DNA Binding Compatibility of the *Streptococcus pneumoniae* SsbA and SsbB Proteins

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**Abstract**

**Background:** *Streptococcus pneumoniae* has two paralogous, homotetrameric, single-stranded DNA binding (SSB) proteins, designated SsbA and SsbB. Previous studies demonstrated that SsbA and SsbB have different solution-dependent binding mode preferences with variable DNA binding capacities. The impact of these different binding properties on the assembly of multiple SsbAs and SsbBs onto single-stranded DNA was investigated.

**Methodology/Principal Findings:** The complexes that were formed by the SsbA and SsbB proteins on dT oligomers of defined lengths were examined by polyacrylamide gel electrophoresis. Complexes containing either two SsbAs or two SsbBs, or mixed complexes containing one SsbA and one SsbB, could be formed readily, provided the dT oligomer was long enough to satisfy the full binding mode capacities of each of the bound proteins under the particular solution conditions. Complexes containing two SsbAs or two SsbBs could also be formed on shorter dT oligomers via a “shared-strand binding” mechanism in which one or both proteins were bound using only a portion of their potential binding capacity. Mixed complexes were not formed on these shorter oligomers, however, indicating that SsbA and SsbB were incompatible for shared-strand binding. Additional experiments suggested that this shared-strand binding incompatibility may be due in part to differences in the structure of a loop region on the outer surface of the subunits of the SsbA and SsbB proteins.

**Conclusion/Significance:** These results indicate that the SsbA and SsbB proteins may co-assemble on longer DNA segments where independent binding is possible, but not on shorter DNA segments where coordinated interactions between adjacent SSBs are required. The apparent compatibility requirement for shared-strand binding could conceivably serve as a self-recognition mechanism that regulates the manner in which SsbA and SsbB interact in *S. pneumoniae*.

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**Introduction**

The naturally transformable Gram-positive bacterium *Streptococcus pneumoniae* has two paralogous, homotetrameric, single-stranded DNA binding (SSB) proteins, designated SsbA and SsbB (Figure 1) [1–3]. The SsbA protein (156 amino acids/17,350 Da per monomer) is expressed constitutively whereas the SsbB protein (131 amino acids/14,926 Da per monomer) is induced specifically during transformational competence. These expression patterns suggest that the SsbA protein may serve as a general SSB protein for routine DNA functions, and that the SsbB protein may be a specialized SSB protein used primarily during natural transformation [1].

The N-terminal domains of the SsbA and SsbB proteins (amino acids 1–105/106) are similar in sequence. The C-terminal domain of the SsbB protein (amino acids 106–131), however, is significantly shorter than that of the SsbA protein (amino acids 107–156) (Figure 1). Structural studies of the corresponding regions of the homotetrameric SSB protein from *Escherichia coli* (SsbEc) and other bacterial SSB proteins have shown that the N-terminal domain contains the DNA binding and subunit tetramerization sites, whereas the C-terminal domain may serve as a binding site for other proteins involved in various DNA functions [4].

Bacterial SSB proteins bind single-stranded DNA in a non-sequence-specific manner. The DNA binding properties of the SsbEc protein have been the most extensively characterized. Two major binding modes have been identified: the SSB35 mode and the SSB65 mode. In the SSB35 mode (favored at lower salt concentrations), two subunits of the SsbEc tetramer interact with the single-stranded DNA (occluding ~35 nucleotides per tetramer), whereas in the SSB65 mode (favored at higher salt concentrations), all four subunits of the SsbEc tetramer interact with the single-stranded DNA (occluding ~65 nucleotides per tetramer) [5].

We previously carried out a comparative analysis of the DNA binding mode properties of the SsbEc, SsbA, and SsbB proteins. In that study, the various SSB proteins were incubated with the oligomer, dT, under different solution conditions and the resulting complexes were examined by polyacrylamide gel electrophoresis. In standard reaction solution (25 mM Tris acetate, pH 7.5), the SsbEc protein was able to bind a single dT oligomer containing ~35 nucleotides per tetramer. In the SSB35 mode (favored at lower salt concentrations), two subunits of the SsbEc tetramer interact with the single-stranded DNA (occluding ~35 nucleotides per tetramer), whereas in the SSB65 mode (favored at higher salt concentrations), all four subunits of the SsbEc tetramer interact with the single-stranded DNA (occluding ~65 nucleotides per tetramer) [5].
molecule, consistent with the SSB35 mode of binding. When Mg^{2+} (10 mM) was included in the reaction solution, however, the SsbEc protein was able to bind two dT35 molecules, consistent with the SSB65 mode of binding. The SsbA protein behaved similarly to the SsbEc protein under all reaction conditions, indicating that it interacted with dT35 in SSB35 and SSB65 modes that were analogous to those of the SsbEc protein. The SsbB protein, in contrast, appeared to bind two dT35 molecules in an SSB65-like mode in the absence of Mg^{2+}, and in an enhanced SSB65-like mode (with positive intersubunit cooperativity) in the presence of Mg^{2+} [2].

