Evolution of the SpoIISABC Toxin-Antitoxin-Antitoxin System in Bacilli

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Abstract: Programmed cell death in bacteria is generally associated with two-component toxin-antitoxin systems. The SpoIISABC system, originally identified in Bacillus subtilis, consists of three components: a SpoIISA toxin and the SpoIISB and SpoIISC antitoxins. SpoIISA is a membrane-bound protein, while SpoIISB and SpoIISC are small cytosolic antitoxins, which are able to bind SpoIISA and neutralize its toxicity. In the presented bioinformatics analysis, a taxonomic distribution of the genes of the SpoIISABC system is investigated; their conserved regions and residues are identified; and their phylogenetic relationships are inferred. The SpoIISABC system is part of the core genome in members of the Bacillus genus of the Firmicutes phylum. Its presence in some non-bacillus species is likely the result of horizontal gene transfer. The SpoIISB and SpoIISC antitoxins originated by gene duplications, which occurred independently in the B. subtilis and B. cereus lineages. In the B. cereus lineage, the SpoIIS module is present in two different architectures.

Keywords: Bacillus subtilis; toxin-antitoxin; Bacillus cereus; SpoIISABC; molecular evolution

1. Introduction

Programmed cell death (PCD) is a genetically-regulated system in which a bacterial cell is able to commit suicide in response to a variety of different stresses. This response includes cell lysis or growth inhibition induced by harsh environmental conditions, such as starvation or antibiotic treatment, active mother cell lysis during sporulation to release the spore or altruistic suicide to release cell content to provide the nutrients required for the normal development of the remaining bacterial population [1]. PCD is usually mediated by a toxin/antitoxin (TA) genetic pair. However, there is controversy about the role of TA systems in PCD, and it was proposed that these systems more likely enhance the formation of dormant persister cells [2]. Toxins are always highly stable proteins. Their antidotes, the antitoxins, are usually labile proteins or small RNAs. TA systems are classified according to the nature of the antitoxin. Antitoxin types I and III are small RNAs, which either inhibit toxin synthesis or capture it. Types II, IV, V and VI, on the other hand, are all proteins that are distinguished based on their mode of action. The type II antitoxins are in many cases small proteins with an N-terminal DNA-binding domain and a C-terminal toxin-bonding domain; the type IV antitoxin is an antagonist of its cognate toxin and competes with it in binding to its target; the type V antitoxin is an endoribonuclease that degrades the toxin-encoding mRNA [3]; and the type VI antitoxin acts as an adaptor by binding to its toxin partner and induces its proteolytic degradation [4].

The SpoIIS system was discovered during a study of Bacillus subtilis cell differentiation and originally appeared to consist of two proteins, the SpoIISA toxin and the SpoIISB antitoxin [5]. SpoIISA is a 248 residue-long protein with three putative transmembrane segments at its N-terminus and a cytosolic domain at its C-terminus. Sporulating B. subtilis cells synthesizing SpoIISA in the absence of SpoIISB exhibit lethal damage to the sporulation septum; the cytoplasmic membrane of vegetatively
growing *B. subtilis* cells is also susceptible to the toxicity of overexpressed SpoIISA [5,6]. Interestingly, the SpoIISA is not toxic to only Gram-positive bacteria. The SpoIISA protein from *B. subtilis* and its homologues from *B. cereus* and *B. anthracis* are also capable of killing Gram-negative, two-membrane *Escherichia coli* cells. This phenotype can be prevented by the formation of a tight complex between the cytosolic domain of the toxin and its small, hydrophilic SpoIISB antidote [5,7]. SpoIISB is a small, 65 residue-long protein with no obvious tertiary structure [7]. The toxin-inhibitor complex adopts a heterotetrameric configuration, with two SpoIISB inhibitors wrapped around a SpoIISA dimer. The entire SpoIISB sequence is involved in extensive interactions with SpoIISA, and each SpoIISB protein interacts with both units of the SpoIISA dimer [7]. Recently, a third component of this system, SpoIISC, was identified, possessing several properties similar to SpoIISB. Both are small proteins and both are capable of neutralizing the toxicity of SpoIISA and, thus, serve as antitoxins [8]. The genes coding for the SpoIISA, SpoIISB and SpoIISC proteins are positioned next to each other on the chromosome, likely forming a single operon. It is thought that both proteins are transcribed together from a common promoter, but SpoIISB is likely to also be transcribed from its own promoter. Because SpoIISB is probably less stable than SpoIISA; two promoters are needed to ensure that it is produced at a high enough level to compensate for its higher rate of degradation [5]. There is presently no evidence for an additional regulatory region for the transcription of *spoIISC*. To date, SpoIISA cell death-inducing activity has been observed only when both SpoIISB and SpoIISC proteins were absent or when SpoIISA was overexpressed. Since the activation of this TA system has never been observed under physiological conditions, the role of this system in programmed cell death remains unknown.

Interestingly, this TA system is widely distributed among Bacilli species. Here, we identified homologues of all *spoIIS* genes on the chromosomes of a wide range of Bacilli species, and we analyzed their possible evolution and genetic transfer. Although the SpoIISB and SpoIISC antitoxins appear to share the same evolutionary origin, they may differ in their functions and biochemical properties.

