RESEARCH ARTICLE

Biofilm-associated toxin and extracellular protease cooperatively suppress competitors in *Bacillus subtilis* biofilms

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Abstract

In nature, most bacteria live in biofilms where they compete with their siblings and other species for space and nutrients. Some bacteria produce antibiotics in biofilms; however, since the diffusion of antibiotics is generally hindered in biofilms by extracellular polymeric substances, i.e., the biofilm matrix, their function remains unclear. The *Bacillus subtilis yitPOM* operon is a paralog of the *sdpABC* operon, which produces the secreted peptide toxin SDP. Unlike *sdpABC*, *yitPOM* is induced in biofilms by the DegS-DegU two-component regulatory system. High *yitPOM* expression leads to the production of a secreted toxin called YIT. Expression of *yitQ*, which lies upstream of *yitPOM*, confers resistance to the YIT toxin, suggesting that YitQ is an anti-toxin protein for the YIT toxin. The alternative sigma factor SigW also contributes to YIT toxin resistance. In a mutant lacking *yitQ* and *sigW*, the YIT toxin specifically inhibits biofilm formation, and the extracellular neutral protease NprB is required for this inhibition. The requirement for NprB is eliminated by Δ*eps* and Δ*bslA* mutations, either of which impairs production of biofilm matrix polymers. Overexpression of biofilm matrix polymers prevents the action of the SDP toxin but not the YIT toxin. These results indicate that, unlike the SDP toxin and many conventional antibiotics, the YIT toxin can pass through layers of biofilm matrix polymers to attack cells within biofilms with assistance from NprB. When the wild-type strain and the YIT-sensitive mutant were grown together on a solid medium, the wild-type strain formed biofilms that excluded the YIT-sensitive mutant. This observation suggests that the YIT toxin protects *B. subtilis* biofilms against competitors. Several bacteria are known to produce antibiotics in biofilms. We propose that some bacteria including *B. subtilis* may have evolved specialized antibiotics that can function within biofilms.

Author summary

Biofilms are multicellular aggregates of bacteria that are formed on various living and non-living surfaces. Biofilms often cause serious problems, including food contamination and infectious diseases. Since bacteria in biofilms exhibit increased tolerance or resistance to antimicrobials, new agents and treatments for combating biofilm-related problems are...
required. In this study, we demonstrated that *B. subtilis* produces a secreted peptide antibiotic called the YIT toxin and its resistant proteins in biofilms. A mutant lacking the resistance genes was defective in biofilm formation. This effect resulted from the ability of the YIT toxin to pass through the biofilm defense barrier and to attack biofilm cells. Thus, unlike many conventional antibiotics, the YIT toxin can penetrate biofilms and suppress the growth of YIT toxin-sensitive cells within biofilms. Some bacteria produce antibiotics in biofilms, some of which can alter the bacterial composition in the biofilms. Taking these observations into consideration, our findings suggest that some bacteria produce special antibiotics that are effective against bacteria in biofilms, and these antibiotics might serve as anti-biofilm agents.

**Introduction**

In the environment, bacteria compete for space and nutrients [1]. Antibiotics are thought to play a critical role in this competition, and antibiotic-producing bacteria are indeed common in various environments [2–4]. However, in the environment, most bacteria are found in sessile multicellular bacterial communities known as biofilms, in which bacteria exhibit increased antibiotic tolerance or resistance [5, 6]. Though alternative environmental roles of antibiotics have been proposed [7], this paradox has not been explained in detail to date.

In biofilms, bacterial cells adhere to each other and to a surface via a mixture of extracellular polymeric substances called the biofilm matrix, which consists of exopolysaccharides, proteins, nucleic acids, and/or lipids [8, 9]. When encased in the biofilm matrix, cells exhibit increased tolerance or resistance to environmental stresses, antibiotics, host defense systems, and predation [8, 9]. Thus, biofilm formation enables bacteria to remain in a favorable niche and to claim territory; however, biofilms are not a utopia for bacteria. The properties of biofilms, including high cell density, decreased internal fluidity, and, in many cases, the presence of multiple species, lead to conditions of harsh competition, especially when nutrients are scarce. Many bacteria secrete biofilm formation-inhibiting molecules, such as biosurfactants, polysaccharides, and molecules that interfere with bacterial quorum sensing, and these secreted molecules help to exclude unfavorable competitors from biofilms [10]. Antibiotics might also play an important role in competition within biofilms. However, since the properties of biofilms, including the protection of member cells by the biofilm matrix, the increased expression of antibiotic resistance genes, and the decreased internal fluidity, reduce the efficacy of antibiotics against biofilm cells [11–16], little attention has been paid to the functions of antibiotics in competition within biofilms. However, some biofilms do indeed produce antibiotics, and several of these antibiotics can alter the bacterial composition of the biofilm [17–23]. These observations suggest that the functions of antibiotics produced in biofilms remain to be investigated. An understanding of how bacteria use antibiotics in biofilms will not only provide insight into bacterial survival strategies within biofilms, it will also lead to the discovery of tactics for combating biofilm-related problems, such as food and beverage safety issues, industrial contamination, and biofilm-related diseases.

The Gram-positive soil bacterium *Bacillus subtilis* is a model organism for biofilm formation. *B. subtilis* forms robust biofilms under laboratory conditions, for example, pellicles on the surface of liquid media under static culture conditions or wrinkled colonies on solid media [24]. *B. subtilis* biofilms are maintained by a biofilm matrix that mainly consists of exopolysaccharides, TasA amyloid fibers, and BslA hydrophobins, which are produced by proteins encoded by the *epsABCDEFGHIJKLMNO* operon, the *tapA-sipW-tasA* operon, and *bslA*,...
respectively [24–29]. These genes are directly or indirectly repressed by the transcriptional repressors AbrB and SinR [30–33]. Phosphorylation of the response regulator Spo0A induces mechanisms that antagonize these repressors, leading to the expression of the biofilm matrix synthesis genes [34, 35].

*B. subtilis* produces a wide array of antibiotics. Many of these antibiotics are non-ribosomally synthesized peptide compounds, such as surfactin, bacillaecine, fengycin, iturin, and bacilysin, which are thought to be important in nature for competition with other organisms, including fungi [4, 36]. Furthermore, *B. subtilis* produces ribosomally synthesized peptide antibiotics, such as bacteriocins and other protein-derived toxins, which are generally effective against other bacteria that are genetically similar and present in similar ecological niches [4, 37–39]. One of these protein-derived toxins is the cannibalism toxin SDP [40], whose function is involved in biofilm formation. The SDP toxin is derived from the internal sequence of SdpC, and it is encoded by the *sdpABC* operon. SdpC is a 203 amino acid protein that contains a typical N-terminal secretion signal and a C-terminal hydrophobic domain. After secretion and cleavage of the signal sequence, SdpC is further processed into the 42 amino acid peptide known as the SDP toxin, which corresponds to the C-terminal hydrophobic domain (C141 to S182) [39–41]. SdpA and SdpB are required for the processing of SdpC to SDP, and this processing is essential for the activity of the SDP toxin [41]. The hydrophobic nature of the SDP toxin enables the SDP toxin to penetrate bacterial membranes, where it then induces cell lysis by collapsing the proton motive force [42]. Downstream of *sdpABC* is the *sdpRI* operon, which encodes its own transcriptional repressor and an anti-toxin protein to the SDP toxin [43]. SdpI is an integral membrane protein that protects cells probably by binding to the SDP toxin. Transcription of the *sdpABC* and *sdpRI* operons is directly or indirectly activated by phosphorylated Spo0A [40, 43]. Spo0A is a master regulator of stationary phase development that is phosphorylated after the onset of stationary phase [44]. However, as the phosphorylation of Spo0A is subject to a bistable regulatory mechanism, a subset of *B. subtilis* cells produce the SDP toxin and the SdpI anti-toxin protein [40, 43]. Consequently, the secreted SDP toxin lyases and kills a fraction of the sibling cells that do not produce the SdpI anti-toxin protein. Since phosphorylated Spo0A also induces biofilm formation in parallel, cells that produce the SDP toxin and the SdpI anti-toxin protein efficiently develop biofilms by using nutrients released from their lysed siblings [45]. Moreover, the SDP toxin is effective not only against *B. subtilis*, but also against many Firmicutes bacteria [39, 41, 46]. Thus, the SDP toxin likely plays an important role in the early phase of biofilm formation by eliminating unnecessary types of cells and closely related competitors in the environment.

