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1	Title
2	Analysis of acid-stressed Bacillus cereus reveals a major oxidative response and inactivation-
3	associated radical formation
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5	Running title
6	Acid-induced radical formation in <i>B. cereus</i>
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1 Summary

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3 Acid stress resistance of the food-borne human pathogen Bacillus cereus may contribute to its 4 survival in acidic environments, such as encountered in soil, food, and the human 5 gastrointestinal tract. The acid stress responses of B. cereus strains ATCC 14579 and ATCC 6 10987 were analysed in aerobically grown cultures acidified to pH values ranging from pH 7 5.4 to pH 4.4 with HCl. Comparative phenotype and transcriptome analyses revealed three 8 acid stress-induced responses in this pH range: growth rate reduction, growth arrest and loss 9 of viability. These physiological responses showed to be associated with metabolic shifts and 10 the induction of general stress response mechanisms with a major oxidative component, 11 including up-regulation of catalases and superoxide dismutases. Flow cytometry analysis in 12 combination with the hydroxyl (OH·) and peroxynitrite (ONOO⁻) -specific fluorescent probe 13 3'-(p-hydroxyphenyl) fluorescein (HPF), showed excessive radicals to be formed in both B. 14 cereus strains in bactericidal conditions only. Our study shows that radicals can indicate acid-15 induced malfunctioning of cellular processes that lead to cell death.

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1 Introduction

2

3 Bacillus cereus is a Gram-positive, spore-forming, facultative anaerobic, rod-shaped food-4 borne human pathogen that appears to be well-equipped to survive in various adverse 5 conditions. The spores and vegetative cells of B. cereus can be found in a range of 6 environments, such as soil (Von Stetten et al., 1999; Vilain et al., 2006), plant rhizosphere 7 (Berg et al., 2005), and various foods (Choma et al., 2000; Rosenquist et al., 2005). Besides 8 being notorious for causing spoilage of dairy products, B. cereus is a food-borne pathogen 9 that can cause two distinct types of disease, i.e., emesis and diarrhoea (Kotiranta et al., 2000). 10 The emetic syndrome occurs upon ingestion of the heat-stable toxin cereulide, which is 11 produced in food by emetic B. cereus strains (Agata et al., 2002). The diarrheal syndrome is 12 associated with the action of enterotoxins, such as non-haemolytic enterotoxin (NHE) and 13 cytotoxin K (CytK) (Granum and Lund, 1997), that are produced by vegetative cells inside 14 the human small intestine (Stenfors Arnesen et al., 2008). Before entering the small intestine 15 and subsequent production of enterotoxins, B. cereus cells have to survive the low pH of the 16 human stomach. Therefore, acid resistance is a key parameter in the pathogenic potential of 17 enterotoxic B. cereus strains. Obviously, the highly resistant dormant spores of B. cereus can 18 pass the stomach unaffected and germination in the acid environment of the small intestine is 19 an important aspect of their pathogenic potential (Wijnands et al., 2007; Hornstra et al., 20 2009). Outside the human host, B. cereus may also be frequently exposed to acidic conditions 21 including a range of low pH foods, where in specific cases organic acids have been added as 22 preservatives (Brul and Coote, 1999). In conclusion, coping with low pH stress is an 23 important feature in the performance of B. cereus in a variety of environments as described 24 above, but also in other ecological niches such as soil and plant rhizosphere (Neumann and 25 Martinoia, 2002).

1 Acid stress responses have mainly been studied in Gram-negative organisms, such as 2 Escherichia coli and Salmonella Typhimurium (Richard and Foster, 2003), and in a select 3 number of Gram-positive bacteria, such as lactic acid bacteria and Listeria monocytogenes 4 (van de Guchte et al., 2002; Cotter and Hill, 2003; Ryan et al., 2008). These reviews highlight 5 the importance of proton pumps, i.e., F₁F₀-ATPase, transcriptional regulators, such as RpoS (Gram-negatives) and σ^{B} (Gram-positives), proteins involved in protection 6 of 7 macromolecules, such as DnaK and GroESL, and enzymes that produce alkaline compounds, 8 such as the ammonium forming enzymes urease and arginine deiminase. In contrast, the acid 9 stress response of B. cereus has not been studied extensively. Available information is limited to alternative sigma factor σ^{B} expression upon exposure to a low pH (van Schaik *et al.*, 10 11 2004), and the acid tolerance response, which includes modulation of intracellular pH and 12 protein synthesis (Browne and Dowds, 2002; Jobin et al., 2002; Thomassin et al., 2006). 13 Additionally, the role of urease in acid resistance of a large number of B. cereus strains has 14 been studied (Mols and Abee, 2008), and revealed that its role in acid resistance of B. cereus 15 was limited.

16 Therefore, we set out to investigate the molecular mechanisms involved in acid stress 17 response of B. cereus and to identify possible acid-induced inactivation mechanisms, by 18 comparing responses of cells exposed to selected pHs leading to mild, bacteriostatic and 19 bactericidal acid stress. To determine both general and phenotype-associated transcriptional 20 responses, two model strains ATCC 14579, isolated from air (Ivanova et al., 2003) and ATCC 21 10987, a food-isolate (Rasko et al., 2004) were investigated. Recently, Kohanski and 22 colleagues (2007) reported that the formation of reactive oxygen species (ROS), such as 23 hydroxyl radicals (OH-), plays a role in antibiotic-induced inactivation of aerobically grown 24 E. coli and Staphylococcus aureus cells. These ROS were suggested to originate from 25 antibiotic-induced perturbation of the electron transfer chain resulting in the production of

1 superoxide (O_2) . O_2 can damage iron-sulphur clusters and subsequently react with the 2 released iron, resulting in OH. formation via the Fenton reaction. Therefore, flow cytometry 3 analysis, in combination with the OH· and peroxynitrite (ONOO⁻) -specific fluorescent probe 4 3'-(p-hydroxyphenyl) fluorescein (HPF) (Setsukinai et al., 2003), was included in our study to 5 detect ROS in (sub)lethally acid-stressed B. cereus cells. Our study provides evidence that 6 radicals can indicate acid-induced malfunctioning of cellular processes and the stress-induced 7 formation of reactive oxygen species as a common theme in bacterial stress response and 8 cellular death is discussed.

