Architecture and Assembly of the *Bacillus subtilis* Spore Coat

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**Abstract**

*Bacillus* spores are encased in a multilayer, proteinaceous self-assembled coat structure that assists in protecting the bacterial genome from stresses and consists of at least 70 proteins. The elucidation of *Bacillus* spore coat assembly, architecture, and function is critical to determining mechanisms of spore pathogenesis, environmental resistance, immune response, and physicochemical properties. Recently, genetic, biochemical and microscopy methods have provided new insight into spore coat architecture, assembly, structure and function. However, detailed spore coat architecture and assembly, comprehensive understanding of the proteomic composition of coat layers, and specific roles of coat proteins in coat assembly and their precise localization within the coat remain in question. In this study, atomic force microscopy was used to probe the coat structure of *Bacillus subtilis* wild type and cotA, cotB, safA, cotH, cotO, cotE, getE, and cotE getE spores. This approach provided high-resolution visualization of the various spore coat structures, new insight into the function of specific coat proteins, and enabled the development of a detailed model of spore coat architecture. This model is consistent with a recently reported four-layer coat assembly and further adds several coat layers not reported previously. The coat is organized starting from the outside into an outermost amorphous (crust) layer, a rodlet layer, a honeycomb layer, a fibrous layer, a layer of “nanodot” particles, a multilayer assembly, and finally the undercoat/basement layer. We propose that the assembly of the previously unreported fibrous layer, which we link to the darkly stained outer coat seen by electron microscopy, and the nanodot layer are cotH-dependent and cotE-specific respectively. We further propose that the inner coat multilayer structure is crystalline with its apparent two-dimensional (2D) nuclei being the first example of a non-mineral 2D nucleation crystallization pattern in a biological organism.

**Introduction**

Spores of bacteria of *Bacillus* species are formed in sporulation and are metabolically dormant and resistant to a large variety of environmental stress factors. While multiple factors contribute to spore resistance, one striking spore feature is the multilayer spore coat that provides protection against many toxic chemicals, as well as digestion by lytic enzymes and being eaten by several types of predatory eukaryotes [1–4]. The spore coat of *Bacillus subtilis* is a fibrous coat consisting of at least 70 proteins. The elucidation of *Bacillus* spore coat architecture, assembly, comprehensive understanding of the proteomic composition of coat layers, and specific roles of coat proteins in coat assembly and function is critical to determining mechanisms of spore pathogenesis, environmental resistance, immune response, and physicochemical properties. Recently, genetic, biochemical and microscopy methods have provided new insight into spore coat architecture, assembly, structure and function. However, detailed spore coat architecture and assembly, comprehensive understanding of the proteomic composition of coat layers, and specific roles of coat proteins in coat assembly and their precise localization within the coat remain in question. In this study, atomic force microscopy was used to probe the coat structure of *Bacillus subtilis* wild type and cotA, cotB, safA, cotH, cotO, cotE, getE, and cotE getE spores. This approach provided high-resolution visualization of the various spore coat structures, new insight into the function of specific coat proteins, and enabled the development of a detailed model of spore coat architecture. This model is consistent with a recently reported four-layer coat assembly and further adds several coat layers not reported previously. The coat is organized starting from the outside into an outermost amorphous (crust) layer, a rodlet layer, a honeycomb layer, a fibrous layer, a layer of “nanodot” particles, a multilayer assembly, and finally the undercoat/basement layer. We propose that the assembly of the previously unreported fibrous layer, which we link to the darkly stained outer coat seen by electron microscopy, and the nanodot layer are cotH-dependent and cotE-specific respectively. We further propose that the inner coat multilayer structure is crystalline with its apparent two-dimensional (2D) nuclei being the first example of a non-mineral 2D nucleation crystallization pattern in a biological organism.
which have drastic effects on overall coat architecture, as these proteins direct the assembly of different subsets of proteins into the coat [2,4,9–11]. In addition, the SpoIID, GerE and GerR proteins have major effects on the expression of genes encoding coat proteins that are transcribed during sporulation, and this in turn has significant effects on coat properties and morphology [4–6].

A variety of studies of the functional repertoire of coat proteins have focused on the determination of the locations of these proteins in the spore coat and their specific roles in spore coat morphogenesis [5,7,12–15]. These studies have been extended and complemented by studies of direct interactions between various coat proteins, both in vitro and in vivo [4,16–20]. All of this work has given a picture of the molecular interactions in the spore coat, as well as the dependencies of the assembly of specific proteins into the coat. However, this type of analysis has not yet been complemented by detailed analysis of the structures of the various spore layers. Atomic force microscopy (AFM) has been used to unravel high-resolution structures of the coats of dormant and germinating spores of various Bacillus [14,21–30] and Clostridium [31] species. However, this analysis has generally been conducted on wild-type spores, with AFM data on only a few mutants lacking specific coat layers. Consequently, in this work we have used high-resolution AFM to analyze the surface structure of spores of wild-type B. subtilis as well as spores of a variety of mutant strains in order to reveal the surface morphology of various layers of the spore coat. The results from these analyses have provided high-resolution visualization of the various protein layers of the spore coat as well as several coat layers not reported previously. This information has allowed the formulation of a model for coat structure and provided further insight into the assembly of the spore coat.

Materials and Methods

Strains used in this study

The B. subtilis strains used in this study (Table 1) except one are isogenic with the wild-type strain PS832, a prototrophic derivative of strain 168. Preparation of strains by transformation with chromosomal DNA was as described [32].

| Strain   | Genotype      | Phenotype* | Source or reference |
|----------|---------------|------------|---------------------|
| PS832    | wild-type     |            | Laboratory stock    |
| PS3394   | ΔcotE::tet   | Kan^r Tet^r | [1]                 |
| PS3735   | ΔspoVID::kan | Kan^r      | [1]                 |
| PS3736   | ΔcotH::cat   | Cm^r       | [1]                 |
| PS3738   | ΔselfA::tet  | Tetr       | [1]                 |
| PS4133   | ΔcotB::cat   | Cm^r       | DL067→PS832         |
| PS4134   | ΔcotO::tet   | Tetr       | PE250→PS832         |
| DL063    | ΔcotA::cat   | Cm^r       | [71]                |
| DL067    | ΔcotB::cat   | Cm^r       | [71]                |
| PE250    | ΔcotO::tet   | Tetr       | [87]                |

*Abbreviations: Cm^r, chloramphenicol resistant; Kan^r, kanamycin resistant; Tetr, tetracycline resistant.

Table 1. B. subtilis strains used in this study.

Spore preparation

B. subtilis strains were grown at 37°C in Luria-Bertani (LB) [33] medium supplemented with the appropriate antibiotics when necessary. Chloramphenicol was used at a final concentration of 5 mg/liter, kanamycin at a final concentration of 10 mg/liter, and tetracycline at a final concentration of 10 mg/liter.