2. Results and Discussion

2.1. Evolution of the SpoIISA Toxin in Bacilli

We found the presence of SpoIISA proteins to be restricted mostly to the Bacillus genus of the Firmicutes phylum (Firmicutes; Bacilli; Bacillales; Bacillaceae, Bacillus) (Table S1). It is notable that there are many Bacillus species whose genomes appear to lack a SpoIISA homologue (Table S1). Most of these are from unfinished genome projects, however (the genome assembly is presently in a form of individual contigs), and it is thus likely that SpoIISA proteins will be found in these genomes in the future. There are still a few species that are almost certainly missing SpoIISA. *Bacillus megaterium* is one example: the complete genomes of multiple strains (*B. megaterium* DSM 319; QM B1551; WSH-002) of this species (including 10 plasmids) are available [9,10], and none of them possess a SpoIISA protein (Table S1). Either *B. megaterium* inherited a SpoIISA protein from the common ancestor of the other SpoIISA-bearing species and subsequently lost it or this species split off from a bacilli subgroup before SpoIISA was obtained. The genomes of the alkaliphilic *B. pseudofirmus* OF4 (one chromosome and two plasmids), *B. selenitireducens* MLS10 (one chromosome) and *B. halodurans* C-125 (one chromosome) all seem to be missing SpoIISA, as well [11–13]. It is notable that *B. selenitireducens* does not form spores. Interestingly, SpoIISA also appears in some species outside of the Bacillus genus and some even appear outside of the Bacillaceae family (Table S1). The SpoIISA sequences from these non-bacillus species are quite dissimilar from each other, however, even if they come from closely related species (e.g., *Lysinibacillus manganicus* and *Lysinibacillus massiliensis*), which is reflected by their failure to group together in the SpoIISA phylogenetic tree (Figure 1).
Moreover, most of them are part of an accessory genome. For example, SpolISA from *Paenibacillus polymyxa* SC2 is positioned on the plasmid PPSC2_p0465. It is thus more likely that some, possibly even all, SpolISA proteins from non-bacillus species were obtained through horizontal gene transfer, rather than from a common ancestor of bacillus and non-bacillus SpolISA-bearing species. This is corroborated by the observation that the non-bacillus species do not form a single separate cluster in the phylogenetic tree, but are intermixed with sequences from bacillus species. Nevertheless, most SpolISA proteins identified in this study come from species that are closely related to *B. cereus* or *B. subtilis* (Table S1), and the SpolISA homologues can be grouped based on whether they originate from *B. subtilis* relatives or *B. cereus* relatives. SpolISA similarity within each group is quite high (more than 90% for species from the *B. cereus* group and more than 80% for species from the more diverse *B. subtilis* group), but inter-group similarity is considerably lower (e.g., the similarity between the SpolISA proteins from *B. cereus* E33L and from *B. subtilis* strain 168 is 45%, and the identity is only 20%). The similarity between the SpolISA proteins from these two groups is thus comparable to the similarity between those located on chromosomes and those on plasmids (e.g., the similarity between *B. cereus* E33L and *Paenibacillus polymyxa* SC2 is 45.3%, and the identity is 19.6%).

Exactly one gene coding for SpolISA was found in every available genome for strains from the *B. subtilis* and *B. cereus* groups. Moreover, the spolISA gene was always found nearby well-conserved genes from the core genome, suggesting that in these species, spolISA genes behave similarly as those
from a very stable part of the core genome. There are two notable exceptions to this, however: *B. cereus* BDRD-ST196 possesses two SpoIISA proteins, one of which (NCBI-Protein ID: EEL06055) shows very high similarity (more than 95%) to SpoIISA proteins from other *B. cereus* strains and is positioned well inside the conserved core region of the genome, while the other one (NCBI-Protein ID: EEL06141) is only weakly similar to SpoIISA proteins from other *B. cereus* strains (about 30% identity and 45% similarity). The closest known relative to this second one is the SpoIISA protein (NCBI-Protein ID: WP_016427205) from *Paenisporosarcina* sp. HGH0030 (54% identity and 71% similarity). The second species with multiple SpoIISA proteins is *B. thuringiensis* Bt407, which contains three SpoIISA proteins. Again, one protein (NCBI-Protein ID: EEM28840) is very similar (more than 95%) to SpoIISA from closely-related strains and species, while the other two (NCBI-Protein ID: EEM25308 and EEM24943) differ considerably (less than 30% identity and less than 55% similarity to SpoIISA proteins from closely-related strains and species). The closest homologue of EEM25308 is again SpoIISA from *Paenisporosarcina* sp. HGH0030 (35% identity and 54% similarity), though they show only weak mutual similarity. The closest homologue of EEM24943 is SpoIISA from *B. simplex* (NCBI-Protein ID: CEG32548), but again, their mutual similarity is quite low (29% identity and 55% similarity). Interestingly, this protein is more similar to the SpoIISA proteins from the *B. subtilis* group (the identity to *B. amyloliquefaciens* LFB112 is 26%, similarity 50%) than to those from the *B. cereus* group (the identity to *B. cereus* E33L is 19%, similarity 46%) (Figure 1). Not only do EEM25308 and EEM24943 show low similarity to proteins from other species from the *B. cereus* group, but the proteins encoded by their neighboring genes are also dissimilar, making it probable that these genes are part of a flexible genome, which arose from horizontal gene transfer. It was recently reported that EEM24943 (locus: AFV22040; NCBI-Protein ID: AAU80350) is also present on plasmid BTB_502p [14]. The role that these multiple SpoIS modules play in a cell is presently unknown, and many outstanding questions remain: Are all of these genes expressed or are the dissimilar ones nonfunctional pseudogenes? Are they all expressed during the same period in the cell cycle? In the same cell compartment? Do they interact with each other?