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10 Results

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12 Physiological response to acid stress

13 The physiological response to acid stress was studied using B. cereus strains ATCC 14579 14 and ATCC 10987 by acidifying aerobically grown cultures to pH values ranging from pH 5.4 15 to pH 4.4 by addition of HCl (Fig. 1). Upon exposure to the different acid shocks, the growth 16 of exponentially growing B. cereus cells was instantly affected. The two strains used showed 17 different phenotypic responses to different levels of acidity. B. cereus strain ATCC 14579 18 showed to continue growth as reflected in an increase of colony forming units upon exposure 19 to pH shocks as low as pH 5.0. This response is hereafter referred to as growth phenotype. 20 Strain ATCC 14579 was inactivated at pH 4.6 and lower as shown by the inability to form 21 colonies on BHI plates incubated at 30°C for 16 hours. This response is hereafter referred to 22 as inactivation phenotype and the condition as bactericidal. Upon exposure to pHs between 23 pH 5.0 and 4.7, ATCC 14579 showed a stable number of viable counts within the first hour of 24 exposure. This response is hereafter referred to as survival phenotype and the condition as 25 bacteriostatic. However, prolonged exposure (overnight) resulted in a decrease of viable

counts (Fig. 2). The growth boundary of the other strain tested, *B. cereus* ATCC 10987, was determined at pH 5.0. Upon exposure to pHs higher than pH 5.0, ATCC 10987 was able to grow and at pHs lower than pH 5.0 this strain was inactivated. There was no apparent survival phenotype in the ATCC 10987 acid shock response using 0.1 pH unit intervals, as observed for ATCC 14579 between pH 5.0 and pH 4.7. The display of an intermediate physiological survival response by strain ATCC 14579 over the pH range 4.7 to 5.0, and the different inactivation boundaries were the main differences between the two strains tested.

8

9 Microarray analysis using hierarchical clustering

10 To investigate the impact of mild, bacteriostatic and bactericidal acid shocks on the gene 11 expression of the two B. cereus strains, four pHs were selected based on the different 12 phenotypic responses displayed by the two strains (Fig. 2A and 2B). At 0, 10, 30 and 60 minutes after the exposure to the different pH shocks, i.e., pH 5.4, pH 5.0, pH 4.8 and pH 4.5, 13 14 RNA samples were collected and subsequent microarray analyses were performed. To 15 compare the transcriptomes of both strains, data obtained of orthologous genes that are 16 present in the genomes of both ATCC 14579 and ATCC 10987 (Ivanova et al., 2003; Rasko 17 et al., 2004; Mols et al., 2007) were collected and subjected to hierarchical clustering. The 18 transcriptome profiles clustered in two different groups, with one cluster including samples 19 obtained of the growth phenotype and the other cluster encompassing samples of the survival 20 (strain ATCC 14579 only) and inactivation phenotypes (Fig. 3). The transcriptome profiles of 21 cultures that showed growth after acid shock exposure clustered together independent of the 22 exposure time and strain. Within these two major groups, the different branches of the 23 hierarchical clustering were separated mostly depending on strain and exposure pH rather 24 than exposure time. In conclusion, the exposure of B. cereus ATCC 14579 and ATCC 10987

to mild, bacteriostatic and bactericidal acid stress led to phenotype specific transcriptome
profiles independent of exposure time.

The pH- and phenotype-specific responses were investigated by analyzing the transcriptome data obtained for genes showing significant differential expression in one or more conditions per strain. Groups of genes with similar expression profiles were identified using hierarchical clustering (*Supplementary material*). The results obtained for a selection of genes putatively involved in low pH or oxidative responses are presented and discussed below.

8

9 Acid shock response of low pH associated genes

10 A selection of genes, based on their putative role in acid stress response of other Gram-11 positive organisms (Cotter and Hill, 2003; Ter Beek et al., 2008), was monitored profoundly. 12 The selection includes transcription regulators, proton pumps, glutamate decarboxylase, 13 production of alkaline compounds, protection of macromolecules, membrane synthesis, and 14 multidrug transporters. The ratios of these genes obtained from cells exposed to pH 5.4 and 15 pH 4.5 for 10, 30 and 60 minutes were averaged and plotted per gene (Fig. 4). In general, the 16 average ratios, showing the up- or down-regulation, were less pronounced in inactivated cells. The genes encoding sigma factors σ^{B} and σ^{H} , involved in the global adaptive response to 17 18 stress, were slightly up-regulated in growing cells of both strains. On the other hand, *codY*, 19 which is a key regulator in the nutrient starvation response of Gram-positive organisms, 20 showed no significant up-regulation. The major oxidative stress response regulator *perR* was 21 one of the most up-regulated transcription regulators in both growing and inactivated cells, 22 indicating an oxidative response upon low pH exposure. Previously, proton pumps, i.e., F₁F₀-23 ATPase, were shown to contribute to pH homeostasis in fermenting Gram-positives exposed to mild acid conditions (Cotter and Hill, 2003). In this study, genes encoding subunits of the 24 F_1F_0 -ATPase (represented by *atpA* in Fig. 4) were highly down-regulated in aerobically 25