For spore preparation, B. subtilis strains were grown for 3 h in LB medium and then spread on 2x Schaeffer’s-glucose medium agar plates without antibiotics [34]. Spores were harvested after incubation at 37°C for 5 d followed by incubation at room temperature for 2 d, and purified as described [34] by brief sonication and repeated washing with distilled water. All spore preparations, except for strain PS3735 (ΔspoVID::kan) (see below) were free (>98%) of vegetative and sporulating cells and germinated spores as determined by phase-contrast microscopy.

Spores of strain PS3735 (ΔspoVID::kan) were generally significantly contaminated with germinated spores and these germinated spores were removed by centrifugation in a one-step Histodenz™ (Sigma, St. Louis, MO) gradient. Four samples, each containing ~3 mg (dry weight) crude spores were suspended in 100 μl of 20% Histodenz™ that was layered on top of 2 ml of 50% Histodenz™ in four Ultra-Clear™ (11×34 mm) centrifuge tubes (Beckman Instruments, Palo Alto, CA) and then centrifuged at 14,000 rpm for 45 min at 20°C in a TLS 55 rotor. After centrifugation, the germinated spores in the supernatant fluid were removed, the pellets containing the dormant spores washed 5 times with 500 μl water and the final pellets were suspended in 500 μl water and combined. These purified spores were free (>98%) from vegetative and sporulating cells as well as germinated spores as determined by phase contrast microscopy.

Chemical decoating of spores

Spores (~6 mg dry weight) were decoated as described previously [35,36]. Briefly, spores were incubated for 90 min at 37°C in 1 ml of 50 mM Tris-HCl (pH 8.0)-8 M urea-10 mM EDTA-1% sodium dodecyl sulfate (SDS)-50 mM dithiothreitol (DTT). After incubation, the spores were centrifuged and the pellets were washed with 1 ml of water 6–10 times. Atomic force microscopy

Droplets of ~2.0 μm of spore suspensions (~3×10⁹ spores/ml) were deposited on plastic cover slips and incubated for 10 min at room temperature and the sample substrate was carefully rinsed...
AFM of Spore Coat Architecture

Results

Surface architecture of wild-type and decoated spore surfaces

As seen previously by AFM [21,22], the prominent surface features of air-dried wild-type B. subtilis spores are surface ridges extending along the long axis of the spore (Fig. 1a,b, light blue arrows). The height of these surface ridges was generally 15–30 nm, occasionally exceeding 40 nm. Similar surface ridges have been observed on spores of Bacillus anthracis [29], Bacillus cereus [22,23], Bacillus atrophaeus [22,24], Bacillus thuringiensis [22,23], and Clostridium novyi NT [31]. This ridge formation appears to be due to coat folding caused by changes in spore size upon dehydration [22,24,30,37,38]. RMS roughness $R_q$ of wild-type coat surfaces measured as described in the Methods section for 20 spores varied between 3.49 nm to 8.71 nm with an average $R_q$ value of ~5.26 nm.

AFM studies of protozoal-digested coat-defective B. subtilis spores [27] showed that the B. subtilis spore's outer surface exhibits a thin layer without prominent structural features, which was defined as an amorphous layer (Fig. 1c,e; green arrows). EM of ruthenium red stained B. subtilis spores demonstrated the presence of an outermost glycoprotein layer, and it was suggested that this layer is an exosporium that is tightly attached to the coat layer [8]. Later, a combination of EM, fluorescence microscopy, and genetic analysis also demonstrated the existence of this outermost glycoprotein layer that was named the spore crust [7]. Thus the outermost layer revealed by AFM and the crust layer correspond to the same spore layer. The thickness of the outermost amorphous layer in B. subtilis spores as measured from AFM images (Fig. 1e) was not uniform and varied between 4–15 nm with an RMS roughness $R_q$ of ~3 nm. Typically, the coverage of surfaces of B. subtilis spores with the amorphous layer was not complete, revealing an underlying rodlet layer, seen on all visualized wild-type spores, with a periodicity of ~7–8.5 nm (Fig. 1c–e; red arrows); note that these rodlets are also seen on the surfaces of the surface ridges. Rodlet structures similar to ones seen in Fig. 1 were previously described in freeze-etching EM [39–41] and AFM studies of both fungal [42,43] and bacterial (B. atrophaeus, B. cereus and B. thuringiensis) [22–25] spores, with rodlet structures on B. atrophaeus, and B. cereus spores exhibiting ~8 nm periodicity. Note, that depending on sporulation conditions for B. thuringiensis, rodlet structures were found either on the spore coat or as extraspinal structures that were present in spore preparations [30].

In order to remove spores’ outer coat, B. subtilis spores were chemically decoated with urea-SDS at slightly alkaline pH as described in Methods. The great majority of the proteins removed by this type of treatment have been well characterized in work from a number of laboratories [44]. This treatment partially or completely removed the amorphous layer, and the outer surface of the decoated spores was now comprised primarily of the intact rodlet layer (Fig. 2b–f; red arrows), which was covered in some cases with remnants of the amorphous layer (Fig. 2a–c; green arrows). The 15–30 nm surface ridges were also seen on the air-dried decoated spores, similar to what was seen on intact spores, and again these ridges appear to contain rodlets (Fig. 2a–d; light blue arrows).

Surface architecture of spores lacking CotA, CotB and SafA

CotA and CotB are two outer coat proteins that are likely localized on or very near the spore’s outer surface [5,13,14]. Loss of either of these proteins has no notable effect on spore resistance.

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with double-distilled water and allowed to dry. Our prior work with spores of other Bacillus spp. [22–25] demonstrated that spore morphological and structural attributes were reproduced both for spores analyzed within the same spore preparation and when multiple spore preparations were analyzed. Thus, in this study for each spore strain a single spore batch was analyzed by AFM with ~50–75 spores being imaged for each spore strain. Detailed experimental procedures for AFM imaging of spores were as described previously [22,24]. Images were collected using a NanoScope IV atomic force microscope (Bruker Corporation, Santa Barbara, CA) operated in tapping mode. For rapid low-resolution analysis of spore samples, fast scanning AFM probes (DMASP Micro-Actuated, Bruker Corporation, Santa Barbara, CA) with resonance frequencies of ~210 kHz were utilized. For high-resolution imaging, SuperSharpSilicon (SSS) AFM probes (NanoWorld Inc, Neuchâtel, Switzerland) with tip radii <2 nm and resonance frequencies of ~300 kHz were used. Nanoscope software 5.30r3sr3 was used for acquisition and subsequent processing of AFM images. In order to successfully assess both overall low-resolution and high-resolution spore features, raw AFM images typically need to be modified. In particular, the contrast enhancement command, which runs a statistical differencing filter on the current image, was typically utilized. This filter can bring all the features of an image to the same height and equalize the contrast among them. This allows all features of an image to be seen simultaneously, and thus a single spore or a group of spores can be imaged at relatively low resolution while visualizing spore coat attributes at high resolution.

