Molecular basis of nucleotide-dependent substrate engagement and remodeling by an AAA+ activator

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ABSTRACT

Binding and hydrolysis of ATP is universally required by AAA+ proteins to underpin their mechano-chemical work. Here we explore the roles of the ATPase site in an AAA+ transcriptional activator protein, the phage shock protein F (PspF), by specifically altering the Walker B motif sequence required in catalyzing ATP hydrolysis. One such mutant, the E108Q variant, is defective in ATP hydrolysis but fully remodels target transcription complexes, the RNAP-σ54 holoenzyme, in an ATP dependent manner. Structural analysis of the E108Q variant reveals that unlike wild-type protein, which has distinct conformations for E108 residue in the ATP and ADP bound forms, E108Q adapts the same conformation irrespective of nucleotide bound. Our data show that the remodeling activities of E108Q are strongly favored on pre-melted DNA and engagement with RNAP-σ54 using ATP binding can be sufficient to convert the inactive holoenzyme to an active form, while hydrolysis per se is required for nucleic acid remodeling that leads to transcription bubble formation. Furthermore, using linked dimer constructs, we show that RNAP-σ54 engagement by adjacent subunits within a hexamer are required for this protein remodeling activity while DNA remodeling activity can tolerate defective ATP hydrolysis of alternating subunits.

INTRODUCTION

In AAA+ proteins a cycle of nucleotide (ATP) binding and hydrolysis allows the formation of at least two different conformational states, each associated with a distinctive functional state (1). AAA+ proteins often exist as hexamers and inter-subunit coupling allows the propagation of conformation from one active site to others within an AAA+ assembly (2). The ATP-dependent conformational coupling can support various activities of AAA+ assemblies, including that of a bimodal switch and a processive machine (3,4). The former is often associated with slow ATP hydrolysis, the latter with multiple rounds of coupled binding and hydrolysis events. AAA+ proteins share conserved functional motifs including those of Walker A for nucleotide binding, Walker B and Arginine fingers for ATP hydrolysis.

Bacterial enhancer binding proteins (bEBPs) are specialized AAA+ proteins that are required to activate the sigma54-dependent transcriptional complexes (5,6). Sigma54 (σ54) recruits RNA polymerase (RNAP) to specific promoter sites and forms a stable closed complex (RPc) unable to proceed to intermediate states (RPi) or the open complex (RPo) used in transcription initiation. As transcriptional activators, bEBPs function to catalyze the conversion of RPc to RPo, causing the RNAP-σ54 holoenzyme to isomerize and promoter DNA to melt, leading to transcription initiation (7,8,9,10). Formation of RPo is associated with large-scale conformational changes in RNAP-σ54, and these are inhibited by sigma54 prior to the remodeling by bEBP activators (11–15).

bEBPs can exist in a number of distinct functional and conformational states including that of nucleotide free (APO), ATP bound, transition state (as mimicked by ATP transition state analogue ADP.AIF), and ADP bound (16–19). Similar to other ATPases, ATP binding and hydrolysis involve distinct kinetic steps including ATP binding, gamma-beta bond cleavage, Pi release and ADP release. Hexameric bEBPs are shown to stably bind sigma54 as well as RPc, only when non-hydrolysable ATP or ATP hydrolysis transition state analogues are present or when the ATP is very slowly hydrolyzed (17,20–22). However, under such conditions either none or an incomplete remodeling of RPc is observed, and the de novo RNA melting characteristic of RPo formation is not observed (23–25). Full remodeling of RPc and formation of RPo was therefore suggested to depend on ADP + Pi formation and/or Pi release (23,26).

A number of important structural and functional motifs within bEBPs have been identified and characterized, including the signature GAFTGA motif, which is almost...
invariant among bEBPs and directly engages with the RPc substrate, and the ‘Glutamate Switch’, a common signature in AAA+ proteins (Figure 1A). The ‘Glutamate Switch’ pair (E-N) was identified originally in PspF as the E108 side chain adopts different conformations in the presence of ATP or ADP, forming or interrupting the interactions with N64 (3,11,16,22,27). These differences are correlated with the distinct conformations of the GAFTGA containing L1 loop inserted into the AAA+ domain, determining with the distinct conformations of the GAFTGA containing 470 nucleotides downstream from the multiple cloning site) and 100 mM holoenzyme (reconstituted in a 1:4 ratio of core RNAP: σ54). For open complex formation 5 mM PspF1–275 and 4 mM ATP or dATP (to limit the role of PspF to RPc formation, since dATP is not an efficient substrate for RNAP) were used. The elongation mixture contained 0.1 mg/ml haperin, 100 nM ATP, CTP and GTP and 50 nM UTP (0.074 μCi/μl of [α-32P] UTP (3000 Ci/mmol)). Reactions were incubated at 37°C and stopped with 4 μl of formamide dye mixture (0.3 mg/ml xylene cyanol, 0.3 mg/ml bromophenol blue, 20 mM EDTA in deionised formamide). 7 μl of the samples were run on a 4% denaturing sequencing gel for 2 h at 50 W. The gels were then dried and quantified and analyzed by PhosphorImager analysis to measure transcriptional activity compared to WT and the varying nucleotide conditions for hydrolysis by PspF variants. All experiments were performed minimally in triplicate.