1 grown and exposed *B. cereus* cells upon exposure to sub-lethal pHs. Upon exposure to lethal 2 acid shocks, genes encoding sodium-proton antiporters napA and nhaC were not down-3 regulated and napA even showed to be up-regulated. In Listeria monocytogenes (Cotter et al., 4 2001) and Lactococcus lactis (Sanders et al., 1998) acid-induced glutamate decarboxylase 5 (gad), which catalyzes the decarboxylation of glutamate with concomitant consumption of 6 protons, was found to play an important role in low pH survival. In B. cereus ATCC 10987, 7 however, the gad gene, that is not present in the genome of ATCC 14579, showed not to be 8 up-regulated upon low pH exposure. This is in line with the notion that B. cereus ATCC 9 10987 lacks a glutamate/GABA exchanger (Mols et al., 2007), that is required to supply 10 glutamate decarboxylase with its substrate (Cotter and Hill, 2003). Alkaline compound 11 forming mechanisms, such as the arginine deiminase (ADI) pathway and the urease enzyme, 12 are involved in acid tolerance of Gram-positive organisms (Cotter and Hill, 2003). Arginine 13 deiminase (arcA), which is involved in acid resistance of streptococci (Curran et al., 1995) 14 and L. monocytogenes (Ryan et al., 2009), showed significant up-regulation in both B. cereus 15 strains upon exposure to sub-lethal acid shocks, whereas exposure to bactericidal acid shocks 16 revealed no significant induction. Urease encoding genes, specific for ATCC 10987 (Mols et 17 al., 2007), were induced upon exposure to pH 5.4 (represented by ureA in Fig. 4A and 4B), 18 but not in bactericidal conditions. Macromolecules are easily damaged during stress exposure, 19 and their protection and repair is crucial for bacterial survival. DnaK and GroES are 20 chaperones, preventing misfolding of proteins, and in Streptococcus mutans deletion of these 21 chaperones resulted in less resistant cells (Lemos et al., 2001). Notably, chaperone encoding 22 genes *dnaK* and *groES* and protease encoding gene *clpC* were up-regulated upon exposure to 23 sub-lethal acid shocks, whereas exposure to lethal pHs did not induce these genes in B. 24 cereus.

1 Mild sorbic acid stress induces the expression of the fatty acid biosynthesis genes (*fab*), *bkdR* 2 and a multidrug transport gene in *B. subtilis* (Ter Beek *et al.*, 2008). Two homologous 3 multidrug systems (*mdr1* and *mdr2*) showed to be also up-regulated in sub-lethal inorganic 4 acid stress in *B. cereus*. In contrast to sorbic acid stressed *B. subtilis*, fatty acid biosynthesis 5 (represented by *fabF* in Fig. 4A and 4B) was down-regulated at pH 5.4 and no significant 6 induction was found for branched-chain fatty acid biosynthesis (*bkdR*) in *B. cereus* upon 7 exposure to (sub)lethal inorganic acid stress.

8

9 Oxidative response and rearrangements in energy metabolism

The response of several genes involved in oxidative stress and energy production were 10 11 investigated in more detail (Fig. 5). Two distinct types of cytochrome oxidases showed 12 different expression patterns. Cytochrome C oxidase, which acts as complex IV in aerobic 13 conditions, was repressed in sub-lethal conditions in both strains. Cytochrome D ubiquinol 14 oxidase, which can act as an alternative complex IV, was also down-regulated upon exposure 15 to pH 5.4. However, it was highly induced in bacteriostatic and bactericidal conditions. Genes 16 involved in oxidative stress, such as sodA, katB (Fig. 5) and perR (Fig. 4) were highly up-17 regulated in all acid shock conditions tested. The induction of these genes indicates that a low pH may induce the formation of oxidative compounds, such as H_2O_2 . Nitric oxide (NO), 18 19 formed from arginine by nitric oxide synthase (nos), putatively protects cells from H₂O₂-20 induced DNA damage by inhibition of the Fenton reaction and activation of catalase (Gusarov 21 and Nudler, 2005; Shatalin et al., 2008). Although nos was only slightly up-regulated in 22 bactericidal conditions, the formation of nitric oxide may be inferred from the induction of 23 nitric oxide dioxygenase (*hmp*) and a nitric oxide dependant transcriptional regulator (*dnrN*, 24 Supplementary material). Nitric oxide dioxygenase facilitates the reaction of nitric oxide with 25 oxygen to form nitrate. Nitrate reductase (nar) and nitrite reductase (nas) are involved in nitrogen metabolism and may serve as an alternative for aerobic respiration. Nitrate reductase
 and nitrite reductase genes are unique for strain ATCC 14579 and cluster together with nitrite
 extrusion protein *narK* (Mols *et al.*, 2007). The cluster, including *nar*, *nas* and *narK*, was
 highly up-regulated upon exposure to all acid shocks tested.

5 Besides genes involved in oxidative responses or energy metabolism, other genes were also 6 up-regulated upon exposure to all the different acid shocks (see Supplementary material). 7 Both strains showed to induce the expression of iron transporting and iron binding proteins, 8 such as feoB (BC0709, BCE0783) and dps (BC2011, BC5044, BCE2092, BC5191). 9 Furthermore, mntH (BC1803, BCE1960), encoding for manganese transport protein, also 10 belonged to the group of up-regulated genes. Manganese and iron ions may play a role in 11 oxidative stress response, conceivably acting as co-factors for superoxide dismutase proteins, 12 and via other redox balancing mechanisms.