The pronounced difference in binding mode preferences raises the question of whether SsbA and SsbB would be able to interact together on single-stranded DNA. To address this issue, we have now examined the assembly of multiple SsbAs or SsbBs on dTn oligomers of various defined lengths. Polyacrylamide gel electrophoresis was particularly well suited for this analysis because the various SSB-dTn complexes were readily resolvable and remarkably stable during electrophoresis, and the effect of solution conditions on complex formation could be assessed by varying the composition of the electrophoresis running buffer. The results indicate that: i) different mechanisms of assembly are available to the SsbA and SsbB proteins, depending on the length of the DNA and the specific solution conditions, and ii) SsbA and SsbB may co-assemble on longer DNA segments where independent binding is possible, but not on shorter DNA segments where coordinated interactions between adjacent SsbBs are required.

Results

Experimental Design

The binding of the Streptococcus pneumoniae SsbA and SsbB proteins to a set of dTn oligomers ranging in length from dT50 to dT130 was examined. Particular attention was placed on determining the shortest dTn oligomer that was able to accommodate the binding of two SsbAs, two SsbBs, or one SsbA and one SsbB, in either the absence or presence of Mg^{2+}. The expectation with this approach was that two SsbBs would have to interact in a coordinated manner to form a complex on a minimal length dTn whereas the SsbBs would be able to bind independently to isolated sites on longer dTn oligomers. All binding reactions were carried out in solutions containing 25 mM Tris acetate (pH 7.5) and either 0 or 10 mM magnesium acetate, and the resulting complexes were analyzed by polyacrylamide gel electrophoresis using a running buffer identical in composition to that of the individual reaction solutions.

SsbA protein assembly

The complexes that were formed by the SsbA protein with the various dTn oligomers in the absence and presence of Mg^{2+} are shown in Figures 2 and 3, respectively (note: in these experiments, the electrophoretic mobilities of the free dTn oligomers exhibit a greater inverse-dependence on length than do the corresponding SsbA-dTn complexes, leading to a progressive decrease in the separation between the free dTn oligomers and the SsbA-dTn complexes with increasing dTn length).

Absence of Mg^{2+}

When increasing concentrations of SsbA were added to dT50 in the absence of Mg^{2+}, a single complex with a gel mobility lower than that of the unbound dT50 was formed (A1 complex). All of the dT50 was converted to this complex at an SsbA concentration that corresponded to approximately one SsbA tetramer per dT50 molecule, and there was no indication of the formation of a second complex at higher SsbA concentrations (Figure 2 and additional data not shown). A similar pattern of binding was obtained with the longer oligomer, dT65 (gel not shown). These results indicated that a single SsbA was able to bind to dT50 and dT65 under these reaction conditions.

When the oligomer length was increased to dT75, an A1 complex was formed with increasing SsbA concentrations in a manner similar to that observed with the shorter oligomers. As the concentration of SsbA was increased further, however, the A1 complex disappeared and a new complex with a lower gel mobility was formed (A2 complex). This result indicated that a second SsbA was able to bind to dT75 under these conditions. A similar pattern of binding was observed with the longer oligomers, dT85, dT95, and dT100, indicating that two SsbAs were able to bind to each of these oligomers as well (Figure 2).

When the oligomer length was increased to dT130, an A1 complex was formed with increasing SsbA concentrations in a manner similar to that observed with the shorter oligomers. As the concentration of SsbA was increased further, however, the A1 complex disappeared and a new complex with a lower gel mobility was formed (A2 complex). This result indicated that a second SsbA was able to bind to dT130 under these conditions (Figure 2).

The results in Figure 2 indicated that the shortest dTn oligomer in the set that was able to bind two SsbAs in the absence of Mg^{2+} was dT75, and the shortest dTn that was able to bind three SsbAs was dT130. These results are summarized in Table 1.

Presence of Mg^{2+}

When increasing concentrations of SsbA were added to dT50 in the presence of Mg^{2+}, a single complex with a gel mobility lower than that of the unbound dT50 was formed (A1 complex), with no indication of the formation of a second complex at higher SsbA concentrations (Figure 3). A similar pattern of binding was obtained with dT65 (gel not shown). These results were similar to those that were obtained in the absence of Mg^{2+}.

Figure 1. Streptococcus pneumoniae SsbA and SsbB proteins. The amino acid sequences of the S. pneumoniae SsbA and SsbB proteins are aligned with that of the E. coli SSB protein, SsbEc. Identical residues are highlighted in black. The division between the N-terminal and C-terminal domains is indicated by an arrow, and the putative Loop 1 region is denoted with a box.

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S. pneumoniae, SsbA, and SsbB proteins are...
and indicated that a single SsbA was able to bind to dT_{50} and dT_{95} in the presence of Mg^{2+}. A_2 complexes were also formed when increasing concentrations of SsbA were added to the longer oligomers, dT_{75} and dT_{105}, but in contrast to the results that were obtained in the absence of Mg^{2+}, A_2 complexes were not detected with these oligomers (Figure 3).

