Purification and Properties of Spore-lytic Enzymes from Clostridium perfringens Type A Spores

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Spores of Clostridium perfringens contain at least two spore-lytic enzymes active in hydrolysing cortical peptidoglycan. One enzyme has been purified 1800-fold and has a molecular weight of 17400 determined from chromatography on Sephadex G-75. Two protein bands were apparent after SDS-PAGE. The isolated enzyme was investigated for response to temperature, pH, ionic strength and enzyme inhibitors, and for mode of action. A second enzyme activity, differing from the first in apparent molecular weight (29800) as determined by gel exclusion chromatography, and also in its pH optimum and activity on cortical substrate, was also isolated, although not purified to the same extent.

INTRODUCTION

Peptidoglycan has structural significance in all bacteria. In vegetative bacteria and bacterial spores peptidoglycan forms the major part of the rigid cell wall and spore cortex respectively. In spores, the cortex must be hydrolysed during germination in order for the spore protoplast to emerge. In Clostridium perfringens, deficiencies in or damage to the cortex-lytic enzymes can result in superdormant spores, or spores which, although viable, cannot germinate and grow out without an exogenous source of lytic agent such as lysozyme (Duncan et al., 1972).

Several researchers have described the cortex-lytic initiation protein (IP) detected in culture filtrates of vegetative and sporulating C. perfringens (Cassier & Ryter, 1971; Duncan et al., 1972; Franceschini & Labbe, 1979; Labbe et al., 1981). Ando (1979) and ourselves (Gombas & Labbe, 1981) have reported a spore-lytic enzyme (SLE) which can be extracted from cleaned C. perfringens spores. Lytic enzymes associated with sporulation and spores of C. perfringens are important for at least two reasons. First, it is the activity of an autolytic system which causes the release (in the intestine) of the mature spore and the concomitant release of an enterotoxin which is responsible for gastroenteritis. Secondly, SLE is involved in the germination of spores of this organism. It is the germination and subsequent vegetative cell multiplication which results in the large numbers of cells associated with food-borne illness due to this organism.

The biochemical properties of partially purified IP have been reported (Labbe et al., 1981). Here we report the isolation and characteristics of SLE from cleaned C. perfringens spores, and present evidence for a second spore-lytic enzyme activity.

METHODS

Organism. The organism used was Clostridium perfringens type A, strain NCTC 8798 (Hobbs serotype 9).

Media and sporulation. Spores were prepared and cleaned as described by Franceschini & Labbe (1979). Spores to be used to prepare SLE extracts were harvested from approximately 300 litres of sporulation medium, resulting

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Abbreviations: IP, initiation protein; SLE, spore-lytic enzyme; UME, urea/mercaptoethanol; CMC, carboxymethyl cellulose; DTT, dithiothreitol; NEM, N-ethylmaleimide; CMPS, chloromercuriphenylsulphonic acid; FDNB, fluorodinitrobenzene.

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in a final washed spore suspension of approximately $3 \times 10^{13}$ spores. Final spore suspensions contained >95% phase-bright spores, as determined by phase-contrast microscopy. Cleaned spores were harvested by centrifuging at 5°C and stored as frozen pellets until use.