13

14 Inactivation associated radical formation

The induction of oxidative stress associated genes and a recent publication that showed that 15 16 hydroxyl radicals (OH-) were formed upon exposure to bactericidal antibiotics in Escherichia 17 coli and Staphylococcus aureus (Kohanski et al., 2007), prompted us to investigate the 18 formation of radicals upon low pH exposure of B. cereus. The formation of OH and/or 19 peroxynitrite (ONOO⁻) in ATCC 14579 and ATCC 10987 cells was tested upon exposure to 20 selected pHs (pH 5.4, pH 5.0, pH 4.8, and pH 4.5) at different intervals using the fluorescent 21 probe 3'-(p-hydroxyphenyl) fluorescein (Fig. 6 and Fig. 7, respectively). Upon exposure to 22 pH 4.5, ATCC 14579 was inactivated and this pH induced an increase of fluorescence 23 indicating the formation of OH· and/or ONOO⁻. The exposure to the other pHs tested, i.e., pH 24 5.4, 5.0, and 4.8, did not result in inactivation of the cells and also did not induce excess 25 radical formation. Strain ATCC 10987 showed excess radical formation corresponding to the inactivation observed at pH 5.0, pH 4.8 and pH 4.5. At pH 5.4, where this strain was able to
resume growth, no excess radical formation was measured.

3 The formation of the oxygen derived radicals OH and ONOO⁻ should be prevented when 4 oxygen is not available during acid exposure. Indeed, no OH and ONOO⁻ formation was 5 observed in both strains anaerobically exposed to similar low pH values. Correspondingly, 6 increased acid resistance was observed in both strains when exposed anaerobically (data now 7 shown). In addition, the generation of superoxide, a key precursor in OH and ONOO 8 formation, was monitored upon low pH exposure using a superoxide-specific fluorescent 9 probe. Detectable levels of superoxide were only found in aerobically exposed cells, and not in anaerobically exposed cells (Mols et al., unpublished results). Taken together, these data 10 11 provide evidence that the formation of hydroxyl radicals and/or peroxynitrite is associated with inactivation of *B. cereus* strains ATCC 14579 and ATCC 10987 exposed to low pH 12 13 environments in the presence of oxygen.

14

15 **Discussion**

16 In this study, we describe the physiological and transcriptional responses of Bacillus cereus 17 strains ATCC 14579 and ATCC 10987 to sub-lethal and lethal acid shocks. The two model 18 strains were subjected to a range of pHs demonstrating that ATCC 14579 was more acid-19 resistant than ATCC 10987. ATCC 14579 survived acid conditions between pH 5.0 and pH 20 4.7 without growth or inactivation in the first hour of exposure. However, a prolonged exposure of ATCC 14579 to pH 4.8 resulted in a decrease of viable cells. In contrast, ATCC 21 22 10987 did not display this survival phenotype and was inactivated within the first hour of 23 exposure to pHs lower than pH 5.0.

The exposure of *B. cereus* to sub-lethal and lethal acid stress resulted in distinct transcriptome profiles related to the physiological response displayed by the cultures. The concurrent

1 analysis of two strains thus enables for distinguishing between phenotype-specific, stress 2 level-specific and strain-specific transcriptome responses. Furthermore, the approach used 3 showed not only the well-studied responses to mild pHs, including the induction of several 4 general stress response genes, but also the response to lethal levels of acidity, an issue that has 5 up to now mostly been neglected, as exemplified in recent studies on mild acid stress response 6 of Bacillus subtilis (Wilks et al., 2009). Cotter and Hill (2003) have reviewed the response of 7 Gram-positive organisms to mild levels of acidity and mechanisms of acid resistance were 8 described for fermentative lactic acid bacteria and L. monocytogenes, including roles of 9 proton pumps, regulators, altered metabolism, protein and DNA repair, cell envelope 10 alterations and alkali production. Using two model strains of B. cereus, we have demonstrated 11 that protein and DNA repair, stress related transcriptional regulators, altered metabolism and 12 alkali production were indeed induced at low pH. In contrast to fermentative lactic acid 13 bacteria, F₁F₀-ATPase was not up-regulated in these respiring *B. cereus* strains upon exposure 14 to acid, indicating that *B. cereus* does not use F₁F₀-ATPase to extrude protons under the 15 conditions tested. Down-regulation of F₁F₀-ATPase is best explained by the cells trying to 16 prevent excessive inward flux of protons via this ATPase upon exposure to acid conditions. 17 Furthermore, no indications were found in the transcriptome analyses for low pH-induced 18 membrane damage or rearrangement of membrane composition. For example, our 19 experiments did not show an induction of fatty acid biosynthesis (fab genes), as was shown 20 for B. subtilis exposed to mild sorbic acid stress (Ter Beek et al., 2008). Furthermore, Ter 21 Beek and colleagues (2008) reported that a putative multidrug resistance (mdr) transporter 22 was induced in B. subtilis exposed to mild sorbic acid stress and they proposed this 23 transporter to export sorbate anions from the cell. Two homologous genes in B. cereus were 24 up-regulated upon exposure to inorganic acid stress at pH 5.4 that were not induced in 25 response to lethal pH exposures. Since there is no apparent connection with sorbic acid stress

and the induction of these putative mdr systems in *B. cereus*, their role in acid resistance of *B. cereus*, if any, remains to be elucidated.