An amino acid sequence comparison of the functionally-characterized SpoIISA proteins from *B. subtilis* PY79 (NCBI-Protein ID: AHA77284) and *B. cereus* ATCC 14579 (NCBI-Protein ID: AAP09399) shows that these two proteins contain 41 identical and 88 similar residues. The similarity between the SpoIISA proteins from *B. subtilis* PY79 and *Lysinibacillus massiliensis* (NCBI-Protein ID: KGR89544) is even higher (Figure 2), with 45 identical and 94 similar residues. Since a multiple sequence alignment of all of the SpoIISA proteins in our set showed that only a very few positions have identical or similar residues (Figure S1), it seems likely that the high level of similarity observed by comparing only these two SpoIISA proteins is not due to purifying selection, but is more likely a remnant of ancestral similarity. This assumes, of course, that all aligned sequences are from functional proteins. Since the presence of nonfunctional SpoIISA proteins could mask functionally-important positions, we aligned only those protein sequences containing the conserved, functionally-important arginine (Arg38 in *B. subtilis*) and the three N-terminal transmembrane helices, which were shown to be indispensable for SpoIISA functionality [5,15]. It is likely, however, that some essential features of SpoIISA are still undiscovered, and thus, we cannot be sure that all nonfunctional proteins were excluded from our final alignment. Comparison of the SpoIISA proteins from 56 different bacterial species revealed only four positions containing invariant residues. According to the *B. subtilis* numbering, they are Arg38, Tyr42, Asp78 and Asp225; eight positions possessed similar residues: Ile37, Lys39, Phe82, Lys89, Ile143, Lys186, Val221 and Leu232 (Figure S1). Most of these conserved residues seem to be located in the second and third transmembrane regions and in the C-terminal helix. Arg38, found near the beginning of the second transmembrane region, is important for oligomerization [15], and its substitution by glutamine results in loss of toxicity [5]. The function of the other conserved residues identified is presently unknown. The conserved aspartate (Asp78) in the third transmembrane region (Figure S1) may have an important functional role. It has been shown that an aspartate found in the transmembrane region of some receptors and transporters plays an important role in their operation [16–19]. Interestingly, a
conserved aspartate (Asp225) is also present in the SpoIISA C-terminal α-helix. Some transmembrane prediction tools suggest that this helix is located in the membrane, but others do not [15]. Structural studies also failed to confirm the transmembrane localization of this helix, but its temporary presence in the membrane still cannot be ruled out [7]. The two conserved aspartates could play a role in SpoIISA assembly or multimerization, or they could be involved in some other process. Further functional and structural studies of the SpoIISA protein with substituted conserved aspartates could throw more light on their function.

2.2. Evolution of the SpoIISB and SpoIISC Antitoxins in Bacilli

The sequence similarity of the SpoIISB and SpoIISC proteins is relatively high within the B. subtilis and B. cereus groups, but quickly decreases with growing evolutionary distance outside of these groups. Consequently, only SpoIISB and SpoIISC from the B. subtilis and B. cereus groups could be reliably identified and only sequences from these two groups were used in our study. We found SpoIISB proteins (hereafter SpoIISBsub) in all B. subtilis strains examined (Table S3), and the identity of SpoIISBsub between all of these strains is more than 96%. The identity of SpoIISBsub from B. subtilis to its homologues from closely-related species from the B. subtilis group (B. vallisnortis, B. tequilensis, B. sp. MSP13, B. mojavensis and various B. atrophaeus strains) is also quite high, above 94%, with nearly all of the differences occurring in the first loop of the N-terminal part, which has been shown to be unnecessary for antitoxin activity [7]. Moving to more distantly-related species, SpoIISBsub identity drops to 85% for various strains of B. amyloquae and is around 70%–75% for SpoIISBsub from B. pumilus, B. altitudinis, B. xiangenensis, B. sp. DW5-4, B. sp. NSP9.1, B. safensis, B. sp. WP8, B. sp. BT1B_CT2, B. sonorensis and B. licheniformis. The N-terminal part again seems to harbor the substitutions. The shared, conserved regions could be identified by comparing the SpoIISBsub proteins from the most distantly-related species (the identity of SpoIISBsub from B. subtilis strain 168 to B. firmus and B. sp. J33 is 57%–59%). The most highly-conserved regions correspond approximately to the SpoIISBsub secondary structure elements (Figure 3). Two of the most highly-conserved regions have been shown to be functionally important for antitoxin activity [7]. The first of these is β-strand β1′, whose three lysines are nearly invariant. This strand forms a two-stranded intermolecular antiparallel β-sheet with strand β1 of SpoIISA, and deletion of this part abolishes SpoIISBsub antitoxin activity [7].

