A spore-lytic enzyme released from *Bacillus cereus* spores during germination

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The exudate of fully germinated spores of *Bacillus cereus*IFO 13597 in 0.25 M sodium phosphate buffer, pH 7.0, was found to contain a spore-lytic enzyme. This enzyme was found to cause loss of absorbance in coat-stripped spore suspensions and phase-darkening of the spores but had minimal activity on isolated peptidoglycan substrates. The enzyme was purified in an active form and identified as a 24 kDa protein which is either an amidase or a peptidase. The amino-terminal 19 residues had the following sequence: FSNQVIQRGASGEKVIELQ. The spore-lytic enzyme retained its activity in a medium of a relatively high ionic strength containing a non-ionic surfactant such as nonaethyleneglycol n-dodecyl ether. This activity was optimum at a salt concentration of about 30 mM in assay buffer at neutral pH. In contrast to the enzyme in a spore-bound form, the enzyme in solution was shown to be heat-sensitive and was readily inactivated by thiol reagents.

Keywords: *Bacillus cereus*, spore-lytic enzyme, germination cortex hydrolysis

INTRODUCTION

The cortical peptidoglycan has been strongly implicated in the maintenance of bacterial spore dormancy. The degradation of peptidoglycan in the spore cortex is an essential event in spore germination (Foster & Johnstone, 1990) and the mechanism of cortex hydrolysis is crucial to an understanding of the germination pathway. A number of spore-lytic enzymes have been isolated from bacterial spores of different origins (Ando, 1979; Brown & Cuhel, 1975; Brown et al., 1978, 1982; Foster & Johnstone, 1987; Gombas & Labbe, 1981, 1985; Gould et al., 1966; Tang & Labbe, 1987). However, in most cases, their involvement in germination has not been unequivocally established.

*Bacillus cereus* spores contain four separate spore-degrading enzymes (Warth, 1972) and spore-lytic enzyme activities are localized in both the core and the coat fraction (Brown & Cuhel, 1975; Brown et al., 1978). Brown et al. extracted a surface-bound spore-lytic enzyme by urea/mercaptoethanol or guanidine HCl/mercaptoethanol treatment (Brown & Cuhel, 1975; Brown et al., 1978, 1982). The enzyme did not induce refractility changes in coat-damaged spores following treatment with this enzyme. However, Gould et al. (1966) found a spore-lytic enzyme which was liberated from a spore-bound form during germination when the ionic strength of the medium was high. The enzyme causes loss of absorbance in coat-stripped spore suspensions and phase-darkening of the spores. These changes are characteristic of normal spore germination. This suggests that the enzyme is involved in the germination pathway. However, the enzyme has not been characterized on a molecular level. In this study we found that a spore-lytic enzyme is released from germinating *B. cereus* IFO 13597 spores into the germination exudate. Furthermore, the enzyme causes refractility changes of coat-stripped spores, and is similar to the enzyme characterized by Gould et al. (1966). This report deals with the purification and the partial characterization of the enzyme.

METHODS

Preparation of spores. Spores of *B. cereus* IFO 13597 were prepared in the following manner. Portions (1 ml) of a nutrient broth culture were inoculated on the surface (600 cm²) of a sporulation medium (1 g Bact-peptone 5 g, meat extract 3 g, yeast extract 1 g, MnSO₄ 0.1 g, agar 15 g, pH 7.0) in culture flasks and inoculated at 32 °C for 3 d. The spores were harvested and washed five times with distilled water. Spores were then incubated in 10 vols 2 M KCl, 10 mM sodium phosphate, pH 7.5, at 37 °C for 40 min and washed twice with water. Afterwards, the spores were further incubated at 37 °C for 30 min in 10 vols 2 M urea, 5 mM CHES, pH 9.3, containing...
25 mM 2-mercaptoethanol. After washing five times with water, the spores were stored at 4 °C. Spores of *Clostridium perfringens* S40 and of *Bacillus steaothermophilus* IAM 1035 were prepared according to the methods of Ando & Tsuzuki (1984) and of Tomida et al. (1991), respectively.