When the oligomer length was increased to dT_{90}, an A_1 complex was formed at lower SsbA concentrations in a manner

Figure 2. Binding of SsbA protein to dT_n oligomers in the absence of Mg^{2+}. The reaction solutions contained 25 mM Tris acetate (pH 7.5), 5 mM dT_n (nucleotide concentration), and the indicated concentrations of SsbA protein (tetramer concentrations). The reactions were analyzed by polyacrylamide gel electrophoresis using a gel running buffer consisting of Tris acetate (pH 7.5). The bands corresponding to the unbound dT_n oligomers, and the A_1, A_2, and A_3 complexes, were visualized by phosphorimaging.

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similar to that observed with the shorter oligomers. As the concentration of SsbA was increased further, however, the A1 complex disappeared and a new complex with an even lower gel mobility was formed (A2 complex). This result indicated that a second SsbA was able to bind to dT90 under these conditions. A similar pattern of binding was obtained with dT100 and dT130, indicating that two SsbAs were able to bind to each of these oligomers as well. In contrast to the results that were obtained in the absence of Mg\(^{2+}\), however, there was no indication of the formation of a third complex with dT_{130}, even at the highest concentrations of SsbA that were examined (Figure 3).

The results in Figure 3 indicated that the shortest dT_n oligomer in the set that was able to bind two SsbAs in the presence of Mg\(^{2+}\) was dT_{90}, and that only two SsbAs were able to bind even when the oligomer length was increased to dT_{130}. These results are summarized in Table 1.

**SsbB protein assembly**

The complexes that were formed by the SsbB protein with the various dT_n oligomers in the absence and presence of Mg\(^{2+}\) are shown in Figures 4 and 5, respectively. In these experiments, the separation between the free dT_n oligomers and the various
SsbB-dTₙ complexes is less than that observed with the SshA protein, owing to the smaller molecular size of the SshB protein and the increased electrophoretic mobility of the SshB-dTₙ complexes.

**Absence of Mg²⁺.** When increasing concentrations of SshB were added to dT₅₀ in the absence of Mg²⁺, a single complex with a gel mobility lower than that of the unbound dT₅₀ was formed (B₁ complex). All of the dT₅₀ was converted to this complex at an SshB concentration that corresponded to approximately one SshB tetramer per dT₅₀ molecule, and there was no indication of the formation of a second complex at higher SshB concentrations (Figure 4 and additional data not shown). A similar pattern of binding was obtained with the longer oligomers, dT₆₅, (gel not shown) and dT₇₅ (Figure 4). These results indicated that only a single SshB was able to bind to these oligomers under these conditions. When the oligomer length was increased to dT₉₀, a B₁ complex was formed at lower SshB concentrations in a manner similar to that observed with the shorter oligomers. When the concentration of SshB was increased further, however, the B₁ complex disappeared and a new complex of even lower gel mobility was formed (B₂ complex). This result indicated that a second SshB was able to bind to dT₁₀₀ under these conditions (Figure 5). A similar pattern of binding was obtained with dT₁₃₀, indicating that two SshBs were able to bind to this oligomer as well. There was no indication of the formation of a third complex with these oligomers, however, even at the highest concentrations of SshB that were examined (Figure 5).

The results in Figure 5 indicated that the shortest dTₙ oligomer in the set that was able to bind two SshBs in the absence of Mg²⁺ was dT₁₀₀, and that only two SshBs were able to bind even when the oligomer length was increased to dT₁₃₀. These results are summarized in Table 1.

| Complexes formed by the SshA and SshB proteins on dTₙ oligomers. |
|------------------------|--------|--------|--------|--------|
| dTₙ       | SshA   | SshB   |
| dT₅₀      | A₁     | B₁     |
| dT₆₅      | A₂     | B₂     |
| dT₇₅      | A₁     | B₂     |
| dT₉₀      | A₂     | B₂     |
| dT₁₀₀     | A₁     | B₂     |
| dT₁₃₀     | A₁     | B₄     |

These results were derived from the experiments shown in Figures 2–5 and additional experiments (gels not shown), and indicate the highest order complexes that were observed when the indicated dTₙ oligomers were mixed with an excess concentration of SshA or SshB protein, in the absence or presence of 10 mM Mg²⁺ (the notations Aₙ and Bₙ indicate the number of SshAs or SshBs bound). doi:10.1371/journal.pone.0024305.t001

**Co-assembly of SshA and SshB proteins.**

The ability of the SshA and SshB proteins to co-assemble on dTₙ oligomers was also investigated. For these experiments, dTₙ oligomers were selected that were long enough to accommodate the binding of either two SshAs or two SshBs, under various solution conditions (see Table 1).

**Absence of Mg²⁺.** The first set of co-assembly experiments in the absence of Mg²⁺ was carried out with dT₉₀ (Figure 6). When an excess concentration of SshA alone was added to dT₉₀, an A₂ complex with two SshAs bound to the dT₉₀ was formed. Similarly, when an excess concentration of SshB alone was added to dT₉₀, a B₂ complex with two SshBs bound to the dT₉₀ was formed. When SshA and SshB were added together to dT₉₀, however, the A₂ and B₂ complexes were again formed, but little if any mixed complexes with one SshA and one SshB to the same dT₉₀ molecule were detected (as judged by the absence of a new band with a mobility intermediate between that of the A₂ and B₂ complexes). These results indicated that although either two SshAs or two SshBs could bind to dT₉₀ in the absence of Mg²⁺, the binding of one SshA and one SshB to dT₉₀ was unfavorable under these conditions.