Heights of spore surface features (i.e. folds, coat layers, etc.) were measured from height images using the section command, which allows measurements of vertical distance (height), horizontal distance, and the angle between two or more points on the surface. Tapping amplitude, phase and height images were collected simultaneously. Height images allow quantitative height determinations, providing precise measurements of spore surface topography. Amplitude and phase images do not provide height information. While amplitude and phase images provide similar morphological and structural information as do height images, they can often display a greater amount of structural detail and contrast compared with height images, often making them a preferred choice for presentation purposes.

Surface roughness of spore surfaces for wild type and cotE gerE spores was evaluated as the root mean square (RMS) value $R_q$ using AFM height images. $R_q$ is the standard deviation of the Z values (height) within the given area and is calculated as: $\Sigma(Z_i-Z_{ave})^2/N$, where $Z_{ave}$ is the average of Z values within the given area, $Z_i$ is the current Z value, and N is the number of points within the given area. $R_q$ was determined for each spore from 4 mm² height images (pixel number ~512²) of multiple spores using a 400 nm² zoomed in area in the center of the spore. In order to eliminate tilt on the spore surface, prior to the measurement of the roughness, the image was flattened using the third flatten order in the flatten command. Step roughness levels were determined by manually digitizing steps’ contour from AFM capture images with a plot digitizer [http://plotdigitizer.sourceforge.net/]. Once the x and y coordinates of the step contours were obtained, the step perimeter length, $S$, was estimated from the sum of all segment lengths given by $S = \Sigma(\Delta x^2 + \Delta y^2)$, where the sum is carried over all digitized contour segments. The sinuosity index, which is a measure of step meandering/roughness, is then calculated by taking the ratio of the contour length $S$ over the shortest path length between the two end points of the step (straight line). Note, that the value of the sinuosity index ranges from 1 (case of straight line) to infinity (case of a closed loop, where the shortest path length is zero).
properties or gross spore coat structure. We found that both cotA and cotB spore morphologies were indistinguishable from wild-type spores by AFM (Fig. 3), as all cotA and cotB spores were encased in the outermost amorphous and rodlet layers (Fig. 3c,d; green and red arrows, respectively) and exhibited 20–40 nm thick surface ridges (Fig. 3a–d; light blue arrows). The cotA and cotB spores also had an undulating surface topography from a subsurface layer (Fig. 3c,d; red circles) that was also seen in wild-type spores (data not shown).

In contrast to CotA and CotB, SafA plays a significant role in the assembly of at least some components of the spore’s outer coat, and much of the coat in safA spores does not adhere tightly and can peel off [4,45]. We observed that the general surface morphology of safA spores as seen by AFM (Fig. 4) appears similar to that of wild-type, cotA and cotB spores, with amorphous and rodlet layers (Fig. 4c; green and red arrows, respectively) forming the outermost safA spores’ coat layer. However, the degree of safA spore coat folding was different from that in wild-type spores. This resulted in the formation of surface ridges in safA spores (Fig. 4a–c; light blue arrows) that appeared shorter (e.g. not running along the whole spore surface as in Fig. 1a) and smaller (ridge heights of 10–20 nm) than in wild-type spores. Furthermore, some safA spores had no or minimal surface ridges (Fig. 4a; dark blue arrow), and ~25% of safA spores had an oversized spore coat sacculus that appeared not to be firmly attached to the body of the spore itself (Fig. 4a,b; spores with adjacent green stars, and data not shown), consistent with previous work [44].

**Surface architecture of spores lacking CotO and CotH**

In addition to SafA, CotO and CotH also play significant roles in outer coat assembly, with perhaps some role in inner coat assembly as well [26,46]. As seen by AFM (Fig. 5), the outer surface of cotO spores was covered either completely or partially by a layer with a grainy appearance (Fig. 5b; brown arrow) and exhibited 15–40 nm thick ridges (Fig. 5a; light blue arrows). The thickness of the grainy layer was 8–20 nm as measured from the AFM images. High-resolution imaging of areas where the grainy structure density was low revealed that this layer actually has a fibrous structure, with the thickness of the thinnest fibers being ~2–4 nm (Fig. 5c; several fibers indicated with light yellow arrows). Thus, high densities of these fibrous structures appear to have assembled on the inner coat to form a layer that has a granular structure (Fig. 5b,c). Underneath the granular structure, multiple structural layers were observed (Fig. 5c; terraces of 3 consecutive layers numbered 1–3, and the edge of one terrace indicated by a purple arrow), and these terraces were decorated with “nanodot” particles (Fig. 5b,c; groups of nanodots indicated with black arrows, and a circle in 5c). While the heights of some
Figure 2. AFM images of decoated *B. subtilis* wild-type spores. Surface ridges extending along the entire length of spores are indicated with light blue arrows in height (a, b) and phase (c, d) images. Patches of rodlet structures are indicated with red arrows in (b–d). The green arrows in (a–c) indicate remnants of the amorphous outermost layer. High resolution height (e) and phase (f) images showing coincidental patches of rodlet structures denoted with red arrows.
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Figure 3. AFM images of *cotA* and *cotB* spores. Height images of *cotA* (a) and *cotB* (b) spores exhibit surface ridges similar to those in wild-type spores (light blue arrows). High-resolution phase images of single *cotA* (c) and *cotB* (d) spores show an irregular outermost amorphous layer (green arrows) as well as underlying rodlets (red arrows). In addition to the amorphous layer and rodlets seen on these spores’ outermost surface, a strong undulating topography from a sub-surface layer is also present (red circles). Surface ridges in (c,d) are indicated with light blue arrows.
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nanodots were as small as 3–4 nm, their typical height was 10–22 nm.

Significant numbers of cotH spores (Fig. 6) were also encased in the outermost amorphous and rodlet layers (Fig. 6b; green arrow and red arrows, respectively) and exhibited 15–40 nm thick surface ridges (Fig. 6a,b; light blue arrows). However, 10–15% of cotH spores were partially (Fig. 6b) or completely (Fig. 6c) devoid of outer spore coat layers. These cotH spores with a defective outer coat exhibited multilayer structures (Fig. 6c; two layers indicated with purple arrows) similar to ones observed on cotO spores (Fig. 5b,c). As seen in Fig. 6b,c, these layers again exhibited high densities of nanodots (Fig. 6b; one group of nanodots indicated with a black arrow) similar to ones seen on cotO spores (Fig. 5). Nanodot heights appeared smaller and more uniform on cotH spores compared with those on cotO spores, varying between 2.5–3.5 nm, and none of the cotH spores exhibited the fibrous/granular structural layer observed on cotO spores.