**Preparation of coat-stripped spores, isolation of cortical fragments and isolation of the vegetative cell wall.** Removal of the spore coat from *B. cereus* and *B. steaothermophilus* spores was performed with a slight modification of the method of Gombar & Labbe (1981). Spores were treated with 30 mM SDS, 0.2 M 2-mercaptoethanol, 0.1 M borate buffer, pH 10.0, at 40 °C. *B. cereus* and *B. steaothermophilus* spores were treated for 8 and 4 h, respectively. Both were extensively washed with distilled water. Coat-stripped spores of *C. perfringens* were prepared according to the method of Ando & Tsuzuki (1984).

Spores or vegetative cells collected from a stationary-phase culture were suspended in distilled water and disrupted at 0–4 °C in a bead-beater (Edmund Buhler) with glass beads (diam. 0.1 mm). Spore cortical fragments and cell walls were isolated according to the methods of Tang & Labbe (1987) and of Kuroda & Sekiguchi (1990), respectively.

**Germination.** A suspension of *B. cereus* spores (0.1 g packed spores/ml water) was heat-activated at 65 °C for 45 min and cooled on ice. After sedimentation by centrifugation (6000 g, 10 min, 4 °C), the spores were germinated at 32 °C in 10 vols 0.25 M sodium phosphate germination buffer containing 10 mM L-alanine and 4 mM adenosine, pH 7.0. Germination was followed by monitoring the decrease in OD$_{600}$ of spore suspension and phase-darkening of the spores. A Jasco UV5 spectrophotometer was used. In some experiments, spores were germinated in the presence of chloramphenicol (100 μg ml$^{-1}$).

**Spore-lytic enzyme assay.** Spore-lytic enzyme activity was assayed by measuring the decrease in OD$_{600}$ of coat-stripped spore suspensions in a cell of 1 mm light path at 32 °C. A Jasco UV5 spectrophotometer was used. The reaction mixture contained coat-stripped spores (OD$_{600}$ of 0.1) and the enzyme in a final volume of 150 μl of 30 mM sodium phosphate buffer, pH 7.0, containing 1 mM EDTA, 1 mM sodium thioglycollate and 0.1% C$_{12}$E$_{9}$ (Nikko Chemicals). One unit of activity is defined as a decrease in OD$_{600}$ of 0.1 min$^{-1}$. At the same time, activity was qualitatively examined through microscopic observation of the change from phase-bright coat-stripped spores to phase-dark ones.

**Purification of spore-lytic enzyme from germination exudate.** A typical purification procedure was as follows. Packed spores (5g) were incubated in 50 ml germination buffer at 32 °C for 45 min. After centrifugation (8000 g, 10 min, 4 °C), the supernatant fluid (50 ml) was brought to 0.5% C$_{12}$E$_{9}$ and then with 40 ml 2-propanol/0.05% TFA, and eluted with a linear gradient of 0-0.4 M NaCl in buffer B. The dialysed solution was applied at 20 °C to a MCI GEL CQK-30S column (75×75 mm), a high performance liquid chromatography column with a sulfopropyl group as an ion exchanger; Mitsubishi Kasei pre-equilibrated with buffer B. The eluted solution was applied at 20 °C to a MCI GEL CQK-30S column (75×75 mm), a high performance liquid chromatography column with a sulfopropyl group as an ion exchanger; Mitsubishi Kasei pre-equilibrated with buffer B. After the column was washed with buffer B, proteins were eluted at 20 °C with a linear gradient of 0–0.4 M NaCl in buffer B. The flow rate was 0.5 ml min$^{-1}$ and the eluate was monitored by measuring $A_{280}$. Fractions (0.75 ml each) were collected and assayed for enzyme activity.

Active fractions were further chromatographed on a µBondapak™ cyanopropyl-300A column (3.9×150 mm, Waters) which had been equilibrated with 5% (v/v) acetonitrile/0.05% TFA, and eluted with a linear gradient of 5% acetonitrile/0.05% TFA and 60% acetonitrile/30% (v/v) 2-propanol/0.05% TFA. The enzyme lost its activity in this purification step. The 24 kDa protein eluted as a major component was used for N-terminal amino acid sequence analysis and preparation of antibodies.

