Analysis of *Bacillus subtilis* 168 prophage-associated lytic enzymes; identification and characterization of CWLA-related prophage proteins

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The CWLA lytic amidase of *Bacillus subtilis* 168 was purified and antisera raised against the purified protein. No expression of *cwLA* could be demonstrated under any conditions by the use of the antisera and *cwLA::ZacZ* fusion analysis. Two lytic enzymes of apparent molecular masses 34 and 30 kDa (as measured by renaturing SDS-PAGE) were found to be mitomycin C-inducible, the larger of which corresponds to a protein immunologically related to CWLA. Both of these inducible lysins were found to be encoded by prophage PBSX. Prophage SPβ was shown by renaturing SDS-PAGE to produce a 43 kDa lytic enzyme unrelated immunologically to CWLA. The smaller of the two PBSX enzymes was purified and found to be an N-acetylmuramyl-L-alanine amidase of 32 kDa (as measured by SDS-PAGE and Coomassie blue staining) which cross-reacts only weakly with the anti-CWLA sera. The potential origin of *cwLA* and its possible relationship to the other phage lytic enzymes are discussed.

Introduction

Autolysins are apparently ubiquitous bacterial enzymes capable of hydrolysing cell wall peptidoglycan (Ghuysen *et al.*, 1966). Many species contain multiple autolysins but their role during growth and division has yet to be established. *Bacillus subtilis* 168 has two major vegetative cell autolysins, a 90 kDa glucosaminidase and a 50 kDa amidase (Herbold & Glaser, 1975; Rogers *et al.*, 1984). The structural genes for these enzymes have been isolated recently, sequenced and inactivated (Kuroda & Sekiguchi, 1991; Lazarevic *et al.*, 1992; Margot, 1992; Margot & Karamata, 1992). The mutant inactivated in the gene encoding the 50 kDa amidase was more resistant to lysis during stationary phase than the parent strain. Two other autolysins of 34 and 30 kDa have been identified during vegetative growth by the use of renaturing SDS-PAGE; however, their hydrolytic bond specificity is unknown (Foster, 1991). During differentiation, three novel enzymes appear, one of which may be responsible for mother-cell lysis at the end of sporulation.

A further *B. subtilis* lytic enzyme has been identified by a direct screening technique from expression libraries of *B. subtilis* genomic DNA (Kuroda & Sekiguchi, 1990; Foster, 1991). The cloned gene encodes a 30 kDa amidase which in *Escherichia coli* is processed to a 21 kDa form (Foster, 1991; Kuroda *et al.*, 1991). A mutant inactivated in the structural gene for this enzyme (*cwLA*) showed the same phenotype as the parent and the inactivation of *cwLA* cannot be correlated with the loss of any of the known autolysins during vegetative growth or differentiation (Foster, 1992).

*B. subtilis* generally contains two prophages, SPβ and PBSX, which can be induced during growth by mitomycin C treatment resulting in cell lysis by the action of phage lytic enzymes (Mauel & Karamata, 1984; Hemp-hill, 1990). Two PBSX lytic enzymes have been identified previously as an amidase of 14.5 kDa and a muramidase of undetermined size (Ward *et al.*, 1982). However CWLA cannot be encoded by SPβ or PBSX as it maps in a different part of the *B. subtilis* chromosome (Piggot, 1989; Foster, 1991).

This paper describes work to try to establish the identity and expression conditions of the CWLA lytic enzyme to elucidate its role during growth and relationship to the other lytic enzymes of *B. subtilis* 168. As part of this study the lytic enzymes of SPβ and PBSX have been characterized by renaturing SDS-PAGE and a PBSX amidase has been purified.

Methods

Bacterial strains and growth conditions. All strains of *Bacillus subtilis* 168 used in this study are shown in Table 1. Growth was carried out at 37 °C in nutrient broth (NB, Oxoid) with shaking (250 r.p.m.) unless otherwise stated. Growth media for strains 1A60 and RB1144 were...
supplemented with uracil and thymine (each 50 pg ml⁻¹), respectively.