A second set of co-assembly experiments was carried out in the absence of Mg²⁺ with the longer oligomer, dT₁₀₀ (Figure 6). When an excess concentration of either SshA or SshB alone was added to dT₁₀₀, the corresponding A₂ or B₂ complexes were formed, as with dT₉₀. In contrast to the results that were obtained with dT₉₀, however, the A₂ and B₂ complexes, and a third complex with an intermediate mobility were formed when SshA and SshB were added together to dT₁₀₀. The intermediate band was excised from the gel, analyzed by SDS-polyacrylamide gel electrophoresis, and found to contain approximately equal amounts of SshA and SshB protein (gel not shown). These results indicated that the intermediate band corresponded to a mixed complex in which one SshA and one SshB were bound to the dT₁₀₀ (A-B complex).

The results in Figure 6 indicated that although the simultaneous binding of SshA and SshB to dT₉₀ was unfavorable in the absence of Mg²⁺, SshA and SshB were able to bind together on dT₁₀₀ under these conditions.

**Presence of Mg²⁺.** The first set of co-assembly experiments in the presence of Mg²⁺ (10 mM) was carried out with dT₁₀₀...
When an excess concentration of either SsbA or SsbB alone was added to dT\textsubscript{100}, the corresponding A\textsubscript{2} and B\textsubscript{2} complexes were formed as expected. When SsbA and SsbB were added together to dT\textsubscript{100}, however, the A\textsubscript{2} and B\textsubscript{2} complexes were again formed, but no mixed complexes with one SsbA and one SsbB bound to the same dT\textsubscript{100} molecule were detected (as judged by the absence of a new band with a mobility intermediate between that of the A\textsubscript{2} and B\textsubscript{2} complexes). These results indicated that although two SsbAs or two SsbBs could bind to dT\textsubscript{100} in the presence of Mg\textsuperscript{2+}, the binding of one SsbA and one SsbB to dT\textsubscript{100} was unfavorable under these conditions.

A second set of co-assembly experiments was carried out in the presence of Mg\textsuperscript{2+} with the longer oligomer, dT\textsubscript{130} (Figure 7). When an excess concentration of either SsbA or SsbB alone was added to dT\textsubscript{130}, the corresponding A\textsubscript{2} or B\textsubscript{2} complexes were formed, as with dT\textsubscript{100}. In contrast to the results that were obtained...
with dT_{100}, however, the A₂ and B₂ complexes, and a third complex with an intermediate mobility were formed when SsbA and SsbB were added together to dT_{100}. The appearance of the intermediate band was consistent with the formation of a mixed complex in which one SsbA and one SsbB were bound to the dT_{130} (A-B complex).

The results in Figure 7 indicated that although the simultaneous binding of SsbA and SsbB to dTₙₙ oligomers was unfavorable in the presence of Mg^{2+}, SsbA and SsbB were able to bind together on dT_{130} under these conditions.

**Figure 5. Binding of SsbB protein to dTₙₙ oligomers in the presence of Mg^{2+}**. The reaction solutions contained 25 mM Tris acetate (pH 7.5), 10 mM magnesium acetate, 5 μM dTₙₙ (nucleotide concentration), and the indicated concentrations of SsbB protein (tetramer concentrations). The reactions were analyzed by polyacrylamide gel electrophoresis using a gel running buffer consisting of Tris acetate (pH 7.5) and 10 mM magnesium acetate. The bands corresponding to the unbound dTₙₙ oligomers, and the B₁ and B₂ complexes, were visualized by phosphorimaging. doi:10.1371/journal.pone.0024305.g005

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**Co-assembly of SsbA protein with SsbA/B and SsbB^{RYTP} proteins**