The undomesticated *B. subtilis* strain NCIB3610 encodes an *sdpABC* paralog known as *yitPOM*. While transcription of *sdpABC* is activated by Spo0A [40], *yitPOM* was previously identified as a member of the group of genes regulated by the DegS-DegU two-component regulatory system [47]. Phosphorylated DegU directly induces transcription of genes for the biofilm matrix protein, BslA, several antibiotic synthetases, and many extracellular degradative enzymes, such as proteases, levansucrase, α-amylase, β-glucanases, and xylanase. These antibiotics and degradative enzymes are thought to play important roles in bacterial competitions and nutrient acquisition in nature [47, 48, and references therein]. Since DegS-DegU is required for biofilm formation, we were interested in determining whether the *yitPOM*-encoded toxin plays a role particularly in biofilms. In this paper, we demonstrate that *yitPOM* encodes a biofilm-associated secreted toxin. Unlike many conventional antibiotics, in particular positive charged antibiotics, this toxin was able to attack cells within biofilms by passing through the layers of the biofilm matrix polymers with the assistance of an extracellular protease. Given that several other bacteria produce antibiotics in biofilms, our results suggest that some bacteria may have evolved specialized antibiotics to suppress competitors in biofilms.
Results

yitPOM encodes a toxin

The undomesticated \textit{B. subtilis} strain NCIB3610 (hereafter referred to as the wild-type strain or 3610) \cite{24} encodes an \textit{sdpABC} paralog known as \textit{yitPOM}. YitP and YitO exhibit approximately 50% sequence similarity to the entire SdpA and SdpB sequences, respectively (Fig 1A and 1B). Like SdpC, YitM has an N-terminal secretion signal; however, the sequence similarity between YitM and SdpC is limited to the N-terminal three quarters of the sequence, which does not include the entire sequence corresponding to the SDP toxin (Fig 1C). Although there is no sequence similarity, like SdpC, the YitM C-terminal region contains a hydrophobic domain (Fig 1C and 1D). These observations suggest the possibility that the C-terminal hydrophobic domain of YitM might be processed to a secreted toxin via a YitP and YitO-dependent mechanism. If this is the case, then \textit{yitPOM} encodes a toxin whose sequence differs from that of the SDP toxin.

To determine whether \textit{yitPOM} encodes a toxin, we examined the effect of \textit{yitPOM} overexpression on growth. We constructed the strain \textit{P\textsubscript{spac-hy}-yitPOM}, which ectopically expresses \textit{yitPOM} from the strong isopropyl \textbeta-D-thiogalactopyranoside (IPTG)-inducible, LacI-repressible \textit{spac-hy} promoter \cite{49} in the \textit{amyE} locus on the chromosome. The wild-type and \textit{P\textsubscript{spac-hy}-yitPOM} strains were grown with vigorous shaking in 2\times Schaeffer’s sporulation medium plus glucose (2\times SG) \cite{50} supplemented with 1 mM IPTG, and the optical density at 600 nm (OD\textsubscript{600}) was measured over time. These strains showed no difference in growth from exponential to stationary phase (Fig 2A). Toxin-producing bacteria normally express cognate anti-toxin proteins against their toxins, and the effects of the toxins do not appear unless the anti-toxin genes are deleted \cite{2, 37, 38, 40}. Since toxin and anti-toxin genes are simultaneously inserted into the genomes as exogenous genes and are frequently located close to each other in the genome \cite{2, 37, 38, 40}, genome comparison is a powerful tool to identify toxin/anti-toxin gene sets. To identify candidates for an anti-toxin gene against the putative toxin encoded by \textit{yitPOM}, we compared the genetic organization of the 3610 and \textit{B. subtilis} var. \textit{natto} BEST195 strains, the latter of which lacks \textit{yitPOM}. This comparison revealed that \textit{yitPOM} appears to be inserted between \textit{yitR} and \textit{yitL} in the 3610 genome, along with \textit{yizB} and \textit{yitQ}, which are predicted to encode a transcriptional regulator and a membrane protein, respectively (Fig 2C). Since the SdpI anti-toxin protein is a membrane protein, YitQ was a candidate for an anti-toxin protein to the putative toxin, although YitQ has no similarity to SdpI. Based on the DNA sequence, \textit{yizB} and \textit{yitQ} are predicted to form an operon with a downstream gene, \textit{yitR}, which also encodes a membrane protein. Although \textit{yitR} is present in BEST195, we kept it as a second candidate for the anti-toxin protein. To test whether these genes encoded anti-YIT toxin, we attempted to disrupt these genes. However, we were concerned that deleting these candidate anti-toxin genes might cause severe growth defects by releasing the activity of the toxin encoded by the genomic \textit{yitPOM} operon. Therefore, we constructed a \textit{ΔyitR-yitM} deletion strain that lacks the entire region from \textit{yitR} to \textit{yitM}, which contains the candidate anti-toxin genes \textit{yitR} and \textit{yitQ}, the unknown repressor gene \textit{yizB}, and the putative toxin-encoding \textit{yitPOM} operon (Fig 2C). Furthermore, since the expression of the secondary resistance mechanism against the SDP toxin is induced by the alternative sigma factor SigW (\textit{σ\textsubscript{W}}) \cite{51}, we also constructed a \textit{sigW} deletion strain. Subsequently, either the \textit{ΔyitR-yitM} and \textit{ΔsigW} mutation alone or both mutations together were introduced into the \textit{P\textsubscript{spac-hy}-yitPOM} strain. We compared the growth of these strains in 2\times SG medium supplemented with 1 mM IPTG. While the \textit{P\textsubscript{spac-hy}-yitPOM} \textit{ΔyitR-yitM} and \textit{P\textsubscript{spac-hy}-yitPOM} \textit{ΔsigW} mutants grew normally, the \textit{P\textsubscript{spac-hy}-yitPOM} \textit{ΔyitR-yitM} \textit{ΔsigW} mutant showed mild cell lysis 3 h after the end of exponential phase (Fig 2A). We confirmed that \textit{yitPOM} expression caused this cell lysis, as cell lysis was only
observed when yitPOM expression was induced with IPTG (Fig 2B). These results indicate that yitPOM expression leads to the production of a toxin that causes cell lysis in a mutant.
strain lacking the putative anti-toxin genes and \( \sigma^W \). The \( \sigma^W \)-regulated genes include multiple antibiotic resistant genes, such as a peptide exporter, an SdpI homolog, fosfomycin resistance proteins, sublancin resistance proteins, and a penicillin-binding protein [51–54]. Some of these genes might contribute to resistance to the putative toxin produced by \( \textit{yitPOM} \).

To further confirm that \( \textit{yitPOM} \) encodes a toxin, we employed a spot-on-lawn assay. We performed this assay in the \( \Delta \text{sdpABC-sdpIR} \) (hereafter referred to as \( \Delta \text{sdpA-sdpR} \) \( \Delta \text{yitR-yitM} \) \( \Delta \sigma^W \)) strain was grown in 2×SG supplemented with or without 1 mM IPTG. (C) Comparison of the genetic organization in NCIB3610 and BEST195. Homologous genes are shown by boxes of the same color. Genes only present in NCIB3610 are shown in red bold. The deleted regions in the \( \Delta \text{yitR-yitM} \) and \( \Delta \text{nprB-yitM} \) mutants are shown below the gene map of NCIB3610.

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Fig 2. \( \textit{yitPOM} \) encodes a toxin. (A) Effect of \( \textit{yitPOM} \) induction on cell growth. \( \textit{B. subtilis} \) strains were grown at 37°C in 2×SG supplemented with 1 mM IPTG with vigorous shaking. Growth profiles were examined at least three time, and the typical examples were shown. (B) Induction of \( \textit{yitPOM} \) caused mild cell lysis. The \( \textit{P}_{\text{spac-hy}} \)-\( \textit{yitPOM} \) \( \Delta \text{yitR-yitM} \) \( \Delta \sigma^W \) strain was grown in 2×SG supplemented with or without 1 mM IPTG. (C) Comparison of the genetic organization in NCIB3610 and BEST195. Homologous genes are shown by boxes of the same color. Genes only present in NCIB3610 are shown in red bold. The deleted regions in the \( \Delta \text{yitR-yitM} \) and \( \Delta \text{nprB-yitM} \) mutants are shown below the gene map of NCIB3610.

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same lawn. These results demonstrate that, like sdpABC, yitPOM encodes a secreted toxin, which we named YIT.

**YitQ is an anti-toxin protein to the YIT toxin**

To determine whether YitQ is an anti-toxin protein to the YIT toxin, we examined the effect of yitQ overexpression on the YIT toxin activity. To this end, the Pspac-hy-yitQ construct was introduced into the amyE locus of the indicator strain (i.e., the ΔsdpA-sdpR ΔyitR-yitM ΔsigW mutant). When spotted on a lawn of the indicator strain expressing YitQ (Pspac-hy-yitQ ΔsdpA-sdpR ΔyitR-yitM ΔsigW), the strain expressing YitPOM (Pspac-hy-yitPOM ΔsdpA-sdpR ΔyitR-yitM) only formed weak halos around its colonies (Fig 3). Thus, yitQ expression conferred resistance to the YIT toxin.