Surface architecture of cotE, gerE and cotE gerE spores

CotE is one of the major morphogenetic proteins in spore coat assembly, and cotE spores lack an outer coat and also have alterations in the inner coat layer [2]. AFM images (Fig. 7) revealed that the outermost surface of cotE spores is also a multilayer structure, composed of ~6 nm thick smooth layers (Fig. 7c; three consecutive layers marked as 1–3). These structures are identical to ones observed for cotO and cotH spores (Fig. 5b,c). Note also that: i) the surface of cotE spores was devoid of nanodots; ii) the vast majority of cotE spores appeared to lack the outermost amorphous and rodlet layers; and iii) cotE spores exhibited no granular/fibrous surface structures.

In contrast to wild-type spores, 20–25% of cotE spores had no surfaces ridges (Fig. 7a; dark blue arrow) or shorter, thinner ridges (Fig. 7a; light blue arrows) that did not extend across the entire spore surface. The thickness of surface ridges that were seen were only 5–13 nm, less than for surface ridges on intact and decoated wild-type spores (Fig. 1,2). The other interesting morphological feature observed on many cotE spores was an oversized spore coat sacculus (Fig. 7a,b; green stars). This was also seen on some safA spores (Fig. 4a,b; adjacent green stars), while the wild-type spore coat was always tightly fitted (Fig. 1). The multilayer outer structure of cotE spores (Fig. 7b,c) exhibited step growth patterns similar to those observed on surfaces of inorganic [47,48] and macromolecular [49–52] crystals. An example of similar structures observed on the surface of a growing trypsin crystal [53] is shown in Fig. 7d. Similar patterns were also observed for the inner coat of C. novyi NT [31] and B. anthracis spores (Plomp and Malkin, unpublished data). As seen in Fig. 7c, layers of structure forming the inner coat of B. subtilis spores are similar in morphology to the surface of trypsin crystals (Fig. 7d), with both showing rough steps with many kinks and a number of 5–10 nm (Fig. 7c) and 70–90 nm (Fig. 7d) wide holes (Fig. 7c,d; purple arrows and circles, respectively). The sinuosity index, which is a further measure of the step roughness (see Methods), was estimated for steps on surfaces of cotE spores (Fig. 7c) and trypsin crystals (Fig. 7d) as 3.84 and 1.49 respectively. Note that high-resolution AFM observations, which allow at least 1 nm resolution for macromolecular crystalline layers [50,54], do not result in molecular scale visualization of the molecular packing within the spore coat layers.

While most cotE spores are encased only in the multilayer coat structure, <5% spores were completely covered by a rodlet layer (Fig. 8a,b; red arrow), with a periodicity of ~7.2 nm (Fig. 8c; insert). Occasionally, as seen in Fig. 8a,b 4–10 nm thick patches of the outermost amorphous layer were observed atop the rodlet layer of <10% of cotE spores (Fig. 8a,b; green arrows). In addition, on <5% of cotE spores a honeycomb-like coat layer with a periodicity of ~8–9 nm was observed atop the inner coat layer (Fig. 9a,b; orange arrow). Note, that loose honeycomb layers with remnants of rodlet structures on top of a honeycomb layer (Fig. 9c, orange and red arrows respectively) were occasionally observed in spore preparations. While >75% of cotE spores lacked a complete rodlet layer, these spores still exhibited patches of rodlet structure of different sizes assembled atop the inner spore coat layer (Fig. 9a; red arrows).

In contrast to CotE and other proteins noted above, GerE is a transcription factor that modulates the expression of some coat proteins genes late in sporulation, including genes that encode proteins in the insoluble fraction of the spore coat [6]. A gerE mutation has drastic effects on overall spore coat structure, as: i) much of the gerE spores’ coat adheres poorly [53]; and ii) some coat component(s) responsible for the strong X-ray scattering by the spore coat is either absent or misassembled on gerE spores.
while this X-ray scattering is observed from cotE spores [56]. As seen by AFM (Fig. 10), gerE spores were devoid of the outer amorphous and rodlet layers, and fibrous structures. Most of these spores were only partially or completely covered by patches of irregular material (Fig. 10a; black stars; Fig. 10b), and 40–45% of gerE spores had only patches of this material (Fig. 10a; grey stars; Fig. 10c; grey arrow), with a thin layer of material covering the spore surface (Fig. 10b,c). The thickness of these patches of coat material was ~6 nm, a value similar to the thickness of the inner coat layers forming the multilayered coat structure (Fig. 5b, 7d).

The combination of cotE and gerE mutations has an even more drastic effect on spore coat structure than either mutation alone, as cotE gerE spores are almost completely devoid of a coat (Fig. 11), except for a thin rind of insoluble material [28]. As reported previously [28], with the exception of small numbers of spores which have remnants of coat material (Fig. 11b; grey arrow), >90% of cotE gerE spores had none of the spore coat structures described above and their outer surface appeared rather smooth (Fig. 11a,b), although high-resolution imaging revealed a slightly bumpy textured outermost surface (Fig. 11c). The RMS roughness $R_q$ of cotE gerE spore surfaces measured for 20 spores as described in the Methods section varied between 0.25 nm to 0.49 nm with average $R_q$ value of ~0.38 nm. These severely coat-defective spores also appeared less rigid than intact spores, as cotE gerE spores within a closely packed monolayer were more deformed compared to ones with fewer near neighbors (Fig. 11a). Approx-

Figure 5. AFM images of cotO spores. (a) Height image of spores with surface ridges extending along the entire length of spores (light blue arrows). (b,c) High-resolution height images of areas on surfaces of single spores showing a dense fiber structure forming a granular structure (b; brown arrow) and individual fibers (c; light yellow arrows). In panel (c), three layers (terraces) of inner coat structure are numbered 1, 2, and 3 in purple. Step edges representing boundaries of each layer (one marked with a purple arrow) are visible. In panels (b) and (c), nanodots are marked with black arrows and one area with a high density of nanodots is circled in panel (c).

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Figure 6. AFM images of cotH spores. (a) Height image of spores with surface ridges extending along the entire length of spores (light blue arrows). (b) High-resolution height image of a spore surface area showing the upper surface area (green rectangle) covered with an amorphous layer (green arrow) and rodlets (red arrows). The lower part of the outermost layer-free area (black rectangle) is covered with nanodots (black arrow). One of the surface ridges in (b) is indicated with a light blue arrow. In panel (c), a two-layer inner coat structure (two purple arrows noting the two layers) decorated with nanodots can be seen.