Additional co-assembly experiments were carried out with the SsbA protein and two SSB variants: the SsbA/B protein and the SsbB^{RYTP} protein. The SsbA/B protein is a chimeric protein in which the C-terminal domain of the SsbA protein (amino acids 106–156) has been replaced with the C-terminal domain from the SsbB protein (amino acids 105–131) [2]. The SsbB^{RYTP} protein is a modified SsbB protein in which a four-amino acid sequence from the N-terminal domain of the SsbB protein (18HKTN21) has...
been replaced with the corresponding sequence from the SsbA protein (18RYTP21) (Figure 1, Material and Methods). The SsbA/B protein (15,039 Da per monomer) and the SsbBRYTP protein (14,963 Da per monomer) are similar in size to the SsbB protein (14,926 Da per monomer), and form complexes on dT90 that are clearly resolvable by gel electrophoresis from the complexes formed by the SsbA protein (17,350 Da per monomer). Experiments analogous to those carried out for the SsbA and SsbB proteins indicated that either two SsbA/Bs ((A/B)2 complex) or two SsbBRYTPs (BRYTP2 complex) were able to bind to dT90 in the absence of Mg2+. SsbA/B protein. The initial set of SsbA and SsbA/B co-assembly experiments was carried out with dT90 in the absence of Mg2+ (Figure 6). When an excess concentration of either SsbA or SsbA/B alone was added to dT90, the corresponding A2 and (A/B)2 complexes were formed, as expected. When SsbA and SsbA/B were added together to dT90, however, the A2 and (A/B)2 complexes, and a third complex with an intermediate mobility, were formed. The appearance of the intermediate band was consistent with the formation of a mixed complex in which one SsbA and one SsbB were bound to dT90 (A·B complex). These results indicated that although the simultaneous binding of SsbA and SsbB was unfavorable (Figure 6), SsbA was able to form a mixed complex with SsbA/B on dT90 in the absence of Mg2+ (Figure 6). Additional experiments indicated that SsbA/B also differed from SsbB in that it was able to form a mixed complex with SsbA on dT100 in the presence of Mg2+ (gel not shown).

**SsbBRYTP protein.** The initial set of SsbA and SsbBRYTP co-assembly experiments was also carried out with dT90 in the absence of Mg2+ (Figure 9). When an excess concentration of either SsbA or SsbBRYTP alone was added to dT90, the corresponding A2 and BRYTP2 complexes were formed, as expected. When SsbA and SsbBRYTP were added together to dT90, however, the A2 and BRYTP2 complexes, and a third complex with an intermediate mobility were formed. The appearance of the intermediate band was consistent with the formation of a mixed complex in which one SsbA and one SsbBRYTP were bound to dT90 (A·BRYTP complex).

These results indicated that although the simultaneous binding of SsbA and SsbB was unfavorable (Figure 6), SsbA was able to form a mixed complex with SsbA/B on dT90 in the absence of Mg2+ (Figure 6). Additional experiments indicated that SsbA/B also differed from SsbB in that it was able to form a mixed complex with SsbA on dT100 in the presence of Mg2+ (gel not shown).

**Discussion**

The results presented here indicate that the shortest dT oligomer that is able to accommodate the binding of two SsbAs or two SsbBs is strictly defined, and depends on whether Mg2+ is...
included in the reaction solution (Table 1). This finding suggests that the minimal oligomer length may be determined by the preferred binding modes and potential binding capacities of the individual SSB proteins under the particular solution conditions.

**SsbA protein assembly**

The shortest dT<sub>n</sub> oligomer that was able to bind two SsbAs in the absence of Mg<sup>2+</sup> was dT<sub>75</sub> (Table 1). Assuming that SsbA interacts with dT<sub>n</sub> oligomers as a tetramer in an SSB<sub>35</sub>-like mode in the absence of Mg<sup>2+</sup> (see Introduction), two SsbAs may assemble on dT<sub>75</sub> under these conditions in a manner in which each of the SsbAs interacts with a 35-nucleotide segment of the oligomer. The observation that only two SsbAs were able to bind even when the oligomer length was increased to dT<sub>100</sub>, whereas three SsbAs were able to bind to dT<sub>130</sub> indicates that at least a 35 nucleotide segment of dT<sub>n</sub> was required for the stable binding of each SsbA (Table 1). In all cases, as the concentration of SsbA was increased, the dT<sub>n</sub> complex with a lesser number of SsbAs bound was replaced completely by the complex with the greater number of SsbAs bound. These results indicate that under SSB<sub>35</sub>-like binding mode conditions, individual SsbAs can organize themselves so as to maximize the number of SsbAs bound to a dT<sub>n</sub> oligomer while satisfying the 35-nucleotide binding requirement of each bound SsbA.

A longer dT<sub>n</sub> oligomer was required for the binding of two SsbAs when Mg<sup>2+</sup> was included in the reaction solution (Table 1). Assuming that SsbA interacts with dT<sub>n</sub> oligomers as a tetramer in an SSB<sub>65</sub>-like binding mode in the presence of Mg<sup>2+</sup> (see Introduction), it is likely that the increased length requirement is due to the higher binding capacity of SsbA under these conditions. The complexes containing two SsbAs that were observed with dT<sub>130</sub> are consistent with an SSB<sub>65</sub>-like mode of binding in that dT<sub>130</sub> is long enough to satisfy the full capacity of 65 nucleotides expected for each of the bound SsbAs (~130 nucleotides total) (Figure 10). However, stable complexes containing two SsbAs could also be formed on oligomers as short as dT<sub>90</sub> under these conditions (Table 1). With these shorter oligomers, one or both of the SsbAs were presumably bound using only a portion of their potential binding capacity. These results suggest that under SSB<sub>65</sub>-like binding mode conditions, two SsbAs are able to assemble onto shorter segments of single-stranded DNA via a coordinated sharing of the DNA strand between the bound proteins.