**Enzyme extraction and purification.** Spores were heat activated (75°C, 15 min), then germinated with 0.1 mM-sodium nitrite, as previously described (Gombas & Labbe, 1981). Germinated spores were extracted for 2 h with 7.2 M-urea plus 10% (v/v) mercaptoethanol, pH 2.85 (UME) at 45°C. The spores were separated from the extract by centrifugation (3020 g, 10 min) at 20°C, and re-extracted for an additional 2 h in fresh UME (secondary UME extract). The two extracts were treated similarly, but separately, through the purification. The extracts were dialysed overnight against approximately 100 vols distilled water at 4°C. Insoluble material was removed from the crude retentate by centrifugation at 3020 g for 10 min at 5°C, resulting in a 1.8-fold increase in purity (Table 1). Phosphate-treated SLE was prepared by adding sufficient 0.5 M-sodium phosphate, pH 6-0, to the supernatant to bring the phosphate concentration to 60 mM. After 5-10 min at 22°C, the precipitate was removed by centrifugation, without measurable loss of SLE activity. Phosphate-treated SLE was used in some studies of crude SLE properties. SLE in solution was concentrated by placing the enzyme preparation in distilled water-washed dialysis tubing, and dehydrating by osmosis on a bed of polyethylene glycol 20000 (Sigma). Activity recovery ranged from 65 to 100%. Optimum recovery required post-concentration dialysis (usually in 60 or 500 mM phosphate buffer, pH 6-0). Any precipitate that formed during the concentration process was collected and redissolved overnight in a small volume of 0-5 M-phosphate buffer. If it contained significant SLE activity, it was added back to the concentrate. The phosphate-treated material was dialysed overnight against run buffer (60 mM-sodium phosphate, pH 6-0, plus 0.2% sodium azide) and then added to a carboxymethyl cellulose (CMC) column.

CMC (0.72 mequiv. g⁻¹, Sigma) was hydrated and charged as described by Himmelhoch (1971). The CMC column (1.5 cm i.d. × 8.3 cm bed height) was washed with at least 10 bed volumes of 0.5 M-sodium phosphate, pH 6-0, then equilibrated with 3-4 bed volumes of run buffer. Phosphate-treated SLE was pumped onto the column at 0.6 ml min⁻¹ (20°C). The column was washed with run buffer until the protein elution was negligible (3-4 bed volumes). SLE was collected by isocratic elution: SLE from the primary UME extract was eluted with 125 mM-NaCl in run buffer (Table 1, IIA); SLE from the secondary UME extract was eluted with 65 mM-NaCl in run buffer (Table 1, IIB); eluted fractions were collected in acid cleaned tubes (3 ml per tube). Fractions containing the highest specific activity were pooled (>10000 U mg⁻¹ for the primary extract, >50000 U mg⁻¹ for the secondary extract). Pooled eluates were then concentrated to less than 10 ml and dialysed against run buffer before fractionation on Sephadex G-75 Superfine.

SLE was mixed with 70% (w/v) sucrose and layered onto the gel column (2.6 cm i.d. × 97 cm bed height). SLE was eluted at 0.5 ml min⁻¹ and 4 ml fractions were collected. Two peaks with spore-lytic activity were recovered and designated β-SLE and α-SLE, for the earlier and later eluting activities, respectively (Fig. 1a). Fractions with the highest specific activity were pooled for each extract: >85000 U mg⁻¹ for α-SLE of the primary UME extract; >100000 U mg⁻¹ for α-SLE of the secondary UME extract; and >26000 U mg⁻¹ for β-SLE of the primary UME extract. These preparations were used for characterization of the properties of SLE.

**Molecular weight determination by gel exclusion chromatography.** The same Sephadex G-75 Superfine column and procedures were used as described above. Molecular weight standards were mixed with 70% sucrose and layered onto the gel under the buffer. Sample volumes were <2% of the gel bed volume. The void volume was determined by using dextran blue 2000 (Sigma). Molecular weight standards used were: ovalbumin (43000, Sigma), chymotrypsinogen (25000, Sigma), horse myoglobin (17200, Sigma) and ribonuclease (13700, Pharmacia). The elution volume of standards was determined from the absorbance at 280 nm; the elution volume of SLE was determined at peak activity, as assayed on coatless spores (see below).

**SDS-PAGE.** This was done by the method of Laemmli & Favre (1973), as modified by Pandey & Aronson (1979). Acrylamide gels (18%, w/v) were run in a slab gel apparatus (Hoefer Scientific Instruments, SE-600) on the day of preparation. The gels were stained using Coomassie Blue R-250 and recorded either by photography or by densitometer measurements using a soft laser scanning densitometer (Biomed Instruments, Fullerton, Calif., USA). The following proteins (8-10 μg of each; all from Sigma) were used as molecular weight standards: phosphorylase b (94000), bovine serum albumin (67000), ovalbumin (43000), alcohol dehydrogenase (37700), myoglobin (17200) and lysozyme (14400).