We were interested in whether there is crosstalk between yitPOM/yitQ and sdpABC/sdpi. To explore this possibility, a strain expressing SdpABC (Pspac-hy-sdpABC ΔsdpA-sdpR ΔyitR-yitM) was spotted onto lawns of the control indicator strain (ΔsdpA-sdpR ΔyitR-yitM Δspo0A) and the indicator strain expressing YitQ (Pspac-hy-yitQ ΔsdpA-sdpR ΔyitR-yitM ΔsigW) (Fig 3). The SdpABC-expressing strain formed clear halos around its colonies on both types of lawns. Thus, yitQ expression did not confer resistance to the SDP toxin. We also tested whether Sdpi expression confers resistance to the YIT and SDP toxins. A strain expressing YitPOM (Pspac-hy-yitPOM ΔsdpA-sdpR ΔyitR-yitM) formed halos around its colonies on a lawn of the indicator strain expressing Sdpi (Pspac-hy-sdpi ΔsdpA-sdpR ΔyitR-yitM ΔsigW), whereas the strain expressing SdpABC (Pspac-hy-sdpABC ΔsdpA-sdpR ΔyitR-yitM) did not. These results indicate that YitQ and Sdpi are anti-toxin proteins specific to the YIT and SDP toxins, respectively.
Thus, the two toxin/anti-toxin gene pairs yitPOM/yitQ and sdpABC/sdpI most likely function independently.

Expression of yitPOM and yitQ

The yitPOM operon was previously identified as a member of the DegS-DegU-regulated genes via a DNA microarray analysis using another B. subtilis strain, ATCC6051 [47]. To confirm this property in strain 3610, we carried out a Northern blot analysis. RNA samples were isolated from wild-type and ΔdegU mutant cells grown for various lengths of time in 2×SG with vigorous shaking (Fig 4A). We detected a single band at a position between the 23S rRNA (2904 nt) and 16S rRNA (1541 nt) on Northern blots with a yitP-specific probe (Fig 4B). The size of the band was consistent with the length of the entire yitPOM locus (2031 bp), confirming that the yitPOM locus is transcribed as an operon (Fig 4C). On the Northern blots, the yitPOM transcript was observed in the stationary phase samples from the wild-type strain but not in those from the ΔdegU mutant (Fig 4B). These results indicate that DegS-DegU directly or indirectly induces yitPOM transcription in stationary phase.

yitQ is predicted to form an operon with its upstream and downstream genes, yizB and yitR. We detected a band below the position of 16S rRNA on Northern blots with a yizB-specific probe (Fig 3B). The size of the band was consistent with the length of the yizB-yitQ-yitR locus (1244 bp), supporting the conclusion that yizB, yitQ, and yitR are transcribed as an operon (Fig 4C). Based on the Northern blots, the yizB-yitQ-yitR operon was transcribed at low levels during exponential phase and then induced during stationary phase in the wild-type

Fig 4. Transcription of yitPOM and yitQ in the ΔdegU mutant. (A) Growth profiles of the wild-type and ΔdegU mutant strains. Strains were grown in 2×SG with vigorous shaking. Arrows indicate the time points at which samples were taken for RNA isolation. (B) Northern blot analysis of yitPOM, yizB-yitQ-yitR, and nprB. Transcripts were detected with genespecific DIG-labeled RNA probes. Lane numbers (time points) under the strain names correspond to the time points shown in panel A. rRNA stained with methylene blue is shown as a loading control. The positions of 23S rRNA and 16S rRNA are indicated by arrows. (C) The transcription map of the yitPOM region. The transcripts are represented as lines with arrows below the gene map, and their estimated lengths are indicated.

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The ΔdegU mutation had no significant effect on the transcription of the yizB-yitQ-yitR operon. The SDP toxin mediates cannibalism between “Spo0A ON” and “Spo0A OFF” cells [40]. The finding that DegS-DegU regulates yitPOM but not yitQ rules out the possibility that the YIT toxin mediates cannibalism between “DegU ON” and “DegU OFF” cells.

To explore the function of the YIT toxin, we asked under what conditions yitPOM expression is induced. The DegS-DegU-regulated gene bpr, which encodes an extracellular protease, is expressed in biofilms [55]. We therefore speculated that yitPOM is also expressed in biofilms. To visualize yitPOM expression in biofilms, the yitP promoter was fused to the green fluorescent protein (GFP) reporter, and the resulting PyitP-gfp reporter construct was introduced to the amyE locus on the chromosome of the wild-type strain. B. subtilis biofilms are wrinkled structures on the surfaces of colonies grown on solid media that support biofilm formation, such as 2×SG [24]; therefore, we attempted to examine the expression level of the PyitP-gfp reporter in colonies grown on 2×SG solid medium. However, we did not detect any fluorescent GFP signal on these colonies, probably because the yitP promoter activity was too low to detect signals in our microscopy. We next examined the expression of the PyitP-gfp reporter inserted into the multi-copy plasmid pHYG2. The expression of the PyitP-gfp reporter on the plasmid was examined at 37°C because the plasmid pHYG2-yitP negatively affected biofilm formation at 30°C. The wild-type strain carrying the multi-copy plasmid pHYG2-yitP (PyitP-gfp) formed wrinkled structures on the surfaces of colonies grown on 2×SG solid medium as the biofilms developed. Weak green GFP signal was observed in these wrinkles with a color digital camera (Fig 5A). By contrast, this strain formed flat colonies on LB medium, which does not support biofilm formation, and produced no detectable green GFP signal. Under the same conditions, the wild-type strain carrying the parental plasmid pHYG2 (promoterless gfp) produced no green fluorescent signal on either 2×SG or LB.

The expression of PyitP-gfp in these colonies was also analyzed at the single-cell level. Flow cytometry analysis revealed that the expression of PyitP-gfp was heterogeneous in the population in colonies grown on 2×SG and that the considerable portion of PyitP-gfp cells exhibited stronger fluorescence in 2×SG than in LB (Fig 5B). Combined with the microscopic observation, these results indicate that biofilms contain cells that highly express the yitPOM operon. Moreover, we confirmed that the expression of PyitP-gfp in these colonies was DegU-dependent as the level of fluorescence decreased to the background level in the ΔdegU mutant (Fig 5B).

To eliminate potential artifacts resulting from multi-copy plasmid-based experiments, we examined the expression of aprE, which is one of the most highly expressed genes among the DegS-DegU-regulated genes [47]. We used a strain carrying a single copy of the PaprE-gfp reporter inserted into the amyE locus on the chromosome. The PaprE-gfp strain produced GFP fluorescent signal in the wrinkles of colonies grown on 2×SG (Fig 5A). By contrast, no detectable GFP signal was observed when the PaprE-gfp strain was grown on LB. Flow cytometry analysis revealed that the considerable portion of PaprE-gfp cells exhibited stronger fluorescence in 2×SG than in LB (Fig 5B). Thus, the expression profiles of PaprE-gfp were quite similar to those of PyitP-gfp. These results indicate that DegS-DegU strongly induces its regulatory target genes, including yitPOM, in biofilms and that the YIT toxin may play a role in biofilms.

The YIT toxin inhibits colony biofilm formation

We hypothesized that if the YIT toxin has an effect on biofilm formation, we expected to detect this effect on biofilm supporting media but not on the media that do not support biofilm formation. First, we examined whether yitPOM overexpression from the spac-hy promoter affects colony biofilm formation on biofilm-supporting media. We used two biofilm-supporting
Fig 5. *yitPOM* expression is induced in biofilms by DegS-DegU. (A) Expression of *yitPOM* and *nprB* in biofilms. The wild-type strain 3610 carrying the multi-copy plasmid pHYG2 (promoterless *gfp*), pHYG2-*yitP* containing the *P_{yitP}-gfp* reporter, or the single copy of *aprE-gfp* was grown at 37˚C for 24 h on 2×SG or LB. GFP fluorescence was analyzed with a digital color camera. GFP fluorescence was observed as green light signals. Some excitation light reflections on the surfaces of colonies and media were observed as blue light signals on GFP images. Strains 3610 pHYG2 (promoterless *gfp*) and 3610 were used as negative controls. Scale bar, 1 mm. (B) Flow cytometry analysis of *gfp* reporter strains. B. *subtilis* strains were grown at 37˚C for 24 h on 2×SG or LB. Expression of *gfp* reporters in the colonies was analyzed using stains 3610 pHYG2 and 3610 as negative controls.