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Figure 7. AFM images of cotE spores. (a,b) Height images of spores that exhibit surface ridges (light blue arrows), and several spores with an oversize sacculus are labeled with green stars. In (a) a spore with no apparent ridges is indicated with a dark blue arrow. (c) Height image of a multilayer inner coat structure. Three layers are indicated with numbers, and a kink on a step edge is marked with a purple arrow. Several holes in the layered structure are also indicated with purple circles. The hole in the middle circle corresponds to a pinning point on the step. (d) Height image of a multilayer layer structure similar to ones seen in Figs. 5b,c, 6c, and 7c, as seen on the surface of a trypsin crystal. Similar to the spore coat layers in (c), three layers, kinks and several holes are indicated with purple numbers, arrows and circles, respectively. The insert in (d) is a larger area of the crystal surface seen in (d). The same holes and three layers seen in (d) are indicated in the insert. The red line in (d) denotes the step contour, which was utilized for the measurement of the sinuosity index. Panel (d) is reprinted with permission from Plomp M, McPherson A, Larson SB, Malkin AJ (2001). Growth mechanisms and kinetics of trypsin crystallization. J Phys Chem B 105: 542–551. [52]. © (2001) American Chemical Society. doi:10.1371/journal.pone.0108560.g007

Figure 8. AFM images of cotE spores. (a) High-resolution height (a) and phase (b) images of the spore surface showing (coincidental in both images) a rodlet structure (red arrows) covered with patches of an amorphous layer (green arrows). (c) High-resolution height image of the spore surface with an insert with a cross section line drawn perpendicular to rodlets showing the periodicity of ~7.2 nm. doi:10.1371/journal.pone.0108560.g008
imately 7% of \textit{cotE} \textit{gerE} spores also exhibited 80–100 nm wide and 30–40 nm deep depressions (Fig. 11a; black circles), which were also observed on some \textit{gerE} spores (data not shown). Note, that neither \textit{gerE} nor \textit{cotE gerE} spores exhibited surface ridges (Fig. 10,11).

Surface architecture of \textit{spoVID} spores

\textit{SpoVID} is another major morphogenetic protein in spore coat assembly. This protein is essential for the adherence and assembly of the coat, and while the peptidoglycan cortex forms relatively normally in \textit{spoVID} spores, the coat largely assembles as swirls in the cytoplasm, giving rise to spores with little coat material [2,10]. Consequently, the surface architecture of \textit{spoVID} spores is drastically different from that of wild-type spores, as a number of \textit{spoVID} spores were again encased in only loosely fitted coat sacculi (Fig. 12a; green stars). Indeed, for a number of \textit{spoVID} spores, the coat sacculi were partially (Fig. 12 c,d; grey stars) or completely (Fig. 12b,d; white stars) sloughed off, releasing empty sacculi (Fig. 12a, insert; dark blue star) and leaving spores encased in what appeared at lower resolution to be a rather smooth structure (Fig. 12a). Note that the shape of a number of the coatless \textit{spoVID} spores was altered significantly compared either to other mutant spores described above or to \textit{spoVID} spores still encased in coat sacculi. The shape of the coatless \textit{spoVID} spores also varied significantly (Fig. 12b; spores with white stars), sometimes having a shape resembling a bowling pin.

The outer and internal surface structures of the coat sacculi released from \textit{spoVID} spores were similar to the outermost surface structure of wild-type spores, as seen in a high-resolution image of a \textit{spoVID} spore sacculus (Fig. 13a), and consisted of rodlet layers (Fig. 13a; red arrows) covered with amorphous material (Fig. 13a; green arrows). As illustrated in Fig. 13b, high-resolution images of the surfaces of the coatless \textit{spoVID} spores revealed a 2–6 nm thick amorphous layer (Fig. 13b; grey arrow) and an underlying pitted surface structure (pink arrow).

![Figure 9. AFM images of \textit{cotE} spores.](image-url)

![Figure 10. AFM height images of \textit{gerE} spores.](image-url)
Figure 11. AFM height images of cotE gerE spores. (a) Spores which appeared to be devoid of spore coat material. Closely packed spores are more deformed than ones that are not surrounded by other spores. Some spores exhibit 80–100 nm wide and 30–40 nm deep depressions (black circles). The insert in (a) is a cross section line (indicated with a white line) drawn across the ~100 nm wide depression showing a depth of ~40 nm. (b) Image showing small patches of coat material (grey arrow) on the spore surface. (c) High-resolution image of a spore devoid of any obvious coat material, and showing a textured outermost surface.

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Figure 12. AFM height images of spoVID spores. (a) Many of the spoVID spores are devoid of obvious spore coat material, although some spoVID spores are encased in loosely fitting coat sacculi (green stars); insert: an empty intact sacculus (blue star) present in a spore preparation. (b,d) Severely deformed spores without any visible coat material are indicated with white stars. (c, d) Spores with partially sloughed off coat sacculi are indicated with grey stars.

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Discussion

Topography of the outer spore surface

The ridges on the surfaces of air-dried *B. subtilis* spores (Fig. 1a) have been seen previously in EM [39,40] and AFM [22–24,29,30] studies of spores of various *Bacillus* species. These ridges have been proposed to form due to the folding of the coat in response to dehydration, likely as a consequence of decreases in spores' internal volume [22,37,38]. Indeed, AFM measurements of the morphology of fully hydrated and air-dried spores demonstrate that surface ridges on dehydrated spores mostly disappear or decrease in size upon hydration [22]. Thus, spore coat flexibility can compensate for decreases in spore surface area upon drying by surface folding and ridge formation [22]. Current work demonstrated that this surface folding takes place in the spore coat, since dry *gerE*, *cotE* *gerE* and *spoVID* spores lacking much of the spore coat exhibit no surface ridges (Fig. 10–12). Note, that while *gerE* and *cotE* *gerE* spores as well as coat rinds produced by protozoal digestion of spores exhibit some coat material [1,28] (Fig. 9,10), the amount of this material is either not sufficient or its proteomic composition is not appropriate to form surface ridges. In contrast, the presence of surface ridges on *cotO*, *cotH* and *cotE* spores lacking the amorphous and rodlet layers (Fig. 5–7) indicates again that surface ridge formation takes place within the multilayer spore coat structure.