Figure 2. Comparison of two SpoIISA proteins. Amino acid sequence of the SpoIISA from B. subtilis PY79 is aligned with the SpoIISA from (A) B. cereus ATCC 14579 and (B) from L. massiliensis. Functionally-important residues are in white on a red background, identical residues are purple and similar ones are green.
The second is a positively-charged C-terminal region comprising 2–4 tandem lysine or arginine residues (in B. subtilis K88K55). Although none of these four residues is invariant, they are always substituted with other hydrophilic residues (Figure 3). The interaction of this region with SpoIISA could not be inferred from the SpoISAB complex structure, since it was not resolved, but its importance was clearly demonstrated by a deletion experiment: deletion of the last four C-terminal SpoISB sub residues obliterates the protein’s antitoxin activity [7]. Other conserved regions, whose function has not presently been identified, include β-strand β2′ and α-helix α1′ (the N-terminal edge of this helix is the most highly conserved region of the whole protein).

Figure 3. Multiple sequence alignment of the SpoISB and SpoISC amino acid sequences from the B. subtilis group. Residues that are identical within either the SpoISB or SpoISC proteins are red, and similar ones are purple. Selection pressure is indicated on the SpoISB and SpoISC reference sequences (from B. subtilis strain 168) shown above each sequence block. Selection pressure intensity is indicated by the given scale bar. SpoISB secondary structure elements (as identified in [7]) are shown above the alignment; the abbreviations are from Table S3.

The identity of the SpoISC proteins (hereafter SpoISC sub) between the various B. subtilis strains is more than 93%. The identity of B. subtilis SpoISC sub to its homologues from closely-related species (B. callismortis, B. tequilensis, B. sp. MSP13, B. mojavensis) is comparable, from 90%–97%. The identity to its homologues from various B. atrophaeus strains, which is also a close relative, is lower, around 90%, but almost all substitutions are for residues that are functionally similar (96% positives). Moving to more distant relatives, the B. subtilis/B. amyloliquefaciens identity is 87%, with most substitutions concentrated in the C-terminal half of the protein, the region corresponding to the middle part of the α1′-helix in SpoISB sub. The identity drops to 75% for SpoISC sub homologues from B. licheniformis WX-02 and B. sp. CPSM8, and the substitutions are now distributed equally throughout the protein (Figure 3). The identity of B. subtilis SpoISC sub to its homologues from the more distantly-related members of the B. subtilis group (B. pumilus, B. sp. M 2–6, B. altitudinis 41KF2b, B. xiamensis) is 68%, and the identity to the most distant SpoISC sub homologs (from B. firmus and B. sp. J33) is 55%–60%.
Since the identities of the SpoIISC\textsubscript{sub} proteins from various species of the \textit{B. subtilis} group do not differ much from the identities observed for the SpoIISB\textsubscript{sub} proteins, it seems likely that both the SpoIISB\textsubscript{sub} and SpoIISC\textsubscript{sub} proteins from all of these organisms were inherited vertically and not obtained through horizontal gene transfer. In contrast to SpoIISB\textsubscript{sub}, only a few SpoIISC\textsubscript{sub} residues are invariant. Two conserved lysines are present at the N-terminus (K\textsubscript{2}K\textsubscript{3} in \textit{B. subtilis} B\textsubscript{5}n\textsubscript{5}), and a conserved serine and tyrosine appear in the central part of the protein (S\textsubscript{19}Y\textsubscript{21}). A C-terminal stretch of 3–4 positively-charged residues (Lys or Arg) is also well conserved (Figure 3).

