unique betahairpin protruding from AAA+ ATPase domain of RuvB motor protein is involved in the interaction with RuvA DNA recognition protein for branch migration of Holliday junctions. *J Biol Chem* **276**: 35024–35028.
- Hanson, P.I., and Whiteheart, S.W. (2005) AAA+ proteins: have engine, will work. *Nat Rev Mol Cell Biol* **6:** 519– 529.
- Iyer, L.M., Leipe, D.D., Koonin, E.V., and Aravind, L. (2004) Evolutionary history and higher order classification of AAA+ ATPases. J Struct Biol 146: 11–31.
- Joly, N., Schumacher, J., and Buck, M. (2006) Heterogeneous nucleotide occupancy stimulates functionality of phage shock protein F, an AAA+ transcriptional activator. *J Biol Chem* **281**: 34997–35007.
- Joly, N., Rappas, M., Wigneshweraraj, S.R., Zhang, X., and Buck, M. (2007) Coupling nucleotide hydrolysis to transcription activation performance in a bacterial enhancer binding protein. *Mol Microbiol* **66**: 583–595.
- Joly, N., Rappas, M., Buck, M., and Zhang, X. (2008) Trapping of a transcription complex using a new nucleotide analogue: AMP aluminium fluoride. *J Mol Biol* **375**: 1206– 1211.
- Jones, T.A., Zou, J.Y., Cowan, S.W., and Kjeldgaard, M. (1991) Improved methods for building protein models in electron density maps and the location of errors in these models. *Acta Crystallogr A* **47:** 110–119.
- Lee, S.Y., De La Torre, A., Yan, D., Kustu, S., Nixon, B.T., and Wemmer, D.E. (2003) Regulation of the transcriptional activator NtrC1: structural studies of the regulatory and AAA+ ATPase domains. *Genes Dev* **17:** 2552–2563.
- Leslie, A.G.W. (1992) Joint CCP4 and ESF-EACMB New-

sletter on Protein Crystallography, No. 26. Warrington, UK: Daresbury Laboratory.

- McGeoch, A.T., Trakselis, M.A., Laskey, R.A., and Bell, S.D. (2005) Organization of the archaeal MCM complex on DNA and implications for the helicase mechanism. *Nat Struct Mol Biol* **12:** 756–762.
- Martin, A., Baker, T.A., and Sauer, R.T. (2008a) Diverse pore loops of the AAA+ ClpX machine mediate unassisted and adaptor-dependent recognition of ssrA-tagged substrates. *Mol Cell* **29:** 441–450.
- Martin, A., Baker, T.A., and Sauer, R.T. (2008b) Pore loops of the AAA+ ClpX machine grip substrates to drive translocation and unfolding. *Nat Struct Mol Biol* **15**: 1147–1151.
- Morett, E., and Buck, M. (1989) *In vivo* studies on the interaction of RNA polymerase-sigma 54 with the *Klebsiella pneumoniae* and *Rhizobium meliloti nifH* promoters. The role of NifA in the formation of an open promoter complex. *J Mol Biol* **210**: 65–77.
- Murshudov, G.N., Vagin, A.A., and Dodson, E.J. (1997) Refinement of macromolecular structures by the maximum-likelihood method. *Acta Crystallog D* **53**: 240– 255.
- Olmea, O., Rost, B., and Valencia, A. (1999) Effective use of sequence correlation and conservation in fold recognition. *J Mol Biol* **293:** 1221–1239.
- Popham, D.L., Szeto, D., Keener, J., and Kustu, S. (1989) Function of a bacterial activator protein that binds to transcriptional enhancers. *Science* **243**: 629–635.
- Rappas, M., Schumacher, J., Beuron, F., Niwa, H., Bordes, P., Wigneshweraraj, S., *et al.* (2005) Structural insights into the activity of enhancer-binding proteins. *Science* **307**: 1972–1975.
- Rappas, M., Schumacher, J., Niwa, H., Buck, M., and Zhang, X. (2006) Structural basis of the nucleotide driven conformational changes in the AAA+ domain of transcription activator PspF. *J Mol Biol* **357**: 481–492.
- Rombel, I., Peters-Wendisch, P., Mesecar, A., Thorgeirsson, T., Shin, Y.K., and Kustu, S. (1999) MgATP binding and hydrolysis determinants of NtrC, a bacterial enhancerbinding protein. *J Bacteriol* **181**: 4628–4638.
- Sallai, L., and Tucker, P.A. (2005) Crystal structure of the central and C-terminal domain of the sigma (54) -activator ZraR. J Struct Biol 151: 160–170.
- Sasse-Dwight, S., and Gralla, J.D. (1988) Probing the *Escherichia coli* glnALG upstream activation mechanism in vivo. *Proc Natl Acad Sci USA* **85**: 8934–8938.
- Schumacher, J., Zhang, X., Jones, S., Bordes, P., and Buck, M. (2004) ATP-dependent transcriptional activation by bacterial PspF AAA+protein. *J Mol Biol* **338**: 863–875.
- Schumacher, J., Joly, N., Rappas, M., Zhang, X., and Buck, M. (2006) Structures and organisation of AAA+ enhancer binding proteins in transcriptional activation. *J Struct Biol* **156**: 190–199.
- Schumacher, J., Joly, N., Rappas, M., Bradley, D., Wigneshweraraj, S.R., Zhang, X., and Buck, M. (2007) Sensor I Threonine of the AAA+ ATPase transcriptional activator PspF is involved in coupling nucleotide triphosphate hydrolysis to the restructuring of {sigma}54-RNA polymerase. *J Biol Chem* **282**: 9825–9833.
- Schumacher, J., Joly, N., Claeys-Bouuaert, I.L., Aziz, S.A., Rappas, M., Zhang, X., and Buck, M. (2008) Mechanism of