**pH optimum determination.** Buffers (30 mM-cation concentration) were prepared at pH values in the range 2.75-11.0 as follows: sodium acetate, pH 4.1-5.6; citrate/phosphate, pH 2.75-6.7 and sodium phosphate, pH 6.1-11.0. Buffer (190-195 ml) was mixed with either coatless spores or cortical fragments to give a final volume of 2.00 ml and a final OD₄₀₀ of 0.4-0.5. The enzyme, dissolved in 60 mM-sodium phosphate, pH 6.0, was mixed into the prewarmed (37°C) substrate/buffer mixture and assayed for activity. The pH of the post-reaction mixture differed from the original pH by <0.1 unit.

**Effects of cations.** The salt solutions used had a final pH of 8.0 ± 0.1 at 25°C. They were mixed with assay spores to give a final volume of 2.00 ml and an OD₄₀₀ of 0.4-0.5. Cobalt chloride samples were assayed at 600 nm due to interfering absorbance at 400 nm. Enzyme (25 μl), in 60 mM-sodium phosphate, pH 6.0, was mixed into the
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prewarmed (37 °C) substrate/salt solution, and assayed for activity. The pH of the post-reaction mixture differed from the original by <0.1 unit.

**Enzyme solubility.** Samples of the enzyme were pipetted into equal volumes of buffer (200 U ml⁻¹, final concentration) and held at 25 °C for up to 15 min. Samples were centrifuged (3020 g, 10 min) to remove precipitated material, and supernatants were collected and assayed. In some cases, pelleted material was redissolved by adding a small volume of 0.5 M-buffer to the centrifuge tube, vortexing, and storing overnight at 4 °C. Material which did not redissolve was removed by centrifugation.

**Temperature stability.** The enzyme was incubated in 60 mM-sodium phosphate, pH 6.0, in a water bath adjusted to various temperatures, ± 1 °C. Temperature was measured with a thermometer placed in an identical tube with a similar volume of water. At various times, 25 μl samples were removed and assayed at 37 °C in 2.0 ml 20 mM-sodium phosphate buffer, pH 7.8, containing assay spores. Activity is reported as the percentage activity remaining as compared with the preincubation activity.

**Protease activity.** Wells (3 mm diam.) were made in 3–5 mm deep plates of 0.2% (w/v) casein in 2% (w/v) agar (buffered to pH 7.8 with 10 mM-sodium phosphate) and were filled with 20–25 μl enzyme. Trypsin (1 mg ml⁻¹; Sigma) was used as a positive control. Plates were stored at 25 °C for up to 5 d. Protease activity appeared as a clear zone surrounding the well. Protease activity was also assayed on Azocoll (Calbiochem) by the method of Cheng & Aronson (1977).

**Enzyme inhibitors.** Dithiothreitol (DTT; Sigma), N-ethylmaleimide (NEM; Sigma), mercuric chloride (Fisher Scientific, Fairlawn, NJ, USA), and potassium ferricyanide (Sigma) were dissolved in 60 mM-Tris/HCl, pH 8.0.

p-Chloromercuriphenylsulphonic acid (CMPS; Sigma) was dissolved in 25 mM-Tris/maleate, pH 6.0. Coatless spores (50 μl) were mixed with each reagent or with buffer alone to give a final volume of 2.0 ml and OΔ₄₀₀ of 0.4–0.5, and prewarmed to 37 °C. The enzyme (25 μl in 60 mM-sodium phosphate) was added and the OΔ₄₀₀ measured. CMPS-treated samples were assayed at 600 nm. The percentage inhibition was calculated from the ratio of activity of SLE in the reagent to activity in buffer alone. In some experiments, enzyme was incubated with 1.0 ml DTT or buffer at 37 °C for 2 min, and then 1.0 ml reagent was added and the mixture incubated for 2 min before addition of assay spores.