**Gel Filtration**

Forty microliters samples were prepared in running buffer (20 mM Tris–HCl pH 8.0, 50 mM NaCl, 15 mM MgCl2, with or without: 0.4 mM ATP; 0.02 mM ATP) and centrifuged at 15 000 rpm for 3 min (4°C) to remove any particulates. The samples were pipetted into 200 μl crimp autosampler vials. Each sample was placed in the refrigerated autosampler of the Thermo Scientific Surveyor HPLC system. The Biosep Sec-S3000 gel filtration column (Phenomenex) was attached to the system in a column oven (Phenomenex) at a temperature of 8°C. The detector was set to detect broad spectrum and UV at 280 nm. The flow rate was set at 0.5 ml/min with a pressure limit of 1500 psi and the injections were set at 15 μl. Security Guard (Phenomenex) was used as the guard column to protect the Biosep column from particulate damage.

**ATPase assay**

* NADH-coupled ATPase assay: The steady-state ATPase activity of PspF1–275 was measured at 37°C using an NADH-coupled ATP regeneration system (Norby, 1988). Reactions were conducted in 100 μl volumes: 25 mM Tris–HCl (pH 8.0), 100 mM KCl, 10 mM MgCl2, 1 mM DTT, 1 mM NADH, 10 mM phospheolpyruvate, 10 U/ml pyruvate kinase, 20 U/ml lactate dehydrogenase, 0.02 mM ATP, ±10 μM σ54 and 1–5 μM PspF1–275. The rate of NADH absorbance decrease at 340 nm is proportional to the rate of steady-state ATP hydrolysis. Assays were performed minimally in triplicate.

**TLC plate ATPase assay** Typically in a 10 μl volume, 4 μM PspF1–275 was preincubated with the ATPase buffer.
Figure 1. (A) Crystal structure of PspF AAA domain with key residues labeled. (B) Cartoon depicting the transition from RP_c to RP_o with intermediates RP_i (either WT PspF trapped with ADP-AIF or PspF E108Q variant in complex with ATP). (C) Transcriptional activities of E108 variants (E108A, E108D and E108Q) and wild-type protein. Different nucleotides are assayed with duplex DNA or DNA with pre-opened transcription bubbles.
(20 mM Tris-HCl pH 8.0, 50 mM NaCl, 15 mM MgCl₂, 0.1 mM EDTA, 10 mM DTT) at 37°C for 5 min. ATP hydrolysis was initiated by addition of 1 mM unlabeled ATP and 0.6 mCi/ml [α-32P] ATP (3000 Ci/mmol) and incubated for various time spans at 37°C. Reactions were quenched by addition of 5 volumes of 2 M formic acid. The [α-32P] ADP was separated from the [α-32P] ATP by thin layer chromatography (Macherey–Nagel) in 0.4 M K₂HPO₄/0.7 M boric acid. Radioactivity was scanned by PhosphorImager and analyzed by Aida software. Assays were performed minimally in triplicate.

Native gel mobility shift assay

Native gel mobility shift assays were conducted in STA buffer in a total reaction volume of 10 μl containing 200 nM RNAP-σ54 (reconstituted using a 1:4 ratio of RNAP-σ54) and 20 nM 32P-labeled probe, which was incubated for 5 min at 37°C. Reactions were analyzed using a 4.5% native (non-denaturing) gel run at 100 V for 55 min. The gels were dried and protein–DNA complexes were visualized and quantified using an FLA-5000 PhosphorImager. These experiments were minimally performed in triplicate.