SpoIISB\textsubscript{sub} proteins show only limited similarity to SpoIISC\textsubscript{sub} proteins. For example, the SpoIISB\textsubscript{sub} and SpoIISC\textsubscript{sub} proteins from \textit{B. subtilis} strain 168 are only 14.3\% identical (39.3\% similar). They also differ in length: SpoIISB\textsubscript{sub} from the \textit{B. subtilis} 168 strain is 56 residues long, while its SpoIISC\textsubscript{sub} is only 45 residues. Since the exact position of the start and stop codons was not identified, however, the real proteins could be slightly longer or shorter than this. SpoIISB\textsubscript{sub} and SpoIISC\textsubscript{sub} do share two stretches of positively-charged residues near their C- and N-terminal ends (according to \textit{B. subtilis} strain 168 numbering, the conserved, shared residues are K\textsubscript{15}XXKIL(LV)KK\textsubscript{22} at the N-terminus and K\textsubscript{52}R(K)K\textsubscript{55} at the C-terminus). The central parts of these proteins are also well conserved (Y\textsubscript{31}XV(L)SXH(Y)T(S)XR(L)\textsubscript{40}), with two invariant residues, a serine (Ser\textsubscript{34}) and an arginine (Arg\textsubscript{39}) (Figure 3). However, comparing SpoIISB\textsubscript{sub} and SpoIISC\textsubscript{sub} from multiple species reveals that only the first three of the four N-terminal positively-charged residues are conserved (according to \textit{B. subtilis} strain 168 numbering K\textsubscript{15}X\textsubscript{sub}XKXXK). Even more strikingly, only one of the conserved positively-charged residues at the C-terminus (K\textsubscript{53}) is completely conserved across all proteins (Figure 3). As noted, deletion of the last four SpoIISB\textsubscript{sub} residues considerably decreased sporulation efficiency. It is thus possible that it was the deletion of just this universally-conserved residue that was responsible for this effect. Measuring the sporulation efficiency of a point mutant in this lysine would be needed to corroborate this hypothesis, however. Of the two remaining residues common to both SpoIISB\textsubscript{sub} and SpoIISC\textsubscript{sub}, Ser\textsubscript{34} was conserved in all studied SpoIISB\textsubscript{sub} and SpoIISC\textsubscript{sub} proteins, but Arg\textsubscript{39} was not; the significance of this is presently unclear. Finally, for whatever reason, conservation of the SpoIISC\textsubscript{sub} proteins is more relaxed than the conservation of the SpoIISB\textsubscript{sub} proteins. This is most clearly seen by comparing the SpoIISB\textsubscript{sub} and SpoIISC\textsubscript{sub} proteins from \textit{B. subtilis} and \textit{B. pumilus}. The similarity between the SpoIISB\textsubscript{sub} proteins from the two species is 81.4\% (65.1\% identity), and the similarity between the SpoIISC\textsubscript{sub} proteins is only 58.1\% (32.6\% identity).

Since \textit{spolIIB}\textsubscript{sub} and \textit{spolIIC}\textsubscript{sub} are positioned adjacently on the chromosome, exhibiting sequence similarity, it is likely that they are the result of a gene duplication event. Further evidence for this comes from the recent report that both proteins have similar functions [8]: they are both able to interact with and neutralize the toxicity of SpoISA. An earlier report [5] showed that a sporulation defect was still observed when only \textit{spolIIB}\textsubscript{sub} (and not \textit{spolIIC}\textsubscript{sub}) was inactivated [5]. The most likely reason for the earlier observation was that the insertion of the tetracycline resistance cassette into codon 17 of \textit{spolIIB}\textsubscript{sub} might have disrupted a potential \textit{spolIIC}\textsubscript{sub} promoter positioned inside the \textit{spolIIB}\textsubscript{sub} gene. The SpoIISC protein also did not compensate for \textit{mut}\textsubscript{14}, which deactivated \textit{spolIISB}\textsubscript{sub}, but \textit{mut}\textsubscript{14} is a frameshift mutation caused by the deletion of 2 bp after codon 52 of \textit{spolIISB}\textsubscript{sub} [5], and this mutation would also be expected to affect a \textit{spolIISC}\textsubscript{sub} promoter or regulatory region. Alternatively, \textit{spolIISC}\textsubscript{sub} might have been deactivated by its own, independent mutation, a possibility that was not explored in the original report [5]. A final possibility is that SpoIISC\textsubscript{sub} is expressed at a different stage of the cell cycle and therefore would not have been available to compensate for the loss of SpoIISB\textsubscript{sub}. As noted earlier, different gene expression profiles could be one reason why there are two \textit{spolII} antitoxin genes present. Generally, after gene duplication, one of the copies becomes redundant and usually goes through a pseudogenization process in which it neutrally accepts nonsynonymous or even deleterious mutations and subsequently becomes nonfunctional or is lost. It may also obtain a new function, different from that of the other copy or the copies may also undergo a division of labor, with each copy retaining a different sub-function of the original, ancestral function. We thus searched for differences between the amino acid sequences of SpoIISB\textsubscript{sub} and SpoIISC\textsubscript{sub}, which could indicate either a division...
of labor or the gain of a new function. We found one position, conserved in both proteins, that contains dissimilar residues. A polar residue (arginine, lysine or glutamine; Arg23 in B. subtilis) in SpoIISBsub is replaced by a nonpolar residue (valine or leucine) in SpoIISCsub. The implication of this change remains to be investigated, but it could make SpoIISCsub bind less (Figure 4). The alignment in Figure 4 shows that the SpoIISBsub proteins contain more (twelve) conserved, non-shared positions than SpoIISCsub (only two). Since these proteins have no reported function other than inhibiting SpoISA, it is difficult to imagine how an ancestral function could be subdivided between them. It is possible (although it has not yet been investigated) that they differ in binding strength to SpoISA and that this difference could explain why SpoIISBsub contains more conserved positions than SpoIISCsub. For example, SpoIISBsub might contain more conserved positions than SpoIISCsub because more of its residues are involved in SpoISA interactions, resulting in stronger binding. It is not clear what advantage the presence of two inhibitors of different binding strengths would give to a cell, suggesting that SpoIISCsub might have an additional function. The most likely possibility is that SpoIISCsub (or SpoIISBsub) when bound to SpoISA can regulate the expression of the spoIS operon, making it serve as the third regulatory component of a three-component TA system [20–22]. However, the SpoIS cluster, with its two antitoxin proteins, would differ from other known three-component systems, which are formed by a toxin, an antitoxin and a regulatory protein, in that no regulatory proteins have yet been found that also exhibit antitoxin capabilities. On the contrary, it has been shown that PasC, the third component of the plasmid addiction system from Thiobacillus ferrooxidans plasmid pTF-FC2, is not capable of neutralizing the toxicity of the PasB toxin on its own [20].

