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Temperature Dependence of the Proteome Profile of the Psychrotolerant Pathogenic Food Spoiler Bacillus weihenstephanensis Type Strain WSBC 10204

Sacha K. Stelder, Siraje A. Mahmud, Henk L. Dekker, Leo J. de Koning, Stanley Brul, and Chris G. de Koster

ABSTRACT: Bacillus weihenstephanensis is a subspecies of the Bacillus cereus sensu lato group of spore-forming bacteria known to cause food spoilage or food poisoning. The key distinguishing phenotype of B. weihenstephanensis is its ability to grow below 7 °C or, from a food safety perspective, to grow and potentially produce toxins in a refrigerated environment. Comparison of the proteome profile of B. weihenstephanensis upon its exposure to different culturing conditions can reveal clues to the mechanistic basis of its psychrotolerant phenotype as well as elucidate relevant aspects of its toxigenic profile. To this end, the genome of the type strain B. weihenstephanensis WSBC 10204 was sequenced and annotated. Subsequently, the proteome profiles of cells grown at either 6 or 30 °C were compared, which revealed considerable differences and indicated several hundred (uncharacterized) proteins as being subproteome- and/or temperature-specific. In this manner, several processes were newly indicated to be dependent on growth temperature, such as varying carbon flux routes and a different role for the urea cycle. Furthermore, a possible post-translational regulatory function for acetylation was suggested. Toxin production was determined to be largely independent of growth temperature.

KEYWORDS: Bacillus weihenstephanensis, genome sequencing, proteomics, mass spectrometry, food pathogen, toxigenicity, psychrotolerance

INTRODUCTION

Bacillus weihenstephanensis is a spore-forming Gram-positive bacterium known for its ability to cause food poisoning and food spoilage. The phenotype that distinguishes B. weihenstephanensis from other members of the Bacillus cereus group is its psychrotolerant growth profile, i.e., its ability to grow at 4–7 °C but not at 43 °C, with an optimum growth temperature of 30 °C. From a food safety perspective, characterization of this species is important, as the psychrotolerant phenotype introduces an increased potential hazard in an environment reliant on refrigerated storage of products (i.e., 4 °C) for microbial inactivation.

The most important vehicle by which B. weihenstephanensis manages to contaminate food is through (endo)spores. Spores are a metabolically dormant, highly resistant survival form of the cell capable of withstanding common processing techniques used in the food industry such as heat or acid treatment. Although spores themselves are harmless, once the bacteria return to their vegetative state through a process called germination they can resume growth and produce toxins. Beyond its ability to grow at lower temperatures as well those of its close relatives, B. weihenstephanensis has been indicated as being able to produce spores, germinate, and produce toxins at lower temperatures as well. Reports of outbreaks of B. weihenstephanensis are rare, as it is not a specifically reportable illness and is not always diagnosed. Furthermore, little effort is made to characterize isolates enough to distinguish it from B. cereus.

Lechner et al. originally distinguished the species from the rest of the B. cereus sensu lato group based on sequence differences in the 16S rDNA, 23S rDNA, 16S–23S rDNA spacer region, and cspA, the gene encoding the major cold-shock protein homologue. Growth characteristics are not enough to classify isolates, as several variants of B. cereus are, in fact, capable of growth at 6 °C. There is still some controversy as to whether or how the B. cereus sensu lato group should be subdivided into different species, as there is no true phylogenetic basis. While other members of the B. cereus group have been studied in great detail, with genome sequences available for many strains, B. weihenstephanensis falls behind, with only one completed genome and until now no sequence for the type strain.

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In order to make *B. weihenstephanensis* more tractable for studies, the type strain *B. weihenstephanensis* WSBC 10204 (DSM11821) has been sequenced. Annotation of the newly created genome sequence allowed for a complete proteome prediction and subsequent analysis by bottom-up mass spectrometry-based proteomics. In order to assess how *B. weihenstephanensis* copes with low growth temperature and to determine its toxigenic profile, the proteome expressed at growth temperatures of 6 and 30 °C was analyzed. Comparison of the cellular proteome and secretome isolated from late-exponential phase cultures indicated the presence of many temperature-specific proteins. These differentially expressed proteins were related to several processes that were expected to be growth temperature-dependent as well as processes newly correlated with growth temperature.