3 The transcriptome analyses of the phenotypic responses to various levels of acidity revealed a 4 major oxidative response. In bactericidal conditions, the oxidative response could be linked to 5 the formation of OH· and/or ONOO⁻ using flow cytometry in combination with the 6 fluorescent probe HPF that specifically targets these reactive oxygen species. The observed 7 oxidative burst in B. cereus may originate in a similar way as described for the formation of 8 OH radicals in Escherichia coli and Staphylococcus aureus upon exposure to bactericidal 9 antibiotics in aerobic conditions (Kohanski et al., 2007). The formation of OH- and ONOO 10 was not observed in anaerobically acid-stressed B. cereus cells, and correspondingly, 11 increased acid resistance was observed under these conditions for both strains (data not 12 shown). Based on phenotype and transcriptome analyses we propose a model for acid-induced 13 radical formation, including OH· and ONOO⁻, in B. cereus (Fig. 8). Acid stress may cause 14 perturbation of the aerobic electron transfer chain (ETC) in B. cereus indicated by the 15 differential expression of several genes potentially involved in ETC activity. This disturbance 16 may cause premature leakage of electrons to oxygen leading to the formation of superoxide 17 (O_2) . Indeed, elevated levels of O_2 could be detected in *B. cereus* cells upon exposure to 18 lethal levels of acidity as indicated by staining of these cells with a superoxide-specific 19 fluorescent probe (Mols *et al.*, unpublished results). Furthermore, the formation of O_2^- can be 20 inferred from the induction of superoxide dismutase and catalase genes. Iron-sulphur clusters 21 may subsequently be damaged by O₂⁻ releasing iron in the cytoplasm (Imlay, 2006). Free iron 22 can react with hydrogen peroxide, originating from the dismutation of O_2^- , forming the highly toxic OH· radicals in the Fenton reaction (Imlay et al., 1988). Furthermore, O2⁻ can rapidly 23 24 react with nitric oxide (NO) to form another highly toxic oxidative compound, ONOO⁻ 25 (Beckman and Koppenol, 1996). NO is formed by a reaction catalyzed by nitric oxide

1 synthase (bNOS). Indirect indications for the formation of NO upon low pH exposure can be 2 inferred from the up-regulation of nitric oxide dioxygenase and nitric oxide dependant 3 regulator *dnrN*. The induction of bNOS activity, which is possibly regulated at protein level 4 (Shatalin et al., 2008), may initially have a positive effect on surviving oxidative stress. 5 bNOS-derived NO may inhibit thiol reduction leading to the inhibition of the OH forming 6 Fenton reaction (Gusarov and Nudler, 2005; Sudhamsu and Crane, 2009). Furthermore, NO 7 induces catalase activity in *B. anthracis* (Shatalin *et al.*, 2008) and inhibits the aerobic ETC 8 (Husain et al., 2008). On the other hand, NO facilitates the formation of ONOO⁻, which may 9 have a damaging effect that could lead to cell death. Nitric oxide dioxygenase and nitrite 10 reductase are described to be possible NO dissipation routes (Payne et al., 1997; Gardner, 11 2005). The genome of ATCC 14579 encodes both mechanisms and this strain showed to be 12 more acid resistant than the nitrite/nitrate reductase deficient ATCC 10987 strain.

13 The phenomenon that exposure to stresses such as salt, heat, acid, and bile, results in 14 secondary oxidative stress, has been described earlier for B. cereus and numerous other 15 bacteria (Aldsworth et al., 1999; Clements et al., 1999; Hecker and Volker, 2001; Airo et al., 16 2004; Banjerdkij et al., 2005; Latifi et al., 2005; Dodd et al., 2007; Kim et al., 2008), but up 17 to now, this secondary oxidative stress response has not been linked to radical-associated cell 18 death. Moreover, our findings are supported by earlier observations in amongst others S. 19 aureus and Vibrio vulnificus, where acid resistance was found to be superoxide dismutase 20 (and catalase) dependant (Clements and Foster, 1999; Kim et al., 2005).

In conclusion, the results obtained in our study provide evidence for the origin of acid stressinduced oxidative stress. In extension to the antibiotic study of Kohanski and colleagues (2007), we now propose that in aerobic conditions, the formation of radicals such as $OH \cdot$ and ONOO⁻ may be a common mechanism of cellular death in bacteria exposed to severe stress conditions.

2 **Experimental procedures**

3

4 Bacterial strains and growth conditions

B. cereus strains ATCC 14579 and ATCC 10987 were obtained from the American Type
Culture Collection (ATCC). Stock cultures, grown in brain heart infusion (BHI, Becton
Dickinson, France) broth, were stored at -80°C in 33% glycerol. To prepare pre-cultures, 10
ml BHI in a 100 ml Erlenmeyer flask was inoculated with a droplet from the glycerol stock
and incubated overnight at 30°C, with shaking at 200 rpm.

10 To study the effect of pH on *B. cereus* cells and the corresponding transcriptome profiles, 100 11 ml BHI in a 500 ml Erlenmeyer flask was inoculated with 0.5 ml pre-culture and incubated at 12 30°C, with shaking at 200 rpm. Upon reaching an optical density of 0.5 measured at 600 nm 13 (OD, Novaspec II, Pharmacia Biotech, Germany), the culture pH was measured (PHM 240 14 pH/ION Meter, Radiometer, Denmark) and serial dilutions were made in peptone 15 physiological salt solution (PPS, 1g/l neutralized bacteriological peptone (Oxoid, England) 16 and 8.5 g/l NaCl in water) and plated with a spiral-plater (Eddy Jet; IUL Instruments, Spain) 17 on BHI agar plates (15 g/l bacteriological agar, Oxoid, England). 20 ml of the culture was 18 used to extract RNA (sample t = 0). The remaining volume of the culture was acidified with 19 hydrochloric acid (HCl 37%, Merck, Germany) to pH 5.4, 5.0, 4.8 or 4.5 and incubated at 20 30°C, with shaking at 200 rpm. At designated time points (10, 30 and 60 minutes), samples 21 were taken to measure the OD, to determine the viable counts, and to extract RNA.