**Lysozyme assay.** Micrococcus lysodeikticus cells (Sigma) were suspended in 2.0 ml of either sodium phosphate buffer, pH 7.8, or citrate/phosphate buffer, pH 4.5, and pre-warmed to 37 °C. Enzyme was added to the suspension, and the OΔ₄₀₀ measured for 25 min. Lysozyme (0.1 mg ml⁻¹; Sigma) was used as a positive control.

**N-Acetylglucosaminidase assay.** p-Nitrophenyl-N-acetylglucosaminide (Sigma) was dissolved in sodium phosphate buffer, pH 7.8, or citrate/phosphate buffer, pH 4.5, and pre-warmed to 37 °C. Enzyme was added to the solution, and the A₄₅₀ measured for 25 min. N-Acetylglucosaminidase (Sigma) was used as a positive control.

**Reducing sugars.** Generation of reducing sugars during enzyme hydrolysis of cortical fragments was measured by the ferricyanide method (Thompson & Shockman, 1968) using N-acetylglucosamine (Sigma) as the standard.

**N-terminal amino groups.** Measurement of N-terminal amino groups generated during enzyme hydrolysis of cortical fragments was by the fluorodinitrobenzene (FDNB) procedure described by Ghuysen et al. (1966).

**SLE activity assay.** SLE was assayed as previously described (Gombas & Labbe, 1981) using as substrate coat-stripped C. perfringens spores (Franceschini & Labbe, 1979), or cortical fragments from C. perfringens spores prepared as described by Hashimoto et al. (1972). Complete removal of spore coats was verified by electron microscopy. Unless otherwise stated, a wavelength of 400 nm was used for all activity assays, which were done in a Beckman 25 spectrometer equipped with a kinetic system. One unit (U) of activity was defined as a decrease in the OΔ₄₀₀ of 0.01 min⁻¹.

**Protein assay.** Protein was measured by the dye-binding method of Bradford (1976), using bovine serum albumin (Sigma) as the standard.

**RESULTS**

**Purification of SLE**

The purification protocol used (Table 1) resulted in a 15–16% recovery of SLE activity from the primary UME extract, with an overall purification of approximately 600-fold and 200-fold for α- and β-SLE respectively. When analysed by SDS-PAGE, α-SLE was found to contain one major band (Fig. 2c; 64% of the total stained protein based on densitometer tracings, Fig. 2d) of molecular weight 29500, a lesser band (11% of total protein, molecular weight 22500), and four lower molecular weight bands. The β-SLE fraction was found to contain bands at all six molecular weights (Fig. 2a), in approximately the same ratio as in α-SLE, and five additional protein bands with apparent molecular weights of 38000, 33000, 32000, 22400, and 17500 (with relative peak intensities of 30.5, 30.5, 22, 33, and 15%, respectively, as compared with the
Purification of the secondary UME extract by the modified protocol (i.e. 65 mM-NaCl instead of 125 mM-NaCl for the isocratic elution of activity from CMC) resulted in approximately 30% less activity eluted from CMC, but an increase in the overall purification to approximately 1800-fold (Table 1). Only a barely detectable β-SLE fraction, less than 2% of the α-SLE activity, was eluted from Sephadex G-75 by this modified protocol (Fig. 1b). When analysed by SDS-PAGE, the α-SLE fraction was found to contain only the 29500 and the 22500 bands (Fig. 2f), in a ratio of 8:1, based on densitometer tracings (Fig. 2g). This α-SLE fraction was used for the characterization of the properties of α-SLE.

When phosphate-treated SLE was assayed on coatless spores, a sharp pH optimum was found at approximately 4.25, with a lesser peak (approximately 70% of the pH 4.25 peak) over the range of pH 6.6-8.4 (Fig. 3a). Negligible activity was found at pH values < 3.1 or > 8.75. Essentially the same profile was found at pH values < 5.0 if activity was assayed on cortical fragments, although the secondary peak at pH 6.2-8.0 was less prominent.