Purification

Purification of the recombinant CWLA protein and preparation of antiserum. A CWLA-containing extract (50 ml) of *E. coli* DH5α with plasmid pSFP102 was prepared as described previously (Foster, 1991) from 51 of culture. The extract was centrifuged (100000 g, 1 h, 4 °C) to remove insoluble material and the supernatant dialysed overnight against 5 l of 10 mM-Tris/HCl, pH 7-5 at 4 °C. After dialysis a white precipitate had formed which contained > 95% of the enzyme activity. This precipitate was collected by centrifugation (10000 g, 5 min, 4 °C) and washed three times in dialysis buffer by resuspension and centrifugation. The final pellet was then resuspended in 50 ml of 300 mM KCl, 50 mM-Tris/HCl, pH 7-5 (resuspension buffer) by gentle mixing at 4 °C overnight. After centrifugation (20000 g, 20 min, 4 °C) to remove insoluble material, the enzyme-containing solution was loaded on a 10 x 16 cm column containing Reactive Red 120-Agarose (Sigma) equilibrated with enzyme resuspension buffer. All the enzyme bound to the column and was eluted with a linear gradient of LiCl (0-3 M) in enzyme resuspension buffer. The enzyme-containing fractions were pooled and the enzyme precipitated by dialysis against two changes of 100 vols of 10 mM-Tris/HCl, pH 7-5 at 4 °C. After dialysis the precipitated enzyme was recovered and resuspended as above. After SDS-PAGE (Laemmli, 1970), the 21 kDa protein corresponding to the processed major recombinant form of CWLA was electroeluted (Extraphor, LKB), extensively dialysed against water and the resulting precipitate recovered. The preparation was resuspended in water and used for the production of anti-CWLA sera in a rabbit. For the initial injections, 100 µl of purified protein was used and 50 µg per boost.

Construction of a *B. subtilis* strain carrying a cwla::lacZ fusion. Plasmid pSFP100 carries a 3 kb EcoRI insert containing the cwla gene and surrounding sequences (Foster, 1991). A 936 bp HindIII fragment was excised from this plasmid, end-filled and purified (Genecelean, Stratex Scientific). This fragment contains the coding sequence for the first 108 amino acids of CWLA and 672 bp of upstream sequence (Foster, 1991). After ligation into SmaI-cut and dephosphorylated pAZ106 (Kemp et al., 1991; A. Zuberi & R. Doi, unpublished data) this construct was used to transform *E. coli* DH5α (Hanahan, 1983). The orientation of the fragment in pAZ106, which created a transcriptional fusion between cwla and lacZ, was confirmed by restriction mapping and the construct pSFP10 used to transform competent *B. subtilis* 168 HR (Anagnostopoulos & Spizizen, 1961). Recombinants were selected by plating on erythromycin (1 µg ml⁻¹) and lincomycin (25 µg ml⁻¹) containing nutrient agar plates. The insertion resulting in the cwla::lacZ fusion was checked by Southern blotting (Sambrook et al., 1989) using the 936 bp HindIII fragment as probe. β-Galactosidase activity was measured by the method of Youngman (1990) using 4-methylumbelliferyl-β-D-galactopyranoside (MUG) as the substrate.

Renaturing SDS-PAGE and Western blotting. SDS extracts of cultures were prepared for SDS-PAGE as described previously (Foster, 1992). Renaturing SDS-PAGE and visualization of lytic enzyme activity was exactly as described previously (Foster, 1992). Samples for Western blot analysis were separated by 11% SDS-PAGE, transferred to nitrocellulose and treated with the anti-CWLA sera by the method of Burnet (1981). Antibody-antigen complexes were detected by the use of goat anti-rabbit IgG coupled to alkaline phosphatase (Sigma). All experiments were repeated at least twice and the results show representative examples.

Mitomycin C and prophase mutant induction. Mitomycin C induction was performed by the addition of mitomycin C (Sigma) to a final concentration of 0.5 µg ml⁻¹ (Buxton, 1980) to a growing culture at 37 °C in nutrient broth. SPF was induced in the strain CU1089 by the addition of 1/3 volume of boiling NB to an exponentially growing culture at 37 °C (Rosenthal et al., 1979). The culture was held at 50 °C for 6 min and then returned to incubate at 37 °C to allow the phase to replicate. PBSX was induced in the strain L4587 by the transfer of an exponentially growing culture at 37 °C to a shaking water bath at 45 °C to allow phage replication and cell lysis.