Protein purification, crystallization and nucleotide soaking

PspF1–275E108Q was over expressed and purified as described previously (11). Crystals of PspF1–275E108Q were grown in a sitting drop vapor diffusion experiment at 18°C with 40 mg/ml protein concentration in a precipitant consisting of 12–16% MPD, 2 M ammonium formate and 100 mM Bis-tris Propane, pH 8.0. For APO datasets, the native crystals were soaked in a buffer containing 30% (w/v) PEG8000, 30% (w/v) glycerol and 100 mM HEPES, pH 8.0 before flash freezing in liquid nitrogen. For Mg-ATP and ADP bound structures, native crystals were soaked in 30% (w/v) PEG8000, 30% (w/v) glycerol, 100 mM HEPES, pH 8.0 with 30 mM MgCl₂ and 25 mM ATP/ADP for 2 h at room temperature and then flash frozen in the same buffer in liquid nitrogen.

Crystallographic data collection and processing

Data were collected under cryogenic conditions (100 K) at the Diamond Light Source (DLS, UK) at beam lines I02 for nucleotide soaked crystals and I04 for APO crystals. All the datasets were processed using XDS (29) and the statistics are summarized in Table 1.

Structure determination

Phases for the datasets were obtained using Phaser (30) resulting from molecular replacement solutions, with the native WT PspF structure as the search model (2BJW) (11). Refinement of the models was carried out using phenix.refine implemented in the software suite PHENIX (31). Cross-validation was performed using 5% of the dataset set aside for Rfree calculations. Electron density for the nucleotide analogues ATP and ADP was clearly visible after an initial round of refinement using the molecular replacement solution of the native structure and the ATP and ADP were then accordingly placed into density. Clear density was also visible for Mg²⁺ in the Mg-ATP structure and was thus placed. Refinement statistics for all the structures are summarized in Table 1.

RESULTS

The effects of the Walker B residue E108 are highly dependent on DNA substrates

In order to investigate the precise functionality of the Walker B residue E108, which was proposed to coordinate the hydrolytic water molecule for in-line nucleophilic attack (2), we carried out transcription assay using WT, E108A, E108D and E108Q variant proteins in the presence of different nucleotides and in the presence of a DNA duplex (mimicking DNA substrate in RPo) or with mismatches from −10 to −1, mimicking the transcription bubble, which is found in a fully formed RPo (Figure 1) (as in (23)). The E to A mutation removes the side chain of the residue, E to D maintains the negative charge property but with a shortened side chain while E to Q change maintains the side chain geometry except the OH of one side chain branch is replaced by a NH₂ group. In addition to different DNA substrates, we also challenged the reactions with different non-hydrolysable nucleotide analogues, in order to probe different functional states during ATP hydrolysis cycle. Remodeling activities of transcriptionally competent complexes were monitored by 5′-UpGGG transcript formation primed with the dinucleotide primer UpG (Figure 1). Notably, with a standard duplex DNA, only E108D behaves similarly to WT protein, and dATP or ATP (we used dATP as well as ATP to maintain a constant pool of hydrolyzable nucleoside triphosphate for PspF NTPase activity during transcription) is required for transcription activation. All other variant proteins tested are defective in transcription activation using dATP or ATP. However, with the pre-opened DNA substrate, all variants showed transcription activation in an (Nucleoside triphosphate) NTP-dependent fashion although E108A has reduced activities. Strikingly, at low ATP concentration, when WT protein is unable to activate transcription, the E108Q variant not only then activates transcription on duplex DNA but this activity is elevated when the DNA is pre-opened.

E108Q mutant supports transcription at lower ATP concentrations compared to wild-type even though it has reduced ATPase activity and enhanced ATP binding

To characterize the ATP dependence of E108Q further, we carried out transcription activation assays for E108Q and WT in a range of ATP concentrations. In agreement with previous reports, the WT has a maximal transcription stimulating activity at around 1–2 mM ATP (Figure 2A). Interestingly, E108Q reaches its maximal activity at 100-fold less ATP concentrations, around 0.02 mM (Figure 2A). Since transcription activation is closely related to ATPase activity, we measured the effects of the Walker B substitution on ATP binding and hydrolysis capability. Not surprisingly, E108Q has severely impaired ATPase activity compared to WT (<1% activity), highlighting the importance of the integrity of Walker B residues. However, it has 30-fold in-
increase in binding affinity to ATP (Figure 2B) (22). These data show that E108Q binds ATP tightly, has poor hydrolysis rate, but can effectively activate transcription using the pre-formed transcription bubble at lower ATP concentration. PspF assemblies have been shown to exist in a mixed nucleotide bound state (32), where both ATP and ADP are bound and proposed to alter the detailed geometry of hexamer favoring its activity in making RPc. Further, high ATP concentrations inhibit PspF functionality. The higher affinity for ATP that E108Q possesses compared to the WT PspF suggest that at the E108Q ATPase activity and other functionality could be inhibited at ATP concentrations that are more optimal for WT PspF activities.

