**Structure-based design of peptides that trigger**
**Streptococcus pneumoniae** cell death

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Toxin–antitoxin (TA) systems regulate key cellular functions in bacteria. Here, we report a unique structure of the *Streptococcus pneumoniae* HigBA system and a novel antimicrobial agent that activates HigB toxin, which results in mRNA degradation as an antibacterial strategy. In this study, protein structure-based peptides were designed and successfully penetrated the *S. pneumoniae* cell membrane and exerted bactericidal activity. This result represents the time during which inhibitors triggered *S. pneumoniae* cell death via the TA system. This discovery is a remarkable milestone in the treatment of antibiotic-resistant *S. pneumoniae*, and the mechanism of bactericidal activity is completely different from those of current antibiotics. Furthermore, we found that the HigBA complex shows a crossed-scissor interface with two intermolecular β-sheets at both the N and C termini of the HigA antitoxin. Our biochemical and structural studies provided valuable information regarding the transcriptional regulation mechanisms associated with the structural variability of HigAs. Our in vivo study also revealed the potential catalytic residues of HigB and their functional relationships. An inhibition study with peptides additionally proved that peptide binding may allosterically inhibit HigB activity. Overall, our results provide insights into the molecular basis of HigBA TA systems in *S. pneumoniae*, which can be applied for the development of new antibacterial strategies.

**Databases**
Structural data are available in the PDB database under the accession number 6AF4.

**Introduction**

*Streptococcus pneumoniae* is a Gram-positive species of bacteria that causes disease mainly through respiratory infections [1]. Diseases caused by pneumococcal species include otitis, sinusitis meningitis, bacteremia, and pneumonia [2]. The infections caused by *S. pneumoniae* resulted in 515 000 death in 2015 [3], with 335 000 of which in children under 5 years of age [4]. Patients infected with *S. pneumoniae* are mainly treated with antibiotics, such as penicillin, cephalosporins, macrolides, and quinolones [5]. However, the advent of...
increasing antibiotic resistance strains of *S. pneumoniae* [6] requires the development of novel therapeutics.

Toxin–antitoxin (TA) systems were first discovered as plasmid genes in the 1980s, which is related to the inheritance of plasmids in daughter cells [7]. TA systems consist of two or more adjacent genes encoding a toxin protein and an antitoxin. Toxins involved in diverse cellular functions, such as protein synthesis inhibition, DNA replication, and cell wall synthesis in response to unfavorable growth conditions. Usually, toxins as ribonucleases degrade mRNA in a specific or non-specific fashion, and some toxins play roles as gyrase inhibitors and kinases. Their cognate antitoxins neutralize the toxins and regulate the transcription levels of TA operons [8]. TA systems are comprised of different kinds of modules and their roles in bacterial cells have been intensively studied [9]. TA systems consist of a noncoding RNA antitoxin and a toxin-encoding mRNA. Type II TA systems consist of a protein antitoxin and its cognate protein toxin. Type III TA systems consist of a small RNA antitoxin that binds directly to the toxin protein as an inhibitor. Similar to the type II TA system, both the toxin and the antitoxin of a type IV TA system are proteins, although their inhibition mechanism is not direct binding. Instead, these proteins play opposing roles in bacterial filamentation. In a type V TA system, a ribonuclease antitoxin degrades the mRNA toxin. The protein toxin of a type VI TA system is neutralized by the protein antitoxin, which is degraded by cellular proteases [10].

The most widely distributed type II TA systems comprise the thermodynamically stable toxin and the unstable antitoxin. The antitoxin is cleaved by some cellular proteases and has a lower half-life than that of the toxin. Therefore, the host depends on a continuous supply of antitoxin to neutralize the toxicities of toxins [11]. Currently, seven type II TA loci have been reported in *S. pneumoniae*: three RelBEs, phd-doc, HicBA, HigBA, and one unidentified family. Among these seven TA systems, HigBA systems share limited sequence homology among HigB toxins and HigA antitoxins exhibit considerable structural variability [12–14].

Here, we present the complex structure of HigBA from *S. pneumoniae* as determined by X-ray crystallography. The crossed-scissor constitutes a new type of interface between HigB and HigA. The binding interface showed critical residues in the formation of the HigBA complex. HigB from *S. pneumoniae* showed in vitro ribonuclease activity, and the mRNA cleavage mechanism was scrutinized through numerous single point mutations. The peptides that mimicked toxin or antitoxin helices could hinder HigBA complex formation. These peptides could penetrate *S. pneumoniae* and exert bactericidal activity. The antitoxin does not block the active site of HigB in the HigBA complex structure, thus revealing that peptide binding may allosterically inhibit HigB activity. HigA has DNA-binding properties through its helix-turn-helix motif (HTH) and regulates HigBA expression. The residues of HigA that are important for DNA binding were investigated by nuclear magnetic resonance (NMR). This approach may contribute to our understanding of the structure–function relationships of TA systems, including the HigBA system. Our biochemical and structural studies will provide valuable information and insights into the molecular basis of TA systems as an antibacterial target.

### Results

#### Overall structure of the HigBA complex

The asymmetric unit of the HigBA complex structure is a heterotetramer containing two HigB toxins and two HigA antitoxins (Protein Data Bank (PDB) code 6AF4) (Fig. 1A and Fig. S1A–D). A secondary structural analysis was conducted using the 2Struc server [18]. The oligomeric state of HigBA was determined to be a heterotetramer based on a molecular weight comparison with various reference proteins (Fig. S2).

The *S. pneumoniae* HigB toxin contains three α-helices, one 3_10-helix (η), and five β-strands in the following order: β1 (residues 4–7), α1 (residues 15–25), α2 (residues 29–48), η1 (residues 49–52), β2 (residues 57–61), β3 (residues 64–67), β4 (residues 72–78), β5 (residues 84–91), and α3 (residues 99–115). The *S. pneumoniae* HigA antitoxin contains five α-helices and two β-strands in the following order: β1 (residues 6–9), α1 (residues 10–17), α2 (residues 20–42), α3 (residues 47–54), α4 (residues 58–66), α5 (residues 73–83), and β2 (residues 85–91). The β-strands of HigB (β1–β5) form antiparallel β-sheets. The interactions of helices α1 and α2 of both HigB and HigA form a heterodimeric interface (Fig. 1B). Helices α3 and α4 of the HigA antitoxin constitute the HTH DNA-binding motif [19] (Fig. 1C). The N-terminal β-strands of both HigB and HigA form intermolecular β-sheets. Because the C-terminal β-strands of proximal HigAs...
bind to form a dimeric interface, two intermolecular β-sheets form at both the N and C termini of HigA (Fig. 1A).

**Comparative analysis of HigBA structures**

The three previously reported structures of HigBA from *Proteus vulgaris* (PDB code 4MCT) (Fig. 2A) [16], *Escherichia coli* (PDB code 5IFG) (Fig. 2B) [17], and *Vibrio cholera* (PDB code 5JAA) (Fig. 2C) [15] were compared with the newly obtained structure of HigBA from *S. pneumoniae* (PDB code 6AF4) (Fig. 2D). These four structures share the features that each HigB has antiparallel β-sheets as its main functional unit and each HigA contains an HTH DNA-binding motif. They form a heterotetrameric assembly, and dimerization is achieved only through the HigAs [HigB-(HigA)2-HigB]. However, significant differences occur in the complex structures, and the interface between HigB and HigA, the conformation of the HigA dimer, and the position of the DNA-binding motif of HigA are quite different from each other.