Purification and characterization of a PBSX lysin. Following temperature induction of a 500 ml culture of L4587 which carries the xin mutation (Buxton, 1976), lysis was allowed to proceed for 3 h. Debris was then removed by centrifugation (15000 g, 20 °C, 10 min; step 1). The supernatant containing the enzyme activity was concentrated by centrifugation (10000 g, 5 min, 4 °C) and washed three times in dialysis buffer by resuspension and centrifugation. The final pellet was then resuspended in 5 ml of 300 mM KCl, 50 mM-Tris/HCl, pH 7-5 (resuspension buffer) by gentle mixing at 4 °C overnight. After centrifugation (20000 g, 20 min, 4 °C) to remove insoluble material, the enzyme-containing solution was loaded on a 10 x 16 cm column containing Reactive Red 120-Agarose (Sigma) equilibrated with enzyme resuspension buffer. All the enzyme bound to the column and was eluted with a linear gradient of LiCl (0-3 M) in enzyme resuspension buffer. The enzyme-containing fractions were pooled and the enzyme precipitated by dialysis against two changes of 100 vols of 10 mM-Tris/HCl, pH 7-5 at 4 °C. After dialysis the precipitated enzyme was recovered and resuspended as above. After SDS-PAGE (Laemmli, 1970), the 21 kDa protein corresponding to the processed major recombinant form of CWLA was electroeluted (Extraphor, LKB), extensively dialysed against water and the resulting precipitate recovered. The preparation was resuspended in water and used for the production of anti-CWLA sera in a rabbit. For the initial injections, 100 µg of purified protein was used and 50 µg per boost.

**Table 1. B. subtilis 168 strains**

<table>
<thead>
<tr>
<th>Strain</th>
<th>Genotype</th>
<th>Source/Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>HR</td>
<td>trpC2</td>
<td>Laboratory stock</td>
</tr>
<tr>
<td>SF2</td>
<td>trpC2 cwLA</td>
<td>Laboratory stock</td>
</tr>
<tr>
<td>SF3</td>
<td>trpC2 cwLA::pSFP10 cwLA</td>
<td>This study</td>
</tr>
<tr>
<td>1A60</td>
<td>metC3 pyrA xin-l</td>
<td>BGSC (Thurm &amp; Garro, 1975)</td>
</tr>
<tr>
<td>RB1144</td>
<td>pyrD ileA1 thyA thyB Δ(metC)</td>
<td>P. Longchamp (Buxton, 1980)</td>
</tr>
<tr>
<td>L4587</td>
<td>metC ile xin-1</td>
<td>P. Longchamp (P. Jaunin, unpublished data)</td>
</tr>
<tr>
<td>1A372</td>
<td>SFβc2 novA1 recF15 trpC2 xin</td>
<td>BGSC</td>
</tr>
<tr>
<td>1A692</td>
<td>metB5 spf</td>
<td>BGSC</td>
</tr>
<tr>
<td>CU2059</td>
<td>[metB3] dal-1 (SFβc2) (SFβc2 int5 dissp44-1)</td>
<td>A. Moir (Lipsky et al., 1981)</td>
</tr>
</tbody>
</table>

* BGSC, Bacillus Genetic Stock Centre, Department of Biochemistry, Ohio State University, Columbus, OH, USA.
The mode of action of the enzyme was determined exactly as described previously (Foster, 1991). Purified 32 kDa lytic enzyme was dialysed against 100 vols of 10 mm-potassium phosphate buffer, pH 7.0. Purified walls were digested at 2 mg ml⁻¹ in 20 mm-MgSO₄, 10 mm-potassium phosphate buffer. pH 7.0 at 37 °C containing 960 U ml⁻¹ of purified enzyme. Periodically 0.5 ml samples were removed for OD₆₆₀ readings and determination of the appearance of new reducing and amino termini. The method of Margot et al. (1991) as described by Foster (1991) was used to determine to which isomer of alanine new amino termini could be attributed.

The N-terminal sequence of the purified 32 kDa amidase was determined using an Applied Biosystems 476A Protein sequencer.