The spore coats of *B. subtilis* mutants lacking morphogenetic coat proteins as well as chemically decoated wild-type spores have different thickness and composition, and these variations could affect the coat's elastic properties and thus change its folding and surface ridges. However, affecting the outer spore coat architecture by mutation or chemical decoating gave no large changes in spore surface ridge parameters or patterns. These results suggest that formation of spore surface ridges originates within multilayer coat structures, which are relatively unaffected by loss of some coat proteins, with the amorphous, rodlet and fibrous layers only following the ridge-associated topography. Our data showing pronounced changes in the surface folding of *safA* spores (Fig. 4) may indicate that these spores’ multilayer coat structure is either thinner or more flexible than in wild-type, decoated wild-type, *cotA*, *cotB*, *cotH*, and *cotO* spores (Fig. 1–3,5,6). However, *cotE* spore coats with lower levels of surface folding typically have the same number of layers as do *cotH* and *cotO* spore coats. Perhaps the decreased surface folding of *cotE* spore coats is due to changes in the elastic properties of inner coat layers because one or more inner coat proteins are not assembled in *cotE* spores. Note, that the wide range of surface ridge parameters and folding patterns observed with spores of different species [22–24] and isogenic strains (this work) makes it problematic to assign these parameters as spore species-specific structural attributes.

Spore coat architecture

In addition to providing information on spore surface topography, AFM images allowed construction of a detailed model of coat architecture (Fig. 14). In this model starting from the outside, the coat consists of an amorphous layer/crust, a rodlet layer, a honeycomb layer, a fibrous layer, a nanodot particle layer, the multilayer assembly, and the undercoat/basement layer just above the pitted surface, which we tentatively assign as the spore cortex.
The existence of an outermost tightly fitting spore layer was initially reported in thin section EM images of *B. subtilis* spores treated with a reducing agent [57]. It was suggested that this layer is an exosporium-like structure, which in EM images of untreated spores is usually indistinguishable from the darkly stained outer spore coat. While the amorphous spore coat layer reported here (Fig. 1) could correspond to this outer coat structure, it does not resemble an exosporium, as this outer layer has no paracrystalline basal layer typical of the exosporium of spores of the *B. cereus* group [41,58]. Rather the amorphous layer likely corresponds to the outer crust layer of *B. subtilis* spores that stains with ruthenium red and is glycoprotein-rich [7,8]. Patches of an outermost amorphous layer are also observed in AFM studies of *B. atrophaeus* spores (Plomp and Malkin, unpublished data).

Rodlet structures, similar to ones seen in Fig. 1b,c, were previously described on the outer surface of a diverse set of microorganisms (see [59]), including Gram-negative bacteria and various fungi. The fungal rodlet layers were resistant to treatment by detergents, organic solvents, enzymes, alkali and mild acids [60,61], and the structural proteins hydrophobins [62,63] and chaplinns [64] were integral components of fungal rodlet structures. In several cases, these rodlets had a cross β-structure similar to that in the amyloid fibrils [64] associated with several neurodegenerative diseases [65]. The amyloid-like rodlet fibrils forming microbial outer surface layers appear to play important roles in attachment, dispersal and pathogenesis [59].

Rodlet structures were also reported in EM [38–40] and AFM studies [22–24,30] of spores of several *Bacillus* species, although the proteins that form these rodlet structures are not known. The structural similarities between *B. atrophaeus* rodlets seen during their germination-induced disassembly [25] and amyloid rodlets found on surfaces of fungi and bacteria suggest that *B. subtilis* coat rodlets may also be amyloids. However, full understanding of the function of rodlet structures in spores awaits elucidation.

Interestingly, micro-etch pits form in the rodlet layer early in spore germination [25], and these could facilitate access of degradative enzymes to their targets in an otherwise tightly packed coat. Characterization of the strength and mechanical stiffness of individual amyloid fibrils of insulin reveals that these parameters are similar to those of steel and silk [66]. Thus, the spore coat' rodlet layer could play a role in protecting spores from mechanical stress, and a combination of rodlet and amorphous structures could provide spores with a wide range of physicochemical properties. Indeed, the existence of both hydrophobic (rodlets) and hydrophilic (glycoproteins) structures on the outermost layer might enable spores' successful dissemination as both as air-borne and fully hydrated particles.

Current results indicate that the assembly of the outermost coat layer does not require CotA and CotB, two of the most abundant spore coat proteins [2,14,67–71]. While interactions between CotB and CotG are critical in guiding assembly of the outer coat layer, no coat assembly defect has been observed in cotA or cotB mutants [69,70]. In addition, cotA and cotB mutations have no effects on spore lysozyme resistance, germination [2] or surface appearance (Fig. 3), and CotB is absent from cotH spores [72] yet the outermost coat structure of CotH spores is similar to that of wild-type spores (Fig. 6), including both the rodlet and amorphous layers. Thus, neither CotA nor CotB appear to play important roles in directing the assembly of spores' outer layers. The similar surface ridges on cotA and cotB spores further suggests that loss of these proteins does not significantly alter the elastic properties of the spore coat.

Loss of SafA also does not affect the high-resolution architecture of spores' amorphous and rodlet layers (Fig. 4), consistent with safA spores' lysozyme resistance [2,45]. However, SafA plays an important role in coat assembly, as in many safA spores the coat is loosely attached to the spore (Fig. 4). SafA is localized in the spore cortex near the inner coat, and SafA may help associate the spore cortex and coat [2,73]. The absence of surface ridges on a large portion of safA spores, along with relatively thinner existing surface ridges, also suggest that safA spores' coat is thinner and/or more flexible. This is consistent with EM analyses that indicate the safA spore coat often has 1–2 layers instead of the typical 3–5 layers [45].

### Inner coat structures

In *B. cereus* [22] and *B. atrophaeus* spores [25] the coat's rodlet layer is underlain by a honeycomb structure also observed in *B. subtilis* spores (Fig. 9). Since disordered microporous inorganic substrates can effectively initiate three-dimensional protein crystallization [74], perhaps the spore coat's honeycomb structure represents a biological example of a microporous matrix that facilitates the ordered self-assembly of the coat's rodlet structure. Note that the 8–9 nm periodicity of the *B. subtilis* coat honeycomb layer is similar to periodicities of honeycomb structures for *B. cereus*, *B. thuringiensis* [22] and *C. novyi* NT [31] spores. This indicates that molecular dimensions of proteins forming these honeycomb structures are similar for different bacterial species, and the molecular composition of the honeycomb layer in different bacterial species may thus be similar.

Studies of cotO, cotH and cotE spores revealed consecutive structural layers of granular/fibrous material (cotO spores; Fig. 5), nanodots (cotO and cotH spores; Fig. 5, 6), and multilayer structures (cotO, cotH and cotE spores; Fig. 5–7). While spores of these mutants had the multilayer structure, only cotO spores retained the granular/fibrous structure and cotE spores lacked the nanodot layer. We propose that the granular/fibrous layer represents an outer spore coat layer that appears as a darkly stained irregular layer in EM images [69]. The thickness of this outer coat layer varies significantly in EM images on both the same spore and between spores, consistent with the range of granular/fibrous layer thickness observed on cotO spores.