Previously, we have shown that the integrity of GAFTGA motif of the L1 is important for PspF to contact RPc (33) while W56 is essential for interacting with its negative regulator PspA, which has been proposed to act through the ‘Glutamate Switch’ pair (E108-N64) that is part of the ‘Glutamate Switch’ pair (E108-N64) that links substrate binding to ATP binding site, we investigated whether its ATPase activity could be stimulated by binding to its substrate. We added RPc to ATPase reactions. Within a 5-min timescale WT PspF shows no discernible difference (at 1 mM ATP) when supplemented with RPc. At 0.1 mM ATP, however, the addition of RPc stimulates ATP turnover by 2-fold (Supplementary Figure S2). It is unclear if this effect is due to a higher affinity e.g. ATP or more efficient phosphate release. Although WT shows a 2-fold stimulation by binding to its substrate, the E108Q variant shows a 4-fold stimulation upon binding to RPc at the lower ATP concentration (0.1 mM) and this stimulation depends on substrate interaction via the GAFTGA motif as mutating T86A, which is shown to impair its interactions with RPc, target remodeling substrate abolishes the stimulatory effects (Figure 2D). The double variants E108Q-T86A and E108Q-W56A double display an increase in ATPase activity compared to E108Q (Figure 2D). These observations are consistent with previous studies that implicate the connection of the L1 loop and W56 to the E108-N64 pair as a means of hydrolysis control (16,27).

### Table 1. Crystallographic data and refinement statistics

|                        | Apo-PspF E108Q | Mg-ATP-PspF E108Q | ADP- PspF E108Q |
|------------------------|----------------|-------------------|-----------------|
| Space group            | P65            | P65               | P65             |
| Unit cell (Å)          | a = b = 113.72, c = 39.45 | a = b = 113.55, c = 39.51 | a = b = 113.38, c = 39.33 |
| A. Data reduction statistics |                |                   |                 |
| λ (Å)                  | 0.9794         | 0.9794            | 0.9794          |
| Resolution (Å)         | 28.43–1.63     | 28.39–1.54        | 28.35–1.42      |
| Total/unique reflections | (1.67–1.63)    | (1.59–1.54)       | (1.47–1.42)     |
| Redundancy             | 8.3 (7.9)      | 9.8 (9.2)         | 9.8 (9.4)       |
| Completeness (%)       | 21.1 (2.08)    | 23.9 (3.2)        | 28.7 (3.3)      |
| R-meas                 | 99.35 (93.34)  | 99.85 (98.54)     | 99.96 (99.63)   |
| CC(1/2)                | 0.06 (0.89)    | 0.052 (0.81)      | 0.042 (0.62)    |
| Wilson B-factor        | 23.35          | 21.67             | 18.58           |
| B. Refinement Statistics |                |                   |                 |
| Reflections (work/free)| 34746/1826     | 41255/2182        | 52102/2752      |
| Number of non hydrogen atoms/water/ligand | 2095/174/21 | 2203/213/59 | 2211/241/48 |
| Rwork (%)              | 0.172 (0.257)  | 0.147 (0.189)     | 0.169 (0.235)   |
| Rfree (%)              | 0.204 (0.283)  | 0.179 (0.265)     | 0.189 (0.261)   |
| Ramachandran favoured (%) | 98            | 98                | 99              |
| Ramachandran outliers (%) | 0.42          | 0                | 0               |
| RMS (bonds) (Å)        | 0.008          | 0.01              | 0.012           |
| RMS (angles) (°)       | 1.17           | 1.14              | 1.35            |
| Average B-factor       | 31             | 29                | 25              |
| (Macromolecule/ligand/water) | (30/51/45) | (28/44/41) | (23/33/40) |

are favored in the E108Q variant. Like W56A, N64S has been shown to escape negative regulation by PspA (27). Interestingly, E108Q-N64S double mutant is still under negative control by PspA, further supporting the notion that E108Q mutation promotes inhibitory routes by PspA that are independent of W56.
E108Q forms constitutive hexamers and alternative competitor resistant complexes

Hexamerization has been linked to nucleotide binding and is one pre-requisite for ATP hydrolysis by PspF (5). To investigate if the reduced ATPase activity of E108Q is due to impaired hexamerisation, we assayed the oligomeric state of PspF variants using gel filtration (Figure 3). For WT proteins, in the absence of nucleotide, PspF exists primarily as dimers at low μM concentrations and upon ATP binding, PspF shifts to a predominately hexameric peak. E108Q, on the other hand, is hexameric irrespective of the nucleotide being present, suggesting that the reduced ATPase activity is due to a reduced hydrolysis reaction per se, and not due to gross defects in hexamerisation or loss of nucleotide binding. The nucleotide analogue ADP-AlF has been traditionally used to capture the RNAP-σ54-DNA intermediate complex (RPc) (Figure 1B). E108Q, however, forms an RP, in the presence of ATP but not in the presence of ADP or in the APO state (Supplementary Figure S3).