When α-SLE was assayed on coatless spores over the same pH range, the pH optimum was 7.2-8.4 (Fig. 3b). When α-SLE was assayed on cortical fragments at pH 7.7 or 4.2, negligible activity was found.

When β-SLE was assayed on coatless spores over the same pH range, the activity profile was essentially the same as that of the phosphate-treated SLE (Fig. 3c): the pH optimum was at approximately pH 4, with a lesser peak at pH 6-75 having 70% of the optimum activity. When β-SLE was assayed on cortical fragments, it had the same activity (U ml⁻¹) on cortical fragments as on coatless spores at pH 3.6, but only 18% of the activity expressed on coatless spores when assayed at pH 7.7. Subsequently, α-SLE activity was assayed at pH 7.8-8.1 and β-SLE activity was assayed at pH 4.2.
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Fig. 2. SDS-PAGE of purified SLE eluted from Sephadex G-75 SF column. (a–d) Material from CMC column eluted with 125 mM-NaCl before separation on Sephadex column: (a) β-SLE, (b) molecular weight standards, (c) α-SLE, (d) densitometer tracing of (c). (e–g) Material from CMC column eluted with 65 mM-NaCl before separation on Sephadex column: (e) molecular weight standards, (f) α-SLE, (g) densitometer tracing of (f).

Table 1. Summary of SLE purification protocol
The values are from a single experiment.

<table>
<thead>
<tr>
<th>Purification step</th>
<th>Protein activity (mg ml⁻¹)</th>
<th>Specific activity (U mg⁻¹)</th>
<th>Recovery (%)</th>
<th>Purification (fold)</th>
</tr>
</thead>
<tbody>
<tr>
<td>I Crude dialysate</td>
<td>0.76</td>
<td>213</td>
<td>100</td>
<td>1.0</td>
</tr>
<tr>
<td>Centrifuged</td>
<td>0.56</td>
<td>380</td>
<td>107</td>
<td>1.8</td>
</tr>
<tr>
<td>Phosphate-treated</td>
<td>0.15</td>
<td>4100</td>
<td>107</td>
<td>19</td>
</tr>
<tr>
<td>Concentrated</td>
<td>0.755</td>
<td>4700</td>
<td>85</td>
<td>22</td>
</tr>
<tr>
<td>IIA Pooled CMC fractions (eluted with 125 mM-NaCl)</td>
<td>0.275</td>
<td>31700</td>
<td>65</td>
<td>150</td>
</tr>
<tr>
<td>Concentrated</td>
<td>ND</td>
<td>ND</td>
<td>47</td>
<td>ND</td>
</tr>
<tr>
<td>α-SLE</td>
<td>0.55</td>
<td>141000</td>
<td>10</td>
<td>660</td>
</tr>
<tr>
<td>β-SLE</td>
<td>0.57</td>
<td>41900</td>
<td>5.6</td>
<td>200</td>
</tr>
<tr>
<td>IIB Pooled CMC fractions (eluted with 65 mM-NaCl)</td>
<td>0.07</td>
<td>75000</td>
<td>ND</td>
<td>350</td>
</tr>
<tr>
<td>Concentrated</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
<td>ND</td>
</tr>
<tr>
<td>α-SLE</td>
<td>0.25</td>
<td>390000</td>
<td>ND</td>
<td>1800</td>
</tr>
</tbody>
</table>

ND, Not determined.