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**Fig. 1.** Overall structure of heterotetrameric *S. pneumoniae* HigBA. PyMOL was used to generate Fig. 1. A ribbon representation of the HigBA heterotetramer. Chains A and C of HigB are shown in green and olive, respectively. Chains B and D of HigA are shown in orange and yellow, respectively. The structure reveals a dimeric interface between the C-terminal β-strands of the HigA subunits. The β-strands of HigB (β1-β5) form antiparallel β-sheets. B Binding conformation of the HigBA complex. Interactions between the N-terminal helices of both HigB and HigA result in a crossed-scissor conformation at the heterodimeric interface. (C) HTH DNA-binding motif consisting of α3 and α4 of HigA.
HigAs from *E. coli* and *S. pneumoniae* wrap the cognate HigB with their two helices. HigA from *V. cholerae* wraps its cognate HigB, with one α-helix and one β-strand. However, two helices from *P. vulgaris* HigA form boundary junctions with two helices from its cognate HigB. The dimerization domains of HigAs are also quite different from each other. In *P. vulgaris* and *E. coli*, one helix is involved in dimerization, and in *V. cholerae*, two helices are packed to form a dimerization domain. In contrast, in *S. pneumoniae*, an antiparallel β sheet is formed between the C termini of the two subunits. The HTH motif of *P. vulgaris* HigA is located at the N terminus, while the motif of *E. coli* HigBA is at the C terminus. In *V. cholerae* and *S. pneumoniae* HigA, the HTH motif occurs in the middle of the proteins. To understand the differences among HigBAs, a full-length sequence comparison and the resulting amino acid identity percentages are presented in Fig. S3. Alignments of amino acid residues were carried out using ClustalW [20] and visualized using ESPript 3.0 [21].

DNA-binding properties of HigBA

To investigate the DNA-binding properties of HigBA and HigA, an electrophoretic mobility shift assay (EMSA) experiment was conducted using the four palindromic DNAs of the HigBA upstream region (Pal-I, Pal-II, Pal-III, and Pal-IV) (Fig. 3A). The palindromic DNAs exhibited a conserved sequence represented as xaxxt(x. .)jaxxa. In the EMSA experiment, the DNA concentration was fixed at 0.01 mm and the protein concentration was increased from 0 to 0.6 mm (HigA dimer and HigBA heterotetramer). As the
amount of added protein increased, the bands of the DNA–protein complex shifted upward, showing that HigBA binds to all four DNAs (Pal-I, Pal-II, Pal-III, and Pal-IV) (Fig. 3B). In the EMSA results, two complexes showing distinct mobility were observed. We surmised that HigBA binds to DNA, which results in the formation of a lower-shifted band. However, because the DNA sequences are relatively long (at least 20mer), two HigBA heterotetramers might be able to bind to one DNA. Therefore, an increase in the fraction of the super-shifted band likely occurred at higher concentrations of HigBA. HigBA showed the highest affinity for Pal-III among the four DNAs. To monitor the binding specificity of HigBA to its upstream promoter, an EMSA experiment was also conducted using the other two palindromic sequences ‘A’ and ‘B’, which did not belong to the promoter region of HigBA. The results showed that band shifts did not occur in the cases of ‘A’ and ‘B’ or the control DNA ‘X’, indicating a lack of DNA–protein complex formation between the proteins and DNA (Supplementary Fig. S4). An isothermal titration calorimetry (ITC) experiment was conducted to estimate the binding affinity of HigBA (Fig. 3C). The DNA-binding reaction of HigBA is endothermic and entropically driven, and the thermodynamic parameters are listed in Table 1. The binding stoichiometry (n) indicates that one HigBA heterotetramer binds to each DNA duplex. The thermodynamic parameters of HigA are provided in Fig. S5. The experimental results showed that the HigBA complex has a higher affinity...
Table 1. Thermodynamic parameters upon the binding of DNA to HigBA

| DNA   | n (the binding stoichiometry) | $K_d$ (μM) | $\Delta H$ (kcal mol$^{-1}$) | $\Delta S$ (kcal mol$^{-1}$) | $\Delta G$ (kcal mol$^{-1}$) |
|-------|------------------------------|-----------|--------------------------|--------------------------|--------------------------|
| Pal-I | 0.53 ± 0.07                  | 0.46 ± 0.03 | 2.8 ± 0.4                | 8.74                     | −5.94                    |
| Pal-II| 0.49 ± 0.06                  | 0.70 ± 0.08 | 3.6 ± 0.3                | 7.37                     | −3.77                    |
| Pal-III| 0.48 ± 0.04                | 0.20 ± 0.01 | 1.7 ± 0.2                | 8.14                     | −6.44                    |
| Pal-IV| 0.48 ± 0.05                  | 1.19 ± 0.14 | 4.3 ± 0.3                | 9.73                     | −5.43                    |

for DNA than the HigA complex based on the $K_d$ values. Other ITC experiments were also performed in the absence of proteins under the same experimental conditions. No calorimetric changes were observed in the control experiments using a protein-free buffer. In addition, no thermal changes were observed in the experimental injections of control DNA ‘X’ or those with the palindromic sequences ‘A’ and ‘B’.

To determine the DNA-binding site of HigA, a NMR titration experiment on HigA was conducted with Pal-III DNA, which had the highest affinity with HigA. Because of the poor quality of the spectrum of full-length HigA (Fig. S6A), HigA$^{19–97}$ was used for this experiment. The 63 N-H peaks in the $^{1}H$–$^{15}N$ heteronuclear single quantum coherence spectroscopy (HSQC) spectra were assigned to individual residues of HigA$^{19–97}$ (Fig. S6B). The oligomeric state of the deletion mutant HigA$^{19–97}$ was determined to be a homodimer and showed an unchanged binding mode (Fig. S2). Then, the HSQC spectra of HigA$^{19–97}$ (0.5 mm) with increasing concentrations of Pal-III (0 to 0.1 mm) were overlaid (Fig. 4A). The $^{1}H$–$^{15}N$ HSQC peaks of HigA showed no significant chemical shift changes with decreased intensity as illustrated in Fig. 4B. The intensity reduction and disappearance of the $^{1}H$–$^{15}N$ HSQC cross-peak of HigA may originate from the increased transverse relaxation rate of HigA$^{19–27}$ by complex formation with Pal-III DNA. Although small chemical shift changes were observed, S34, T69, Q72, and A88 showed relatively distinct chemical shift changes compared with the other residues (Fig. 4B) as well as large intensity reductions. Although the overall cross-peaks exhibited intensity reduction, the largest effects for $^{1}H$–$^{15}N$ HSQC cross-peak intensity reduction were found in helices α3, α4, and α5 and their connecting loops. Helices α3 and α4 form the HTH motif, which is strongly related to DNA binding. In addition, α5 and its connecting loop are close to the HTH motif.

Based on these results, the binding pattern between DNA and HigBA was deduced by HADDOCK. Multiple solutions of the in silico docking model of the HigBA-DNA complex were clustered automatically to obtain the best single cluster of docking results. The best solution showed a score of $−207.9 ± 2.7$, the smallest root mean square deviation (RMSD) (2.2 ± 0.3 Å), and the largest buried surface area (2458.2 ± 38.0 Å$^2$) (Fig. 4C). Helices α3 and α4 of HigA constituting the HTH motif bind mainly to the major groove of the DNA. In the electrostatically generated interface, α3 exhibited a negative charge while α4 exhibited a positive charge.