E108Q clearly has altered properties in ATP binding and hydrolysis, hence altered kinetics of the ATP hydrolysis cycle. It is therefore potentially capable of forming altered intermediate transcriptional complexes with its RPc. To probe for these, we challenged the transcription reactions with the competitor heparin. Normally RPc is heparin sensitive while RPo is resistant. Interestingly, two distinct complexes formed for E108Q protein depending on the level of ATP, and these complexes are not observed for WT protein (Figure 4A). The magnitude of the shifts and the ability of ATP to stabilize complexes between σ54 and PspF E108 variants suggest that these gel mobility results were due to the presence of E108Q within the new complexes. At the non-inhibitory low ATP levels, as lower migrating complex containing E108Q is the predominant species and its formation is dependent on an intact GAFTGA motif (which contacts σ54) and that the new co-complex is stable under heparin challenge (100 μg/ml). Notably these complexes are distinct from RPc (by their stability to heparin) and the RPo (in terms of gel mobility) and so are most likely transcription intermediates containing the E108Q variant and lying between RPc and RPo on the pathway to transcription initiation. At higher ATP concentrations, a complex corresponding to RPo dominates although other fast migrating bands are also seen.
Figure 3. E108Q forms constitutive hexamers. Gel filtration profile of E108Q mutant protein compared to wild-type protein. WT PspF AAA domain predominantly exists as dimers in the absence of ATP, which is required for hexamer formation. PspF AAA domain E108Q variant exists as hexamers irrespective of nucleotide and the hexamers are not dependent on the integrity of RNAP-σ54 binding site (T86A in Loop1) nor the PspA inhibitor binding site (W56A).
which are consistent with PspF oligomers. This suggests that at inhibitory ATP concentrations, significant portion of PspF E108Q mutant protein dissociates from RNAP-σ54-DNA complex, consistent with the inhibitory effects at high ATP concentrations.

We considered whether the accumulation of transcription intermediates with E108Q might hinder steps after RP0 formation such as promoter escape. However E108Q at low ATP concentrations supported full length transcript formation, and did not prevent promoter escape (Figure 4B). Full length transcript assays showed that although E108Q exhibited a delayed activation compared to WT protein, and that E108Q displayed a persistent transcription stimulating activity as shown in the time course experiments (Figure 4B). Furthermore, our data show that the E108Q mutant protein could support full length transcript formation at least at the pspA promoter site, without addition of ATP after 120 min incubation with 0.02 mM initial ATP whereas WT protein did not with 4 mM initial ATP (Figure 4C, compare the boxed areas). Together these data support the idea that E108Q is a slow but stable productive ATPase compared to the faster WT PspF.

Previously, using pre-formed transcription bubble substrate, the ADPAlF nucleotide analogue was shown to sup-
port a partial remodeling of the RNAP-σ^{54} holoenzyme but not the extensive de novo DNA melting as seen in RPo, nor full length transcription, suggesting that RNAP-σ^{54} engagement at the ATP hydrolysis transition state can partially overcome transcription suppression imposed by RNAP-σ^{54} (23). Our results here with E108Q show that RNAP-σ^{54} engagement can also remodel the RNAP-σ^{54} holoenzyme to such an extent that it can overcome suppression to support full transcription persistently. Unlike the ADPAIF bound pre-initiation complex, which can only partially activate transcription, E108Q variant can fully activate transcription (comparing the activities between WT and E108Q) when DNA is pre-melted, suggesting that E108Q variant represents a functional state further along the activation pathway compared to those represented by ADPAIF bound complex. The slow ATPase activity of E108Q allows the detection of such an intermediate state that might be short-lived with WT PspF. Our data here thus support a model where the set of distinct kinetic steps in transcription activation intimately involve the functional states of the activator protein, driving the conversion of RPo to RPo.