**Preparation of antiserum and immunoprecipitation.** A mouse was intraperitoneally injected with 10 μl cyanopropyl-column-purified 24 kDa protein emulsified in Freund's complete adjuvant. After 3 weeks, the mouse was injected with the same amount of protein in Freund's incomplete adjuvant. A second boost was performed 5 weeks later, and the mouse was bled 10 days later, after the final injection. The blood was brought to 0.15 M NaCl, 5 mM sodium phosphate, 0.1% NaN$_{3}$, pH 7.0, via the addition of a concentrated buffer and then stored at 4 °C for 20 h. After centrifugation (14000 g, 5 min, 4 °C), the antiserum was stored at −20 °C.

For immunoprecipitations, the antiserum (50 μl), or non-immune control serum of the immunized mouse (50 μl), was incubated with 100 μl protein A-agarose (Pharmacia) at 4 °C for 16 h. After centrifugation (5000 g, 5 min, 4 °C), the gel was washed with a 1:5 glycine–NaOH buffer, pH 7.0, containing 3 M NaCl, and equilibrated with buffer A. The spore-lytic enzyme (100 μl, 0.35 units), partially purified by a BSA-Sepharose 4B column, was mixed with serum-treated protein A-agarose gels. This mixture was incubated at 4 °C for 16 h. After centrifugation (6000 g, 5 min, 4 °C), the spore-lytic activity and the protein of the supernatant were examined.

**Mode of action of the spore-lytic enzyme.** A coat-stripped spore suspension (1-2 ml in 30 mM sodium phosphate buffer, pH 7.0, containing 1 mM EDTA, 1 mM sodium thioglycollate, and 0.1% C$_{12}$E$_{9}$, with an OD$_{600}$ of 1/4 in a cell of 1 mm light path) was incubated with the enzyme (50 μl, 27 units) at 32 °C. The change in OD$_{600}$ was then monitored. After hydrolysis (30 min), the suspension was centrifuged (8000 g, 10 min, 4 °C). The free amino groups and reducing sugars of the supernatant were determined. A control experiment was performed with the enzyme boiled for 10 min.

**Analytical methods.** Protein concentrations were determined according to the methods of Lowry and/or Groves et al. (1960), using BSA as a standard. Free amino groups were identified by their reactions with trinitrobenzenesulfonate using L-alanine as a standard, according to the method of Fields (1972). Reducing sugars were determined by the ferricyanide reduction method of Park & Johnson (1949), using N-acetyl-D-glucosamine as a standard. SDS-PAGE was carried out on 12.5% or 13.3% (w/v) slab gels using a Laemmli buffer system (Laemmli, 1970) at a constant current of 20 mA. Proteins were silver-stained
(Oakley et al., 1980). Sequencing of N-terminal amino acids was carried out on an Applied Biosystems 477A/120A sequence analyser.

**RESULTS**

**Action of spore-lytic enzyme released during germination on spores and isolated peptidoglycan**

Coat-stripped spores were phase-bright, but did not respond to germinants such as L-alanine, adenosine, inosine or a mixture thereof. This suggested that the spores were non-viable. When incubated with the germination exudate of *B. cereus*, the coat-stripped spores became phase-dark as the extinction of the suspensions decreased. The response was quite similar to changes characteristic of normal spore germination. However, the enzyme did not cause germination of spores which had not previously been coat-stripped. The release of the enzyme occurred in the presence of chloramphenicol, an inhibitor of protein synthesis, suggesting that the enzyme is not synthesized *de novo* during germination. Table 1 shows lytic activity of the germination exudate for a variety of substrates. Under our experimental conditions used here, changes in the extinction of coat-stripped spore suspensions were completed within 10 min. However, the exudate did not cause a detectable loss in OD of isolated spore cortex and isolated cell wall suspensions during incubation for 1 h.

**Stability of spore-lytic enzyme activity**

It was found that subtilisin-like protease activity is presented at the spore surface, which could not be removed from spores by washing with water. When protease-carrying spores were germinated, spore-lytic activity released into the germination exudate rapidly declined. This is most likely due to proteolytic degra-

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**Table 1. Lytic activity of the germination exudate for a variety of substrates**

The lytic activity was measured by the decrease in OD at 600. The control value (0.25 units for coat-stripped *B. cereus* spores) was taken as 100%. Values were rounded off to integers. These data represent the mean of three independent experiments (SD < 12% of the mean). The same substrate specificities were also observed when the BSA-Sepharose 4B column-purified enzyme was used.