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media, the rich complex medium 2×SG and the synthetic medium MSgg [24]. On 2×SG solid medium, the wild-type strain formed whitish wrinkled colonies (Fig 6A). Induction of yitPOM did not affect the colony morphologies of the wild-type, ΔyitR-yitM mutant, or ΔsigW mutant strains; these Pspac-hy-yitPOM strains formed similar whitish wrinkled colonies in the presence or absence of IPTG. By contrast, yitPOM induction altered the colony morphology of the ΔyitR-yitM ΔsigW mutant; the Pspac-hy-yitPOM ΔyitR-yitM ΔsigW mutant formed brown flat colonies in the presence of IPTG (Fig 6A). Magnified images showed that the whitish wrinkled layers (biofilms) were completely absent on the surfaces of the Pspac-hy-yitPOM ΔyitR-yitM ΔsigW mutant colonies in the presence of IPTG (Fig 6B). Similar results were obtained on MSgg medium. The wild-type strain formed light brown wrinkled colonies on MSgg (Fig 6A). Induction of yitPOM altered the colony morphology of the ΔyitR-yitM ΔsigW mutant. The Pspac-hy-yitPOM ΔyitR-yitM ΔsigW mutant formed brown flat colonies in the presence of IPTG (Fig 6A), and these colonies completely lacked the light brown wrinkled layers (biofilms) on their surfaces (Fig 6B). Unlike on rich 2×SG medium, induction of yitPOM also altered the colony morphology of the ΔyitR-yitM mutant when it was grown on MSgg (Fig 6A). In the presence of IPTG, the Pspac-hy-yitPOM ΔyitR-yitM mutant formed colonies covered with attenuated wrinkles at 96 h post-inoculation; however, these wrinkles faded over time (S1 Fig). This phenotype suggests that YitQ may play a major role in resistance to the YIT toxin under low nutrient conditions, such as B. subtilis natural habitats, soils. We next examined the colony morphologies on the complex medium LB and on the synthetic medium Spizizen minimal medium (SMM) [56]. B. subtilis forms flat colonies rather than biofilms on these media. On these media, yitPOM induction had little or no effect on colony morphology, even in the ΔyitR-yitM ΔsigW mutant (Fig 6A). We compared the effect of yitPOM overexpression on colony morphology with that of sdpABC overexpression. For this, sdpABC was expressed under the control of the same promoter (spac-hy) in the ΔsdpA-sdpR ΔsigW mutant. The Pspac-hy-sdpABC ΔsdpA-sdpR ΔsigW mutant formed normal colonies in all four media in the absence of IPTG, but it did not form colonies in the presence of IPTG (Fig 6A). Thus, the SDP toxin inhibited overall cell growth independently of the medium conditions.

We investigated the relationship between the expression levels of yitPOM and sdpABC and colony morphology. For this purpose, Pspac-hy-yitPOM ΔyitR-yitM ΔsigW and Pspac-hy-sdpABC ΔsdpA-sdpR ΔsigW mutants were grown on 2×SG medium supplemented with various IPTG concentrations (0 to 1000 μM) (Fig 6C). The effect of yitPOM expression on colony morphology appeared when the Pspac-hy-yitPOM ΔyitR-yitM ΔsigW mutant was grown in the presence of 30 μM or higher IPTG concentrations. At 30 μM IPTG, attenuated wrinkles appeared on the colonies at 24 h post-inoculation; however, these wrinkles failed to grow further. At 100 μM IPTG, the colonies completely lacked the whitish wrinkled layers on their surface and became flat. Higher IPTG concentrations did not further alter the colony morphology. Despite having an obvious effect on colony morphology, yitPOM induction did not affect colony size. By contrast, when grown on 2×SG medium supplemented with various IPTG concentrations, the Pspac-hy-sdpABC ΔsdpA-sdpR ΔsigW mutant formed small colonies in the presence of 10 or 30 μM IPTG but did not form colonies at 100 μM or higher IPTG concentrations (Fig 6C). Thus, sdpABC expression exerted a stronger effect on colony formation as its expression levels increased. These results demonstrate that the YIT and SDP toxins have different effects on colony growth and that the YIT toxin specifically inhibits biofilm formation in the absence of its resistance genes.

We considered how the YIT toxin inhibits biofilm formation. As described above, yitPOM induction caused mild cell lysis only in ΔyitR-yitM ΔsigW mutant cells grown in 2×SG medium with shaking. Induction of yitPOM in the ΔyitR-yitM ΔsigW mutant did not cause cell lysis in cultures grown in LB medium with shaking (S2 Fig). Thus, yitPOM induction
Fig 6. Expression of yitPOM inhibits biofilm formation. (A) $P_{\text{spac-hy}}$-yitPOM strains with the indicated mutations were grown at 30°C for 48 h on biofilm formation media ($2\times$SG and MSgg) and non-biofilm formation media (LB and SMM) with or without 1000 μM IPTG. Colonies of the wild-type and $P_{\text{spac-hy}}$-sdpABC ΔsdpA-sdpR ΔsigW strains are also shown as references. (B) Magnified images of the $P_{\text{spac-hy}}$-yitPOM ΔyitR-yitM ΔsigW mutant colonies shown in panel A. (C) Comparison of the effects of yitPOM and sdpABC overexpression on colony morphology. The $P_{\text{spac-hy}}$-yitPOM ΔyitR-yitM ΔsigW and $P_{\text{spac-hy}}$-sdpABC ΔsdpA-sdpR ΔsigW mutant strains were grown at 30°C for 48 h on $2\times$SG with various IPTG concentrations. Colony morphology analysis was done at least three time, and the typical examples were shown in the figure. Scale bar, 2 mm.
caused cell lysis and inhibition of biofilm formation only in the ΔyatR-yitM ΔsΔgW mutant cells grown on biofilm formation media, indicating that these two phenotypes represent different aspects of one phenomenon. Moreover, induction of yitPOM led to formation of halos in the spot-on-lawn assays. We propose that the YIT toxin likely inhibits biofilm formation by killing biofilm-forming cells rather than by preventing expression of biofilm formation genes.

**NprB allows the YIT toxin to attack cells within biofilms**

Induction of yitPOM exerted its effects only in cells grown on biofilm formation media. However, the spac-hy promoter is active in rich and poor media, including LB and SMM, as observed for the Pspac-hy-sdpABC ΔsdpA-sdpR ΔsΔgW mutant (Fig 6A). These observations suggest the involvement of other factor(s) in the functions of the YIT toxin. A comparison of the genetic organization of the 3610 and BEST195 strains revealed that, in addition to yizB and yitQ, nprB appears to be inserted into the 3610 genome along with yitPOM (Fig 2C). Like yitPOM, nprB, which encodes an extracellular neutral protease, was transcribed in a DegU-dependent manner (Fig 4B), and its expression was induced in biofilms (Fig 5A and 5B). To determine whether nprB is involved in the YIT toxin function, we introduced a deletion of the nprB-yitM region (Fig 2C) into the Pspac-hy-yitPOM ΔsΔgW mutant and examined the colony morphology of the resulting strain. Unlike in the Pspac-hy-yitPOM ΔyatR-yitM ΔsΔgW mutant, yitPOM induction did not inhibit biofilm formation in the Pspac-hy-yitPOM ΔnprB-yitM ΔsΔgW mutant. This strain formed whitish wrinkled colonies like those of the wild-type strain in the presence or absence of IPTG (Fig 7A). Because nprB deletion was the only genetic difference between Pspac-hy-yitPOM ΔyatR-yitM ΔsΔgW and Pspac-hy-yitPOM ΔnprB-yitM ΔsΔgW mutants (Fig 2C), this result suggests that the NprB protease is required for the production or function of the YIT toxin.

To distinguish these possibilities, we examined the production of the YIT toxin in these mutants via spot-on-lawn assays. The Pspac-hy-yitPOM ΔyatR-yitM ΔsΔgW and Pspac-hy-yitPOM ΔnprB-yitM ΔsΔgW mutants were spotted on the lawn of the ΔyatR-yitM ΔsΔgW mutant. Although both mutants formed halos around their colonies in the presence of IPTG, the Pspac-hy-yitPOM ΔnprB-yitM ΔsΔgW mutant formed clearer halos than did the Pspac-hy-yitPOM ΔyatR-yitM ΔsΔgW mutant (Fig 7B). The Pspac-hy-yitPOM ΔyatR-yitM ΔsΔgW mutant formed smaller colonies on the lawn than did the Pspac-hy-yitPOM ΔnprB-yitM ΔsΔgW mutant, likely due to loss of biofilm formation. Therefore, we compared the YIT toxin production between the Pspac-hy-yitPOM ΔyatR-yitM and Pspac-hy-yitPOM ΔnprB-yitM mutants. Although these mutants formed similar colonies on the lawn of the ΔyatR-yitM ΔsΔgW mutant, the Pspac-hy-yitPOM ΔnprB-yitM mutant formed clearer halos than did the Pspac-hy-yitPOM ΔyatR-yitM mutant (S3 Fig). Thus, the ΔnprB mutation increased YIT toxin production or activity. These results suggest that NprB is not required for YIT toxin production. Given that NprB is an extracellular neutral protease, these results suggest that the YIT toxin is probably a substrate for NprB.

We next examined the alternative possibility that NprB might be required for the function of the YIT toxin. To test this idea, we spotted the Pspac-hy-yitPOM ΔyatR-yitM ΔsΔgW and Pspac-hy-yitPOM ΔnprB-yitM ΔsΔgW mutants on the lawn of the ΔnprB-yitM ΔsΔgW mutant. Both mutants failed to form clear halos around their colonies (Fig 7B), supporting this idea.