### E108Q variants support functional asymmetry required for PspF activities

To explore further subunit specializations within hexameric βEBP assemblies, we constructed single chain forms of the PspF AAA+ domain with alternating WT and E108Q, E108D or E108A substitutions (36). A linked dimer form of PspF was constructed by adding a linker sequence between the C and N terminal residues of two PspF AAA domains and expressing the linked coding sequences as a single polypeptide chain (36). By assaying these constructs against fully duplexed or open DNA templates bound by RNAP-σ^{54} we show that E108Q, E108D and E108A, even when coupled to WT subunits, greatly favors transcription activation with pre-formed transcription-bubble over the duplex DNA substrates (Figure 5, Supplementary Figure S4). E108Q, E108D and E108A can all support transcription activation with pre-formed transcription bubble (Figure 1). E108Q, E108D and E108A are shown to have reduced ATPase activity but can still use ATP to engage with RPo. T86A in Loop1, on the other hand, does not alter ATPase activity but fails to engage with RPo (11). As expected, T86A cannot efficiently support transcription even when a subset of the subunits are WT (Figure 5) because two adjacent L1 loops need to be fully functional for binding to RPo. The ‘Glutamate Switch’ variant N64S, which is shown to be able to engage with the RNAP-σ^{54}, can also support transcription and greatly favors pre-formed transcription bubble over the duplex DNA. These outcomes imply that transcription activation can be supported by a fraction of the ATPase subunits, some of which may have defective ATPase activity but can associate with the target RPo, such as in WT/E108Q variant. However, subunits that are relatively fast for ATPase activity but do not stably associate with RPo, as shown in WT/T86A, are insufficient for transcription activation. These results highlight that multiple subunits with intact L1 loops are required to engage with the RNAP-σ^{54} (as shown in WT/T86A) for efficient remodeling although stable association with a subset of subunits (WT/E108Q) showing slow ATPase are sufficient. Interestingly, despite the defective ATPase activity of PspF E108Q, the ATPase activity of WT/E108Q linked dimer is WT-like at the higher ATP concentration (1 mM) but E108Q-like at lower ATP. This is in agreement with a model in which mixed nucleotide states exist but in a highly coordinated fashion within a hexameric PspF. It is possible that E108Q subunits adopt certain conformations that are coupled to adjacent subunits, which then adapt conformations competent for using ATP in remodeling the RPo. This model can also explain the elevated ATPase activity of the linked dimer at the lower ATP concentration (0.1 mM) compared to WT/WT (Supplementary Figure S5) as the increased ATP affinity of E108Q would promote efficient conformational coupling to its adjacent WT subunits.

### Crystal structures of PspF_{1–275} E108Q mutant in APO, ATP and ADP states

In order to understand the biochemical properties of E108Q and to provide a structural basis for the ‘Glutamate Switch’ pair, we determined the crystal structures of E108Q mutant of PspF AAA+ domain (PspF_{1–275},E108Q) in the absence of nucleotide (APO), and in the presence of Mg-ATP or Mg-ADP. The crystals diffracted to 1.6–1.8 Å and have the same space group and unit cells as those of WT proteins and are thus subsequently rebuilt and refined using WT protein structures as templates. The crystallographic data and refinement statistics are summarized in Table 1.

We first compared the conformations surrounding the ‘Glutamate Switch’. In WT protein, when ADP is bound, E108 does not interact with N64 albeit in the ATP bound or APO state, E108 interacts with N64 (Figure 6A, (16)). In the mutant structures, on the contrary, E108Q in the ADP bound state as well as in the APO state also interacts with N64 (Figure 6B). This conformation, which enables the ‘Glutamate Switch’ pair to interact, is linked to the ability of PspF being able to engage with its target, the RPo. The crystal structures of the E108Q therefore suggest that PspF E108Q variant could engage with RPo, irrespective of its nucleotide bound states. Such a conformation could contribute to the ability of E108Q to bind RPo and move RPo along the pathway of transcription activation despite its slow ATPase activity.

To further investigate the structural basis for these distinct conformations, we analyzed the detailed interaction networks surrounding these key residues in the different nucleotide bound states, both in the WT and in the mutant protein structures.

The WT and E108Q mutant structures are almost identical surrounding the ‘Glutamate Switch’ pair (Figure 7 and also (16)). The ‘Glutamate Switch’ pair is located at the interface of two neighboring protomers (Figure 5 and S7A). Similar residues are involved in the interaction networks and the interactions are mainly electrostatic involving E108, N64 as well as residues from adjacent protomer including the R finger residue R162’ and D164’ (‘denotes adjacent protomer). Specifically, the oxygen atom (O2E in Figure 7B) of E108 interacts with NH2 groups of N64 and R162’ from adjacent protomer. R162’ also interacts with D164’ (Figure 7A–
Figure 5. Activities of single chain forms of PspF variants. Alternating wild-type with glutamate switch mutants (E108 or N64 variants) within the hexamer result in forms that favor protein remodeling over DNA remodeling.