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Lytic activity (percentage of control)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Normal <em>B. cereus</em> spore</td>
<td>&lt; 1</td>
</tr>
<tr>
<td>Coat-stripped <em>B. cereus</em> spore</td>
<td>100</td>
</tr>
<tr>
<td>Coat-stripped <em>B. stearothermophilus</em> spore</td>
<td>65</td>
</tr>
<tr>
<td>Coat-stripped <em>C. perfringens</em> spore</td>
<td>56</td>
</tr>
<tr>
<td>Isolated <em>B. cereus</em> spore cortex</td>
<td>&lt; 1</td>
</tr>
<tr>
<td>Isolated <em>B. cereus</em> cell wall</td>
<td>&lt; 1</td>
</tr>
<tr>
<td>Isolated <em>C. perfringens</em> cell wall</td>
<td>&lt; 1</td>
</tr>
</tbody>
</table>

**Table 2. Purification of the spore-lytic enzyme**

The spore-lytic enzyme was purified from 50 g packed weight of spores as described in Methods. ND, Not determined.

<table>
<thead>
<tr>
<th>Procedure</th>
<th>Total activity (units)</th>
<th>Total protein (mg)</th>
<th>Specific activity [units (mg protein)-1]</th>
<th>Yield (percentage total original activity)</th>
<th>Purification factor</th>
</tr>
</thead>
<tbody>
<tr>
<td>Germination exudate</td>
<td>2445</td>
<td>45.0</td>
<td>54.3</td>
<td>100</td>
<td>1</td>
</tr>
<tr>
<td>SP-Sepahex C-25 column</td>
<td>1338</td>
<td>0.94</td>
<td>142.3</td>
<td>54.7</td>
<td>26.2</td>
</tr>
<tr>
<td>BSA-Sepharose 4B column</td>
<td>512</td>
<td>ND</td>
<td>ND</td>
<td>20.9</td>
<td>ND</td>
</tr>
<tr>
<td>MCI GEL CQK-30S column</td>
<td>428</td>
<td>0.028</td>
<td>15.285.7</td>
<td>17.5</td>
<td>281.5</td>
</tr>
</tbody>
</table>
Retention time (min) 1 2 3 4 5

Fig. 3. Immunoprecipitation of the spore-lytic enzyme with anti-24 kDa protein antibody. Spore-lytic enzyme (100 μl, 0.35 units, partially purified by BSA-Sepharose 4B column chromatography) was incubated with protein A-agarose gels which were treated with antiseraum or control serum as described in Methods. After the suspensions were centrifuged, proteins in the supernatants were analysed by SDS-PAGE and enzyme activities of the supernatant were measured. (a) SDS-PAGE was run on a 13.5 % gel and 60 μl of the supernatant was electrophoresed. Lanes: 1, supernatant treated with control serum; 2, supernatant treated with antiserum. The arrow indicates the 24 kDa protein. (b) Changes in OD₅₆₀ of coat-stripped spore suspensions (initial OD₅₆₀ was 0.1 in a cell of 1 mm light path) by the addition of supernatants (15 μl supernatant/50 μl spore suspension). Curve 1, supernatant treated with control serum; curve 2, supernatant treated with antiserum.

Fig. 2. Chromatography of spore-lytic enzyme on a MCI GEL CQK-305 column and SDS-gel electrophoretic profiles showing purification of the enzyme. (a) Fractions retarded on a BSA-Sepharose 4B column which contain spore-lytic activity (30 ml, from 4 g packed spores) and fractions unretarded on the column (40 ml) were applied to a MCI GEL CQK-305 column. The elution profiles are shown in curves 1 and 2, respectively. Proteins were monitored by A₂₅₀. (b) The spore-lytic enzyme was purified as described in Table 2 and analysed by 0.1 % SDS-12.5 % PAGE. Lanes: 1, germination exudate; 2, components containing the spore-lytic activity eluted from a SP-Sephadex C-25 column; 3, components unretarded on a BSA-Sepharose 4B column; 4, components retarded on a BSA-Sepharose 4B column which contain spore-lytic enzyme activity; 5, MCI GEL CQK-305 column-purified spore-lytic enzyme [a major peak fraction of curve 1 in (a)]. Approximately 3-30 μg of proteins were electrophoresed. Migration positions of standard proteins are indicated; BSA (67000), ovalbumin (44000), subtilisin BPN' (27500), chymotrypsinogen (25700) and lysozyme (14300).