Results

Production and use of anti-CWL A sera to identify CWLA-related proteins during growth and differentiation

The recombinant CWLA protein was purified from an E. coli strain carrying the plasmid pSF102 as described in Methods by selective precipitation and reactive red 120 dye affinity chromatography. The enzyme was eluted at about 1 M-LiCl from the reactive red column. The activity-containing eluate was analysed by 11% SDS-PAGE and the 21 kDa CWLA protein electroeluted. Antiserum was raised in rabbits to this recombinant protein and used to probe various cell extracts by Western blotting (Fig. 1a). The antiserum reacts with two proteins of 30 and 21 kDa in an extract of E. coli(pSF102) (Fig. 1a, track 2) which match the sizes of the active precursor and processed forms of CWLA in E. coli (Fig. 1b, track 2). The specificity of the sera is shown by the inability of an E. coli(pUBS) extract (plasmid without cwlA-containing insert), which has no autolytic activity against B. subtilis walls (Fig. 1b, track 1), to cross-react with the sera (Fig. 1a, track 1). CWLA is not related to any proteins in an SDS extract of vegetative or stage VI sporulating cells which show the characteristic autolysin profiles (Fig. 1a, b, tracks 3 & 4). Either CWLA or related proteins are not present in these samples or are at too low a level to be detected using the antisera.

Analysis of cwlA expression by the creation of a cwlA::lacZ fusion

Plasmid pSF10 contains a 936 bp HindIII, end-filled fragment from pSF100 ligated into pAZ106. Transformation of this plasmid into B. subtilis HR resulted in the creation of strain SF3 containing a cwlA::lacZ fusion. Analysis of the expression of the reporter gene β-galactosidase revealed no increase in expression as compared to B. subtilis HR background levels either during vegetative growth or differentiation (results not shown). The cwlA::lacZ transcriptional fusion was expressed in E. coli DH5α because colonies of a clone harbouring plasmid pSF10 were blue on LB ampicillin plates which had been spread with 80 μl of a 20 mg ml⁻¹ solution of X-gal in dimethylformamide.

Identification of mitomycin C-inducible lytic enzymes and their antigenic relationship to CWLA

Mitomycin C exposure is commonly used for the induction of prophage in B. subtilis 168. Treatment of an exponentially growing culture of strain HR (OD₆₀₀ = 0.2) resulted in the cessation of growth as measured by the increase in OD₆₀₀ (0.72) after about 1 h, followed by cellular lysis (Fig. 2a). A renaturing gel of SDS extracts of samples taken during mitomycin C treatment is shown in Fig. 2(b). After 60 min, a large increase appears in the amounts of the A3 and A4 lytic bands of molecular masses approximately 34 and 30 kDa, respectively (Fig. 2b, track 4). These further increase during the time of cellular lysis. Neither of these bands corresponds to the product of the cwlA gene, as strain SF2, which is inactivated in cwlA, still shows the same mitomycin C-induced lytic profile (Fig. 2b, track 8). Also, no expression of the cwlA gene occurs during mitomycin C treatment as measured by the cwlA::lacZ fusion created above. Interestingly, however, if these SDS extract samples were analysed by Western blotting using the anti-CWLA sera as the probe, a mitomycin C-induced CWLA-cross-reactive protein appeared concomitant with the increase in lytic activity (Fig. 2c). This protein
Fig. 2. Identification of mitomycin C-induced lytic enzymes and a CWLA-related protein. (a) The effect of mitomycin C (0.5 μg ml⁻¹) on growth (OD₆₀₀) of B. subtilis 168 HR was measured (■) as compared to an untreated control culture (□). The results are from a representative experiment from which the samples were taken for further analysis. Samples were taken at various times after addition of mitomycin C to both an HR and an SF2 (cwlA-inactivated) culture. SDS-extracts were prepared as described previously (Foster, 1992) and separated by 11% SDS-PAGE. All samples contained 10 μg protein. (b) Renaturing SDS-PAGE. (c) Western blot probed with anti-CWLA sera. Tracks: 1–7, SDS extracts of B. subtilis 168 HR taken at 0, 20, 40, 60, 80, 100 and 120 min, respectively, after the addition of mitomycin C; 8, SDS-extract of B. subtilis 168 SF2 120 min after the addition of mitomycin C. The position of molecular mass standards and the major lytic bands are indicated.

Identification of B. subtilis 168 prophage lytic enzymes

B. subtilis generally contains two mitomycin C-inducible prophages, PBSX and SPβ. To determine if the mitomycin C-induced lytic enzymes were prophage-encoded, the lytic profiles of various phage mutants were examined.