![Figure 4](image_url)

**Figure 4.** Conserved positions in the SpoIISB and SpoIISC proteins from the B. subtilis group. Positions containing identical or similar residues in both groups are colored red; positions containing identical or similar residues in one, but not the other group are green. Positions conserved in both groups, but containing dissimilar residues, are cyan. Selection pressure intensity and secondary structure elements are indicated as in Figure 3. The abbreviations are from Table S3.
SpoIISBcer proteins are present in all members of the B. cereus group. Some B. cereus strains possess two SpoIISBcer (SpoISB1cer, SpoISB2cer) and two SpoIISCcer (SpoIISC1cer, SpoIISC2cer) proteins. The differences between them will be explored in Section 2.3 below. The identity between SpoIISBcer from various B. cereus strains and from closely-related species like B. anthracis and B. thuringiensis is more than 80% (similarity is more than 90%). The identity between SpoIISBcer proteins from more distantly-related members of the B. cereus group (e.g., between B. cereus E33L and B. mycoides Rock1-4 SpoIIBcer) is more than 70% (the similarity is more than 85%). The most conserved part is an eight-residue central region, including residues D19FSLIKGD26 (B. cereus m1293 numbering) and the C-terminal region. The N-terminal region, except for the highly-conserved K3 and a F9FK11 motif, is less conserved (Figure 5). Similarly, SpoIISCcer proteins are present in all members of the B. cereus group. The identity between SpoIISCcer from various B. cereus strains and from closely-related species like B. anthracis and B. thuringiensis is more than 80% (similarity is more than 90%). The identity between SpoIISCcer proteins from more distantly-related members of the B. cereus group (e.g., between B. cereus E33L and B. mycoides Rock1-4 SpoIISCcer) is more than 60% (the similarity is more than 80%). The SpoIISCcer proteins are all about 16 residues longer than their corresponding SpoIISBcer proteins (e.g., B. cereus m1293 SpoIISBcer is 41 residues long, while its SpoIISCcer is 58 residues). The conservation pattern in SpoIISCcer is very similar to that of SpoIISBcer: the lower similarity at the N-terminus and the higher similarity at the C-terminus. Both SpoIISBcer and SpoIISCcer proteins have an N-terminal polar residue (K3 in B. cereus m1293), an FFK motif (F9FK11), a DFSLI(V)KG motif (D19FSLIKGD25) and a conserved C-terminal region. The SpoIISCcer proteins contain an additional 12 residues at the C-terminus (Figure 5), but this region is not especially well conserved. Like their B. subtilis counterparts, the SpoIISCcer proteins are less tightly conserved than the corresponding SpoIISBcer proteins. For example, the similarity between the SpoIISBcer proteins from B. cereus m1293 and B. cereus AH620 is 97.6% (87.8% identity), while the similarity between the corresponding SpoIISCcer proteins is 89.7% (84.5% identity). Since the genes coding for SpoIISBcer and SpoIISCcer proteins are positioned next to each other on the chromosome and show a high degree of sequence similarity, it is likely that they are the result of a gene duplication event. A search for signs of either division of labor or gain of function yielded three interesting positions. In SpoIISBcer, a conserved aspartate (Asp26 in B. cereus m1293) appears after the DFSLI(V)KG motif, but in SpoIISCcer, an alanine is found instead (Figure 6). The following residue in SpoIISBcer, a conserved threonine (Thr27 in B. cereus m1293), is replaced by leucine in SpoIISCcer. Finally, a negatively-charged glutamate appears 10 positions downstream of this threonine in SpoIISBcer (Glu37 in B. cereus m1293), but is replaced by the aromatic phenylalanine (or the aliphatic leucine) in SpoIISCcer (Figure 6). We also examined the selection pressure acting upon the SpoIISBcer and SpoIISCcer proteins, but no significant differences could be inferred (Figures 5 and 6). Only a weak negative selection signal was detected in the highly-conserved central region. The extremely high level of conservation in this region, which is almost invariant even at the DNA level, provides only a low level of statistical support for negative selection, which is the most likely reason for this weak signal.
**Figure 5.** Multiple sequence alignment of the SpoIISB and SpoIISC amino acid sequences from the *B. cereus* group. Residues that are identical in either SpoIISB or SpoIISC are red, and similar ones are purple. Selection pressure is indicated on the SpoIISB and SpoIISC reference sequences (from *B. anthracis* A0248) shown above each sequence block. Selection pressure intensity is indicated by the given scale bar. The abbreviations are from Table S3.
2.3. Evolution of the SpoIIS Module