C). These interactions are similarly maintained in the mutant structure since the mutation does not affect the oxygen (OE2) of residue 108.

A direct superposition of the mutant and WT ADP bound structures reveal two major differences surrounding the ‘Glutamate Switch’ pair. In addition to the distinct conformation of E108 and E108Q, D164′ from adjacent protomer, another highly conserved residue in bEBPs, also adapts distinct conformations (Figure 7D). In the WT structure, there are a number of water molecules in the nucleotide binding pocket, mediating a network of hydrogen bonding interactions involving E108, N64 and the sensor II R227 as well as residues from the adjacent protomer including R162′ and D164′ (see Figure 7E). In particular, E108 and D164′ only form hydrogen bond interactions with water molecules. However, in the mutant protein, the OH group in E108 (OE1 in Figure 7E) that is involved in hydrogen bonding is replaced by an NH2 group in E108Q, no longer able to maintain a stable hydrogen bond network with these water molecules. Instead, E108Q returns to a conformation optimal for interacting with N64 through OE2 of E108Q (Figure 7F). The loss of water mediated hydrogen bonding network also destabilizes the D164′ side chain conformation. D164′ side chain rotates to interact with R162′. Consequently, in this conformation, R162′ makes charge-charge interactions with both E108Q and D164′ (Figure 7F), resulting in distinctive side chain conformations in ADP bound states between WT and E108Q mutant proteins.

DISCUSSION

A structural basis for the precise role of E108 and the distinct properties of E108 variants

Our data on the formation of a distinct transcriptional intermediate complex with E108Q mutant show that a range of RNAP-σ54 engagements can be obtained when ATP hy-
Figure 6. Structural changes of the ‘Glutamate Switch’ pair. (A) In wild-type (WT) protein, the E108 adapts different conformations depending on the nucleotide states. (B) In E108Q variant protein, irrespective of nucleotide bounds states, E108Q adapts a conformation similar to the APO/ATP bound conformation in the WT protein, where the Glutamate Switch pair is engaged. Gray—apo, orange—ATP, cyan—ADP. OE1 and OE2 refer to the two different glutamate oxygen atoms.

Figure 7. Structural comparisons of the nucleotide binding pockets between wild-type (WT) and E108Q mutant proteins. Red—ATP bound states, blue—ADP bound states. (A) Superposition of WT and E108Q ATP bound states. (B) WT ATP states showing key interactions involving E108 OE2. (C) E108Q ATP state showing key interactions. (D) Superposition of WT and E108Q ADP states. (E) WT ADP state showing key interactions involving E108 OE1 which is mutated in E108Q. (F) E108Q ADP state showing interactions involving OE2 instead.
drolysis is slowed down. The significantly impaired ATPase activity of E108Q mutant and its ability to activate transcription when a transcription bubble is pre-formed imply that efficient RNAP-σ\(^{54}\) engagement is sufficient for removing transcription inhibition but not sufficient for extensive \textit{de novo} DNA opening. ATP hydrolysis and the associated conformational changes in the AAA domains of at least two adjacent subunits with a hexameric assembly are required for this DNA remodeling. Strikingly, competitor resistant transcriptional intermediates are evident with E108Q (Figure 4A), and these intermediates can be correlated with persistent transcription of E108Q at lower ATP concentration (Figure 4B) or even without further ATP addition after a prolonged incubation with R\(_{\alpha}\). (Figure 4C).

It appears that the E108Q variant is competent at engaging R\(_{\alpha}\) in a nucleotide bound state prior to ADP + Pi release. This engagement property of E108Q can be explained by considering the altered interaction networks in its nucleotide bound forms compared to the WT. These changes are mainly due to the hydrogen bonding networks mediated via water molecules and involve the OH group of one of the E108 side chain branches. Mutating this OH to NH\(_2\) would thus destabilize the water-mediated hydrogen bonding network, irrespective of the nucleotide bound state. This explains its ability to engage stably with RNAP-σ\(^{54}\), hence its favored ability in activating transcription from a pre-opened transcription bubble and persistent transcription under competitor challenge conditions.