Active site of HigB deduced from homologs

HigB toxins contain several conserved active site residues that are essential for the cleavage of mRNA. For a detailed comparison of S. pneumoniae HigB (Fig. 5A and E) with other HigBs, the structural similarities of HigBs were analyzed using Dali [22], and the reported active site residues were arranged. Despite low statistical similarity, the active sites of HigBs were found to be located in their antiparallel β-sheets. The detailed Dali search results with HigB from Vibrio cholera [15] (Fig. 5B), HigB from E. coli [17] (Fig. 5C), and HigB from Proteus vulgaris [16] (Fig. 5D) are shown in Table 2. According to the structural alignment, the putative active residues of S. pneumoniae HigB for ribonuclease activity may be D61, E66, R68, R73, F90, and K92. Compared with other HigBs, the active site residues of S. pneumoniae HigB are sterically hindered by a flexible loop between β5 and α3.

In addition, the overall structures of ribonucleases from similar families were compared using Dali, which showed high statistical values for ParE from Mesorhizobium opportunistum [23] (Fig. 5F), RelE from Pyrococcus horikoshii [24] (Fig. 5G), and YoeB from E. coli [25] (Fig. 5H) (Table 3). These ribonucleases showed significant structural similarity despite their low sequence identity. The C-terminal α-helix in S. pneumoniae HigB is not conserved in other structural homologs.

Intermolecular interaction of HigBA

Substantial intermolecular networks between HigB and HigA are formed at the cross-scissor interface. Helices α1 and α2 of both HigB and HigA underwent hydrophilic and hydrophobic interactions to form binding interfaces. In the interface, the residues of HigB involved in
hydrophilic interactions are R21, R32, N39, D40, E43, and H48, and they interact with W10, R14, E16, E22, S26, R29, and S34 of HigA. D40 of HigB has the closest interactions with S26 and R29 of HigA (Fig. 6A). Hydrophobic interactions are contributed by F17, M20, L35, Y41, I42, and L44 of HigB and by W10, V13, L17, F18, V30, and M33 of HigA. F17, M20, and L44 of HigB have hydrophobic interactions with multiple counterparts (Fig. 6B). These binding networks are summarized in Fig. 6A and B.

Ribonuclease activities of HigB and its mutants

We performed ribonuclease activity assays on HigB using fluorescence-labeled RNA substrates. According to amount of the substrate cleaved by HigB, fluorescence is proportionally emitted. The ribonuclease activity of *S. pneumoniae* HigB is verified through the increase in resulting fluorescence (RFU) as a function of time and concentration (Fig. 7A). The control containing only RNA substrate without proteins showed no ribonuclease activity.

In addition, to determining the mutation effects of active site residues and the main residues interacting with HigA, ribonuclease activity was also measured for a total of 10 single amino acid mutants (mutations in active site residues D61A, E66A, R68A, R73A, F90A, and K92A and HigA-interacting residues D40A, M20A, F17A, and L44A) (Fig. 7B). R73A showed approximately 70% less ribonuclease activity than the wild-type (WT) HigB, and F90A and R68A also showed significant reductions in ribonuclease activity.
activity. The mutation effect of K92 (K92A) was relatively reduced, and the mutations of D61, E66, and the HigA-interacting residues had almost negligible effects on ribonuclease activity. The secondary structural integrity of HigB and the HigB mutants was confirmed by circular dichroism (CD) spectroscopy (Supplementary Fig. S7A).

Based on these in vitro results, four efficient mutants (R68A, R73A, F90A, and K92A) showing significant reductions in ribonuclease activity were validated through an in vivo toxicity neutralization test. To perform this kill-and-rescue assay using E. coli, HigB and its mutants were cloned into the L-arabinose-inducible vector pBAD33 and HigA was cloned into the IPTG inducible vector pET-21a (+). As expected from the in vitro test results, the toxicity of HigB was efficiently rescued by HigA and HigB mutations of residues R68, R73, and F90. The K92A mutant exhibited slightly

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**Fig. 5.** Active site residues of *S. pneumoniae* HigB compared to those of other HigBs, and overall structures of ribonucleases from similar families. PyMOL was used to generate Fig. 5. (A–D) Active site residues of *S. pneumoniae* HigB compared to those of other HigBs. The HigB active site residues of (A) *S. pneumoniae*, (B) *V. cholerae*, (C) *E. coli*, and (D) *P. vulgaris*. The active sites of HigBs are oriented to show the antiparallel β-sheets in frontal view. (E–H) Overall structures of ribonucleases from similar families. The C-terminal α-helix observed only in *S. pneumoniae* HigB is highlighted in a green dashed box. (E) *S. pneumoniae* HigB, (F) *M. opportunistum* ParE, (G) *P. horikoshii* RelE, and (H) *E. coli* YoeB

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**Table 2.** Structural similarity comparison of HigB with its homologs using the Dali server

| Protein name | Source        | PDB code (used chain) | Z-score | RMSD (Å) | Number of aligned Cα | Sequence identity (%) |
|--------------|---------------|-----------------------|---------|----------|----------------------|----------------------|
| HigB         | *V. cholera*  | 5JAA (D)              | 6.8     | 3.6      | 87                   | 10                   |
| HigB         | *E. coli*     | 5IFG (A)              | 6.9     | 2.9      | 83                   | 4                    |
| HigB         | *P. vulgaris* | 4MCT (D)              | 6.4     | 2.7      | 77                   | 14                   |

**Table 3.** Structural similarity comparison of HigB with its homologs using the Dali server

| Protein name | Source        | PDB code (used chain) | Z-score | RMSD (Å) | Number of aligned Cα | Sequence identity (%) |
|--------------|---------------|-----------------------|---------|----------|----------------------|----------------------|
| ParE         | *M. opportunistum* | 5CEG (D)             | 9.8     | 2.4      | 86                   | 3                    |
| RelE         | *P. horikoshii* | 1WMI (A)              | 9.8     | 1.8      | 80                   | 11                   |
| YoeB         | *E. coli*     | 2A6R (D)              | 9.8     | 2.0      | 79                   | 16                   |
weak toxicity compared to the WT HigB, showing that the toxicity of HigB was not strongly affected by the mutation of K92 compared to mutations of the other residues (R68, R73, and F90) (Fig. 8). This assay demonstrated that HigB is a \textit{bona fide} toxin with cytotoxicity and that the active site of HigB has been successfully deduced.

**HigBA complex disruption by mimicking peptide**

Peptides mimicking the binding region between HigB and HigA were added to disrupt the HigBA complex. When the HigBA complex is disrupted by peptides and free HigB is released, fluorescence-labeled RNA substrates emit fluorescence in proportion to the

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**Fig. 6.** Crossed-scissor heterodimeric interface between HigB and HigA. The same colors as in Fig. 1 are employed. PyMOL was used to generate Fig. 6. (A) Residues involved in hydrophilic interactions at the interface. Hydrogen bonds and salt bridges are shown as black dotted lines. The residues participating in hydrophilic interactions are also shown in stick representation. (B) Residues involved in hydrophobic interactions at the interface. The residues participating in hydrophobic interactions are also shown in stick models. (A,B) Schematic diagrams of the hydrophilic interactions (left) and hydrophobic interactions (right) contributed by residues in the heterodimeric interface of HigBA.

**Fig. 7.** \textit{In vitro} ribonuclease activity assays of \textit{S. pneumoniae} HigB with its mutants. Each experiment was performed in triplicate. (A,B) \textit{In vitro} ribonuclease activity assays of \textit{S. pneumoniae} HigB and comparison of the ribonuclease activity with that of HigB mutants. (A) Fluorescence measurements as a function of time. The control did not show activity, and HigB showed concentration-dependent ribonuclease activity. (B) Comparison of the ribonuclease activity of the WT HigB with those of the HigB mutants. A WT HigB at 2 \(\mu\)M was used as a reference for comparison. The concentration of each HigB mutant was the same as that of the WT HigB. The RFU obtained with the reference was taken as 100%. Arg73 of HigB showed the strongest effect on the ribonuclease activity of HigB. Error bars represent the standard deviations of the three replicate reactions.
amount of the substrate cleaved by HigB. The α-helical nature of one HigB-mimicking peptide and two HigA-mimicking peptides was confirmed by CD spectroscopy (Fig. S7B).