Various arrangements can be envisioned for the DNA strand in a "shared-strand binding" mechanism (Figure 10). With dT<sub>90</sub> for example, one SsbA could be bound to a ~65-nucleotide segment, with the second SsbA bound to the remaining ~25-nucleotide segment of the oligomer. Alternatively, the dT<sub>90</sub> might be equally shared between the two SsbAs, with each binding to a ~45-nucleotide segment of the oligomer. These possibilities are not necessarily mutually exclusive and a combination of different binding arrangements may also occur. In any case, the observation that complexes with two SsbAs were not formed on dT<sub>n</sub> oligomers shorter than dT<sub>90</sub> when Mg<sup>2+</sup> was included in the reaction solution, but could be formed on oligomers as short as dT<sub>75</sub> in

![Figure 7. Binding of SsbA and SsbB proteins to dT<sub>100</sub> and dT<sub>130</sub> in the presence of Mg<sup>2+</sup>.](https://doi.org/10.1371/journal.pone.0024305.g007)
absence of Mg$^{2+}$ (Table 1) suggests that the shared-strand arrangement that is adopted when two SsbAs bind in the SSB65-like mode may be different from the binding arrangement that is used when two SsbAs bind in the SSB35-like mode.

SsbB protein assembly

Longer dT$_n$ oligomers were required for the binding of two SsbBs than were needed for two SsbAs, both in the absence and presence of Mg$^{2+}$ (Table 1). Assuming that SsbB interacts with dT$_90$ oligomers as a tetramer in an SSB65-like mode in the absence of Mg$^{2+}$ and in an enhanced SSB65-like mode in the presence of Mg$^{2+}$ (see Introduction), the longer length requirement may reflect the higher binding capacity of SsbB, relative to that of SsbA, under these reaction conditions. The complexes containing two SsbBs that were observed with dT$_{130}$ in both the absence and presence of Mg$^{2+}$ are consistent with either the SSB65-like mode or the enhanced SSB65-like mode of binding in that dT$_{130}$ is long enough to satisfy the full ~65 nucleotide binding capacity expected for each of the bound SsbBs in either case (~130 nucleotides total) (Figure 10). However, stable complexes containing two SsbBs could also be formed on oligomers as short as dT$_{85}$ (in the absence of Mg$^{2+}$) or dT$_{100}$ (in the presence of Mg$^{2+}$). These results suggest that two SsbBs can bind to the shorter oligomers using a shared-strand binding mechanism analogous to that proposed for SsbA in...
the presence of Mg$^{2+}$ (SSB$_{35}$-like binding mode conditions) (Figure 10). In the case of SsbB, the observation that dT$_{100}$ was able to accommodate the binding of two SsbBs in absence of Mg$^{2+}$, whereas at least dT$_{100}$ was required for the binding of two SsbBs in the presence of Mg$^{2+}$, may reflect a difference in the arrangement of the shared strand between the two SsbBs under normal SSB$_{65}$-like mode versus enhanced SSB$_{65}$-like mode binding conditions.

SsbA and SsbB protein co-assembly

The shortest dT$_n$ oligomer that was able to accommodate the simultaneous binding of one SsbA and one SsbB was also strictly defined and depended on the solution conditions. In the absence of Mg$^{2+}$, complexes containing either two SsbAs or two SsbBs were readily formed on dT$_{100}$. However, little or no mixed complexes with one SsbA and one SsbB were detected when both proteins were added together to dT$_{90}$ (Figure 6). If the binding capacity of SsbA under these conditions is assumed to be $\sim 35$ nucleotides (SSB$_{35}$-like mode) and the binding capacity of SsbB is assumed to be $\sim 65$ nucleotides (SSB$_{65}$-like mode), a dT$_{100}$ molecule would not be long enough to satisfy the full binding capacities of one SsbA and one SsbB ($\sim 100$ nucleotides total). Therefore, the simultaneous binding of SsbA and SsbB to dT$_{100}$ would presumably require one or both of these proteins to bind using only a portion of their potential binding capacity. The absence of mixed complex formation on dT$_{90}$ thus suggests that SsbA and SsbB are not able to engage in shared-strand binding in the absence of Mg$^{2+}$.

The apparent incompatibility in shared-strand binding does not appear to preclude SsbA and SsbB from binding independently on longer dT$_n$ oligomers where strand sharing would not be required. Although they were unable to co-assemble on dT$_{90}$, SsbA and SsbB were able to form a mixed complex on dT$_{100}$ in the absence of Mg$^{2+}$ (Figure 6). In this case, a dT$_{100}$ molecule could potentially provide a $\sim 35$-nucleotide segment for the SsbA and a $\sim 65$-nucleotide segment for the SsbB, and thereby satisfy the full binding capacities of both proteins under the reaction conditions.