We explored why the ΔnprB mutation impaired the function of the YIT toxin. Cells in biofilms are covered with and protected by biofilm matrix polymers, a key reason why cells in biofilms exhibit increased antibiotic tolerance or resistance [5, 6]. We hypothesized that a similar mechanism might work against the YIT toxin and that the NprB protease might enable the YIT toxin molecules to pass through the layers of the biofilm matrix polymers to attack cells
within the biofilms. If this were true, then disrupting the biofilm matrix would enable the YIT toxin to inhibit biofilm formation even in the ΔnprB mutant. The biofilm matrix of *B. subtilis* biofilms mainly consists of exopolysaccharides (synthesized by the products of the eps operon) and polymers of the TasA (produced by the tapA-tasA operon) and BslA proteins [24–29].
test our hypothesis, we introduced Δeps, ΔtapA-tasA, and ΔbslA deletion mutations into the P_{spac-hy}-yitPOM ΔyitR-yitM ΔsigW and P_{spac-hy}-yitPOM ΔnprB-yitM ΔsigW mutants and examined their colony morphologies. The P_{spac-hy}-yitPOM ΔyitR-yitM ΔsigW Δeps mutant formed whitish mucoid colonies in the absence of IPTG, while it formed flat brown colonies in the presence of IPTG (Fig 7C). The difference in colony morphology depending on the presence or absence of IPTG indicates that the induced YIT toxin can function in these colonies even though the Δeps mutation impaired biofilm formation and led to the formation of mucoid colonies. Unlike the P_{spac-hy}-yitPOM ΔnprB-yitM ΔsigW mutant, the P_{spac-hy}-yitPOM ΔnprB-yitM ΔsigW Δeps mutant also formed whitish mucoid colonies in the absence of IPTG and flat brown colonies in the presence of IPTG (Fig 7C). Thus, the ΔnprB mutation did not interfere with the function of the YIT toxin in the Δeps mutant. Similar results were obtained with the ΔbslA mutant. The P_{spac-hy}-yitPOM ΔyitR-yitM ΔsigW ΔbslA and P_{spac-hy}-yitPOM ΔnprB-yitM ΔsigW ΔbslA mutants formed whitish mucoid colonies in the absence of IPTG and flat brown colonies in the presence of IPTG (Fig 7C). These results demonstrate that the Δeps and ΔbslA mutations eliminate the requirement for NprB in the function of the YIT toxin. Thus, our idea that NprB enables the YIT toxin to pass through the layers of the biofilm matrix polymers to attack cells in the biofilm is very likely. On the other hand, the ΔtapA-tasA mutation did not restore the YIT toxin activity in the ΔnprB-yitM mutant. The P_{spac-hy}-yitPOM ΔnprB-yitM ΔsigW ΔtapA-tasA mutants formed whitish mucoid colonies in the presence or absence of IPTG (Fig 7C). These results indicate that exopolysaccharides, BslA polymers or molecules associated with these polymers probably trap the YIT toxin in ΔnprB mutant biofilms.

We further examined the effect of overexpression of biofilm matrix polymers on the YIT toxin activity. SinR is a major repressor of the biofilm matrix synthesis genes [32], and a ΔsinR mutant formed large swollen colonies due to overproduction of biofilm matrix polymers (Fig 7D). We introduced the ΔsinR mutation into the P_{spac-hy}-yitPOM ΔyitR-yitM ΔsigW mutant and examined the colony morphology of the resulting strain. In the absence of IPTG, the P_{spac-hy}-yitPOM ΔyitR-yitM ΔsigW ΔsinR mutant formed large swollen colonies, like those of the ΔsinR mutant. Induction of yitPOM inhibited biofilm formation in this mutant. The P_{spac-hy}-yitPOM ΔyitR-yitM ΔsigW ΔsinR mutant formed flat brown colonies in the presence of 100 or 1000 μM IPTG (Fig 7D), as was also observed in the P_{spac-hy}-yitPOM ΔyitR-yitM ΔsigW mutant (Fig 6C). We also examined the effect of the ΔsinR mutation on the SDP toxin. The P_{spac-hy}-sdpABC ΔsdpA-R ΔsigW ΔsinR mutant formed large swollen colonies in the absence of IPTG. Induction of sdpABC did not inhibit colony formation and only partly suppressed the swollen colony phenotype even in the presence of 1000 μM IPTG (Fig 7D). These results indicate that overproduction of biofilm matrix polymers interferes with the activity of the SDP toxin but not with that of the YIT toxin. Based on these results, we conclude that the YIT toxin can function within mature biofilms with the assistance of the extracellular neutral protease NprB.

We examined the colony morphology of the ΔnprB mutant. The ΔnprB mutant formed wrinkled colonies on 2xSG medium similar to those of the wild-type strain (Fig 8A). We extracted the extracellular proteins and cell surface-associated proteins from these colony biofilms and analyzed them via SDS-PAGE. We detected little or no difference in the protein composition between the wild-type and ΔnprB mutant strains in the gels after Coomassie brilliant blue (CBB) staining (Fig 8B). These results indicate that, despite a clear effect on the function of the YIT toxin, the ΔnprB mutation does not significantly alter biofilm structure.

**The YIT toxin is present in biofilms of the wild-type strain**

So far, we have reported the results of experiments designed to uncover the function of the YIT toxin via yitPOM expression from the strong spac-hy promoter. We asked whether
yitPOM expression from its own promoter is sufficiently high to exhibit the phenotypes observed above. As described above, the action of the YIT toxin was antagonized by YitQ and unidentified $\sigma^W$-regulated gene product(s) in the wild-type strain. If the YIT toxin is present in biofilms of the wild-type strain, its effect should appear in $\Delta{yitQ}$ and $\Delta{sigW}$ mutants. Therefore, we examined the colony morphologies of mutants lacking $yitQ$ and/or $sigW$ on 2×SG medium (Fig 9A). While the $\Delta{yitQ}$ and $\Delta{sigW}$ single mutants formed whitish wrinkled colonies like those of the wild-type strain, the $\Delta{yitQ}\Delta{sigW}$ double mutant formed colonies with attenuated wrinkles (biofilms). The $\Delta{yitR}\Delta{yitM}\Delta{sigW}$ mutant, which lacks both the toxin and anti-toxin genes, formed whitish wrinkled colonies like those of the wild-type strain. Thus, the phenotype of the $\Delta{yitQ}\Delta{sigW}$ mutant was caused by the YIT toxin. However, the phenotype of the $\Delta{yitQ}\Delta{sigW}$ mutant was slightly less noticeable than that of the $P_{spac-hy} yitPOM \Delta{yitR}\Delta{yitM}\Delta{sigW}$ mutant in the presence of IPTG. Likewise, when grown in shaking culture, the $\Delta{yitQ}\Delta{sigW}$ mutant did not display the culture lysis phenotype as observed for the $P_{spac-hy} yitPOM \Delta{yitR}\Delta{yitM}\Delta{sigW}$ mutant; instead, the $\Delta{yitQ}\Delta{sigW}$ mutant reached slightly lower OD<sub>600</sub> in the stationary phase than that of the wild-type strain (Fig 9B).
As mentioned above, the P<sub>aprE</sub>-gfp reporter is induced in biofilms by DegS-DegU. The wild-type strain and the ΔyitR-yitMΔsigW mutant with the P<sub>aprE</sub>-gfp reporter displayed bright GFP signal at wrinkles on colonies. However, the ΔyitQΔsigW mutant with the P<sub>aprE</sub>-gfp reporter displayed no detectable GFP signal on its colonies with attenuated wrinkles (Fig 9C). Thus, the ΔyitQΔsigW mutant showed decreased biofilm formation and decreased aprE expression. These results indicate that the YIT toxin reduces the number of biofilm-forming cells that express DegS-DegU-regulated genes, including yitPOM, in the ΔyitQΔsigW mutant. In other words, the action of the YIT toxin reduces the number of cells producing the YIT toxin in the ΔyitQΔsigW mutant. This effect can explain why the phenotype of the ΔyitQΔsigW mutant was slightly less obvious. These results demonstrate that the YIT toxin is present within biofilms of the wild-type strain.

**The YIT toxin can mediate intercellular competition within biofilms**

We hypothesized that the YIT toxin might mediate intercellular competition within biofilms. To test this hypothesis, we designed the following experiment. Dilutions of cultures of the wild-type and ΔyitR-yitMΔsigW mutant strains were mixed, and the mixtures were spotted on...
2×SG solid medium. The inoculated cells grew and formed biofilms in which the wild-type cells were expected to produce the YIT toxin. The YIT toxin then exerted its effect in biofilms with the assistance of NprB, which expressed in wild-type and ΔyitR-yitM ΔsigW mutant biofilms. If the YIT toxin suppressed the growth of ΔyitR-yitM ΔsigW mutant cells within biofilms, the ratio of these strains within the biofilms would change from the initial ratio. To estimate the ratio of two strains within biofilms, the P_{aprE-gfp} reporter was introduced into one strain to detect its cells within biofilms.