The addition of mimicking peptides resulted in the effective disruption of HigBA, showing regeneration of the toxicity of peptide-treated HigBA compared to the activity of HigB (Fig. 9A). When the HigB α2 helix-mimicking peptide was added to HigBA, the ribonuclease activity reached approximately 80% of that of HigB. Subsequently, the complex disruption of the HigB-mimicking peptide was also verified by chromatograms using a Superdex 75 10/300 prepacked column (GE Healthcare, Chicago, IL, USA) (Fig. 9B).

**Artificial activation of toxin by designed peptides can exert bactericidal effects in vivo**

To support the above results in an in vitro enzyme-level experiment, diverse in vivo studies were performed. An antimicrobial activity test of the mimicking peptides against *S. pneumoniae* TIGR4 revealed that cell growth was inhibited at concentration of 6.3 μm HigA α1-, HigA α2-, or HigB α2-mimicking peptide. A cell growth assay of the designed peptides at 6.3 μm showed the highest growth defects in the presence of the HigB α2-mimicking peptide (Fig. 9C). Scrambled peptides with the same amino acid sequence as the HigB α2-mimicking peptide did not inhibit *S. pneumoniae* cell growth.

We then tested the effect of HigB α2-mimicking peptide addition on *S. pneumoniae* TIGR4 cells by the flow cytometry analysis of cells labeled with LIVE/DEAD BacLight stains (Fig. 9D). In the absence of peptides, 22.1% of *S. pneumoniae* TIGR4 cells were permeable to propidium iodide (PI). In contrast, for *S. pneumoniae* TIGR4 cells treated with peptides, the proportion of PI-permeable cells increased to 78.1% in the presence of 6.3 μm peptides after 10 h of incubation, indicating the bactericidal activity of the structure-based designed peptides. To support the suggested mode of killing, the data after 1 h of incubation with peptides are provided (Fig. 9E and F). The results show that most cells lived after 1 h of incubation. Thus, the peptides might not have caused the loss of membrane potential and membrane barrier function.

The successful intracellular localization of peptides was also demonstrated by confocal microscopy using fluorescein-labeled peptides (Fig. 9G). Fluorescein-only (left) and bright-field images of *S. pneumoniae* TIGR4 (middle) are presented. An overlaid image (right) indicates the successful intracellular localization of the peptide. Because surface-bound peptides and extracellular peptides were degraded by trypsin, the images indicate that the fluorescence signals are indeed mediated by peptide internalization. Because confocal images were taken after 1 h of incubation with the peptides, the peptides were introduced much earlier than PI and peptide passage might have allowed the
cytoplasmic membrane to maintain its barrier and energy-generating function for quite some time after introducing the peptide.

For further proof of the suggested killing mechanism, *S. pneumoniae* D39 without *higBA* and *S. pneumoniae* D39 containing intact *higBA* or active site-mutated *higBA* were treated with 6.3 μm peptides. After 1 h, the peptides entered the *S. pneumoniae* D39 cells containing active site-mutated *higBA*, and the results were consistent with that of *S. pneumoniae* TIGR4 (Fig. 9H). However, after 10 h, *S. pneumoniae* D39s containing active site-mutated *higBA* presented longer lifespans than *S. pneumoniae* D39s containing intact *higBA* (Fig. 9I). *S. pneumoniae* D39s that did not contain *higBA* seemed to be unaffected by peptides.

**Discussion**

**Insights into the transcription regulation mechanism of the HigBA system**

Although many other proteins present HTH DNA-binding motifs, antitoxins with HTH motifs in the HigBA system not only reveal differences in the location of their DNA-binding motifs but also show functional differences. HigA antitoxins have almost no structural similarity to other proteins with an HTH motif, such as the domain position, dimerization mechanism, and binding pattern to the cognate HigB toxin. The structural variability of these HigA antitoxins will cause differences in their transcriptional regulation. In the case of *S. pneumoniae* HigA, the binding between DNA and HigA is dominated by certain residues belonging to the HTH motif and their neighboring residues. When DNA was added to HigA through titration, these residues showed an overall reduction in NMR peak intensity, which shows that these residues are affected by DNA binding.

HigBA and HigA from *S. pneumoniae* seem to have different binding affinities for the four palindromic sequences tested. HigA has been suggested to regulate transcription by sequence-specific interactions with DNA and by multisite binding [26]. In addition, our experimental data show that the HigBA complex has higher binding affinity to DNA than to HigA antitoxin alone. Because the affinities of HigA to Pal-I, Pal-II, and Pal-III are weak compared with those of HigBA, the DNA-binding modes of HigA and the HigBA complex will differ. Additionally, an operator appears to be preferentially recognized when the HigBA complex is formed. Therefore, there might be more than one transcriptional regulation mechanism. In general, the diversity of transcriptional regulation mechanisms due to the structural variability of HigA antitoxins is likely a key aspect of the transcription regulation mechanism governed by the HigBA TA system.

**Application of structural studies to the catalytic mechanism of HigB**

HigB from *S. pneumoniae* is structurally homologous to the ribosome-dependent toxin RelE [27]; however, experimental results showed possible similarities to the ribosome-independent toxin YoeB [25]. Since HigBA system proteins share very low sequence similarity, many proteins belonging to the HigBA system have been elucidated via structural studies [15–17]. Therefore, the deposition of structural information regarding additional HigBA proteins in the database may allow for confirmation of whether the C-terminal helix present in HigB is related to target specificity. Knowledge about the regulation of type II toxin-mediated cleavage could explain the mechanism underlying the cellular responses of bacteria to environmental changes [27,28]. Furthermore, recent studies demonstrated that type II toxins play crucial roles in bacteria and participate in various functions, such as drug resistance and persistent cell formation [29,30]. Therefore, understanding the molecular mechanism of RNA recognition can provide fundamental principles for understanding toxin function. Residues that have a major role in the activity of HigB have been proposed according to the results of structural and functional studies; however, data on other types of HigB are needed to understand the detailed mechanisms, such as substrate recognition and catalytic stabilization.

The active site of HigB from *S. pneumoniae* is positively charged, suggesting possible interactions with the negatively charged RNA phosphate backbone (Fig. S8). As evidenced by the mutational study, the following four key residues participate in catalysis and constitute the active site: R68, R73, F90, and K92. Individual active site residues can significantly contribute to the activity of HigB. The basic nature of the positively charged residues Arg68 and Arg73 is thought to contribute to the charge stabilization of the negatively charged transition state [27]. The HigBs may also have specificity depending on the type of residue containing the phenyl ring. F90 might be important for the catalytic efficiency of HigB and facilitate the positioning of nucleotide substrates [28]. Taken together, the evidence shows that arginine and phenylalanine, which constitute the active site, will participate in the RNA cleavage reaction by organizing the active site and substrate geometry through an acid-base mechanism. Lysine may only be related to...
additional charge stabilization in the catalysis without greatly affecting the activity of *S. pneumoniae* HigB in this study [31].