### MATERIALS AND METHODS

#### DNA Isolation

An overnight culture of *B. weihenstephanensis* WSBC 10204 (available at DSMZ under DSM-11821) grown in 25 mL of tryptic soy broth medium was harvested by centrifugation, and the pellet was resuspended in 2 mL of resuspension buffer (10 mM Tris-HCl, pH 7.5, 50 mM EDTA, 100 mM NaCl). Cells were lysed for 30 min at 37 °C by addition of 2 mg of lysozyme. Two-hundred microliters of 2% SDS was added, and the solution was incubated on ice for 5 min. DNA was extracted by addition of 1 volume of phenol/chloroform. The phases were separated by centrifugation (15 min, 13,000 rpm), and 1/10th volume of 3 M sodium acetate was added to the aqueous phase. DNA was precipitated with 1 volume of cold 100% ethanol, washed once with 70% ethanol, and air-dried before resuspension in water.

#### Genome Sequencing and Annotation

Illumina HiSEQ2500 sequencing and assembly into scaffolds was performed by BaseClear BV, Leiden, The Netherlands. This resulted in a roughly 5.6 MB sequence with a coverage of 400x. Scaffolds were aligned to the *B. weihenstephanensis* KBAB4 sequence [GenBank no. NC_010184.1] to produce a draft genome using MAUVE.10 Residual gaps were closed by PCR amplification and 454 GS FLX+ sequencing, with 1 out of 250 reads aligned to the *B. weihenstephanensis* WSBC 10204 sequence. Annotation of the newly generated draft genome using MAUVE.10 Residual gaps were closed by PCR amplification and 454 GS FLX+ sequencing, with 1 out of 250 reads aligned to the *B. weihenstephanensis* WSBC 10204 sequence. Annotation was compared with the published genome annotation of *B. weihenstephanensis* KBAB4 in order to improve the quality. Annotations were manually verified by comparing against nonredundant databases of NCBI and UniProt. In order to associate a function with a predicted gene, a minimum cutoff of 30% identity and 80% coverage of the gene length were used, checking at least two best hits in nonredundant protein databases. The BLASTp algorithm was used to search for protein similarities to other deposited sequences, following these criteria: >50% similarity at the amino acid level and >50% coverage of protein length.

#### Culturing and Protein Sample Preparation

Duplicate batch cultures of *B. weihenstephanensis* WSBC 10204 were grown in TSB medium at either 6 or 30 °C until late-exponential phase, as determined by optical density at 600 nm (OD600 = 10), synchronizing their growth phase but not growth rate. Cells were harvested by centrifugation and freeze-dried. Fifty milligrams of dried cell pellet was resuspended in 100 mM dithiothreitol, 100 mM Tris-Cl, pH 6.8, and 4% SDS, disrupted by three cycles of sonication consisting of 3 min sonication (50% duty cycle, Output Control 7) and 5 min rest on ice in a Branson Sonifier 250 (Branson Ultrasonics, Danbury, USA), and freeze-dried.

For secretome preparation, the supernatants of duplicate cultures grown as described above were filtered through a 0.2 μm poly(ether sulfone) filter to remove remaining cells. Secretr proteins were precipitated from the supernatant by addition of up to 10% trichloroacetic acid and incubation overnight at 4 °C. Precipitated proteins were washed three times with 95% acetone and freeze-dried.

#### Preparation for Mass Spectrometry (MS) Analysis

Five milligrams of lyophilized cellular extracts was dissolved in sample buffer, loaded onto a 12% SDS-PAGE gel, and run for 2 h at 100 V until the dye ran off the gel. Gels were stained with Coomassie brilliant blue to confirm separation. Gel lanes were excised and cut into approximately 20 slices of equal size. Slices were diced into small pieces and dehydrated by vortexing in 25 mM NH4HCO3, 50% acetonitrile twice for 30 min. After drying fully in a centrifugal vacuum concentrator (miVac DNA concentrator, Genevac Ltd., Ipswich, UK), proteins were reduced by incubation with 10 mM dithiothreitol in 25 mM NH4HCO3 for 60 min at 55 °C followed by alkylation with 55 mM iodoacetamide for 45 min in the dark. Gel pieces were washed with 25 mM NH4HCO3, dehydrated, and dried fully as above. Proteins were subjected to in-gel tryptic digestion by rehydration with 2.5 μg of trypsin in 25 mM NH4HCO3 per fraction and incubation at 37 °C overnight. After digestion, the supernatant containing peptides was collected and peptides were further extracted from the gel by vortexing in 50% acetonitrile/5% formic acid twice for 30 min. The combined supernatants were partially dried to reduce the acetonitrile concentration before desalting, as described below.