22

23

24 RNA isolation

1 RNA isolation was performed by transferring 20 ml of the cultures into a 50-ml Falcon tube 2 (Greiner Bio-one, Germany) at the designated time points. Subsequently, the cultures were 3 pelleted at maximum speed at 4°C for 30 s (Eppendorf centrifuge 5804 R, Eppendorf, 4 Germany). After decanting the supernatant, the cell pellets were frozen in liquid nitrogen. 5 Within 10 min after freezing the cell pellets, 1 ml TRI-reagent (Ambion, United Kingdom) 6 was added to the pellets. The samples were stored at -80°C until RNA extraction. RNA was 7 extracted as described previously (van Schaik et al., 2004). Residual chromosomal DNA was 8 removed by treating the samples with DNA-free (Ambion, United Kingdom). The RNA 9 concentration was measured in 2 ml cuvettes (UVettes, Eppendorf, Germany) with a 10 BioPhotometer (Eppendorf, Germany) by determining the OD_{260} and OD_{280} . The quality of 11 the RNA was monitored using the RNA 6000 Nano Assay (Agilent, United States) and the 12 Agilent 2100 Bio-analyzer (Agilent, United States) according to the provided protocol. The 13 extracted RNA samples were stored in 70% ethanol with 83 mM sodium acetate buffer (pH 14 5.2) at -20°C.

15

16 cDNA synthesis, labelling and microarray hybridization and design

17 Complementary DNA with amino-allyl-labelled dUTP (Ambion, United Kingdom) from the 18 extracted RNA was prepared in reverse transcription reactions using Superscript III 19 (Invitrogen, The Netherlands). Cy3 and Cy5 labelling of the cDNAs was performed with the 20 CyScribe Post-Labeling kit (GE Healthcare, Belgium) as previously described (den Hengst et 21 al., 2005). The labelled cDNAs were purified using the CyScribe GFX purification kit (GE 22 Healthcare) according to the provided protocol. To conduct the microarray hybridization, the 23 Cy5-labelled cDNA samples were combined with the corresponding Cy3-labelled t0 reference 24 samples (1:1 ratio). The microarray experiments for the comparison of the transcriptomes of 25 the cultures exposed to various pHs were performed in two independent biological replicates, where the replicate was performed with the dyes swapped. *B. cereus* ATCC 14579 and *B. cereus* ATCC 10987 microarrays (details below) were hybridized with 200 to 300 ng labelled
 cDNA following the 60-mer oligo microarray processing protocol (Agilent, United States).

4 The microarrays used in this study were custom-made B. cereus ATCC 14579 and custom-5 made B. cereus ATCC 10987 microarrays developed by Agilent Technologies (United 6 States). The B. cereus ATCC 14579 microarray design was based on the 11K platform of 7 Agilent Technologies (GEO accession number GPL7680). A total of 10,262 spots represented 8 5,131 chromosomal open reading frames, meaning that 98.0% of the predicted chromosomal 9 open reading frames (NCBI accession number NC_004722) are represented on the 10 microarray. 99.6 % of the open reading frames for which probes could be designed were 11 represented by two non-overlapping probes on the array. The remaining 0.4% of the open 12 reading frames was represented by a single oligonucleotide spotted in duplicate on the array. 13 The B. cereus ATCC 10987 microarray design was based on the 22K platform of Agilent 14 Technologies (GEO accession number GPL7681). A total of 17,697 spots represented 5,578 15 chromosomal open reading frames, 240 plasmid open reading frames and 81 putative small-16 RNAs, meaning that 99.6% of the predicted chromosomal and the plasmid open reading 17 frames (NCBI accession numbers NC_003909 and NC_005707, respectively) were represented on the microarray. All features (chromosomal and plasmid open reading frames 18 19 and small-RNAs) were represented by three individual spots. For 4,914 features three non-20 overlapping probes were designed, for 488 features two probes were designed (one probe 21 spotted in duplicate) and for 497 only one oligonucleotide could be designed (one probe 22 spotted in triplicate).

After hybridization at 60°C for 17 hours, the microarrays were washed with $6 \times SSC$ (0.9 M NaCl and 0.09 M sodium citrate) supplemented with 0.005% Triton X-102 at room

1 temperature for 10 min. Subsequently, the microarray slides were washed at 4° C with 2 prechilled 0.1 × SSC with 0.005% Triton X-102 for 5 min and dried with nitrogen gas.

3

4 Microarray scanning and data analysis

5 The microarray slides were scanned using an Agilent microarray scanner (G2565BA), and 6 data were extracted from the scanned microarrays with Agilent's Feature Extraction software 7 (version 8.1.1.1), which includes a LOWESS (locally weighted scatterplot smoothing) 8 normalization step for the raw data. After removal of the data for the control spots, the 9 normalized data for each spot from the microarrays were analyzed for statistical significance 10 using the web-based VAMPIRE microarray suite (Hsiao et al., 2005). A spot was found to be 11 differentially expressed between two samples when the false discovery rate was smaller than 12 0.05. Subsequently, the data for the single spots were integrated to obtain expression ratios for 13 a corresponding feature (i.e., open reading frame or sRNA). A feature was found to be 14 differentially expressed when all spots representing the feature were significantly 15 differentially expressed between samples.