**HigBA peptide derivatives are promising new antimicrobial agents**

The active site of HigB is not occluded, although the formation of the HigBA complex can block the toxicity of HigB. We focused on the neutralization of HigB toxin via binding with HigA and designed a peptide that mimics the binding site between HigB and HigA to estimate the potential of an antibiotic peptide through free toxin release by competition with HigB or HigA at the binding site. In our mutational study, the single point mutation of any interacting residue in the tight binding region of the HigBA complex did not result in HigB toxicity. However, the HigBA complex was fully disrupted by mutation of the binding region-mimicking peptides containing numerous interacting residues; thus, a large increase in RFU was observed in the *in vitro* test. The active site of HigB is not blocked by HigA in the HigBA complex structure. And the active site and HigA binding region of HigB are distant from each other. Therefore, it can be inferred that the peptide blocks the toxicity of HigB allosterically. These notable characteristics are also observed in other HigBA complex structures. In *P. vulgaris* HigBA complex, *P. vulgaris* HigA does not cover the *P. vulgaris* HigB active site and only forms a minimal boundary junction [16]. Similarly, *V. cholerae* HigA does not interact with the active site of *V. cholerae* HigB but sufficiently neutralizes the *V. cholerae* HigB [15]. In general, HigB inhibition likely is not solely mediated by active site obstruction by HigA.

In our *in vivo* experiments, the peptides designed based on structural information about HigBA also successfully penetrated the cell membrane of *S. pneumoniae* and proved its growth arrest effect on *S. pneumoniae*. Therefore, we suggest the possibility of new antimicrobial agents based on structural and functional studies of the HigBA TA system, which has a completely new mechanism that does not exist in current antibiotics. HigB toxin does not exhibit toxicity when it is part of a TA complex because the HigA antitoxin completely blocks the active site of the HigB. However, when a peptide binds to a TA complex and interacts with its binding partners, the complex is disrupted by the peptide inhibitor and the toxin is released from the complex. Additionally, in the case of the VapBC and HicBA systems, the activity of the toxin is regenerated when an active site of the free toxin is exposed by the addition of a peptide mimicking the binding site [32,33].

Taken together, our study establishes that the structurally designed peptides can function as bactericidal agents in *S. pneumoniae*. Furthermore, improvements in activity and stability obtained by modifying current candidate peptides using hydrocarbon α-helix stapling or hydrogen bond surrogates will enable these antibiotic peptides as a new type of antimicrobial agent that targets specific pathogenic bacteria [14,34,35].

**Materials and Methods**

**Cloning, mutation, expression, and purification**

The genes encoding *S. pneumoniae* (strain TIGR4) HigB (sp_1143) and HigA (sp_1144) were amplified by polymerase chain reaction (PCR) using the primers HigB-F, HigB-R, HigA-F, and HigA-R (Table S1). The restriction enzymes *Nde1* and *Xhol* were used to cut the PCR products and vectors [pET-28b(+)] for HigB and [pET-21a(+)] for HigA (Novagen, Madison, WI, USA). The cloned plasmids of HigB and HigA were cotransformed into *E. coli* Rosetta (DE3) pLysS for the overexpression of the HigBA complex. The cells were grown in Luria-Bertani (LB) medium containing 50 µg·mL<sup>−1</sup> ampicillin and 30 µg·mL<sup>−1</sup> kanamycin at 310 K until an OD<sub>600</sub> of 0.5 was reached. The overexpression of the target proteins was induced by 0.5 mm isopropyl-β-D-thiogalactopyranoside (IPTG) with an additional incubation for 4 h. The cells were harvested at 11,355 x g. The harvested cells were lysed by ultrasonication in buffer A (20 mm Tris·HCl, pH 7.9, 500 mm NaCl) containing 10% (v/v) glycerol. The entire lysate was centrifuged at 28,306 g for 1 h at 277 K. The supernatant was loaded onto an open Ni<sup>2+</sup>-affinity column (Qiagen, Hilden, Germany), pre-equilibrated with buffer A, and washed with buffer A containing 50 mm imidazole. The protein bound to the column was eluted by an imidazole gradient (50–700 mm). For further purification using ion-exchange chromatography, the buffer containing the target proteins was changed to buffer B (50 mm MES, pH 6.0). Proteins were loaded onto a HiTrap SP column (GE Healthcare) pre-equilibrated with buffer B and eluted with a NaCl gradient (0–1.5 m). As a final step, size-exclusion chromatography was conducted using an HiLoad 16/600 Superdex 200 prep-grade column (GE Healthcare) in a final buffer (50 mm MES, pH 6.0, 500 mm NaCl). The purified proteins were confirmed by sodium dodecyl sulfate–polyacrylamide gel electrophoresis and concentrated using an Amicon Ultra centrifugal filter unit (Merck Millipore, Burlington, MA, USA) in preparation for crystallization. The HigBA complex displaced by selenomethionine (SeMet) (Calbiochem, San Diego, CA, USA) was expressed and purified by the same procedure, except M9 medium containing extra essential amino acids was used for cell growth.
Design of toxin activator based on HigBA structure

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For the ribonuclease activity assay of HigB, 8 M urea was added to purified HigBA complex as a denaturant. The final buffer containing 8 M urea and HigBA complex was incubated at 293 K for 4 h. Size-exclusion chromatography was conducted on the denatured HigBA complex using a HiLoad 16/600 Superdex 200 prep-grade column (GE Healthcare). The eluted fraction containing HigB was diluted to remove the denaturant and enable refolding, and the denatured HigB was diluted by a factor of one thousand with the final buffer. Mutated forms of HigB were constructed and subjected to a ribonuclease activity assay. Ten residues of HigB were mutated to alanine (F17A, M20A, D40A, L44A, D61A, E66A, R68A, R73A, F90A, and K92A) (Table S1) and cloned into pET-28b (Novagen), resulting in an EZ Change Site-Directed Mutagenesis Kit (Enzymics, Daejeon, South Korea). The protein complexes of HigB mutants with the WT HigA were expressed and purified by the same procedure used for the WT HigBA. Each HigB mutant was generated by denaturation and refolding by the same procedure used for the WT HigB.

A HigA construct consisting of residues 19 to 97 was used for the NMR experiments. The genes encoding HigA19-97 were amplified by PCR using primers HigA-F19 19-97 and HigA-R (Table S1) and cloned into pET-28b (+) (Novagen), resulting in a twenty-residue tag (MGSSHHHHHHSSGLVPRGSH) at the N terminus. The plasmid encoding HigA19-97 was transformed into E. coli Rosetta (DE3) pLysS for overexpression, and the expression was induced with 1 mm isopropyl β-D-1-thiogalactopyranoside (IPTG). The expressed HigA construct was purified using a Ni-NTA column (GE Healthcare). The eluted fraction containing HigA was diluted by a factor of one thousand with the final buffer. Mutated forms of HigA were constructed and subjected to a ribonuclease activity assay.