Properties of SLE

Temperature stability. α-SLE in 60 mM-sodium phosphate, pH 6.0, retained 90% of its activity after incubation at 42.5 °C for 45 min (Fig. 4). However, incubation of the isolated enzyme at higher temperatures resulted in rapid inactivation, with 88% loss of activity after 2.5 min at 57.5 °C. Both purified α-SLE and phosphate-treated crude SLE were stable at 4 °C for several weeks. Purified α- and β-SLE each retained >70% activity after several weeks of frozen storage.
** Ionic strength effects.** Optimum activity of α-SLE occurred at a monovalent cation concentration of approximately 25 mM, plus 5 mM-Tris/maleate, pH 7.8, with a rapid decrease in activity at lower monovalent cation concentrations and a gradual decrease at higher concentrations. The activity profile was identical for the cations sodium, potassium and ammonium. Activity in 5 mM-buffer alone was 11–12% of optimum, while activity in 100 mM-salt was approximately 35% of optimum.

Maximum activity in the presence of the divalent cations calcium or magnesium occurred at the lower concentration of 2.5–10 mM. Activity dropped off rapidly at higher concentrations, with 40% of maximum at 25 mM-cation, and <10% of maximum at >50 mM-cation. No detectable activity was found in the presence of the divalent cation cobalt in the range of 1–100 mM, and only 35% of maximum at 0.1 mM.

At pH 7.9, α-SLE responded to the sodium phosphate buffer concentration as it did to other monovalent cations, with maximum activity at a buffer concentration of 20–25 mM, falling off rapidly at higher or lower concentrations (Fig. 5). Negligible activity was found at buffer concentrations >100 mM. At buffer concentrations of <10 mM solubility of α-SLE was <20% of the maximum. Solubility gradually increased at higher buffer concentrations until reaching >80% at buffer concentrations of >50 mM. Enzyme solubility in 20–25 mM-sodium phosphate was approximately 50%. Activity which was lost due to precipitation could be quantitatively recovered after resolubilization of pelleted material in 0.5 M-buffer.

**Molecular weight.** Analysis by gel exclusion chromatography on Sephadex G-75 indicated a Stokes radius equivalent to a molecular weight of 17400 for α-SLE, and 29800 for β-SLE (Fig. 6).
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Fig. 5. Effect of sodium phosphate, pH 7.8, on activity (▪) and solubility (○) of α-SLE. Activity was assayed in 20 mM-phosphate as described in Methods. The values are representative of those obtained in duplicate experiments.

1 a). Analysis by SDS-PAGE indicated a molecular weight for the major protein band in both α- and β-SLE of 29500 (Fig. 2). The reasons for the apparent discrepancy in the molecular weight measured by the two procedures are discussed below.

Enzyme inhibitors. SLE activity on coatless spores was inhibited by thiol-active agents to various degrees: mercuric chloride (0.05 mM) inhibited α-SLE at pH 7.8 by >97% and CMPS (10 mM) inhibited phosphate-treated SLE at pH 6.0 by 80%. NEM concentrations as high as 50 mM were only 40–60% inhibitory to α-SLE, while DTT was slightly stimulatory at 0.1–10 mM. Potassium ferricyanide (50 mM) was >90% inhibitory to α-SLE. All inhibitory effects of thiol-active agents were partially prevented by pre-incubation of enzyme with equimolar DTT, except for mercuric chloride, which required a 1000:1 ratio for 50% retention of activity. Potassium ferricyanide inhibition was not prevented by equimolar DTT.

Proteolytic activity. Neither phosphate-treated SLE, α-SLE, nor β-SLE showed detectable proteolytic activity on casein plates after 5 d. Proteolysis assays on Azocoll at pH 4.5 and 7.8 were also negative.

Mode of action

α- and β-SLE were both negative for muramidase and N-acetylglucosaminidase activity when each was incubated with M. lysodeikticus or p-nitrophenyl-N-acetylglucosaminide, respectively. Incubation with cortical fragments indicated β-SLE to have approximately equal activity on germination of coatless spores and hydrolysis of cortical fragments. At equivalent levels of spore-lytic activity, the α-SLE preparation had <2% of the corticolytic activity of the β-SLE preparation. However, α-SLE was able to slowly hydrolyse cortical fragments to about the same final percentage change in OD₄₀₀ as β-SLE (32% and 37% for α- and β-SLE respectively). When cortical fragments, incubated with α-SLE or β-SLE until the maximum decrease in OD₄₀₀ had occurred, were assayed for reducing power, the α-SLE sample was not significantly different from untreated cortex, while β-SLE samples were strongly positive. Samples analysed for FDNB-reactive groups were not significantly different from untreated cortical fragments, indicating the absence of additional free amino groups following treatment with α- or β-SLE.