16 Hierarchical clustering (Eisen et al., 1998) was performed per strain to identify groups of 17 genes showing similar expression patterns. In Genemaths XT (version 1.6.1, Applied Maths, 18 Belgium) genes that were significantly differentially expressed in one or more conditions 19 were log₂ transformed and clustered using the complete linkage method and the Euclidian 20 distance matrix. The groups identified from the hierarchical clustering were based on an 21 arbitrary cut-off value. To identify relevant biological processes significantly overrepresented 22 in a group, the genes corresponding to a group were analyzed using FIVA (Blom et al., 2007). 23 To visualize the relation between the different acidic conditions independent of the strains, the 24 log₂ transformed expression ratios from genes present on both microarrays were hierarchically

clustered using the average linkage method and the Euclidian distance matrix (Genemaths
 XT, 1.6.1).

3

4 *Flow cytometry and radical measurements*

5 To detect radical formation, the fluorescent reporter dye 3'-(p-hydroxyphenyl) fluorescein 6 (HPF, Invitrogen, The Netherlands) was used (Setsukinai et al., 2003). At designated time 7 points (0, 10, 30, 60 minutes) after adding HCl to the exponentially growing cultures 8 (OD_{600nm} ~0.5), samples were obtained by centrifuging $(15,000 \times g, 30 \text{ s})$ 1 ml of culture and 9 resuspension of the cell pellet in 1 ml filtered phosphate buffer saline (PBS). The samples 10 were washed once and diluted with filtered PBS to obtain a concentration of approximately 10⁶ cells per ml, subsequently HPF was added at a final concentration of 5 mM. Samples were 11 12 run on a Becton Dickinson FACSCalibur flow cytometer with the following photomultiplier 13 tube (PMT) voltage settings: E00 (FSC), 360 (SSC) and 825 (FL1). Data were obtained from 20,000 events (cells) at medium flow rate using Cellquest Pro (version 4.0.2), subsequently 14 15 analyzed with WinMDI 2.9 (Joseph Trotter, Salk Institute for Biological Studies, La Jolla, 16 California, USA; http://facs.scripps.edu/software.html) and graphically presented using 17 Adobe Illustrator CS2 (version 12.0.1).

18

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20

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9	
10	Figure legends
11	
12	Fig. 1. Physiological response of <i>B. cereus</i> ATCC 14579 (filled squares) and ATCC 10987
13	(open diamonds) upon exposure to a range of acidity levels. The colony forming units were
14	determined after 0 and 60 minutes of exposure, the difference between the t0 and t60 is
15	depicted. Data points represent single experiments, indicating a large variability of the
16	responses of ATCC 10987 exposed to pH 5.0. The grey line at 0 corresponds to no growth
17	(growth above the line) and no inactivation (inactivation below the line). The filled area
18	between the dotted lines depicts the pH values where ATCC 14579 displayed no growth and
19	no inactivation, i.e., survival, and where ATCC 10987 already showed to be inactivated.
20	
21	Fig. 2. Physiological response of <i>B. cereus</i> ATCC 14579 (A) and ATCC 10987 (B) upon
22	exposure to low pH. The colony forming units determined at different time points upon
23	exposure to pH 5.4 (squares), pH 5.0 (diamonds), pH 4.8 (triangles) and pH 4.5 (circles) are
24	depicted. At 0, 10, 30 and 60 minutes samples were taken for microarray analysis, indicated

with arrows and the error bars represent the standard deviation between duplicate
 experiments.

3

Fig. 3. Hierarchical clustering of the transcriptome profiles of different pH exposures based on the common genes of *B. cereus* ATCC 14579 and ATCC 10987. Samples with similar expression patterns were clustered using Euclidean distance and complete linkage. The corresponding phenotypic responses are shown at the right. Samples obtained from growing cultures are depicted in black, samples from non-growing ("survival" and "inactivation" phenotypes) cultures are shown in grey. Relative distance in similarity between the branches is shown at the top and bootstrap values are indicated at each branch.

11

12 Fig. 4. Average ratios of low pH and stress associated genes from B. cereus ATCC 14579 13 (closed bars) and ATCC 10987 (open bars) upon exposure to pH 5.4 (A) and pH 4.5 (B). The 14 global adaptive response is represented by sigB (BCE1086 and BC1004), sigH (BCE0093 and BC0114) and *codY* (BCE3869 and BC3826), encoding σ^{B} , σ^{H} and CodY respectively. 15 16 Additionally, perR (BCE0592 and BC0518), a major oxidative stress response regulator is 17 shown. atpA (BCE5432 and BC5308), napA (BCE1729 and BC1612), and nhaC (BCE1840 18 and BC1709) represent F₁F₀-ATPase and two proton antiporters. gad (glutamate 19 decarboxylase, BCE2691), arcA (arginine deiminase, BCE0472 and BC0406), and ureA 20 (urease, BCE3664) represent systems that are described to be involved in alkaline production. 21 Glutamate decarboxylase and urease are specific for ATCC 10987 and are indicated with an 22 asterisk. The general stress response chaperones and proteases are depicted by dnaK 23 (BCE4395 and BC4312), groES (BCE0288 and BC0294), and clpC (BCE0079 and BC 24 BC0100). Additionally, mechanisms involved in sorbic acid stress of B. subtilis (Ter Beek et 25 al., 2008), such as, multidrug transporters (mdr1, BCE4699 and BC4568 and mdr2, BCE1943 and BC1786), fatty acid biosynthesis (*fabF*, BCE1294 and BC1174), and branched-chain fatty
 acid biosynthesis (*bkdR*, BCE4239 and BC4165) are shown.