Crystallization, X-ray data collection, and structure determination

Crystals of the HigBA complex were grown using the sitting-drop vapor diffusion method at 293 K by mixing equal volumes (0.5 μL each) of the protein solution (11 mg·mL−1 in 50 mm MES, pH 6.0, 500 mm NaCl) and the reservoir solution (20% (w/v) PEG 1000, 0.1 m potassium phosphate monobasic/sodium phosphate dibasic, pH 6.2, and 0.2 m sodium chloride). A cryoprotectant solution consisting of 25% (v/v) glycerol was added to the reservoir solution for data collection under a liquid nitrogen stream. The crystal was vitrified in the cold nitrogen gas stream, and the data were collected using an ADSC Quantum Q270r CCD detector at beamline 5C of the Pohang Light Source, Republic of Korea. Crystals of HigBA belonged to the monoclinic space group P21, with unit cell parameters of a = 74.903 Å, b = 74.403 Å, c = 98.321 Å, α = γ = 90.00°, β = 90.08° for the native complex and a = 74.575 Å, b = 67.038 Å, c = 87.771 Å, α = γ = 90.00°, β = 94.22° for the SeMet-labeled complex. The structure of the HigBA complex was first phased at 2.80 Å resolution by single-wavelength anomalous dispersion using SeMet. The structure of the HigBA complex was calculated using 2.30 Å resolution diffraction data of the native complex with a molecular replacement based on the SeMet protein model. The detailed statistical information is described in Table S2. PHENIX[37] was first used to automatically build a residue, and the remaining residues were built in Coot [38] to provide the final model. The Rwork and Rfree [39] values were calculated by REFMAC and PHENIX for refinement [37,40]. The overall geometry was validated using molprobity [41]. Figures were generated using pymol [42]. The electrostatic potential surfaces were calculated using the Adaptive Poisson-Boltzmann solver (APBS) method [43].
Electrophoretic mobility shift assay

The binding affinities of HigA and HigBA to the palindromic sequence in their own operator region were estimated by EMSA. Typically, type II antitoxins interact with DNA via symmetric palindromic stretches that are approximately 15–30 nucleotides long called inverted repeats. For higBA, we were able to locate several inverted repeats within the promoter region. Four DNA duplexes were selected and termed Pal-I, Pal-II, Pal-III, and Pal-IV (Table S3). Control experiments were also performed with DNA ‘X’ and the two other palindromic sequences ‘A’ and ‘B’ (Table S3). Purified DNA duplexes were purchased from Bioneer (Daejeon, South Korea). Reaction mixtures (10 µL) containing 0.01 mm of duplex DNA and different protein concentrations (a series of protein concentrations, with the HigA dimer and HigBA heterotetramer ranging from 0 to 0.6 mm) were prepared in binding buffer (50 mm MES, pH 6.0) and incubated for 30 min at 293 K. The reaction mixtures were electrophoresed on a 0.5% agarose gel with 0.5x TBE for 20 min. The results were visualized using a Printgraph 2M (ATTO, Tokyo, Japan).

Isothermal titration calorimetry measurements

Isothermal titration calorimetry (ITC) experiments were conducted using an iTC200 calorimeter (Malvern Instruments, Malvern, Worcestershire, UK) at 25 °C. The proteins and promoter dsDNAs were prepared in a buffer consisting of 50 mm MES, pH 6.0, and 100 mm NaCl for both HigA and HigBA. The DNA duplexes used in the ITC measurements were the same as those used in the EMSA (Table S3). Affinity experiments were conducted with the protein solution (10 µm; HigA dimer and HigBA heterotetramer, 320 µL) in the cell and the dsDNA solution (600 µm) as the injected titrant. In total, 19 injections performed at 180-s intervals were used for data collection. The MicroCal Origin software was used for curve fitting to calculate the binding affinity ($K_d$), the enthalpy of binding ($\Delta H$), the entropy of binding ($\Delta S$), and the stoichiometry ($n$). The raw data were fitted using a one-site binding model. The Gibbs-free energies ($\Delta G$) were calculated using the standard equation $\Delta G = \Delta H - T\Delta S$.

NMR spectroscopy

For the NMR analysis, HigA$^{19-97}$ was uniformly labeled with $^{13}$C and $^{15}$N and prepared in 20 mm MES, pH 6.0, 50 mm NaCl, 1 mm EDTA, 0.1 mm PMSF, and 10% D$_2$O. All NMR measurements were conducted at 308 K using an AVANCE 800 MHz spectrometer equipped with a cryogenic probe (Bruker BioSpin, Billerica, MA, USA). $^1$H, $^13$C, and $^{15}$N chemical shift assignments for the backbone nuclei were obtained using 2D $^1$H-$^15$N HSQC and a set of triple resonance experiments to obtain the HNCO, HN(CO)CA, HN(CA)CO, HN(CA)CN, HN(CA)CO, HN(CA)CN, and HN(CA)CN spectra. Data were processed using NMRPipe and nmrDraw [44] and further analyzed using NMRviewJ [45].

NMR titration experiment

The NMR titration experiments on $^{15}$N-labeled HigA$^{19-97}$ with the promoter dsDNA Pal-III (Table S3) were conducted by acquiring the $^1$H-$^15$N HSQC spectra. Throughout the NMR titration experiments, the concentration of the HigA$^{19-97}$ dimer was maintained at 0.5 mm, and the concentration of the promoter dsDNA was increased from 0 to 0.1 mm. The chemical shift perturbation (CSP) was calculated and analyzed using Equation 1, where $\Delta\delta_N$ and $\Delta\delta_H$ are the CSP values of the amide nitrogen and proton, respectively. The intensity ratio was calculated by dividing the peak intensity of the last spectrum by the peak intensity of the first spectrum:

$$\Delta\delta_{avg} = \frac{\text{intensity of last spectrum}}{\text{intensity of first spectrum}}.$$

In silico HigBA-DNA docking

In silico molecular docking to examine the interaction between Pal-III and HigBA was performed using the high ambiguity-driven protein–protein DOCKing algorithm (HADDOCK) [46]. The 20-base pair promoter DNA was modeled using a 3D-DART server [47]. Residues S57, T69, and Q72 were defined as ‘active residues’ based on the NMR titration results. Residues A2, C12, T13, and T14 of Pal-III-F, T7, T8, A9, and C19 of Pal-III-R, Q47, R63, and T75 of HigA were additionally designated to generate a rational interface for HTH recognition contact between the protein and DNA. Passive residues were defined automatically as the residues surrounding the active residues.

In vitro ribonuclease assay

The ribonuclease activity of HigB and HigB mutants was verified using an RNase Alert Kit (IDT, Coralville, IA, USA) as previously reported (32).

Kill-and-rescue assay

Genes encoding HigB mutants (R68A, R73A, F90A, and K92A) were amplified and cloned into pBAD33. The compatible plasmid pairs (pBAD33 and pET-21a(+) or pET-21b(+) with and without each HigB mutant and HigA were cotransformed into BL21 (DE3) competent cells and grown overnight at 37 °C. The overnight cultures were diluted by 100-fold in 20 ml of fresh M9 minimal medium containing 25 µg mL$^{-1}$ chloramphenicol and 50 µg mL$^{-1}$ ampicillin. The bacteria were then streaked onto LB medium plates supplemented with inducer L-arabinose (0.2% w/v) or 0.5 mm IPTG. The plates were incubated overnight at 37 °C before examination of the cell growth.
Circular Dichroism spectroscopy

Circular Dichroism (CD) measurements were conducted to verify the structural integrity of the HigB and HigB mutants using a Chirascan plus spectropolarimeter (Applied Photophysics, Leatherhead, UK) at 20°C in a cell with a 1-mm light path. CD scans were performed from 260 nm to 190 nm with a 1 nm bandwidth and a scan speed of 100 nm·min⁻¹. Three scans were averaged, followed by subtraction of the solvent signal.

Peptide design and addition to the HigBA complex

Three peptides that mimicked the binding interface region between HigB and HigA were designed and purchased from AnyGen (GwangJu, South Korea, www.anygen.com). Theoretically, mimicked peptides compete with the original protein for binding, and if peptides occupy the binding site, the toxin is freely released to digest RNA, which results in the detection of fluorescence in the fluorescence quenching assay. To disrupt the binding interface between HigB and HigA and restore the toxicity of HigB, peptides were added to the HigBA complex (Table S4).