Secretr proteins were resuspended in 8 M urea, 0.5 M ammonium bicarbonate, pH 8.0, and 4 M dithiothreitol and incubated at 55 °C for 60 min. Afterward, 25 μL of 20 mM iodoacetamide was added, and the samples were incubated in the dark for 45 min. The samples were diluted 4 times with water to reduce the concentration of urea to avoid inhibition of trypsin, after which 2.5 μg of trypsin was added and proteins were digested at 37 °C overnight.

Tryptic digests were desalted using Omix μC18 tips (80 μg capacity, Varian, Palo Alto, CA) according to the manufacturer’s protocol, with the final elution of peptides in 50% acetonitrile, 0.1% trifluoroacetic acid. Prior to analysis by liquid chromatography mass spectrometry, the peptide concentration was estimated on a Nanodrop ND1000 spectrophotometer (Isogen Life Sciences, De Meern, The Netherlands) at 205 nm.

#### Liquid Chromatography–Fourier Transform–Ion Cyclotron Tandem Mass Spectrometry (LC–FT–ICR MS/MS) Analysis

LC–MS/MS data were acquired with a Bruker ApexUltra Fourier transform ion cyclotron resonance mass spectrometer (Bruker Daltonics, Bremen, Germany) equipped with a 7 T
magnet and a nanoelectrospray Apollo II DualSource coupled to an Ultimate 3000 (Dionex, Sunnyvale, CA) high-performance liquid chromatography system. For each digest, samples containing up to 200 ng of tryptic peptides were injected as a 20 µL 0.1% trifluoroacetic acid, 3% acetonitrile aqueous solution and loaded onto a PepMap100 C18 (5 µm particle size, 100 Å pore size, 300 µm inner diameter × 5 mm length) precolumn at a flow rate of 50 µL/min. Following injection, the peptides were eluted via an Acclaim PepMap 100 C18 (3 µm particle size, 100 Å pore size, 75 µm inner diameter × 250 mm length) analytical column (Thermo Scientific, Etten-Leur, The Netherlands) to the nanoelectrospray source. For analyses of the third biological replicate, a similar 500 mm column at 60 °C was used. LC gradient profiles of up to 120 min were obtained using 0.1% formic acid/99.9% H2O (A) and 0.1% formic acid/80% acetonitrile/19.9% H2O (B) (0 min, 97% A/3% B; 2 min, 94% A/6% B; 110 min, 70% A/30% B; 120 min, 60% A/40% B; 125 min, 100% B) at a flow rate of 300 nL/min. Data-dependent Q-selected peptide ions were fragmented in the hexapole collision cell at an argon pressure of 6 mbar (measured at the ion gauge), and the fragment ions were detected in the ICR cell at a resolution of up to 60 000. In the MS/MS duty cycle, three different precursor peptide ions were selected from each survey MS. The MS/MS duty cycle time for one survey MS and three MS/MS acquisitions was about 2 s. Instrument mass accuracy was better than 5 ppm over an m/z range of 250 to 1500.

Raw FT-MS/MS data were processed with the MASCOT DISTILLER program, version 2.4.3.1 (64 bit), MDRO 2.4.3.0 (MATRIX science, London, UK), including the Search toolbox and the Quantification toolbox. Peak picking for both MS and MS/MS spectra were optimized for the mass resolution of up to 60 000. Peaks were fitted to a simulated isotope distribution with a correlation threshold of 0.7 and with a minimum signal-to-noise ratio of 2. For each replicate, the processed data of gel fraction digests were combined and searched in a MudPIT (multidimensional protein identification technique) scoring approach with the MASCOT server program 2.3.02 (MATRIX science, London, UK) against the newly created B. weihenstephanensis WSBC 10204 predicted proteome database. Processed data from the secretome digests were searched separately. The database was complemented with its corresponding decoy database for statistical analyses of peptide false discovery rate. Trypsin/P was used as enzyme, and one missed cleavage was allowed. Carbamidomethylation of cysteine was used as a fixed modification. The peptide mass tolerance was set to 40 ppm, and the peptide fragment mass tolerance was set to 0.03 Da. MASCOT MudPIT peptide identification score was set to a cutoff of 20. At this cutoff and based on the number of assigned decoy peptide sequences, the significance threshold was adjusted to obtain a peptide false discovery rate of 2% for all analyses. The MASCOT protein identification reports were exported as XML and then imported in a custom-made VBA software program running in Microsoft Excel. The program facilitates organization and data mining of large sets of proteomics data.