First, wild-type cells carrying the P_{aprE-gfp} reporter (the P_{aprE-gfp} strain) were mixed with wild-type cells at various ratios from 10:0 to 0:10, and the mixtures were spotted on 2×SG solid medium. After 2 days of incubation, we observed the colonies with a fluorescence stereomicroscope. A bright GFP fluorescent signal was detected on colonies grown from the 10:0 mixture, and the GFP signal decreased as the proportion of the P_{aprE-gfp} strain decreased (Fig 10A). When P_{aprE-gfp}-expressing cells were mixed with ΔyitR-yitM ΔsigW mutant cells and grown on 2×SG medium, the GFP signal on the colonies also decreased as the proportion of the P_{aprE-gfp} strain decreased; however, its decrease was moderate compared with that in the former experiment. For example, at a ratio of 3:7, obvious GFP signal was observed on colonies grown from the mixture of the aprE-gfp and ΔyitR-yitM ΔsigW mutant strains but not on colonies grown from the mixture of the aprE-gfp and wild-type strains. We also introduced the P_{aprE-gfp} reporter into the ΔyitR-yitM ΔsigW mutant. When ΔyitR-yitM ΔsigW P_{aprE-gfp} mutant cells were mixed with wild-type cells, no GFP signal was observed, even on the colonies grown from the 9:1 mixture (Fig 10A). By contrast, when ΔyitR-yitM ΔsigW P_{aprE-gfp} mutant cells were mixed with ΔyitR-yitM ΔsigW mutant cells, the GFP signal on the colonies decreased as the proportion of the ΔyitR-yitM ΔsigW P_{aprE-gfp} mutant cells decreased, as was also the case with the mixture of the aprE-gfp and wild-type strains. Thus, the YIT toxin is likely to mediate intercellular competition within biofilms.

To confirm these results, we analyzed the population of cells carrying the aprE-gfp reporter in those colonies over time by determining colony forming units. In this experiment, aprE-gfp reporter strains were mixed with competitors at the ratio of 1:1. In the colonies of aprE-gfp and wild-type cells, the population ratio of the aprE-gfp cells did not change drastically from the initial ratio by 72 h after inoculation (Fig 10B). By contrast, in the colonies of P_{aprE-gfp} and ΔyitR-yitM ΔsigW cells, a drastic increase in the population ratio of the aprE-gfp cells was observed 24 h after inoculation, and its ratio increased to 93% 72 h after inoculation. In colonies of P_{aprE-gfp} ΔyitR-yitM ΔsigW and wild-type cells, the population ratio of the P_{aprE-gfp} ΔyitR-yitM ΔsigW cells decreased to 4% by 72 h after inoculation. However, such a decrease was not observed in colonies of P_{aprE-gfp} ΔyitR-yitM ΔsigW and ΔyitR-yitM ΔsigW cells. These results demonstrate that the YIT toxin can mediate intercellular competition within biofilms. In conclusion, we propose that the YIT toxin functions within biofilms from YIT toxin-sensitive unfavorable competitors.

**The variety of SDP toxin homologs**

Genome comparison revealed that sdpABC homologs are widely conserved among _B. subtilis_ strains (S1 Table). sdpABC homologs, including sdpABC itself and yitPOM, can be classified into five groups based on their genome positions and SdpC homolog sequences (S1 Table, S4 and S5 Figs). Many _B. subtilis_ strains have one or two sdpABC homologs, and yitPOM appears to be more widely conserved among _B. subtilis_ strains than the other sdpABC homologs (S1 Table). Homologous genes encoding membrane proteins are found downstream of sdpABC homologs 2 and 4 (S4 Fig). The sdpIR homolog are found upstream of the sdpABC homolog 3.
### A

| Competitor | WT | Δ\(\text{yitR-yitM}\) Δ\(\text{sigW}\) | WT | Δ\(\text{yitR-yitM}\) Δ\(\text{sigW}\) |
|------------|----|--------------------------------|----|--------------------------------|
| 10:0       | ![Image](image1) | ![Image](image2) | ![Image](image3) | ![Image](image4) |
| 9:1        | ![Image](image5) | ![Image](image6) | ![Image](image7) | ![Image](image8) |
| 8:2        | ![Image](image9) | ![Image](image10) | ![Image](image11) | ![Image](image12) |
| 7:3        | ![Image](image13) | ![Image](image14) | ![Image](image15) | ![Image](image16) |
| 6:4        | ![Image](image17) | ![Image](image18) | ![Image](image19) | ![Image](image20) |
| 5:5        | ![Image](image21) | ![Image](image22) | ![Image](image23) | ![Image](image24) |
| 4:6        | ![Image](image25) | ![Image](image26) | ![Image](image27) | ![Image](image28) |
| 3:7        | ![Image](image29) | ![Image](image30) | ![Image](image31) | ![Image](image32) |
| 2:8        | ![Image](image33) | ![Image](image34) | ![Image](image35) | ![Image](image36) |
| 1:9        | ![Image](image37) | ![Image](image38) | ![Image](image39) | ![Image](image40) |
| 0:10       | ![Image](image41) | ![Image](image42) | ![Image](image43) | ![Image](image44) |

### B

**GFP strain in colonies (%)**

| Time after inoculation (h) | 0 | 24 | 48 | 72 |
|----------------------------|---|----|----|----|
| GFP strain                 | 0 | 50 | 50 | 50 |
| P\(_{\text{apre}}\)gfp    | 0 | 25 | 25 | 25 |
| Δ\(\text{yitR-yitM}\) Δ\(\text{sigW}\) P\(_{\text{apre}}\)gfp | 0 | 25 | 25 | 25 |
Some strains possess only *sdpRI* homologs in the *sdpABC* homolog 3 locus. These observations suggest functional similarities between the *sdpABC* homolog 2 and the *sdpABC* homolog 4 and between *sdpABC* and the *sdpABC* homolog 3. However, each group of SdpC homologs has a unique C-terminal hydrophobic domain, the sequence of which differs from those of the others (S5 Fig); therefore, each group of SdpC homolog-derived toxins may have different sequences. If the differences in these sequences impart their functional differences, as observed for SdpC and YitM, then SdpC homolog-derived toxins are likely to play more diverse roles.

**Discussion**

Biofilms often contain a high-density bacterial community consisting of a mixture of various species, and under these conditions, the bacteria compete with their own siblings and other species for limited space and nutrients. Since cells in biofilms exhibit increased antibiotic tolerance or resistance, the functions of antibiotics in the competition between biofilm bacteria remain unclear. In this study, we demonstrated that *B. subtilis* produces a biofilm-associated toxin, and that this toxin can attack toxin-sensitive cells within the biofilm by passing through the protective barriers of the biofilm with assistance from an endogenously produced extracellular protease.

The *yitPOM* operon is a paralog of the *sdpABC* operon; however, these operons are under different control and play distinct roles. Transcription of *sdpABC* is induced by low levels of Spo0A-P [44]. Given that low levels of Spo0A-P also induce the expression of biofilm matrix synthesis genes including the *eps* operon [44, 45], it is likely that the SDP toxin is induced during the early phase of biofilm formation. By contrast, *yitPOM* is induced by DegS-DegU in biofilms. Induction of these operons had different effects on *B. subtilis* cells lacking genes whose products confer resistance to the SDP and YIT toxins, i.e., *sdpABC* induction prevented colony formation independently of the medium conditions, whereas *yitPOM* specifically inhibited biofilm formation. Like SdpC, YitM contains a C-terminal hydrophobic domain that seems to be processed to produce the YIT toxin. The hydrophobic nature of the YIT toxin probably enables the YIT toxin to penetrate bacterial membrane, as observed for the SDP toxin. However, the sequence of the hydrophobic domain of YitM is different from that of SdpC. We assume that this difference contributes to the differences in the roles of these toxins. The genomes of many *B. subtilis* strains have both the *sdpABC* and *yitPOM* operons (S1 Table), suggesting that having both *sdpABC* and *yitPOM* may provide survival advantages in the environment.

Overproduction of biofilm matrix polymers interfered with the function of the SDP toxin but not with that of the YIT toxin, suggesting that the YIT toxin has a mechanism to pass through the layers of the biofilm matrix polymers, and we showed that this mechanism involves the extracellular neutral protease NprB. The YIT toxin could not inhibit biofilm formation in the absence of NprB even though the ΔnprB mutation increased the production of the YIT toxin. The requirement of NprB was eliminated by Δeps and ΔbslA mutations, either of which impairs biofilm matrix formation and, thus, biofilm formation [24, 28]. Cells in biofilms are encased in the biofilm matrix, which functions as a physical and chemical barrier...
against antibiotics by limiting their penetration [11–16]. Our results suggest that similar mechanisms might contribute to resistance to the YIT toxin in *B. subtilis* biofilms and that the extracellular protease NprB is required for the YIT toxin to pass through these defense barriers. The ΔnprB mutation had no significant effect on the composition of the extracellular and cell surface-associated proteins of biofilms nor on biofilm formation. We speculate that NprB may degrades the exopolysaccharides- or BslA polymer-associated protein that interacts with the YIT toxin, or that NprB may digest the YIT toxin smaller. By either or both actions, NprB may enable the YIT toxin molecules to pass through the layers of the biofilm matrix polymers. Moreover, we showed that the ΔnprB mutation increased activity of the YIT toxin, suggesting the possibility that the YIT toxin is a substrate for NprB protease. We speculate that NprB may also play a role in controlling YIT toxin levels in biofilms to avoid self-intoxication.