Antimicrobial activity test

The antimicrobial activity of the mimicking peptides was evaluated as previously reported (32). The activity of the peptides against Gram-positive *S. pneumoniae* TIGR4 (ATCC® BAA-334™) was tested. Several scrambled peptides with the same amino acid sequence as the HigB α2 helix-mimicking peptide were also tested (Table S4).

In vivo cell growth assay

For the cell growth assay, *S. pneumoniae* TIGR4 was grown for 10 h without agitation at 310 K in Todd Hewitt Broth (THB) in air containing 5% CO₂ until the mid-exponential phase (OD₆₀₀ of 0.5). The 10 h of cultures were inoculated into fresh medium for assays. The cells were incubated at 37°C for 10 h, and growth was monitored at 1 h of intervals. The average OD₆₀₀ values, along with the standard error of the mean, were plotted in MicroCal Origin software.

Generation of *S. pneumoniae* D39 containing mutated HigBA

For the electroporation experiments, *S. pneumoniae* D39 cells were grown at 37°C in THB until the early-exponential phase (OD₆₀₀ of 0.1–0.2). They were harvested and washed twice using electroporation medium containing 0.5 M sucrose, 7 mm potassium phosphate, pH 7.5, and 1 mm MgCl₂. Subsequently, they were concentrated ten times in the same medium mentioned above. Then, 0.1 mL of the concentrated cell suspension was poured into a Bio-Rad cuvette along with 1 μg·mL⁻¹ plasmid DNA. The mixture was incubated on ice for 10 min. A single impulse of current was supplied with the gene pulse apparatus set at 6.25 kV and 25 μF. The electro-transformed cells were then incubated at 37°C with constant shaking for approximately 1 h. Finally, the desired cells were obtained using selective media.

Flow cytometry

*Streptococcus pneumoniae* TIGR4 cells were cultured in a 96-well flat-bottom plate containing THB (MBcell) with various concentrations of peptide using a serial dilution method. At two time points (1 h later and 10 h later), bacteria cells were harvested and stained with a LIVE/DEAD BacLight Kit (Thermo Fisher Scientific, Waltham, MA, USA) containing SYTO® 9 and PI. A positive control was prepared by measuring fresh cells, and a negative control was prepared by incubating cells with 70% isopropyl alcohol for 15 min before washing. For staining, the cells were incubated with 5 μm SYTO® 9 and 30 μm PI at room temperature and protected from light for 15 min. The cells were analyzed with an LSRFortessa cell analyzer (BD Bioscience, San Jose, CA, USA), and the data were analyzed using FlowJo software (TreeStar, Ashland, OR, USA). *S. pneumoniae* D39 cells and four active site mutants containing *S. pneumoniae* D39 cells were subjected to the same treatment used for the *S. pneumoniae* TIGR4 cells after 10 h of incubation with peptides.

Confocal images

*S. pneumoniae* cells were cultured in THB during exponential growth (OD₆₀₀ between 0.4 and 0.6). The cells were diluted with Difco™ Mueller Hinton Broth (BD Bioscience) to obtain a 10% cell suspension. The cell suspension was incubated with 1 μm peptide samples conjugated with fluorescein for 30 min, and then, to remove the peptides bound to the cell surface, 1 mg·mL⁻¹ trypsin from bovine pancreas (Sigma Life Sciences) was added for quenching. After 30 min, the cells were washed with PBS and then analyzed with a Leica TCS SP8 (Leica Microsystems, Wetzlar, Germany) confocal microscope.

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Conflict of Interest
The authors declare no conflict of interest to declare.

Author contributions
SMK, CJ, and DHK conceived the study. SMK, CJ, DHK, and SWH performed the experiments. SMK, CJ, and DHK analyzed the data. SMK, CJ, DHK, and SWH wrote the manuscript. SMK, DHK, CJ, SJP, and BJL edited the manuscript.

References
1 Henriques-Normark B & Tuomanen EI (2013) The pneumococcus: epidemiology, microbiology, and pathogenesis. Cold Spring Harb Perspect Med 3, a010215.
2 Pletz MW, Maus U, Krug N, Welte T & Lode H (2008) Pneumococcal vaccines: mechanism of action, impact on epidemiology and adaption of the species. Int J Antimicrob Agents 32, 199–206.
3 Wahl B, O’Brien KL, Greenbaum A, Majumder A, Liu L, Chu Y, Lukic I, Nair H, McAllister DA, Campbell H et al. (2018) Burden of Streptococcus pneumoniae and haemophilus influenzae type b disease in children in the era of conjugate vaccines: global, regional, and national estimates for 2000–15. Lancet Glob Health 6, e744–e757.
4 von Mollendorf C, Tempia S, von Gottberg A, Meiring S, Quan V, Feldman C, Cloete J, Madhi SA, O’Brien KL, Klugman KP et al. (2017) Estimated severe pneumococcal disease cases and deaths before and after pneumococcal conjugate vaccine introduction in children younger than 5 years of age in South Africa. PLoS One 12, e0179905.
5 Bridy-Pappas AE, Margolis MB, Center KJ & Isaacman DJ (2005) Streptococcus pneumoniae: description of the pathogen, disease epidemiology, treatment, and prevention. Pharmacotherapy 25, 1193–1212.
6 Aliberti S, Cook GS, Babu BL, Reyes LF, H. Rodriguez A, Sanz F, Soni NJ, Anzueto A, Faverio P, Sadud RF, et al. (2019) International prevalence and risk factors evaluation for drug-resistant Streptococcus pneumoniae pneumonia. J Infect 79, 300–311.
7 Ogura T & Hiraga S (1983) Mini-F plasmid genes that couple host cell division to plasmid proliferation. Proc Natl Acad Sci U S A 80, 4784–4788.
8 Hayes F & van Melderen L (2011) Toxins-antitoxins: diversity, evolution and function. Crit Rev Biochem Mol Biol 46, 386–408.
9 Otsuka Y (2016) Prokaryotic toxin-antitoxin systems: novel regulations of the toxins. Curr Genet 62, 379–382.
10 Harms A, Brodersen DE, Mitarai N & Gerdes K (2018) Toxins, targets, and triggers: an overview of toxin-antitoxin biology. Mol Cell 70, 768–784.
11 Fernandez-Garcia L, Blasco L, Lopez M, Bou G, Garcia-Contreras R, Wood T & Tomas M (2016) Toxin-antitoxin systems in clinical pathogens. Toxins 8, 227.
12 Kedzierska B & Hayes F (2016) Emerging roles of toxin-antitoxin modules in bacterial pathogenesis. Molecules 21, E790.
13 Lee KY & Lee BJ (2016) Structure, biology, and therapeutic application of toxin-antitoxin systems in pathogenic bacteria. Toxins (Basel) 8, E305.
14 Kang SM, Kim DH, Jin C & Lee BJ (2018) A systematic overview of type II and III toxin-antitoxin systems with a focus on druggability. Toxins (Basel) 10, E515.
15 Hadzi S, Garcia-Pino A, Haesaerts S, Jurenas D, Gerdes K, Lah J & Loris R (2017) Ribosome-dependent Vibrio cholerae mRNAse HigB2 is regulated by a beta-strand sliding mechanism. Nucleic Acids Res 45, 4972–4983.
16 Schureck MA, Maehigashi T, Miles SJ, Marquez J, Cho SE, Erdman R & Dunham CM (2014) Structure of the Proteus vulgaris HigB-(HigA)2-HigB toxin-antitoxin complex. J Biol Chem 289, 1060–1070.
17 Yang J, Zhou K, Liu P, Dong Y, Gao Z, Zhang J & Liu Q (2016) Structural insight into the E. coli HigBA complex. Biochim Biophys Res Commun 478, 1521–1527.
18 Klose DP, Wallace BA & Janes RW (2010) 2Struc: the secondary structure server. Bioinformatics 26, 2624–2625.
19 Xiong W, Li T, Chen K & Tang K (2009) Local combinational variables: an approach used in DNA-binding helix-turn-helix motif prediction with sequence information. Nucleic Acids Res 37, 5632–5640.
20 Thompson JD, Higgins DG & Gibson TJ (1994) CLUSTAL W: improving the sensitivity of progressive multiple sequence alignment through sequence weighting, position-specific gap penalties and weight matrix choice. Nucleic Acids Res 22, 4673–4680.
21 Robert X & Gouet P (2014) Deciphering key features in protein structures with the new ENDscript server. Nucleic Acids Res 42, W320–324.
22 Holm L & Rosenstrom P (2010) Dali server: conservation mapping in 3D. Nucleic Acids Res 38, W545–W549.
23 Aakre CD, Herrou J, Phung TN, Perchuk BS, Crosson S & Laub MT (2015) Evolving new protein-protein interaction specificity through promiscuous intermediates. *Cell* 163, 594–606.