Strain CU2059 bears the SPβC2 mutation which confers SPβ temperature-sensitive induction. Fig. 3(a) shows the optical density of a culture grown at 37 °C until OD₆₀₀ = 0.33 and then heat-shocked at 50 °C for 6 min before continued incubation at 37 °C. About 20 min after heat shock the culture stopped growing and began to lyse. Concurrent with this lysis was the induction of a 43 kDa SPβ-specific lytic enzyme as shown by renaturing SDS-PAGE (Fig. 3b). This 43 kDa enzyme did not cross-react with the anti-CWLA sera as shown by Western blotting (data not shown). No lytic enzyme induction occurred upon heat shock of strain HR and mitomycin C-induction of a strain cured of SPβ (1A692) showed an identical pattern of lytic enzyme induction to strain HR.

Strain L4587 carries the xhi-1479 mutation which allows specific heat shock-induction of PBSX (P. Jaunin, University of Lausanne, Switzerland, unpublished data). Conversely RB1144 has had the PBSX prophage deleted (Buxton, 1980). The optical density of a mitomycin C-treated culture of RB1144 and a heat-shocked L4587 is shown in Fig. 4(a). The PBSX-deleted strain was no longer mitomycin C-sensitive and carried on growing (Fig. 4a) although at a reduced rate compared with a non-treated control (results not shown). The PBSX temperature-sensitive strain stops growing after about 60 min at 45 °C and then begins to lyse. Renaturing SDS-PAGE revealed the induction concomitant with lysis, of the two lytic enzymes A3 and A4 (Fig. 4b) the larger of which corresponded to the CWLA cross-reactive protein (Fig. 4c). Thus, both of these mitomycin C-induced lysins are PBSX-associated. Mitomycin C treatment of the PBSX-deleted strain (RB1144) showed no increase in the two lytic enzymes or induction of the CWLA cross-reactive protein (Fig. 4b, c). Strains 1A372 and 1A60 which have the xin mutation, such that PBSX cannot be induced, are mitomycin C-resistant and
no lytic enzyme or CWLA-related proteins are induced in response to mitomycin C (data not shown).

**Purification of a PBSX-encoded lytic enzyme**

PBSX has been shown to produce two lytic enzymes, a 14.5 kDa amidase (as measured by gel filtration) and a muramidase of unknown size (Ward *et al*., 1982). To determine if the PBSX lytic enzymes shown in this study correspond to the previously identified enzymes, purification of the proteins from a lysate of the temperature-sensitive strain L4587 was attempted. Only one lytic activity could be purified from the lysate (Fig. 5a; Table 2). The amount of lytic activity of this enzyme in the
Fig. 5. Purification of a PBSX-associated lytic enzyme. For details of the various samples, see Table 2 and Methods. (a) Coomassie blue-stained SDS-PAGE gel of samples taken during the purification procedure. All samples contained 10 μg protein. Track 1, molecular mass standards of sizes indicated; 2, ammonium sulphate precipitate; 3, Econo-Pac Q unbound material; 4, Sephadex G75 eluate; 5, Econo-Pac Q eluate (purified enzyme). (b) Renaturing SDS-PAGE. Track, 1, ammonium sulphate precipitate (5 μg protein); 2, Econo-Pac Q eluate (5 μg purified protein). (c) Western blot probed with anti-CWLA sera. All samples contained 5 μg protein. Track 1, SDS-extract of HR 120 min after mitomycin C treatment (see Fig. 2c, track 7); 2, ammonium sulphate precipitate; 3, Econo-Pac Q eluate (purified protein). The sizes of molecular mass standards are indicated.