If NprB mediates structural changes in biofilms, that can increase the risk that biofilms become susceptible to antibiotics produced by other bacteria. However, the capability of the YIT toxin to attack sensitive cells within biofilms must be important for maintaining biofilm communities. Biofilms often consist of multiple types and multiple species of bacterial cells, some of which exploit others as free-loaders or cheaters that do not produce biofilm matrix polymers and other public goods [57]. The production of biofilm matrix polymers is metabolically costly; however, biofilm matrix polymers are extracellular products that are accessible even to non-producing cheater cells from which they receive protection [57]. An increase in the number of cheater cells, therefore, can disturb the cooperative relationships within biofilm communities and lead to instability within biofilm communities. The production of antibiotics that can diffuse through the biofilm offers the great advantage of being able to eliminate cheater cells and other unfavorable competitors present in the biofilm. Further work is required to determine which types of *B. subtilis* cells or what bacterial species are susceptible to the YIT toxin.

Previous studies showed that positively charged antibiotics interact with negatively charged matrix components, such as extracellular DNA and exopolysaccharides, and impede their penetration into biofilms [12, 15]. The SDP toxin contains two positively charged amino acid residues, whereas the hydrophobic region of YitM contains four negatively charged amino acid residues but no positively charged amino acid residues. These observations suggest that both NprB function and the amino acid sequence of the YIT toxin may be important for the ability of the YIT toxin to pass through the layers of biofilm matrix polymers; however, we have not yet determined the relevant sequence of the YIT toxin.

Bacterial competition is mediated by multiple factors [58]. In addition to the YIT toxin, DegS-DegU directly or indirectly induces non-ribosomally synthesized peptide antibiotics, e.g., bacilysin, fengycin, iturin, difficidin, and bacillomycin, although the repertoire of antibiotic synthesis genes differs from strain to strain and no *B. subtilis* strain produces all of these [59, 60, 61, 62]. DegS-DegU also induces a wide array of extracellular degradative enzymes, including six extracellular proteases [47 and references therein]. Although these proteases have been thought to play roles in nutrient acquisition from the surrounding environment, our results suggest that these degradative enzymes may also play roles in competition within biofilms. Indeed, several proteases were previously shown to disrupt biofilms of heterologous bacteria by degrading critical protein components in biofilms [63, 64, 65, 66]. Simultaneously producing multiple antibiotics and degradative enzymes within biofilms affords *B. subtilis* the ability to attack competitor cells protected by their own biofilm matrixes and to exclude them from biofilm communities. We expect that antibiotics and degradative enzymes cooperate extensively in *B. subtilis* biofilms.

The YIT toxin/NprB system seems to have evolved to be specifically adapted to the *B. subtilis* biofilm environment. Some bacteria produce specific antibiotics in biofilms [17–21].
Among them, the *Escherichia coli* ROAR029 strain produces the bacteriocin colicin R in biofilms, and colicin R is more active against biofilms than against planktonic cultures, as is the YIT toxin [20]. *Pseudomonas aeruginosa* produces bacteriocins pyocins and can suppress the growth of pyocin-sensitive bacteria in biofilms [21, 22]. Based on our results and these previous observations, we propose that bacteria may have evolved specialized antibiotics that function in biofilms as biofilm-specific competition mechanisms. The properties of these antibiotics may differ from those of conventional antibiotics. Biofilm-associated antibiotics might serve as anti-biofilm agents, especially in combination with degradative enzymes.

**Materials and methods**

**Bacterial strains and culture condition**

*B. subtilis* strain NCIB3610 and its derivatives used in this study are listed in Table 1. Construction of the *B. subtilis* mutants is described in S1 File. Primers used for the strain construction are listed in S2 Table. *B. subtilis* strains were maintained in LB (LB Lennox; BD Difco, Franklin Lakes, NJ, USA). For colony morphology observation, *B. subtilis* strains were grown at 30°C on LB plates overnight. A small single colony was suspended in 100 μl of LB, and 2 μl of the suspension was spotted onto 2×SG [49], MSgg [24], LB, and SMM media [55]. The plates were incubated at 30°C. Colony morphology was observed after 48 h of incubation on 2×SG and LB, after 72 h on SMM, and after 144 h on MSgg. Colony morphology observation was carried out at least three times and typical examples were shown in figures.

**Comparison of genetic organization and protein sequence analysis**

A comparison of the genetic organization in different *B. subtilis* strains was carried out using the MBGD website (http://mbgd.genome.ad.jp/) [67]. Protein alignments were constructed using the Protein BLAST program on the NCBI website (https://www.nlm.nih.gov/) and GENETYX ver.14 (GENETYX, Tokyo, Japan). Kyte & Doolittle hydropathy plots were constructed using the ExPASy website (https://web.expasy.org/protscale/) with a window size of 19 and default settings (hydropathy scale values of amino acids; A, 1.8; R, -4.5; N, -3.5; D, -3.5; C 2.5; Q, -3.5; G, -0.4; H, -3.2; I, 4.5; L, 3.8; K, -3.9; M, 1.9; F, 2.8; P, -1.6; S, -0.8; T, -0.7; W, -1.3; V, 4.2).

**Comparison of growth profiles**

*B. subtilis* strains grown at 30°C on LB plates overnight were inoculated into 5 ml of 2×SG and were grown at 37°C to the mid-exponential phase with vigorously shaking. These cultures were then added to 50 ml of warm 2×SG in a 500 ml baffled flask to give an OD₆₀₀ of 0.01. These cultures were shaking at 37°C, and OD₆₀₀ of these cultures was measured over time. The experiments were performed at least three times, and the typical results were shown in figures.

**Spot-on-lawn assay**

Indicator strains (lawn strains) were grown at 28°C overnight in LB with vigorous shaking. Culture (1 μl) was mixed with 12 ml of 50°C 2×SG 1.2% agar with brief vortexing, and the mixture was immediately poured into a 9 cm plate. The lawn plates were dried for 20 min in a laminar flow cabinet. The strains tested for antibiotic production were grown at 28°C overnight in LB with vigorous shaking. These cultures were diluted 10 times with LB, and 2 μl of the dilutions were spotted onto the dried lawn plates. The plates were incubated at 37°C for 24 to 30 h until halos appeared around the colonies. The experiments were performed at least three times, and the typical example was shown in figures.
Table 1. *B. subtilis* strains used in this study.

| Strain name | Genotypes | References or construction* |
|-------------|------------|-----------------------------|
| NCIB3610   | prototroph |                             |
| N1285      | amyE::P_{pacb}−yitPOM (erm) | This study |
| N1263      | ΔyitR−yitM::tet | This study |
| N1286      | amyE::P_{pacb}−yitPOM (erm) ΔyitR−yitM::tet | N1263 → N1285 |
| NTF88      | ΔsigW::cat | 67 |
| N1356      | amyE::P_{pacb}−yitPOM (erm) ΔsigW::cat | NTF88 → N1285 |
| N1357      | amyE::P_{pacb}−yitPOM (erm) ΔyitR−yitM::tet ΔsigW::cat | NTF88 → N1286 |
| N1333      | amyE::P_{pacb}−sdpABC (erm) | This study |
| N1458      | ΔsdpA−sdpR::spc | This study |
| N1335      | amyE::P_{pacb}−sdpABC (erm) ΔsdpA−sdpR::spc | N1458 → N1333 |
| N1337      | amyE::P_{pacb}−sdpABC (erm) ΔsigW::cat | NTF88 → N1333 |
| N1340      | amyE::P_{pacb}−sdpABC (erm) ΔsdpA−sdpR::spc ΔsigW::cat | N1458 → N1335 |
| N741       | ΔsdpA−sdpR::spc ΔyitR−M::tet | N1458 → N1263 |
| N764       | ΔsdpA−sdpR::spc ΔyitR−M::tet ΔsigW::cat | NTF88 → N741 |
| N1498      | amyE::P_{pacb}−yitQ (erm) ΔsdpA−sdpR::spc ΔyitR−M::tet ΔsigW::cat | amyE::P_{pacb}−yitQ (erm) → N764 |
| N1497      | amyE::P_{pacb}−sdpI (erm) ΔsdpA−sdpR::spc ΔyitR−M::tet ΔsigW::cat | amyE::P_{pacb}−sdpI (erm) → N764 |
| N942       | amyE::P_{pacb}−yitPOM (erm) ΔsdpA−sdpR::spc ΔyitR−yitM::tet | N1285 → N741 |
| N776       | amyE::P_{pacb}−sdpABC (erm) ΔsdpA−sdpR::spc ΔyitR−yitM::tet | N1333 → N741 |
| NTF28      | ΔdegU::cat | 62 |
| N1443      | pHYG2 (promoter-less gfp, tet) | pHYG2 → NCIB3610 |
| N1444      | pHYG2−yitP (P_{cat}−gfp, tet) | pHYG2−yitP → NCIB3610 |
| N1446      | pHYG2−nprB (P_{npr}−gfp, tet) | pHYG2−nprB → NCIB3610 |
| N345       | ΔdegU::kan | This study |
| N1515      | pHYG2−yitP (P_{cat}−gfp, tet) ΔdegU::kan | N345 → N1444 |
| N1516      | pHYG2−nprB (P_{npr}−gfp, tet) ΔdegU::kan | N345 → N1446 |
| N1382      | amyE::P_{apre}−gfp (cat) | W740 (amyE::P_{apre}−gfp (cat)) [68] → NCIB3610 |
| N1268      | ΔsigW::neo | This study |
| N355       | ΔepsA−O::spc | This study |
| N1230      | amyE::P_{pacb}−yitPOM (erm) ΔyitR−yitM::tet ΔsigW::neo | N1268 → N1286 |
| N1253      | amyE::P_{pacb}−yitPOM (erm) ΔyitR−yitM::tet ΔsigW::neo ΔepsA−O::spc | N355 → N1230 |
| N1500      | amyE::P_{pacb}−yitPOM (erm) ΔyitR−yitM::tet ΔsigW::neo ΔtapA−tasA::cat | N11 (ΔtapA−tasA::cat) [62] → N1230 |
| N1255      | amyE::P_{pacb}−yitPOM (erm) ΔyitR−yitM::tet ΔbslA::spc | N254 (ΔbslA::spc) [28] → N1230 |
| N924       | ΔnprB−yitM::tet | This study |
| N1290      | amyE::P_{pacb}−yitPOM (erm) ΔnprB−yitM::tet | N924 → N1285 |
| N1238      | amyE::P_{pacb}−yitPOM (erm) ΔnprB−yitM::tet ΔsigW::neo | N1268 → N1290 |
| N1293      | amyE::P_{pacb}−yitPOM (erm) ΔnprB−yitM::tet ΔsigW::neo ΔepsA−O::spc | N355 → N1238 |
| N1503      | amyE::P_{pacb}−yitPOM (erm) ΔnprB−yitM::tet ΔsigW::neo ΔtapA−tasA::cat | N11 (ΔtapA−tasA::cat) [62] → N1238 |
| N1294      | amyE::P_{pacb}−yitPOM (erm) ΔnprB−yitM::tet ΔsigW::neo ΔbslA::spc | N254 (ΔbslA::spc) [28] → N1238 |