The identified cellular and secretome fraction-associated proteins for the three replicates grown at 6 and 30 °C are listed in Supporting Information Table S1, together with their protein MASCOT scores and the number of peptide spectrum matches. The mass spectrometry proteomics data have been deposited to the ProteomeXchange Consortium via the PRIDE partner repository with the data set identifier PXD002007 and 10.6019/PXD002007.

RESULTS AND DISCUSSION

Sequencing and Annotation

Illumina sequencing resulted in a 5.6 MB sequence with an overall GC content of 35.3%. The genome carries 5678 coding sequences (CDSs), with 77 tRNA and 9 rRNA genes. Of the predicted CDSs, 4697 (82.7%) have a predicted biologically known function. The rest are either similar to hypothetical proteins in other genomes or have no substantial similarity to other predicted proteins. The CDSs with known biological functions were also classified into TIGR (The Institute for Genomic Research) biological roles for further analysis.

The draft sequence contained 44 contigs and 35 scaffolds. Mapping the scaffolds against B. weihenstephanensis strain KBAB4 (GenBank no. NC_010184.1) using MAUVE allowed for PCR primers to be designed for genome closing. Using LR-PCR, fragments were obtained for all of the assumed gaps. Further sequencing and annotation of the resulting fragments revealed the presence of tRNA operons. However, one gap between two scaffolds remained resistant to sequencing, likely due to a high number of repeats and/or secondary structure of the DNA or a possible insertion of a phage sequence. Alignment with the published sequence of B. weihenstephanensis strain KBAB4 indicated that this region of the genome consists exclusively of tRNA sequences. This allowed for a full proteome prediction of the type strain of B. weihenstephanensis. Automatic annotation by IGS Annotation Engine11 predicted 5723 encoded proteins. Manual curation and start codon correction were executed using the Manatee annotation tool.13 The high coding density (82.7%) also accounted for the higher number of protein coding sequences in this strain as compared to those in strain KBAB4 (5155), which indicates that genomic rearrangements have taken place as a result of the acquisition of mobile genetic elements. This insertion might be involved in niche adaptation and contribute to the pathogenicity of the current strain.17

Proteome Profiling and Categorization

To assess how B. weihenstephanensis responds to growth at low temperatures, the proteome profile was determined at 6 and 30 °C. Proteins were identified in three replicates of cellular isolates and three replicates of secreted protein isolates for each temperature. A total of 2112 individual proteins were detected at least once, across all isolates and replicates. Predicted proteins were grouped to TIGR role categories automatically by the Manatee annotation tool,13 with proteins not attributed to any category being listed as uncategorized. The number of detected proteins per category is displayed in Figure 1. This indicated that the obtained proteome profiles covered a very large variety of processes. In particular, the majority of proteins detected were ascribed to growth-related processes, as is to be expected from late-exponential phase cell cultures. These categories included Nucleotide Synthesis, covering purine and pyrimidine metabolism; Protein Synthesis, which covers for instance ribosomal proteins and tRNA ligases; and Energy Metabolism, which includes glycolysis, the TCA (tricarboxylic acid) cycle, the pentose phosphate pathway, etcetera.

A protein was considered to be positively identified if it was detected in at least two replicates. Specific localization of identified proteins, either cellular, secreted, or both, is indicated...
in both subproteomes. A total of 976 proteins were specifically identified, with 88 proteins being secretome-specific. Proteins that were not assigned a category are listed as uncategorized. 2112 proteins were detected spanning all TIGR categories. Proteins that were not appointed a category are listed as uncategorized.

Figure 2. Subproteome localization of identified proteins. Identified proteins are indicated according to the isolate(s) in which they were detected. A total of 976 proteins were specific to the cellular proteome, 88 proteins were secretome-specific, and 262 proteins were identified in both subproteomes.