It might be speculated that SpoIISA acts as a holin, forming a pore in the cytoplasmic membrane through which some lytic protein, e.g., an amidase, is released to lyse the cell wall peptidoglycan. Intriguingly, the *B. subtilis* spoIIS cluster partially overlaps the *N*-acetylmuramoyl-L-alanine amidase (XlyA) of prophage PBSX (although in the opposite direction). Inspection of the gene arrangement in various species reveals, however, that the spoIIS cluster is positioned near the PBSX prophage only in species from the *B. subtilis* group and that this connection is missing in the more distantly-related spoIIS cluster-bearing species. Although an evolutionary relationship, or some past or present interaction, between the spoIIS cluster and the PBSX prophage cannot be ruled out, expression of the spoIIS cluster from *B. subtilis* and *B. cereus* in *E. coli* cells shows that SpoIISA is able to induce cell lysis without the help of XlyA or any other prophage protein [6].
The overall similarity between SpoIISB and SpoIISC from species of *B. cereus* and *B. subtilis* groups is very low, with no invariant residues and only three positions containing relatively similar residues (Figure 7). Thus, they either shared a common ancestor and afterwards evolved beyond recognition or they originated independently. Common ancestry seems the more probable explanation, since it is hard to imagine that so potent of a toxin as SpoIISA could exist in a viable cell without an inhibitor. A given SpoIISB or SpoIISC from a species belonging to one of the two groups is more similar to another SpoIISB or SpoIISC (either one) from the same group than it is to its corresponding orthologue from a species of the second group. This can be most clearly seen in the phylogenetic tree of SpoIISB and SpoIISC (Figure 8) where the SpoIISB<sub>sub</sub> and SpoIISC<sub>sub</sub> proteins from the *B. subtilis* group are positioned on a common branch, which is isolated from the branch occupied by the SpoIISB<sub>cer</sub> and SpoIISC<sub>cer</sub> proteins from the *B. cereus* group. It thus seems very likely that the SpoIISB and SpoIISC proteins are the result of two independent gene duplication events, which took place after the *B. cereus* and *B. subtilis* groups separated (Figure 9, upper part). Although the SpoIISB and SpoIISC proteins in a given organism are clearly paralogues, because the gene duplication events appear to have happened after the *B. subtilis* and *B. cereus* groups separated, the SpoIISB and SpoIISC proteins from species belonging to different groups are not clearly orthologues. Interestingly, species (and strains) from the *B. cereus* group have two different architectures for the SpoIS module. There is a shorter one, consisting of the *spoIISA* gene followed by the *spoISB<sub>cer</sub>* and *spoIISC<sub>cer</sub>* genes, and a longer one comprising the *spoIISA* gene followed by the *spoIISB<sub>1cer</sub>* and *spoIISC<sub>1cer</sub>* genes, then two genes for unknown hypothetical proteins and, finally, the *spoIISB<sub>2cer</sub>* and *spoIISC<sub>2cer</sub>* genes. Although one might expect that the longer form is the result of duplicating the shorter form, the opposite seems to be true. As the phylogenetic tree (Figure 8) shows, SpoIISB<sub>cer</sub> from the short form corresponds to SpoIISB<sub>1cer</sub> from the long form, while SpoIISC<sub>cer</sub> from the short form corresponds to SpoIISC<sub>2cer</sub> (or at least the C-terminal part of it) from the long form. The most parsimonious explanation for these observations is that the middle part of the longer form was deleted (Figure 9, lower part). Since the intergenic regions between *spoIISB<sub>1cer</sub>* and *spoIISC<sub>1cer</sub>* and *spoIISB<sub>2cer</sub>* and *spoIISC<sub>2cer</sub>* are identical in the shorter form, it is difficult to determine the exact position of the cleavage site. If the cleavage site was positioned inside the intergenic region, the deleted part would comprise the entirety of the *spoIISB<sub>1cer</sub>* and *spoIISC<sub>1cer</sub>* genes, leaving the complete sequences of the *spoIISB<sub>1cer</sub>* and *spoIISC<sub>2cer</sub>* genes in the short form. Alternatively, if the cleavage site was positioned inside *spoIISC<sub>1cer</sub>* then the C-terminal part of *spoIISB<sub>1cer</sub>* are all of *spoIISB<sub>2cer</sub>* and the N-terminal part of *spoIISB<sub>2cer</sub>* would be deleted. *SpoIISB<sub>cer</sub>* of the short form would then correspond to *spoIISB<sub>1cer</sub>* and *spoIISC<sub>cer</sub>* would have been formed by the fusion of the C-terminal part of *spoIISB<sub>1cer</sub>* and the N-terminal part of *spoIISB<sub>2cer</sub>* from the longer form (Figure 9). Notably, we found signs of nonfunctionalization in the *spoIISB<sub>1cer</sub>* and *spoIISB<sub>2cer</sub>* genes in some strains of *B. cereus*, including deletions, nonsense mutations and reading-frame shifts. It is thus possible that these two genes were either pseudogenes (or were on their way to becoming ones) even before they were deleted. The high mutual sequence similarity between the *spoIISB<sub>1cer</sub>*/*spoIISB<sub>1cer</sub>* and *spoIISB<sub>2cer</sub>*/*spoIISB<sub>2cer</sub>* pairs and the presence of pseudogenes in the middle part of this region thus poised it for deletion.