On wild-type spores and spores of some mutants lacking specific coat proteins (i.e. cotA and cotB), the grainy/fibrous outer coat layer was largely obscured by the rodlet and amorphous layers. However, the force exerted by the AFM probe tip on the outermost spore layer allows visualization of underlying structures, as in the AFM visualization of a cytoskeleton beneath a cellular plasma membrane [75]. AFM phase imaging can probe micro-mechanical properties of sample materials (e.g. viscoelasticity) [76] and map surface inhomogeneity of these properties. Furthermore, when mechanical properties of two layers are significantly different, phase imaging can provide structural information on layers beneath the topmost layer [77]. Thus an irregular grainy layer can often be seen in AFM phase images beneath the outer rodlet structure (Fig. 3c,d), and we suggest that this underlying layer corresponds to a grainy/fibrous outer coat layer (Fig. 5, 414). Note, that an undulating surface morphology similar to that seen on cotA and cotB spores (Fig. 3c,d) was also observed on the surface of wild-type spores (data not shown).

Typically, multilayer structures on cotO, cotH and cotE spores contained 3–5 layers, consistent with the appearance of the lightly stained lamellar inner coat of *B. subtilis* spores seen by EM [68], and thus these multilayer structures may correspond to the *B. subtilis* spores' inner coat. We further suggest that the nanodots between the outer and inner coat layers but absent on cotE spores, might be CotE molecules that facilitate the assembly of the grainy/granular outer coat layer. The height of the smallest nanodots seen
on cotH spores was ~3 nm, consistent with CotE’s mol wt of 20.9 kDa [2], and this suggestion is consistent with current models of B. subtilis spore coat assembly that have CotE positioned between the inner and outer coat layers [2, 26]. However, these nanodots could also be small coat protein aggregates, and further experiments, perhaps using AFM-based immunolabeling techniques [29], will be needed to identify the protein(s) forming the nanodots.

The cotO spores have no amorphous or rodlet layers, which could explain the partial lysozyme sensitivity of cotO spores [26]. However, the presence of these outer layers on the majority of cotH spores (Fig. 6) is consistent with their relatively normal lysozyme resistance [1]. The outer coat of cotO spores often appears disorganized and missing in EM thin sections [26] and is generally indistinguishable from that of cotH spores. CotO and CotH are suggested to be localized below the coat surface [13, 26] and to participate in a late phase of coat assembly. However, our AFM analyses showed pronounced differences between cotO and cotH spore coats. In particular, CotO plays a critical role in the assembly of the amorphous and rodlet layers, while assembly of the fibrous outer coat requires CotH and CotE. AFM studies also indicated that these proteins play a role in assembly of the coat’s amorphous and rodlet layers, consistent with biochemical, genetic and EM studies [26, 46, 72]. It has been suggested [26] that CotO and CotH also play an important role in inhibiting the tendency of outer coat protein layers to stack up resulting in the polymerization of the coat layers into closed shells. However, AFM demonstrates that cotO and cotH spore coats self-assemble to form contiguous shells rather than disorganized coats. At the same time, many cotE spores exhibited only a loose coat sacculus (Fig. 7), indicating that CotE plays an important role in the assembly of the inner coat and/or its attachment to the cortex as noted above.

The crucial role for CotE and GerE in proper coat assembly was further highlighted by the AFM of gerE and cotE gerE spores. First, loss of gerE prevented formation of the outer coat, rodlet, and amorphous layers. Second, while most gerE spores are encased in a loose structure formed by what appeared to be patches of the inner coat (Fig. 10), these structures do not resemble the inner coat multilayer structures described above. The cotE gerE spores were devoid of the amorphous and rodlet layers, and both complete inner and outer coats, and these spores’ surface exhibited some roughness (Fig. 11). This surface likely corresponds to the basement/undercoat layer [4]. Thus, both CotE and GerE are crucial in proper assembly of the inner coat. Note, also that cotE gerE spores are less rigid than cotE or gerE spores. This increased deformability is due either to the loss of the inner coat or a role for CotE in the assembly and elastic properties of the basement layer. The nature of the 80–100 nm wide and 30–40 nm deep depressions seen in Fig. 11a is unclear, but we speculate that these holes may facilitate germinant access to the spore inner membrane, and are perhaps associated with the GerP proteins important in germinant movement through spores’ outer layers [78].

Another coat protein important for proper spore coat assembly and attachment to the cortex is SpoVID, as a large percentage of spoVID spores lacked obvious coat structures, with some encased in a misassembled sacculus composed of amorphous and rodlet structures (Fig. 12). The thickness of the sacculi walls varied between 15–30 nm, indicating that the sacculi could contain coat material in addition to the rodlet and amorphous layers (Fig. 13a). Note that none of the spoVID spores visualized in this study exhibited the multilayer inner coat structures seen on cotO, cotH and cotE spores indicating that the inner coat is absent on spoVID spores. Most spoVID sacculi were only loosely attached to the spore body and were partially sloughed off, exposing a relatively smooth spore surface (Fig. 12). These AFM data are consistent with observations of swirls of spore coat in spoVID mother cells [10] (Fig. 12a, inset) and that SpoVID is required for the stable attachment of the coat. High-resolution imaging of the surface of spoVID spores indicated the existence of two prominent layers (Fig. 13b). One layer (13b; square) could correspond to the basement layer [4] and a pitted layer (Fig. 13b; black arrow) could correspond to either a subbasement coat layer or the cortex. Note, that spoVID spores lacking sacculus exhibit very high deformability (Fig. 12).

During wild-type B. subtilis sporulation, proteins forming honeycomb and rodlet coat layers self-assemble on the outer spore coat layer. Based on AFM results with cotE spores, the complete outer coat layer is not essential for formation of patches of the honeycomb and rodlet coat layers. Thus, the underlying integument is not crucial for assembly of the rodlet and honeycomb layers. Proteins that form honeycomb and rodlet spore coat structures must therefore be present during cotE spore formation, and self-assemble on the outer spore surface producing amorphous and rodlet layers (Fig. 8). Indeed, during B. thuringiensis sporulation rodlet proteins can self-assemble on the underlying spore coat, or in either the mother cell cytoplasm or the sporulation medium [23]. Hydrophobins, which form fungal rodlet layers, also self-assemble into rodlet fibrils in vitro (for review see [59]).

The multilayer structure forming the inner coat of B. subtilis spores exhibits patterns similar to ones described for the inner coats of spores of C. novyi NT [31] and B. anthracis (Plomp and Malkin, unpublished data). These patterns are also similar to those observed on surfaces of inorganic and macromolecular crystals. In addition to growth steps, these patterns include two-dimensional (2D) nuclei and screw dislocations that are major growth sources of inorganic, organic, and macromolecular crystals [79]. The presence of these growth patterns plus the smooth appearances of coat layers strongly point to a crystalline nature [79] of B. subtilis inner coat layers. While no screw dislocation sources similar to ones observed on the C. novyi NT inner spore coat [31] were seen on B. subtilis spores, on some spores with a low density of the grainy/fibrous outer layer, circular 2D nuclei were observed on the inner coat (Fig. 15a; dark blue arrows). This indicates that B. subtilis spores could represent the first case of non-mineral 2D nucleation growth patterns in a biological organism.