Identified proteins divided into the same categories as above are indicated according to their growth temperature predominance and/or specificity in Figure 3. A total of 505 proteins showed temperature-specific behavior, being identified in isolates from either 6 or 30 °C exclusively. Another 129 proteins showed a predominance at 6 or 30 °C, determined as follows. In label-free mass spectrometric quantification methods, more abundant peptides are assumed to be selected for fragmentation more often, thereby increasing the number of times peptides are identified for abundant proteins in a sample. It therefore serves as an indication of temperature-dependent expression of proteins. Using this spectral counting approach, the number of peptide spectrum matches for a given protein was normalized to the total number of matches in each singular analysis to correct for variation in sample quality. Subsequently, the sum spectral count was averaged for all three replicates and compared between the two experimental conditions. Proteins with a 3-fold difference in their associated spectral count were considered to be predominant at a certain temperature. Individual proteins with MASCOT scores and number of peptide spectrum matches are listed in Supporting Information Table S1. Proteins from across all categories were influenced by growth temperature to a certain extent.

Growth Temperature-Dependent Processes

The batch culture setup used for acquiring cell material allowed for a difference in growth rate as a result of the imposed difference in growth temperature. In order to separate a growth temperature effect from a growth rate effect, it is imperative to find some sort of normalization and establish whether cells are in the same growth phase. A good indicator for this would be the ribosomal protein content, which can be considered to be proportional to the growth rate. If the ribosomal proteins are present in similar amounts, then the inclination of the cell to grow and divide can be considered to be the same as well, although actual growth rate may still differ as a result of the temperature dependence of ribosomal activity. By harvesting all cell cultures in the late-exponential phase and determining the ribosomal protein content to be largely temperature-independent (Supporting Information Table S1), it was concluded that cells were in the same growth phase, i.e., maximum growth rate for that respective temperature. This allowed for an appreciation of growth temperature-dependent effects despite the growth rate difference.

Upon encountering a reduction in temperature, bacteria need to adjust in order to maintain viability and/or resume growth. Toward this end, several hurdles must be overcome, including hampered translation, increased superhelicity of DNA, reduced membrane fluidity, and others. Several proteins related to these processes were identified as being expressed in a growth temperature-dependent manner. Major cold-shock proteins CspA (Gene ID: bwe1_3875, GenBank accession no. AIW86483) and CspE (bwe1_2608, AIW85234) were found to be predominantly or exclusively present at 6 °C and are known to aid in preventing premature folding, which can occur during translation at low temperatures. DNA gyrase subunit B (bwe1_4278, AIW86876) was predominant at 6 °C as well, allowing for restoration of correct superhelicity of DNA. Furthermore, enzymes responsible for membrane remodelling, such as fatty acid desaturase (bwe1_2028, AIW84669) and a short-chain specific acyl-CoA hydrolase (bwe1_1913, AIW84555) were identified in 6 °C isolates exclusively,
indicating a tendency toward shorter unsaturated fatty acids, which should result in a more fluid membrane. These findings indicated that known candidate proteins related to a cold shock or cold adaptation response were indeed detected.

Beyond known candidates, several processes were newly indicated to be correlated to growth temperature. Looking at Energy Metabolism, the proteome profiles showed a few differences. Fermentative pathways, in the form of several lactate dehydrogenases (bwei_0009, AIW82688; bwei_3088, AIW85707; bwei_4768, AIW87353) and pyruvate formate lyase (bwei_5272, AIW87817), were identified exclusively or predominantly in 30 °C isolates, with pyruvate formate lyase showing the clearest semiquantitative difference among all proteins identified. However, enzymes pertaining to glycolysis and the TCA cycle were present irrespective of temperature. This points to the notion that oxygen limitation is likely not a major factor, as aerobic metabolism presumably takes place under the same conditions. Rather than a direct growth temperature effect, induction of fermentative pathways may be a response to alleviate higher levels of pyruvate. This, in turn, may be caused by a higher flux through glycolysis as a result of higher enzyme activity at 30 °C. In essence, the data indicated that a bacterial Crabtree effect occurs at 30 °C, where glucose consumption cannot be matched by full aerobic conversion of pyruvate, diverting carbon flux to lactate instead.

Looking at cold-induced Energy Metabolism on the other hand, a glyoxylate bypass of the TCA cycle in the form of isocitrate lyase (bwei_3878, AIW86486) and malate synthase (bwei_3879, AIW86487) was identified in 6 °C isolates only. Isocitrate can cleave isocitrate into succinate and glyoxylate, after which succinate is readily used in the remainder of the TCA cycle. The glyoxylate, however, first needs to be condensed by malate synthase using an acetyl group from acetyl-CoA to form malate before this can re-enter the TCA cycle. During adjustment of the membrane to low temperatures, fatty acid chains are shortened to increase fluidity, as mentioned above. The glyoxylate bypass could be induced to enable the cell to partition the acetyl fragments released during fatty acid chain degradation for either anabolic processes or full oxidation.