24 Takagi H, Kakuta Y, Okada T, Yao M, Tanaka I & Kimura M (2005) Crystal structure of archaeal toxin-antitoxin ReE-ReLB complex with implications for toxin activity and antitoxin effects. *Nat Struct Mol Biol* 12, 327–331.

25 Kamada K & Hanaoka F (2005) Conformational change in the catalytic site of the ribonuclease YoeB toxin by YeF antitoxin. *Mol Cell* 19, 497–509.

26 Schureck MA, Meisner J, Hoffer ED, Wang D, Onuoha N, Ei Cho S, Lollar P 3rd & Dunham CM (2019) Structural basis of transcriptional regulation by the HigA antitoxin. *Mol Microbiol* 111, 1449–1462.

27 Neubauer C, Gao YG, Andersen KR, Dunham CM, Kelley AC, Hentschel J, Gerdes K, Ramakrishnan V & Brodersen DE (2009) The structural basis for mRNA recognition and cleavage by the ribosome-dependent endonuclease RelE. *Cell* 139, 1084–1095.

28 Schureck MA, Repack A, Miles SJ, Marquez J & Dunham CM (2016) Mechanism of endonuclease cleavage by the HigB toxin. *Nucleic Acids Res* 44, 7944–7953.

29 Page R & Peti W (2016) Toxin-antitoxin systems in bacterial growth arrest and persistence. *Nat Chem Biol* 12, 208–214.

30 Michiels JE, van den Bergh B, Verstraeten N & Michiels J (2016) Molecular mechanisms and clinical implications of bacterial persistence. *Drug Resist Updates* 29, 76–89.

31 Griffin MA, Davis JH & Strobel SA (2013) Bacterial toxin RelE: a highly efficient ribonuclease with exquisite substrate specificity using atypical catalytic residues. *Biochemistry* 52, 8633–8642.

32 Kang SM, Kim DH, Lee KY, Park SJ, Yoon HJ, Lee SJ, Im H & Lee BJ (2017) Functional details of the *Mycobacterium tuberculosis* VapBC26 toxin-antitoxin system based on a structural study: insights into unique binding and antibiotic peptides. *Nucleic Acids Res* 45, 8564–8580.

33 Kim DH, Kang SM, Park SJ, Jin C, Yoon HJ & Lee BJ (2018) Functional insights into the *Streptococcus pneumoniae* HicBA toxin-antitoxin system based on a structural study. *Nucleic Acids Res* 46, 6371–6386.

34 Chapman RN, Dimartino G & Arora PS (2004) A highly stable short alpha-helix constrained by a main-chain hydrogen-bond surrogate. *J Am Chem Soc* 126, 12252–12253.

35 Cromm PM, Spiegel J & Grossmann TN (2015) Hydrocarbon stapled peptides as modulators of biological function. *ACS Chem Biol* 10, 1362–1375.

36 Kruger NJ (1994) The bradford method for protein quantitation. *Methods Mol Biol* 32, 9–15.

37 Adams PD, Afonine PV, Bunkoczi G, Chen VB, Davis IW, Echols N, Headd JJ, Hung LW, Kapral GJ, Grosse-Kunstleve RW et al. (2010) PHENIX: a comprehensive Python-based system for macromolecular structure solution. *Acta Crystallogr D Biol Crystallogr* 66, 213–221.

38 Emsley P, Lohkamp B, Scott WG & Cowtan K (2010) Features and development of Coot. *Acta Crystallogr D Biol Crystallogr* 66, 486–501.

39 Brunger AT (1992) Free R value: a novel statistical quantity for assessing the accuracy of crystal structures. *Nature* 355, 472–475.

40 Murshudov GN, Skubak P, Lebedev AA, Pannu NS, Steiner RA, Nicholls RA, Winn MD, Long F & Vagin AA (2011) REFMAC5 for the refinement of macromolecular crystal structures. *Acta Crystallogr D Biol Crystallogr* 67, 355–367.

41 Chen VB, Arendall WB, Headd JJ, Keedy DA, Immormino RM, Kapral GJ, Murray LW, Richardson JS & Richardson DC (2010) MolProbity: all-atom structure validation for macromolecular crystallography. *Acta Crystallogr D Biol Crystallogr* 66, 12–21.

42 Delano WL (2002) The PyMOL Molecular Graphics System. DeLano Scientific, San Carlos, CA.

43 Baker NA, Sept D, Joseph S, Holst MJ & McCammon JA (2001) Electrostatics of nanosystems: application to microtubules and the ribosome. *Proc Natl Acad Sci U S A* 98, 10037–10041.

44 Delaglio F, Grzesiek S, Vuister GW, Zhu G, Pfeifer J & Bax A (1995) NMRPipe: a multidimensional spectral processing system based on UNIX pipes. *J Biomol NMR* 6, 277–293.

45 Johnson BA (2004) Using NMRView to visualize and analyze the NMR spectra of macromolecules. *Methods Mol Biol* 278, 313–352.

46 Dominguez C, Boelens R & Bonvin AM (2003) HADDOCK: a protein-protein docking approach based on biochemical or biophysical information. *J Am Chem Soc* 125, 1731–1737.

47 van Dijk M & Bonvin AM (2009) 3D-DART: a DNA structure modelling server. *Nucleic Acids Res* 37, W235–W239.

### Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.
Fig. S1. Ribbon representation and 90°-rotated diagrams of (A) HigBA heterotetramer, (B) HigBA dimer, (C) HigB and (D) HigA. PyMOL was used to generate Fig. S1.

Fig. S2. Molecular weight estimates of HigBA and HigA19−97 obtained by size-exclusion chromatography.

Fig. S3. Full-length sequence comparison and percentages of amino acid identities of (A) HigBs and (B) HigAs.

Fig. S4. EMSA study using other palindromic sequences ‘A’ and ‘B’ and control DNA ‘X’.

Fig. S5. ITC results with palindromes in the higBA promoter region.

Fig. S6. 2D 1H-15N HSQC spectra of full-length HigA and HigA19-97.

Fig. S7. Validation tests for toxin mutants and peptide mimetics.

Fig. S8. Active site of HigB from S. pneumoniae.

Table. S1. Primers used for cloning.

Table. S2. Data collection and refinement statistics for SeMet-substituted and native structures.

Table. S3. DNA used in EMSA, ITC and NMR titration.

Table. S4. Peptides used to disrupt the binding interface of HigBA.