The single chain form of PspF shows that when only a subset of E108Q subunits is present in the hexamer, its ability to remodel DNA is similar to that of WT (Figure 5), implying that DNA opening does not require an equivalent active ATPase activity of adjacent subunits. In low ATP concentration conditions, WT/E108Q does not activate transcription to the same level as E108Q (Supplementary Figure S4), suggesting that even though active ATPase activity is not required for adjacent subunits, ATP binding to adjacent subunits is required for transcription activation. WT has 30-fold lower affinity for ATP, correlating with its optimal ATP concentration being 100-fold higher than that of E108Q variant. These results support a model in which PspF does not function as a processive motor. Rather a bimodal switch function of PspF requiring the cooperation of several neighboring subunits, at least for ATP binding, seems to operate. This correlates with the requirement of stable RNAP-σ\(^{54}\) engagement by several adjacent subunits to re-model R\(_{\alpha}\) and activate transcription. Furthermore, defects in ‘Glutamate Switch’ mutants (such as those E108 or N64 mutants, Figure 5) results in a form of PspF that favors protein remodeling over DNA remodeling, supporting the idea that stable RNAP-σ\(^{54}\) engagement is sufficient to remove the suppression imposed by RNAP-σ\(^{54}\) on protein organization for R\(_{\alpha}\) formation while full ATP hydrolysis and associated conformational changes are required for the DNA melting and transcription bubble formation seen in an R\(_{\alpha}\).

**Implications for isomerisation**

Our results reveal and explain the differences in the ATP binding and hydrolysis properties between the WT and the E108Q variant proteins. The preferred conformation of E108Q in the ‘inactivated’ form could also explain the significantly enhanced affinity for ATP (32-fold increased compared to WT) and the ability to readily form hexamers. However, due to its reduced stability in the ‘for hydrolysis activated’ form, the ability to hydrolyse ATP is significantly impaired (<1% compared to WT). The constitutive hexamer formation in E108Q and reduced ATPase activity could also explain the observed stimulatory effects on ATPase activity in E108Q upon RNAP-σ\(^{54}\) binding, which re-orient the E108Q to be in the ‘for hydrolysis activated’ form. In WT proteins, the ATPase activity is affected by hexameration which in turns is affected by ATP and RNAP-σ\(^{54}\) binding. Furthermore, the high base level of the ATPase activity in the WT protein suggests a preferred orientation of E108 in the ‘for hydrolysis activated’ form, hence the limited stimulatory effect upon RNAP-σ\(^{54}\) binding.

Interestingly, in WT, the optimal ATP concentration for ATP hydrolysis as well as transcription activation is at 1–2 mM while the E108Q variant requires significantly less ATP (0.02 mM) for its optimal function even though the ATP hydrolysis rate is much slower. This reduced ATPase is partially restored due to the stimulatory effects upon binding to the R\(_{\alpha}\). These results also re-enforce that R\(_{\alpha}\) engagement, not remodeling due to hydrolysis \textit{per se}, is sufficient in removing the inhibition to protein conformational change in R\(_{\alpha}\), which forms a major kinetic barrier to isomerisation to make the R\(_{\alpha\theta}\). Due to the 30-fold increased affinity for ATP, significantly lower ATP concentrations are thus required for stable R\(_{\alpha}\) engagement. Interestingly, mutations in the ‘Glutamate Switch’ N64 have similar phenotypes to those of E108 in favoring the ‘pre-opened transcription bubble’ as a transcription substrate (Figure 5), suggesting that the ability for the ‘Glutamate Switch’ to form and dissociate are key to DNA opening. Target engagements can be achieved when ‘Glutamate Switch’ pair is formed as demonstrated with E108Q mutant protein, which readily forms hexamers and binds to ATP, and is able to engage with the R\(_{\alpha}\) complex for a prolonged period of time ahead of ATP turnover. The reduced hydrolysis rate would reduce the speed for the final conformation changes required for transcription bubble formation but favor protein remodeling (23). The particular properties of the E108Q variant provide evidence for a set of distinct kinetic steps along the activation process. The altered dynamics between the different steps due to the E108Q mutation seems to result in an overall more efficient use of ATP in this process, but as noted above the penalties are defects in DNA opening and a strong ATP inhibition and a slow conversion of R\(_{\alpha}\) to R\(_{\alpha\theta}\). In physiologically conditions, the 2–3 mM concentrations of ATP in \textit{Escherichia coli} suggests that E108Q, though a functional ATPase variant, is unlikely to be viable to support transcription.

**PDB ACCESSION CODES**

4QNM: PspF1-275E108Q mutant; 4QNR: PspF1-275E108Q mutant bound to ATP; 4QOS: PspF1-275E108Q mutant bound to ADP.
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