As displayed in Figure 4, most urea cycle related proteins were detected exclusively in 30 °C isolates. In fact, these proteins were among the most dominant proteins identified at 30 °C and, together with pyruvate formate lyase mentioned above, showed the clearest growth temperature dependence (Supporting Information Table S1). Of the proteins involved in this cycle, only arginase (bwei_5574, AIW88091) deviated from this behavior, being identified exclusively in 6 °C isolates. The mechanistic rationale behind this remains unclear, but it appears that the urea cycle is used in a different manner depending on the growth temperature. The net result of the activity of the proteins identified at 30 °C would be production of fumarate, whereas at 6 °C, the suggested end product is urea. Perhaps, in line with the oversaturation of the TCA cycle mentioned above, at 30 °C the urea cycle is used to feed fumarate into the TCA cycle to allow it to keep up better with high pyruvate production. Conversely, urea is known for its chaotropic capabilities, which might be beneficial at low temperatures. As a so-called compatible solute, it may counteract the tendency of macromolecules to rigidify and/or aggregate at lower temperatures, thereby increasing the bacterium’s viable growth temperature range. To put this in perspective, in the related species *Bacillus subtilis*, Budde et al. have shown that arginine biosynthetic pathways were in fact chill-induced at the transcriptional level. Here, an opposite effect is observed at the translational level for arginosuccinate synthase and lyase in *B. weihenstephanensis*. Apart from the species differences, this may have resulted from a difference in growth medium, as a minimal growth medium was used in the Budd study compared to the rich culture medium used here. Interestingly, conversion of urea to ammonium and CO2 was shown to be chill-induced in *B. subtilis* as well, yet *B. weihenstephanensis* lacks the urease required for this process. This may signify the alternative strategies available to these organisms that allow one to be psychrotolerant while the other is mesophilic.

A very striking discovery was the fact that a substantial number of acetyltransferases with undetermined substrate specificity were detected (Table 1). Acetylation of proteins by acetyltransferases can occur at α- or ε-amino groups and is known to be a key post-translational regulatory mechanism in eukaryotes. In prokaryotes, this has only recently gained some traction. In *Escherichia coli*, studies have shown that certain stimuli (hypoxic growth) can induce altered acetylation profiles, as monitored by immunoblotting with an acetyl-lysine-specific antibody. Furthermore, changes in growth phase were also linked to changes in acetylation. Although the specific role of acetylation remains unclear, there have been indications that carbon source utilization and metabolic flux can be regulated by acetylation. As most of the identified acetyltransferases were temperature-specific (Table 1), this indicated a possible regulatory function. MASCOT searches of the data set using protein N-terminal acetylation and lysine acetylation as a variable modification indeed revealed acetylated peptides. However, no clear relation to growth temperature was apparent, as both modified and unmodified versions of the same peptides were detected at both temperatures (data not shown). Modified peptides are suggested to be of low abundance, in addition to the modification being transient, thus requiring dedicated methods for analysis beyond the profiling approach used in this study. The role that acetylation and the temperature-specific acetyltransferases play in *B. weihenstephanensis* remains to be determined.
Toxicogenic Profile Is Largely Temperature-Independent

From a food safety perspective, it is important to highlight the toxicogenic profile. *B. weihenstephanensis* can potentially produce a variety of toxins associated with diarrheal disease: hemolysin BL (Hbl), nonhemolytic enterotoxin (Nhe), and cytotoxin K. In this strain, two copies of the genes encoding the homologous three-component pore-forming cytotoxins Hbl and Nhe have been found. As displayed in Table 2, component proteins of both copies of Hbl were identified in the secretome regardless of the growth temperature, with copies of hemolysin BL lytic components L1 and L2 being exclusive or predominant at 30 °C. On the other hand, not all components of Nhe were detected, with only one of the copies of NheC (*bwei_3144, AIW85763*) being identified at 6 °C.