DISCUSSION

Although studies have shown that spores of Bacillus cereus contain at least four different cortex-degrading enzymes (Warth, 1972), no previous reports have indicated more than one spore-lytic activity from spores of C. perfringens. As that activity is undoubtedly involved in germination of spores, it was considered of interest to compare some of the properties of the isolated enzyme with the observed characteristics of the spore. In order to minimize misinterpretation of enzyme properties because of competing or co-activities in the enzyme
preparation, steps were taken to purify the enzyme from any potentially interfering proteins. It was only during the final steps of the purification that the SLE activity resolved into two fractions, which were designated α-SLE and β-SLE. Because of the higher concentration of α-SLE activity in the preparation, efforts were focused on elucidating its properties. Preliminary trials indicated that the protocol used for purification of the primary UME extract would yield a preparation with a single protein component, as determined by SDS-PAGE. However, in scaling up from the 10^10 spores per extraction of the preliminary trials to the 10^13 spores used for this study, several previously undetected proteins became evident in the final α-SLE fraction of the primary extract. Decreasing the ionic strength of the eluent for the CMC elution of the secondary UME extract decreased SLE recovery but yielded a purer α-SLE preparation. However, it is not known if the improved purification was due to the lower ionic strength eluent, or because the contaminating proteins were not present in the secondary extract.

After purification of α-SLE to two protein bands as determined by SDS-PAGE, its properties were investigated. Like the phosphate-treated preparation and many other spore enzymes, α-SLE is not inherently heat resistant, being inactivated by temperatures far below those used in thermal activation, such as used on the spores from which the SLE was extracted. Consequently, both SLE activities must be stabilized or otherwise protected in vivo. By contrast, Srivastava & Fitz-James (1981) reported that partially purified cortex-lytic enzyme and purine nucleoside hydrolase from Bacillus subtilis spores were stable for 20 min at 75 °C and 50 °C, respectively.

An N-acetylmuramyl-L-alanine amidase, extracted from sporulating cells of C. perfringens (Traci, 1977), was reported to be stimulated by cobalt ions. In contrast, α-SLE was totally inactive in millimolar concentrations of cobalt, and was inhibited by concentrations which were reported to be stimulatory to the amidase.

Germination of bacterial spores, including those of C. perfringens, has been found to be greatly stimulated by various cations, including potassium and calcium (Ando, 1978; Levinson & Feeherry, 1975). By contrast, activity of α-SLE, as well as the more crude preparation, was very sensitive to ionic strength in vitro, showing activity only in a narrow range of monovalent cation concentrations, and an even narrower range of divalent cations.

Like the cruder preparation, α-SLE demonstrated an unexpected solubility profile in various ionic strength buffers. The property of insolubility in the narrow ionic range in which SLE has activity, and the loss of enzyme activity as solubility increases, may be related; e.g. low ionic strength may cause the enzyme to precipitate and adsorb to its substrate, while high ionic strength may preclude contact between the two. The isoelectric points of both SLE activities are > pH 8 (data not shown), so at a physiological pH, SLE is positively charged. As its substrate, cortical peptidoglycan, is negatively charged (Gould & Dring, 1974), the attraction between the two could be electrostatic, and ions in solution might interfere with the interaction.