DNA binding compatibility also appears to govern the co-assembly of SsbA and SsbB on dT$_n$ oligomers when Mg$^{2+}$ is included in the reaction solution. Although complexes with two SsbAs or two SsbBs were readily formed on either dT$_{100}$ or dT$_{130}$ in the presence of Mg$^{2+}$, mixed complexes with one SsbA and one SsbB were detected only with dT$_{130}$ (Figure 7). If the binding capacity of SsbA under these conditions is assumed to be $\sim 65$ nucleotides (SSB$_{65}$-like mode) and the binding capacity of SsbB is also assumed to be $\sim 65$ nucleotides (enhanced SSB$_{65}$-like mode), a dT$_{130}$ molecule would be able to satisfy the full binding capacities of one SsbA and one SsbB ($\sim 130$ nucleotides total), whereas a dT$_{100}$ molecule would only be able to partially satisfy the binding capacities of the two proteins. Thus, the formation of mixed complexes on dT$_{130}$ but not on dT$_{100}$, indicates that SsbA and SsbB are able to bind independently, but are not able to engage in shared-strand binding, in the presence of Mg$^{2+}$.

SsbA/B and SsbBRytp proteins

Although SsbA and SsbB appeared to be incompatible for shared-strand binding, SsbA was able to form mixed complexes under some shared-strand binding conditions with two SSB protein variants: the SsbA/B protein and the SsbBRytp protein.

The SsbA/B protein, in which the C-terminal domain of the SsbA protein has been replaced with the C-terminal domain from the SsbB protein, was prepared previously to assess the contribution of the C-terminal domains to the DNA binding properties of the SsbA and SsbB proteins [2]. The DNA binding mode preferences of the SsbA/B protein were found to be similar to those of the SsbA protein, suggesting that the primary structural determinants of DNA binding may be contained within the N-terminal domains of the various SSB proteins [2]. Moreover, the SsbA/B protein was able to form a mixed complex with SsbA on dT$_{90}$ in the absence of Mg$^{2+}$ and on dT$_{100}$ in the presence of Mg$^{2+}$ (Figure 8, and gel not shown). These findings suggest that the inability of SsbA to engage in shared-strand binding with SsbB may not be due to the dissimilar C-terminal domain of the SsbB protein, inasmuch as the C-terminal domain of the SsbA/B protein is identical to that of the SsbB protein. It is possible, however, that the C-terminal domain functions differently in the SsbB protein than in the chimeric SsbA/B protein, and contributes to the incompatibility of SsbA and SsbB in shared-strand binding.

The SsbBRytp protein, in which the $^{15}$HKTN$^{21}$ sequence of the SsbB protein has been replaced with the corresponding $^{15}$RYTP$^{21}$
sequence from the SsbA protein, was also prepared in an effort to determine the structural basis for the differential DNA binding properties of the SsbA and SsbB proteins. The DNA binding properties of the SsbB<sup>R</sup><sup>RYTP</sup> protein were found to be similar to those of the SsbB protein, indicating that the<sup>18</sup>R<sup>RYTP</sup><sup>21</sup> sequence may not be responsible for the distinctive DNA binding mode preferences of the SsbB protein (see Experimental Procedures). The SsbB<sup>R</sup><sup>RYTP</sup> protein differed from the SsbB protein, however, in that it was able to form a mixed complex with SsbA on dT<sub>100</sub> in the absence of Mg<sup>2+</sup> (Figure 9). These results suggest that the shared-strand binding incompatibility that was observed with the SsbA and SsbB proteins in absence of Mg<sup>2+</sup> was not due to the difference in their preferred DNA binding modes, but may be attributable to the divergent<sup>18</sup>R<sup>RYTP</sup><sup>21</sup> and<sup>18</sup>H<sup>HKTN</sup><sup>21</sup> sequences of these proteins. The observation that SsbB<sup>R</sup><sup>RYTP</sup> was unable to form a mixed complex with SsbA on dT<sub>100</sub> in the presence of Mg<sup>2+</sup>, however, indicates that this sequence difference is not sufficient to account for the shared-strand binding incompatibility that was observed with SsbA and SsbB in the presence of Mg<sup>2+</sup>. Thus, the introduction of the<sup>18</sup>R<sup>RYTP</sup><sup>21</sup> sequence into the SsbB protein has the effect of uncoupling the Mg<sup>2+</sup>-independent shared-strand binding incompatibility from the Mg<sup>2+</sup>-dependent incompatibility.

The<sup>18</sup>R<sup>RYTP</sup><sup>21</sup> sequence of the SsbA protein is at least partially conserved in the SsbEc protein and in a number of other homotetrameric bacterial SSB proteins whose x-ray crystal structures have been determined (Figure 11). An inspection of the various structures shows that the tertiary folds of the N-terminal domains are similar in all cases, and that the<sup>18</sup>R<sup>RYTP</sup><sup>21</sup> sequence of the SsbA protein, and the divergent<sup>18</sup>H<sup>HKTN</sup><sup>21</sup> sequence of the SsbB protein, correspond to the Loop 1 region on the outer surface of the subunits of the SSB tetramers (Figure 11) [6–7]. The differences in the composition of the Loop 1 region may account for the inability of the SsbA and SsbB proteins to engage in shared-strand binding in the absence of Mg<sup>2+</sup>. For example, the Loop 1 variations could conceivably affect the precise orientation of the DNA strand as it winds around the individual SSB tetramers, or influence the manner in which two SSB tetramers interact when bound in close proximity on a DNA strand. Other molecular determinants are apparently required, however, for shared-strand binding compatibility in the presence of Mg<sup>2+</sup>. A definitive determination of the molecular basis for shared-strand binding and DNA binding compatibility will require further structural analysis of the various SSB-dT<sub>n</sub> complexes. These studies are currently underway in our laboratory.