### Table 2. Purification of the 32 kDa PBSX-associated lytic enzyme

<table>
<thead>
<tr>
<th>Purification step</th>
<th>Volume (ml)</th>
<th>Activity (U ml⁻¹)</th>
<th>Specific activity (U mg⁻¹)</th>
<th>Yield (%)</th>
<th>Purification (-fold)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. Cell lysate</td>
<td>460</td>
<td>70</td>
<td>1400</td>
<td>100</td>
<td>1</td>
</tr>
<tr>
<td>2. Ammonium sulphate precipitate</td>
<td>11</td>
<td>3250</td>
<td>1800</td>
<td>111</td>
<td>1.3</td>
</tr>
<tr>
<td>3. Econo-Pac Q wash-through</td>
<td>17</td>
<td>2530</td>
<td>2400</td>
<td>133</td>
<td>1.7</td>
</tr>
<tr>
<td>4. Sephadex G75 Superfine eluate</td>
<td>18</td>
<td>2440</td>
<td>11100</td>
<td>136</td>
<td>7.8</td>
</tr>
<tr>
<td>5. Econo-Pac Q eluate</td>
<td>14</td>
<td>1260</td>
<td>12000</td>
<td>52</td>
<td>8.5</td>
</tr>
</tbody>
</table>

Lysate could not be accounted for by vegetative cell activity. Table 2 shows the results from one representative purification of the lytic enzyme which was repeated three times. The increase in apparent yield seen after some steps during the purification procedure may be due to the removal of an endogenous inhibitor or the activation of a latent proform (Table 2). No fractions containing lytic enzyme activity were found at any point during the procedure apart from those associated with the enzyme which was purified. The enzyme was purified and found to have a molecular mass of 32 kDa as measured by Coomassie blue staining after SDS-PAGE, and 30 kDa by renaturing SDS-PAGE (Fig. 5a, b). SDS-PAGE and Western blotting revealed that protein redissolved after ammonium sulphate precipitation of the lysate contained different CWLA-related proteins than the SDS extract from the lysing cells (Fig. 5c, tracks 1 and 2). Instead of the 34 kDa band, the lysate contained a cross-reactive 18 kDa band. The 18 kDa antisera-cross-reactive band had no lytic activity as measured by renaturing SDS-PAGE (Fig. 5b, track 1). The purified 32 kDa enzyme also reacted weakly with the antisera (Fig. 5c, track 3).

### Determination of the hydrolytic bond specificity of the purified 32 kDa lytic enzyme

A suspension of purified B. subtilis 168 vegetative cell walls (5 mg ml⁻¹) was hydrolysed by the purified 32 kDa PBSX autolysin (results not shown). After 60 min, more than 75% of the initial OD₄₉₀ of the wall suspension had been lost. Concomitant with this decrease in OD₄₉₀ was a large increase in the number of free amino termini [430 nmol (mg cell wall)⁻¹]. After 60 min digestion with the purified enzyme there was no increase in the number of wall-reducing termini. Acid hydrolysis of the FDNB
labelled samples followed by TLC analysis identified only an increase in \(N\)-2,3-dinitrophenyl-L-alanine (DNP-alanine) on hydrolysis of the walls with purified lysis. The use of \(L\)-[\(\text{\textsuperscript{14}C}\)]alanine-labelled walls (Margot et al., 1991) identified this DNP-alanine increase to be solely accounted for by DNP-L-alanine and thus the enzyme is an \(N\)-acetylmuramyl-L-alanine amidase (amidase).

**N-terminal sequencing of the 32 kDa PBSX amidase**

The N-terminal sequence of the purified 32 kDa amidase protein was determined to elucidate its relationship to other known lytic enzymes. The 16 amino acid sequence MVNIIQDFIPVGANRP determined did not match CWLA or any other sequences in the database. This sequence does, however, match exactly that of the cloned and sequenced major amidase of PBSX (P. Longchamp, P. Jaunin & D. Karamata, unpublished data).

**Discussion**

The gene encoding the CWLA amidase has been cloned and inactivated but the function and expression pattern of this enzyme has remained elusive (Kuroda & Sekiguchi, 1990; Foster, 1991). The antisera raised to the purified CWLA protein in this study is specific in its reaction to the 30 and 21 kDa recombinant forms of CWLA and no other *E. coli* proteins. As far as can be ascertained, *cwlA* is never expressed in *B. subtilis* even under mitomycin C treatment in which a CWLA-related protein appears. This protein of 34 kDa is larger than CWLA (30 kDa) and is present in a strain inactivated in the *cwlA* gene. To gain a further insight into the identity of this CWLA-related protein, the resident prophage-associated lytic enzymes of *B. subtilis* 168 were characterized.