As shown in Fig. 1, spore-lytic enzyme activity was found to decrease with lowering of the ionic strength of the buffer. Addition of the non-ionic surfactant C₁₄₃₄ suppressed the loss of enzyme activity. Therefore, purification of the spore-lytic enzyme was performed in a buffer of relatively high ionic strength, > 75 mM, containing 0.1 % C₁₄₃₄.

Purification, N-terminal amino acid sequence and mode of action of spore-lytic enzyme

The steps in the purification procedure and the yields obtained for the spore-lytic enzyme are shown in Table 2. The spore-lytic enzyme was separated from most protein...
components in the first chromatography step using a SP-Sephadex C-25 column. The spore-lytic enzyme activity was retarded on a BSA-Sepharose 4B column, indicating an affinity between BSA and the enzyme. The affinity was found to be stronger at 4 °C than at 20 °C. Fractions containing enzyme activity were further purified by MCI GEL CQK-30s column chromatography and the enzyme was recovered in the major peak of protein (Fig. 2a, curve 1). Fractions comprising this peak contained a single predominant protein with an apparent molecular mass of 24000, as determined by SDS-PAGE (Fig. 2b, lane 5). The yield of enzyme activity was 17.5%. For comparative purposes, components unretarded on a BSA-Sepharose 4B column were also chromatographed on the same column, as shown in curve 2 of Fig. 2(a). The results shown in Fig. 2 indicate that separation of the spore-lytic enzyme from other protein components by a BSA-Sepharose 4B column is a crucial step for purification of the enzyme.

The fraction containing spore-lytic activity, which was obtained by BSA-Sepharose 4B column chromatography, was treated with antiserum raised against the 24 kDa protein as previously described. As shown in Fig. 3, the treatment led to a complete loss of enzyme activity, in parallel with a disappearance of the 24 kDa protein in the BSA-Sepharose 4B column-purified fraction. This strongly supports our view that the 24 kDa protein possesses the spore-lytic activity.

The N-terminal amino acid sequence of the spore-lytic enzyme was determined as FSNQVIQRGASGEKVIE-LQ (19 residues). We could not find any protein sharing this amino-terminus (computer analysis by using Genetyx homology search program and the SWISS-PROT protein database).

Degradation of coat-stripped spores by the enzyme resulted in the release of a large amount of free amino groups (5.3±0.3 μmol ml⁻¹ under the conditions described in Methods) and only a minor amount of reducing groups (0.03±0.006 μmol ml⁻¹). No free amino groups and reducing groups were liberated when the enzyme was boiled for 10 min before the assay. The molar ratio of free amino groups to reducing groups was 175, suggesting that no cleavage of the cortex polysaccharide had occurred.

**Effects of surfactant, temperature, pH, ionic strength and chemicals on spore-lytic enzyme activity**

The spore-lytic enzyme (0.4 units), which was partially purified by a BSA-Sepharose 4B column, was used in these experiments.

Besides C₁₂E₆, non-ionic surfactants such as sucrose monolaurate, octyl glucoside and dodecyl polyglycoside (final concn 0.1%) were also effective in preventing enzyme inactivation. SDS (0.02%), however, inactivated the enzyme.

In contrast to the heat-resistance of the enzyme in situ, the isolated enzyme was completely inactivated after incubation at 60 °C for 2 min or 45 °C for 10 min. Temperature sensitivity of the enzyme in solution is shown in Fig. 4(a).

The pH dependence of activity of the enzyme in solution
Table 3. The effects of various chemicals on germination and spore-lytic enzyme activity

Normal spores were incubated in 75 μl 0.25 M sodium phosphate buffer, pH 7.0, containing the indicated concentration of chemicals at 32 °C for 10 min. The above buffer (75 μl) containing 20 mM L-alanine and 8 mM adenosine was added to the spore suspension to induce germination. The spore-lytic enzyme (15 μl, 0.4 units) was mixed with 130 μl assay medium, to obtain the desired concentration listed. After being incubated at 32 °C for 10 min, 5 μl of coat-stripped spores were added and the enzyme activity measured. The final OD₅₅₀ of these spore suspensions was 0.1 in a cell of 1 mm light path. Sodium thioglycollate and EDTA were removed from the assay medium when effects of thiol reagents and ZnCl₂ were tested. Values are rounded off to integers. These data represent the mean of two (for germination) and three (for lytic activity) independent experiments (sd < 10% of the mean).