The observations above strongly suggest that assembly of inner spore coat layers proceeds by formation of 2D nuclei and their subsequent growth, similar to the birth-and-spread growth mechanism of conventional and macromolecular crystals [51, 79]. In this model, 2D crystal growth takes place by generation and subsequent spread of 2D nuclei that provide a new crystalline layer on crystalline surfaces. Subsequent formation and growth of new 2D nuclei on this layer result in the formation of a new crystalline layer. An example of such growth, showing 2D nuclei on the surface of a crystal of satellite tobacco mosaic virus that are similar to ones seen in Fig. 15a, is presented in Fig. 15b (dark blue arrows). Typically, 2D growth takes place at high supersaturation (e.g. protein and precipitant concentrations used in macromolecular crystallization) [51, 52, 79], suggesting that relatively high concentrations of inner coat protein(s) are present during B. subtilis sporulation.

Step edges seen on the inner coat of B. subtilis spores showed significant roughness with many kinks (Fig. 7b), suggesting that formation of the inner coat was strongly affected by impurities. Similar patterns have been described for a wide range of crystalline surfaces (illustrated in Fig. 7d), where adsorption of
impurities (ones present in solution), but not forming a layer at the step terraces and edges results in step roughening and cessation of growth [80–82]. Indeed the roughness and sinuosity of step edges on the inner coat of *B. subtilis* spores (Fig. 7c) are higher than observed for step edges on the surface of the trypsin crystal (Fig. 7d). This may indicate [80–83] that higher levels of impurities are adsorbed on the inner coat surface of *B. subtilis* spores compared to ones on the surface of trypsin crystals. Growth steps stop at sites of contact with impurity particles [indicated as small balls in Fig. 15c,d] that are adsorbed to the surface. However, portions of steps between neighboring impurity particles continue to grow, resulting in pinning of growth steps (Fig. 15c) as seen in Fig. 7c,d. Step advancement ceases (Fig. 15d) when at increased impurities’ concentration, the distance between impurities/pinning points $d_{\text{imp}}$ becomes smaller than the diameter of critical nuclei $d_c$, necessary for step advancement [80]. One interesting feature of the inner coat is a number of $\sim 5–10$ nm holes (Fig. 7c), which may indicate locations of clusters of impurities [53,83]. Note, that in general the size of such holes is a function of the size of impurities or their clusters adsorbed on the surface. As described for a number of systems [80–83], these clusters of impurities may be responsible for pinning the advancement and cessation of growth of spore coat layers observed in Fig. 7c. Alternatively, such holes that were also observed on inner coat layers of *C. novyi* NT spores [31] and *B. anthracis* spores (Plomp and Malkin, unpublished data) could be an intrinsic feature of spore inner coat layers having a particular function. These results, combined with prior observation of screw dislocations on the inner coat of *C. novyi* NT spores [31], strongly suggest that inner spore coat assembly is governed by two crystallization mechanisms – growth on dislocations and 2D nucleation. These observations suggest that while spore coat proteins are produced enzymatically [84], the assembly of these proteins into coat layers may be a self-assembly process similar to crystallization, and may be influenced by the sporulation conditions (protein and salt concentrations, pH, temperature, impurities) when these proteins assemble.

The lack of high-resolution crystalline lattice structures of the *B. subtilis* inner coat layers is similar to prior observations of *C. novyi* NT [31] and *B. anthracis* inner spore coat layers (Plomp and Malkin, unpublished data). It was suggested that proteins forming the *C. novyi* NT inner coat layers [31] are not globular, but rather peptides ‘standing upright’ in the layers, similar to peptide arrangements found in several organic crystals [85,86]. This hypothesis was based on the fact that for globular proteins, the $\sim 6$ nm height of the inner spore coat layers would not be considerably different in either perpendicular or lateral unit cell parameters, with the latter being amenable for AFM visualization [54]. Based on the lack of molecular scale AFM resolution of the crystalline lattice forming the *B. subtilis* inner coat layer, it is reasonable to suggest that proteins forming the inner coat might be also “standing upright” peptides [31,85,86].

In conclusion, the results presented in this communication provide further understanding of the structure and assembly of the *B. subtilis* spore coat. Furthermore, morphological and structural attributes of *B. subtilis* spores described here could thus serve as a baseline for future studies of effects of sporulation conditions on these structures. In addition, the similarities of some of the new findings with *B. subtilis* spores to findings with spores of *C. novyi* NT and other *Bacillus* species, suggest that the coat structure proposed in this work may generally be similar for spores of all of these species. While there is extensive knowledge of the individual proteins in the spore coat, as well as their location and assembly, there is much less knowledge of precise coat structure. In
particular, the new high-resolution AFM studies have identified a number of new coat structural features, including the nanodots, the fibrous layer, and the terraced multilayer inner spore coat. Based on these results, we propose that the amorphous/crust layer and rodlets form the outermost spore structure, the fibrous layer and multilayer structure correspond to the outer coat and the inner coat respectively, with honeycomb and nanodot structures sandwiched between the outermost layer and the inner coat and the inner and the outer coats respectively.

Note, that high-resolution studies of fully hydrated B. atrophaeus [22,25] and Clostridium novyi NT spores [31] demonstrated that rodlet, honeycomb, and inner coat layer structures, similar to ones described here for B. subtilis, maintained the same patterns, lattice periodicities, and step heights as seen on air-dried spores.

Finally, the striking similarity between the appearance of the terraces and likely 2D nuclei in the multilayer inner coat and in inorganic and macromolecular crystals suggest that at least this part of the coat may assemble by crystallization mechanisms. A consequence of a crystallization spore coat assembly mechanism is that coat structure will be influenced by conditions during which these proteins self-assemble. In particular, variations in rates of 2D nucleation on spores could change the growth rate and hence the thickness of the spore coat, and this could influence spore properties such as their resistance and germination. The challenge now will be to correlate spore coat features identified in this work with specific coat proteins, and to understand how individual proteins contribute to these coat features, in particular, by using AFM-based immunolabeling techniques [29].

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Author Contributions

Conceived and designed the experiments: AJM PS. Performed the experiments: MP AMC PS AJM. Analyzed the data: MP AMC AMC PS. Contributed reagents/materials/analysis tools: MP AMC PS AJM. Wrote the paper: MP PS AJM.

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