(Continued)
Northern blot analysis

Wild-type and ΔdegU mutant cells were grown at 37˚C in 2×SG with vigorous shaking, and samples were taken from the cultures at various time points for RNA isolation. Total RNA was prepared as previously described [47]. The Northern blot analysis was carried out as previously described [47]. Primers used for RNA probe synthesis are shown in S2 Table.

Microscopic observation

The strains carrying gfp reporters were grown on 2×SG or LB solid medium. The expression of the GFP reporters on the colonies was analyzed with a SZX7 stereomicroscope (Olympus, Tokyo, Japan) equipped with an AdvanCam-E3Rs digital color camera (Advan Vision, Tokyo, Japan). For colony observation, the plates were tilted slightly using a small piece of cardboard (1.5 mm thickness) under the microscope to avoid detecting excitation light reflections on the surfaces of colonies and solid media. However, we could not remove reflected light completely, and some reflected light was observed as blue color signals on GFP images. Images were obtained and processed with AdvanView (Advan Vision) and Photoshop Elements (Adobe, San Jose, CA, USA). The experiments were done at least three times, and the typical examples were shown in figures.

Flow cytometry analysis

Strains harboring promoter-gfp fusions were grown on 2×SG or LB solid medium. A whole single colony was scraped from the surface of the solid medium and suspended in 1 ml of PBS buffer. Cells in the biofilms were then dispersed by repetitive pipetting and were fixed in 4% paraformaldehyde for 7 min [69]. Prior to flow cytometry analysis, the cells were subjected to mild sonication [69]. Single-cell fluorescence was measured on an Accuri C6 flow cytometer (BD Biosciences, Franklin Lakes, NJ, USA). The number of recorded events was 50,000. The experiments were done twice, and the typical examples were shown in figures.

Competition assay

Strains were grown at 28˚C overnight in LB with vigorous shaking. Cultures of the strains used for the assay contained 3.0 × 10⁸ cells/ml on average. The cultures were diluted 100-fold in LB.

Table 1. (Continued)

| Strain name | Genotypes | References or construction* |
|-------------|-----------|-----------------------------|
| N1358       | amyE::Pspac-hy-yitPOM (erm) ΔyitR-yitM::tet ΔsigW::neo ΔsinR:cat | WTF92 (ΔsinR:cat) [68] → N1238 |
| N999        | amyE::Pspac-hy-sdpABC (erm) ΔsdpA-sdpR::spc ΔsigW::neo | N1268 → N1335 |
| N1334       | amyE::Pspac-hy-sdpABC (erm) ΔsdpA-sdpR::spc ΔsigW::neo ΔsinR:cat | WTF92 (ΔsinR:cat) [68] → N999 |
| N1264       | nprB::cat | W115 (nprB::cat) [47] → NCIB3610 |
| N1287       | ΔyitQ::cat | This study |
| N1288       | ΔyitQ::cat ΔsigW::neo | N1268 → N1287 |
| N1234       | ΔyitR-yitM::tet ΔsigW::neo | N1268 → N1263 |
| N1413       | ΔyitQ::spc ΔsigW::neo amyE::PaprE-gfp (cat) | N1382 → N1288 |
| N1388       | ΔyitR-yitM::tet ΔsigW::neo amyE::PaprE-gfp (cat) | N1382 → N1234 |

*Arrows indicate B. subtilis transformation: donor strain name → recipient strain name.

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and two dilutions were mixed at the indicated ratios. Aliquots (2 μl each) of the mixtures were spotted onto 2×SG solid medium, and the plates were incubated at 30˚C for 48 h. GFP fluorescence on the colonies was analyzed as described above. The experiments were done three times, and the typical examples were shown in figures. The population ratios of aprE-gfp cells in these colonies were also analyzed by determining the number of cells per colonies. A whole single colony was scraped with an inoculation loop and was dissolved in 1 ml of LB in a test tube. After 10 times pipetting, the cell suspension was left for a while until the unsolved cell aggregates went down to the bottom of the tube. The dissolved cells were then serially diluted with LB, and these dilutions were plated on LB or LB plus chloramphenicol (Cm). Since aprE-gfp cells exhibited Cm<sup>r</sup>, the population ratio of aprE-gfp cells was calculated as Cm<sup>r</sup> CFUs/total CFUs. Each population ratio was the average of 4 independent colony measurements.

**Supporting information**

S1 Fig. The colony morphology of P<sub>pac</sub>-hy<sup>yitPOM</sup> mutants on MSgg medium with 1 mM IPTG. The strains were grown at 30˚C on MSgg. Scale bar, 2 mm. (TIF)

S2 Fig. The growth profile of the P<sub>pac</sub>-hy<sup>yitPOM ΔyitR-yitM ΔsigW</sup> mutant in LB with or without 1 mM IPTG. (TIF)

S3 Fig. The production of the YIT toxin. ΔyitR-yitM ΔsigW and ΔnprB-yitM ΔsigW cells were added to 1.2% 2×SG agar with or without 1000 μM IPTG, and the mixtures were poured into plates. P<sub>pac</sub>-hy<sup>yitPOM ΔyitR-yitM</sup> and P<sub>pac</sub>-hy<sup>yitPOM ΔnprB-yitM</sup> cells were spotted on the lawn plates. The plates were incubated at 37˚C for 24 h. (TIF)

S4 Fig. Comparison of the genetic organization of the sdpABC homologs in different B. subtilis strains. The genetic organization of sdpABC and sdpABC homologs in the indicated B. subtilis strains was compared with that of the corresponding locus in strains that do not have sdpABC or sdpABC homologs. Homologous genes are shown by patterned boxes of the same color. Strain names are shown to the right of the gene maps. (TIF)

S5 Fig. Comparison of SdpC homologs. (A) The alignment of SdpC homologs. The sequences of SdpC homolog 1 (YitM), homolog 2, homolog 3, and homolog 4 are derived from B. subtilis strains NCIB3610, ATCC13952, BEST195, and OH131.1, respectively. The signal sequences and the SDP toxin sequence are shown in blue and red, respectively. Hydrophobic amino acid residues in the C-terminal regions of SdpC homologs are shown in green. Identical and similar amino acids among all of the homologs are indicated by asterisks and dots, respectively. (B) Hydropathy plots of SdpC homologs. The plots were constructed using the ExPASy website (https://web.expasy.org/protscale/) with a window size of 19. (TIF)

S1 Table. The distribution of sdpABC and its homologs in selected B. subtilis strains. (DOCX)

S2 Table. Primers used in this study. (DOCX)

S1 File. Construction of B. subtilis mutants. (PDF)
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