Lindbäck et al. have previously shown maximal cytotoxic activity when NheA, NheB, and NheC were incubated in a molar ratio of 10:10:1. This implies that NheC is likely to be present at much lower levels than the other components, thereby evading the detection methods used in this study. Gohar et al. were also unable to detect NheC in the secretome of *B. cereus* using two-dimensional gel electrophoresis combined with MALDI-TOF MS. NheA and NheB were shown to have low toxic activity in the absence of NheC, implying that, regardless of whether NheC is not present or simply not detected due its low abundance, *B. weihenstephanensis* WSBC 10204 is indeed toxicogenic at low growth temperatures.

### Table 1. Growth Temperature Dependence of Identified Acetyltransferases

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### Table 2. Toxins Produced by *B. weihenstephanensis* WSBC 10204

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<td>Hemolytic enterotoxin NheA</td>
<td><em>bwei_3146</em></td>
<td>AIW85765</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>Hemolytic enterotoxin NheB</td>
<td><em>bwei_3145</em></td>
<td>AIW85764</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>Hemolytic enterotoxin NheC</td>
<td><em>bwei_3144</em></td>
<td>AIW85763</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Nhe2</td>
<td>Hemolytic enterotoxin NheA</td>
<td><em>bwei_5368</em></td>
<td>AIW87907</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>Hemolytic enterotoxin NheB</td>
<td><em>bwei_5367</em></td>
<td>AIW87906</td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td></td>
<td>Hemolytic enterotoxin NheC</td>
<td><em>bwei_5366</em></td>
<td>AIW87905</td>
<td>–</td>
<td>–</td>
</tr>
</tbody>
</table>

†, identified; ‡, predominant; §, not identified.

### CONCLUSIONS

The psychrotolerant food spoiling pathogenic bacterium *B. weihenstephanensis* poses a significant health risk due to its propensity to infect and survive in food processing lines. This, combined with its ability to grow and potentially produce toxins at lower temperatures than related species, begets the questions of how it manages to maintain growth in a refrigerated environment and if it indeed produces toxins under such conditions. To open up molecular investigations into the behavior of *B. weihenstephanensis*, the type strain WSBC 10204 (DSM11821) has been sequenced and annotated. Complete proteome prediction followed by analysis of the cellular proteome and secretome profiles at 6 and 30 °C revealed differential expression of proteins related to processes known to be induced at low temperatures, such as the cold-shock response, homeoviscous adaptation of the membrane, or adjustment of superhelicity of DNA. Furthermore, new processes have been indicated to be dependent on growth temperature, in the form of varying carbon flux as well as a different role for the urea cycle. Finally, a possible post-translational regulatory function via protein acetylation has been suggested as a result of growth temperature-dependent expression of a large variety of acetyltransferases. Toxin production was found to be largely independent of growth temperature, with encoded tripartite toxins showing only partial differential expression of component proteins. Regardless, toxins were produced at refrigerated temperatures, once again stressing the importance of proper control and further characterization of this species.

### ASSOCIATED CONTENT

#### Supporting Information

Table S1: All proteins identified in cellular isolates and secretome isolates of *Bacillus weihenstephanensis* type strain WSBC 10204 grown at 6 and 30 °C with the accumulated MASCOT scores and peptide spectrum matches of each protein per isolate. This material is available free of charge via the Internet at http://pubs.acs.org.

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**Author Contributions**

§S.B. and C.G.d.K. contributed equally to this work and participated in the design and coordination of the study and helped to draft the manuscript. S.K.S. isolated the DNA for *B. weihenstephanensis* WSBC 10204 grown at 6 and 30 °C. S.B. and C.G.d.K. contributed equally to this work and participated in the design and coordination of the study and helped to draft the manuscript. S.K.S. isolated the DNA for *B. weihenstephanensis* WSBC 10204 grown at 6 and 30 °C.
sequencing, performed the proteomics part of the study, and wrote the manuscript. S.A.M. closed the genome and performed the annotation and curation. H.L.D. assisted in setting up the protein isolation and fractionation strategy. L.J.d.K. advised in devising the experimental setup and helped with MS analyses, interpretation of the data set, and drafting the manuscript.

**Notes**
The authors declare no competing financial interest.

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**ABBREVIATIONS**
MS, mass spectrometry; CDS, coding sequence; TIGR, The Institute for Genomic Research; TCA, tricarboxylic acid; Hbl, hemolysin Bl; Nhe, nonhemolytic enterotoxin; LC−MS, mass spectrometry; CDS, coding sequence; TIGR, The Prokaryotes; FC-MassSpec, Fourier transform–ion cyclotron tandem mass spectrometry; MudPIT, multidimensional protein identification technique

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