homotropic control to coordinate hydrolysis in a hexameric AAA+ ring ATPase. *J Mol Biol* **381:** 1–12.

- Storoni, L.C., McCoy, A.J., and Read, R.J. (2004) Likelihoodenhanced fast rotation functions. *Acta Crystallogr D Biol Crystallogr* **60**: 432–438.
- Studholme, D.J., and Dixon, R. (2003) Domain architectures of sigma54-dependent transcriptional activators. *J Bacteriol* **185**: 1757–1767.
- Thomsen, N.D., and Berger, J.M. (2008) Structural frameworks for considering microbial protein- and nucleic aciddependent motor ATPases. *Mol Microbiol* 69: 1071–1090.
- Wedel, A., and Kustu, S. (1995) The bacterial enhancerbinding protein NTRC is a molecular machine: ATP hydrolysis is coupled to transcriptional activation. *Genes Dev* **9**: 2042–2052.
- Wigneshweraraj, S.R., Nechaev, S., Bordes, P., Jones, S., Cannon, W., Severinov, K., and Buck, M. (2003) Enhancerdependent transcription by bacterial RNA polymerase: the beta subunit downstream lobe is used by sigma 54 during open promoter complex formation. *Methods Enzymol* **370**: 646–657.
- Wigneshweraraj, S.R., Burrows, P.C., Severinov, K., and Buck, M. (2005) Stable DNA opening within open promoter complexes is mediated by the RNA polymerase beta'-jaw domain. *J Biol Chem* **280**: 36176–36184.

- Wigneshweraraj, S., Bose, D., Burrows, P.C., Joly, N., Schumacher, J., Rappas, M., *et al.* (2008) Modus operandi of the bacterial RNA polymerase containing the sigma54 promoter-specificity factor. *Mol Microbiol* **68**: 538–546.
- Wittinghofer, A. (1997) Signaling mechanistics: aluminum fluoride for molecule of the year. *Curr Biol* **7:** R682–R685.
- Yamada, K., Miyata, T., Tsuchiya, D., Oyama, T., Fujiwara, Y., Ohnishi, T., *et al.* (2002) Crystal structure of the RuvA-RuvB complex: a structural basis for the Holliday junction migrating motor machinery. *Mol Cell* **10**: 671–681.
- Zhang, X., Chaney, M., Wigneshweraraj, S.R., Schumacher, J., Bordes, P., Cannon, W., and Buck, M. (2002) Mechanochemical ATPases and transcriptional activation. *Mol Microbiol* **45:** 895–903.

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