**SsbA and SsbB binding compatibility**

SsbA is a constitutively-expressed protein, and presumably functions as the primary SSB protein during the routine replication and maintenance of chromosomal DNA in <i>S. pneumoniae</i> (analogous to the SsbEc protein in <i>E. coli</i>). SsbB, in contrast, is induced specifically during natural transformation, and associates transiently with a single-stranded form of the exogenous DNA before the DNA is incorporated into a homologous region of the <i>S. pneumoniae</i> chromosome (there is no analog of the SsbB protein in <i>E. coli</i>). [1,8]

The extent to which the SsbA and SsbB proteins are functionally interchangeable in these various activities is not clear. Our results, however, indicate that the SsbA and SsbB proteins will be able to bind together on longer single-stranded DNA segments where independent binding is possible, but suggest that they may not co-assemble on shorter single-stranded DNA segments where coordinated interactions between adjacent SSBs are required. The compatibility requirement for shared-strand binding could conceivably serve as a self-recognition mechanism that regulates the manner in which SsbA and SsbB interact in <i>S. pneumoniae</i>.

**Materials and Methods**

Materials

<i>S. pneumoniae</i> SsbA protein [9], SsbB protein [10], and SsbA/B protein [2] were prepared as previously described. Gel-purified dT<sub>n</sub> oligomers were from Invitrogen. <sup>32</sup>P-end-labeled dT<sub>n</sub> oligomers were prepared using γ-[<sup>32</sup>P]ATP (PerkinElmer) and T<sub>4</sub> polynucleotide kinase (New England Biolabs).
Preparation and characterization of the SsbB<sup>RYTP</sup> protein

The SsbB<sup>RYTP</sup> protein coding sequence, in which the nucleotide sequence of the SsbB protein corresponding to amino acids 189HKTN<sup>21</sup> was replaced with a sequence coding for the amino acids 189RYTP<sup>21</sup>, was generated by overlap-extension PCR mutagenesis. The initial mutagenesis template was our previously described pETssbB construct, which contains the wild type SsbB sequence cloned into the NdeI/HindIII site of the pET21a expression vector (Novagen) [10]. Primer a (5′-GGATACGCCG-GATCTCAGTGG-3′) flanked the ssbB gene, and primer b (5′-CTTGCATTTGGAGTGTACGCAATTCGCTGTA-GAC-3′) and primer c (5′-GAAATTCGTTAGCATCCAAATGACAAGTCCGTAGC-3′) were the internal overlapping mutagenic primers (mutagenic bases are underlined). The final SsbB<sup>RYTP</sup>-coding PCR product was digested with NdeI and HindIII and then cloned into the NdeI/HindIII site of a pET21a expression vector to yield the construct pETssbB<sup>RYTP</sup>. The insert was sequenced and found to be identical to the expected SsbB<sup>RYTP</sup> protein coding sequence.

The pETssbB<sup>RYTP</sup> expression plasmid was introduced into E. coli strain Rosetta(DE3)pLysS (Novagen), and the SsbB<sup>RYTP</sup> protein was purified from the resulting Rosetta(DE3)pLysS/pETssbB<sup>RYTP</sup> cells using a procedure analogous to that described previously for the wild type SsbB protein [10]. The final fraction of SsbB<sup>RYTP</sup> protein was greater than 95% pure as judged by SDS-polyacrylamide gel electrophoresis.

The purified SsbB<sup>RYTP</sup> protein was characterized using the dT<sub>35</sub> binding assay that was described previously for the SsbB protein [10]. The final fraction of dT<sub>35</sub> binding reaction solutions (30 μl) contained 25 mM Tris acetate (pH 7.5), 5% glycerol, 1 mM dithiothreitol, and the concentrations of magnesium acetate, dT<sub>35</sub> (32P-end-labeled), and SSB protein given in the figure legends. The reactions solutions were incubated at 25°C for 15 min, and then 3 μl of gel loading solution (0.25% bromophenol blue, 40% sucrose) was added. An aliquot (20 μl) of the final solution was analyzed by electrophoresis on 5% native polyacrylamide gels using a buffer system consisting of 25 mM Tris acetate (pH 7.5) and the same concentration of magnesium acetate as in the reaction solutions. The bands corresponding to unbound and SSB-bound dT<sub>35</sub> oligomers were visualized using a Fuji FLA-7000 imager. The specific protein concentrations for the individual gels in Figures 2–9 were selected to illustrate the concentration-dependent formation of the various SSB-ddT<sub>n</sub> complexes.

Author Contributions

Conceived and designed the experiments: FRB BS. Performed the experiments: BS GA. Analyzed the data: FRB BS. Contributed reagents/materials/analysis tools: FRB BS GA. Wrote the paper: FRB BS.

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