3

4 Fig. 5. Average ratios of selected genes associated with respiration and oxidative responses 5 from B. cereus ATCC 14579 (closed bars) and ATCC 10987 (open bars) upon exposure to pH 6 5.4 (A), pH 5.0 (B), pH 4.8 (C) and pH 4.5 (D). ETC-4a and ETC-4b depict two distinct types 7 of complex IV of the aerobic electron transfer chain. Cytochrome C oxidase polypeptide I 8 gene ctaD (BCE3990 and BC3943) represents ETC-4a and cytochrome D ubiquinol oxidase 9 subunit I gene cydA (BCE4949 and BC4792) represent ETC-4b. The genomes of ATCC 14579 and ATCC 10987 harbour four different superoxide dismutase genes and three 10 11 different catalase genes, the data of sodA (BCE5579 and BC5445) and katB (BCE1261 and 12 BC1155) are shown here. Nitric oxide synthase and nitric oxide dioxygenase are represented 13 by the corresponding genes, nos (BCE5578 and BC5444) and hmp (BCE1571 and BC1448), 14 respectively. Nitrate and nitrite reductases are encoded by multiple genes on the genome of 15 ATCC 14579, therefore narI (BC2121), nasD (BC2136) and nark (BC2128) are shown. This 16 nitrate and nitrite reductase cluster of ATCC 14579 is absent in ATCC 10987 and therefore 17 indicated with an asterisk.

18

Fig. 6. Radical formation in *B. cereus* ATCC 14579 upon exposure to pH 5.4, pH 5.0, pH 4.8 and 4.5. Samples were taken at 0 (green), 10 (light blue), 30 (blue) and 60 (dark blue) minutes. The pH and corresponding physiological response are indicated at each graph. The shift in fluorescent signal to the right indicates the formation of hydroxyl and/or peroxynitrite radicals.

Fig. 7. Radical formation in *B. cereus* ATCC 10987 upon exposure to pH 5.4, pH 5.0, pH 4.8 and 4.5. Samples were taken at 0 (green), 10 (light blue), 30 (blue) and 60 (dark blue) minutes. The pH and corresponding physiological response are indicated at each graph. The shift in fluorescent signal to the right indicates the formation of hydroxyl and/or peroxynitrite radicals.

6

7 Fig. 8. Low pH induced oxidative stress response and radical forming mechanisms in B. 8 cereus ATCC 14579 and ATCC 10987. Schematic representation of radical formation 9 conceivably induced upon exposure to lethal acid stress. Acid stress may cause perturbation 10 of the electron transfer chain and an excess of superoxide radicals (O_2) may be formed. Superoxide radicals can be converted to hydrogen peroxide and water by superoxide 11 12 dismutase (sod) and catalase (kat). However, when the capacity to dismutate superoxide is not 13 sufficient, free superoxide radicals can cause damage to iron-sulphur (Fe-S) cluster containing 14 enzymes supplying unbound iron ions. These free iron ions and hydrogen peroxide can react 15 (Fenton reaction) and produce hydroxyl radicals (OH·). Another possible route in forming 16 highly damaging radicals may occur via nitric oxide. Nitric oxide can be formed by nitric 17 oxide synthase (nos) and can react with superoxide radicals to form peroxynitrite (ONOO⁻). 18 Nitric oxide can be converted to nitrate by nitric oxide dioxygenase (*hmp*). Subsequently, 19 nitrate can be converted to nitrite and ammonium by nitrate (nar) and nitrite (nas) reductase, 20 respectively. Nitrite can also be transported outside the bacterial cell by a Nitrite extrusion 21 protein (nark). nar, nas and narK are ATCC 14579 specific, the reactions they catalyze are 22 indicated with dotted lines.

23

24 Supplementary material

1 The following supplementary material is available for this article online:

2 14579_Allgroups.tar. A compressed folder containing the results of the hierarchical 3 clustering of the significant differentially expressed genes of strain ATCC 14579. Opening the 4 htmlOutputFiva.html file shows the graphical output of all used annotation modules and can 5 be used to navigate through the results. By clicking the cluster names, the corresponding 6 transcriptome profiles of the groups are revealed. The size of each group is displayed in blue 7 underneath the group name and by clicking the list of genes will appear. Numbers in each 8 rectangle represent absolute values of occurrences. The significance of occurrences is 9 visualized in a colour gradient and with symbols which are displayed at the bottom of the plot. 10 On the left the categories showing over representation in one or more groups are listed with 11 the corresponding size. Clicking the categories will reveal the genes present in the 12 corresponding category per group. The description of each category is placed at the right and 13 more information about the categories can be displayed by clicking the description.

14 10987_Allgroups.tar. A compressed folder containing the results of the hierarchical
clustering of the significant differentially expressed genes of strain ATCC 10987. A detailed
description is given above.

Supplementary results.pdf. Supplementary results based on the microarray data obtained for *B. cereus* strains ATCC 14579 and ATCC 10987 exposed to pH 5.4, pH 5.0, pH 4.8 and pH 4.5. The data of each strain was clustered hierarchically and different groups of genes were identified based on their common transcriptome profile. The FIVA analysis showed that several function categories were overrepresented in certain groups.

22

Raw and processed microarray data are available for reviewers following the link to the GEOdatabase below.

25 http://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?token=hfsfreqageqakzi&acc=GSE13773



Fig 1





5 Fig 2



2 Fig 3



2 Fig 4



- 5 Fig 5



1 Fig 7





3

4 Fig 8