<table>
<thead>
<tr>
<th>Chemicals</th>
<th>Conc (mM)</th>
<th>Inhibition of decrease in OD₅₅₀ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Germination</td>
<td>Spore-lytic activity</td>
</tr>
<tr>
<td>D-Alanine</td>
<td>5</td>
<td>100</td>
</tr>
<tr>
<td>DFP</td>
<td>9</td>
<td>100</td>
</tr>
<tr>
<td>PSMF</td>
<td>5</td>
<td>35</td>
</tr>
<tr>
<td>HgCl₂</td>
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<td>100</td>
</tr>
<tr>
<td>ZnCl₂</td>
<td>1</td>
<td>95</td>
</tr>
<tr>
<td>NAM</td>
<td>0.5</td>
<td>-*</td>
</tr>
<tr>
<td>NEM</td>
<td>0.3</td>
<td>95</td>
</tr>
<tr>
<td>PCMB</td>
<td>0.5</td>
<td>100</td>
</tr>
<tr>
<td>PNAG</td>
<td>1</td>
<td>-*</td>
</tr>
</tbody>
</table>

* The assay medium contained 5% (v/v) DMSO, which is a solubilizer of NAM and PNAG. The medium containing 5% DMSO inhibited germination, although the enzyme activity was not affected.

Fig. 5. Effects of NEM on spore-lytic enzyme activity and germination. Germination and spore-lytic enzyme activity were measured at 32 °C, after incubation for 10 min in the presence of the indicated concentration of NEM, as described in the footnote of Table 3. Values are expressed as a percentage of the decrease in OD₅₅₀ and are the mean of two determinations (sd < 0.8% of the mean). ●, Spore-lytic enzyme activity; ○, germination.

In addition to the effect of ionic strength on the stability of the spore-lytic enzyme shown in Fig. 1, activity of the enzyme depended strongly on the salt concentration of the assay medium. Fig. 4(c) shows the effect of various concentrations of sodium phosphate on enzyme activity. When NaCl or KCl was added to a 5 mM solution of sodium phosphate, pH 7.0, optimal activity occurred at approx 30 mM, with a rapid decrease in activity at both lower and higher salt concentrations.

The effects of a variety of chemicals, including enzyme inhibitors, were examined on spore-lytic activity and germination (Table 3). D-Alanine, a competitive inhibitor of germination, had no effect on the enzyme activity. Serine protease inhibitors such as DFP and PMSF inhibited germination without affecting the spore-lytic enzyme activity, suggesting involvement of serine protease or serine protease-dependent reactions in the germination pathway other than cortex degradation. Although D- nitrophenyl-N-acetyl-β-D-glucosamide (PNAG) was not a substrate of the spore-lytic enzyme purified here, partial inhibition of the enzyme with the chemical may be due to structural homology between the spore cortex peptidoglycan and PNAG. Germination and spore-lytic enzyme activity were inhibited with thiol reagents such as HgCl₂, N-(9-acridinyl)maleimide (NAM), N-ethylmaleimide (NEM) and p-chloromercuribenzoate (PCMB). Sodium thioglycollate (10 mM) fully reversed the inactivation caused by the mercuric compounds. Fig. 5 shows the effects of NEM on spore-lytic enzyme activity and on germination. The results indicate involvement of multiple sites sensitive to thiol reagents in the germination process of B. cereus spores. ZnCl₂ inactivated the enzyme, but no effect was observed with MnCl₂, MgCl₂ and CoCl₂ (1 mM each).

DISCUSSION

We have purified a spore-lytic enzyme of molecular mass 24 kDa in an active form from the germination exudate of spores of B. cereus IFO 13597. The solubilized enzyme completely lost its activity when heated at 45 °C for 10 min (Fig. 4a). Prior to germination, the spores were extensively washed with 2 M KCl and 2 M urea, and incubated at 65 °C for 45 min. It is most likely that the enzyme released into the germination exudate is not a spore surface-bound enzyme, which might have been liberated from the spores or heat-denatured during urea or heat treatment of the spores.