Three prophage-encoded lytic enzymes of *B. subtilis* 168 were identified during this work. The prophage SP\(\beta\) encodes a 43 kDa enzyme of unknown hydrolytic bond specificity. This enzyme is not present during vegetative growth, differentiation or mitomycin C prophage induction in measurable amounts as judged by renaturing SDS-PAGE. This is not surprising as although SP\(\beta\) is spontaneously induced at a low level during vegetative growth and this induction is greatly increased upon mitomycin C treatment, the titre is still far less than one infectious particle per bacterium (Warner et al., 1977). Mitomycin C treatment induces two lytic enzymes of apparent molecular masses 34 and 30 kDa as judged by renaturing SDS-PAGE. Both of these enzymes were found to be PBSX-associated. Only one enzyme could be purified from a lysate of a PBSX temperature-sensitive strain. This was the smaller of the two PBSX enzymes in SDS extracts and the purified protein has a molecular mass of 32 kDa as measured by SDS-PAGE and Coomassie blue staining but 30 kDa by renaturing SDS-PAGE. A PBSX amidase was partially purified by Ward et al. (1982) and found to have a molecular mass of 14.5 kDa as measured by gel filtration. SDS-PAGE, however, showed the preparation to contain two proteins of 30 and 14.5 kDa (Ward et al., 1982). The amidase purified in this study may correspond to the 30 kDa component of the previously partially purified amidase. The major CWLA cross-reactive lytic enzyme of 34 kDa may be the unpurified PBSX muramidase of Ward et al. (1982).

Interestingly, although it is not apparent in SDS extracts of lysing cells, the 32 kDa PBSX amidase does cross-react to a low level with the CWLA antisera as can be seen using the purified protein. In fact the 32 kDa PBSX amidase and CWLA show 50% amino acid sequence identity (P. Longchamp, P. Jaunin & D. Karamata, unpublished data). The 34 kDa major CWLA cross-reactive band, which is probably the mitomycin C-inducible 34 kDa lytic enzyme, was not recovered from the PBSX-containing lysate after temperature induction of strain L4587. Instead of the 34 kDa cross-reactive band there was an 18 kDa protein which reacted with the antisera. Thus it seems likely that the 34 kDa protein is quite unstable and subject to proteolysis, even after PMSF treatment, and was lost during purification. The PBSX late operon encodes two proteins of 31.5 and 32.5 kDa which are comparable in size to the lytic enzymes identified in this study (Mauel & Karamata, 1984; Wood et al., 1990).

From the results it can be seen that CWLA is related immunologically to two lytic enzymes of the defective prophage PBSX. PBSX produces defective non-infectious particles as it is unable to package its own DNA. PBSX-like defective prophages are found in many *Bacillus* and cured strains have not been isolated. Thus it seems that the maintenance of these phages is in some way advantageous for the cell (Buxton, 1980). It is possible that CWLA is also phage-encoded on another redundant copy of PBSX or a closely related phage. The map locations are such that *cwlA* cannot be encoded for by PBSX itself and may therefore be on a cryptic prophage which can no longer be induced even by the action of mitomycin C and thus *cwlA* is never expressed. The *cwlA* gene seems to be expressed from its own promoter in *E. coli* and so is in itself functional (Foster, 1991). A regulatory mutation analogous to the *xin* mutation of PBSX (Thurm & Garro, 1975) may be responsible for the lack of expression of *cwlA*. The four open reading frames flanking the *cwlA* gene show no significant homology to any protein sequences in the databases (Foster, 1991); however, further sequencing of the PBSX
genome may well identify a homologous region to that containing \textit{cwLA}.

A further lysozyme structural gene has been isolated by the direct screening method used to clone \textit{cwLA} and the recombinant protein shows low level cross-reactivity with the anti-CWLA sera (Foster, 1991; Greene & Foster, unpublished data). The map location and specificity of this enzyme are at present unknown. This new lytic enzyme may also be of prophage origin.

To understand the individual and cooperative roles of the lytic enzymes of \textit{B. subtilis} one must first identify the complement of enzymes present. PBSX produces two lytic enzymes of between 30 and 34 kDa. By the use of renaturing SDS-PAGE it seems there is at least one other enzyme present in this size range in vegetative cells of \textit{B. subtilis} 168 (Foster, 1992). The creation of isogenic multiple and single lytic enzyme inactivated mutants to try to understand the function of lytic enzymes must take into account the presence of spontaneously induced prophage enzymes.

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References