B. cereus NV0597 B1 --MIKNLOWDDDK-EDNEOTDASHLGGDTLQQGELED----RKRK---------
B. cereus NV0597 B2 --LTKKKKEQFEQVQVKTSDFSLKGDTRQQGEELE--RKRK---------
B. cereus NV0597 C1 MAEVWQKEFSKKEKEPMILDDSFSLKGKLRTQQGLRKE-----FYKIPYKVKEGVMKKA
B. cereus NV0597 C2 MAEVWQKEFSKKEKEPMILDDSFSLKGKTEENHLEK-----RNSSSYTVTKEKKN
B. subtilis FY79_B -------MERAQFQCEFRAAKKFPIKRRKSTASSVQPSMTAPKPKERLR1DEYKRKKA
B. subtilis FY79_C -------MGDTVKRLRERVKSTVGFG--GSYSLF------PGRERRAG11KLLKEK

**Figure 7.** Multiple sequence alignment of SpoIISB and SpoIISC amino acid sequences from two representatives from the *B. cereus* and the *B. subtilis* groups. Similar residues are colored purple.
We also found another group of SpoIISA proteins in the Bacillus genus of the Firmicutes phylum, we found exactly one copy of the spoIISA gene, which was positioned next to well-conserved core genes, indicating that they behave as the stable part of the core genome. We also found another group of SpoIISA proteins in

3. Conclusions

In most species of the Bacillus genus of the Firmicutes phylum, we found exactly one copy of the spoIISA gene, which was positioned next to well-conserved core genes, indicating that they behave as the stable part of the core genome. We also found another group of SpoIISA proteins in
non-bacillus species, which were probably obtained from Bacillus species through horizontal gene transfer. Comparing the SpoIISA proteins from 56 different bacterial species revealed that only four positions carried invariant residues and eight positions possessed similar residues (with the caveat that some important residues might have been masked by the possible presence of non-functional proteins in our dataset). Interestingly, one of these conserved residues is an aspartate located in the third transmembrane region. All Bacillus species studied possess two adjacent antitoxin genes, which are likely the result of gene duplication. This gene duplication occurred independently in the \textit{B. subtilis} and \textit{B. cereus} lineages after they had separated. Although the reason for the presence of two antitoxin genes still needs to be investigated, it is clear that the SpoISB protein contains more conserved residues than the SpoISC protein, whose conservation seems to be more relaxed. In the \textit{B. cereus} lineage, the \textit{spolIS} module is present in two different architectures. The shorter of the two consists of the \textit{spolISA} gene followed by the \textit{spolISB} and \textit{spolISC} genes, while the longer one comprises the \textit{spolISA} gene followed by the \textit{spolISB1} and \textit{spolISC1} genes, then genes for two unknown hypothetical proteins and, finally, \textit{spolISB2} and \textit{spolISC2}. A phylogenetic analysis suggests that the short form evolved from the long form by deletion of the middle part.

4. Experimental Section

The amino acid sequences of the SpoIISA, SpoISB and SpoISC proteins from various bacterial species were obtained using protein BLAST [23] against the non-redundant database. Amino acid sequences of the characterized SpoIISA (NCBI-Protein ID: AHA77284), SpoISB \textsubscript{sub} (NCBI-Protein ID: AHA77283) [5,6] and SpoISC \textsubscript{sub} (NCBI-Protein ID: AHA77282) proteins from \textit{B. subtilis} PY79 and the SpoIISA (NCBI-Protein ID: AAP09399) and SpoISC \textsubscript{cer} (NCBI-Protein ID: AAP09400) proteins from \textit{B. cereus} ATCC 14579 [8] were used as queries. Since the amino acid sequence of SpoISB \textsubscript{cer} from \textit{B. cereus} ATCC 14579 is not found in the NCBI protein database, it was translated directly from the genomic sequence [24] and then used as the BLAST search query. If the amino acid sequences of the SpoIS proteins from a given bacterial species were not found using BLAST, they were produced by translating from the appropriate genomic sequence. The resulting amino acid sequences were then aligned using ClustalX [25]. Nucleic acid sequences of particular \textit{spolIS} genes were obtained from the bacterial genomes available in GenBank [26]. A neighbor-joining (NJ) tree [27] was calculated using ClustalX using default parameters, and the tree topology reliability was evaluated using the bootstrap test [28] with 1000 replications. Selection pressure was estimated using the SELECTON tool [29] using the M8 codon-substitution model [30] and a neighbor-joining tree. For calculating selection pressure, the sequences listed in Table S3 were used. Sequence similarities were estimated using MatGAT [31] using the BLOSUM62 matrix.

Supplementary Materials: The following are available online at www.mdpi.com/2072-6651/8/6/180/s1.
Figure S1: Multiple sequence alignment of SpoIISA amino acid sequences from various bacterial species. Functionally-important residues are white on a red background; identical residues are purple; similar ones are green. Substitutions of functionally-important residues with dissimilar residues are colored cyan. Secondary structure elements (as identified in [7]) are shown above the alignment. Table S1: The SpoIISA proteins from Bacillus and non-Bacillus species and species with available genome sequences lacking the SpoIISA proteins. Table S2: The SpoIISA proteins used in phylogenetic analysis. Table S3: SpoISB and SpoISC proteins from the species of the \textit{B. subtilis} and \textit{B. cereus} groups.

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