Although the 24 kDa spore-lytic enzyme remains resistant to heat-treatment to which spores are exposed before germination, the enzyme in solution lost its resistance. The enzyme freed from spores was unstable and readily inactivated at low ionic strength. The presence of non-ionic surfactants effectively prevented the enzyme from inactivation. It has been indicated that reduction of ionic strength decreases hydrophobic interactions (Kennedy,
1990) and that non-ionic surfactants bind to hydrophobic regions on the protein without causing an appreciable change in its native structure (Makino, 1979). The enzyme was found to interact with BSA, which possesses binding sites for hydrophobic ligands (Makino, 1979). Taken together, it is reasonable to assume that protection of the hydrophobic surface on the 24 kDa spore-lytic enzyme by association with hydrophobic ligands is of primary importance to maintain the isolated enzyme in its native structure. In situ, the spore-lytic enzyme probably forms a complex with a hydrophobic substance(s), which leads to a stabilization of the enzyme in a spore-bound form. In addition to the proposed hydrophobic interaction, the enzyme activity also depends on ionic strength of the medium (Fig. 4c) and has minimal activity at salt concentrations > 0.1 M, at which the enzyme was stabilized. This may imply that the catalytic site of the enzyme is sensitive to an ionic environment, which might be of importance for the formation of the precise three-dimensional structure of the site.

The 24 kDa spore-lytic enzyme induced germination-like changes in coat-stripped spores, whereas the enzyme lacked the ability to lyse isolated cortical fragments and isolated cell walls. Such substrate specificities and the sensitivities of the 24 kDa enzyme to temperature, pH, Hg²⁺ and Zn²⁺ (Fig. 4a, b and Table 3) are similar to those of a germination-specific 29 kDa spore-lytic enzyme found in spores of B. megaterium KM, which is most probably an amidase-catalysing formation of muramic acid 3-lactam (Foster & Johnstone, 1987). Our preliminary results showed that exposure of the coat-stripped spores to the 24 kDa enzyme resulted in a release of free amino groups. This suggested that the 24 kDa enzyme is either an amidase or a peptidase. Unfortunately, it is impossible to compare the 24 kDa enzyme with the 29 kDa enzyme on the basis of amino acid alignment, because the sequence of the 29 kDa enzyme is not available. If the 24 kDa enzyme were an amidase, the enzyme would differ from a vegetative cell amidase from Bacillus subtilis (Foster, 1991; Kuroda & Sekiguchi, 1990), at least, in their N-terminal sequences.

Proteolytic cleavage activates the germination-specific 29 kDa spore-lytic enzyme in B. megaterium KM spores (Foster & Johnstone, 1988). Similarly, in spores of B. cereus T, spore-lytic enzyme activation by release from an inactive bound form has previously been suggested to occur during germination (Gould et al., 1966) and protease inhibitors are known to arrest germination of this organism (Boschwitz et al., 1985, 1991). This implies involvement of proteolytic activity in the germination pathway. The present experiments also suggested involvement of protease in the germination pathway of spores of B. cereus (Table 3). Furthermore, it has been demonstrated that two Hg²⁺-sensitive sites are involved in the germination process of spores of B. megaterium KM and that the germination-specific 29 kDa spore-lytic enzyme represents the second sensitive site (Foster & Johnstone, 1986). The most probable candidates for the Hg²⁺-sensitive sites are the thiol groups. The results shown in Fig. 5 suggest that, in addition to the 24 kDa spore-lytic enzyme, another thiol reagent-sensitive site must participate in the germination pathway of the B. cereus spores studied here. It is apparent that NEM can block germination at a stage prior to the inhibition of the 24 kDa enzyme. This suggests the location of a second Hg²⁺-sensitive site on the 24 kDa enzyme. Thus, it appears that B. cereus IFO 13597 shares a germination mechanism common to B. megaterium KM, which was proposed by Foster & Johnstone (1990). However, there is still much to be elucidated about the details of the germination mechanism. Studies are in progress to solve several questions such as whether a precursor of the 24 kDa enzyme is synthesized, whether modification of the enzyme by protease occurs, and the location of the enzyme in the dormant spore.

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