An optimal UME extraction of a spore pellet yielded about 52000 SLE units per 10^12 spores. A specific activity of 3.9 × 10^5 U mg^-1 has been attained for α-SLE, which has an apparent molecular weight of approximately 17000. Assuming near 100% efficiency in the extraction of α-SLE, and an even distribution of active enzyme from the spores extracted, these values indicate a maximum of 4700 molecules per spore, or about 0.05% of the total spore dry weight. Higher actual values for molecular weight (see below) or specific activity, or separate consideration of β-SLE, would decrease the estimated number of molecules per spore.

The resolution of α- and β-SLE on Sephadex G-75 was initially thought to be an artefact, since only one activity peak was resolved by other purification methods, notably Sephadex G-100. However, no such discrepancy was observed with other proteins on Sephadex G-75 and G-100 columns. Further, β-SLE has a very different pH optimum from α-SLE, and a different degree of activity on cortical fragments (although both are apparently capable of lysing cortical fragments). On the other hand, SDS-PAGE of both fractions indicated that not only the major protein band but also the minor protein band of α-SLE appeared to be the same as in the β-SLE fraction. This might be easily understandable if α-SLE were a subunit of β-SLE; however, the major protein fraction of α-SLE on SDS-PAGE was 1.7 times larger than indicated by the elution volume on Sephadex G-75. β-SLE, on the other hand, had the same apparent molecular weight on Sephadex G-75 as the major protein fraction on SDS-PAGE. Moreover, considering
the good resolution of α-SLE and β-SLE on Sephadex G-75, none of the molecular weight 30000
β-SLE protein should have been included in the α-SLE fractions. This apparent paradox would
be expected if α-SLE had more than a typical affinity for Sephadex G-75, and was retarded on
the column, thus only appearing to have a smaller molecular weight, while β-SLE, for reasons
unknown, did not have this affinity for Sephadex, and so eluted normally. Such an affinity for
Sephadex gels has been reported for amidases (Herbold & Glaser, 1975; Williamson & Ward,
1979) and glucosaminidases (Taylor et al., 1980; Williamson & Ward, 1981). Whether this is the
case or not, elution on Sephadex G-75 Superfine has been the only technique we have found to
resolve the two activities.

Another lytic enzyme which appears in the culture filtrate during growth of C. perfringens has
been called initiation protein (IP), so termed for its ability to initiate germination in sensitized
spores (Cassier & Sebald, 1969; Duncan et al., 1972). This enzyme also attacks both coatless
spores and isolated cortical fragments (Labbe et al., 1981). Partially purified IP has been found to
have an apparent molecular weight of 100000 (Labbe et al., 1981), which is three to six times
larger than the enzymes described here. It is apparent, therefore, that C. perfringens does
produce more than one spore-lytic enzyme. Synthesis of more than one lytic enzyme is not
unusual for spore-forming bacteria. For example, Kingan & Ensign (1968) reported three
autolytic enzymes from sporulating cells of Bacillus sphaericus. Brown et al. (1977, 1978) reported
two separate lytic enzymes present in B. cereus spores: a 'surface' or coat-associated enzyme
and a core-associated enzyme; while both enzymes lysed cortical fragments, only the core-associated
enzyme could germinate coat-stripped spores.

The relationship between α-SLE and β-SLE is unknown. Exhaustive efforts were made using
non-denaturing PAGE and isoelectric focusing to determine which of the protein bands were
actually associated with SLE activity. Thus far, we have been unsuccessful in eluting or
otherwise detecting any SLE activity from electrophoretic gels; this precludes the use of
immunochemistry to test the identity of the two activities when neither is in a homogeneous
preparation. Although α-SLE and β-SLE enzyme activities are corticolytic, the specific nature of
each was not determined. Cortical fragments reacted with β-SLE had a substantially higher
concentration of reducing sugars than non-reacted fragments, but showed no increase in free
amino groups. Cortical fragments reacted with α-SLE had neither more reducing sugars nor
more terminal amino groups than the untreated substrate, although it is possible that the latter
was due to a high background of amino groups in the cortex (Ghuysen et al., 1966). The results,
however, suggest that α-SLE and β-SLE lack amidase and peptidase activities.

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