BACTERIAL PATHOGENICITY

Purification and properties of a novel glycosaminoglycan depolymerase from Streptococcus intermedius strain UNS 35

H. SHAIN, K. A. HOMER and D. BEIGHTON

Joint Microbiology Research Unit, King’s College School of Medicine and Dentistry, Caldecot Road, Denmark Hill, London SE5 9RW

A glycosaminoglycan (GAG) depolymerase that acts on chondroitin sulphate A (CS-A), chondroitin sulphate C (CS-C) and hyaluronic acid (HA) was purified to apparent homogeneity from a culture of Streptococcus intermedius, strain UNS 35, grown in minimal medium supplemented with CS-A as the sole carbon source. The enzyme was purified by ammonium sulphate precipitation followed by serial chromatography on DEAE Trisacryl M, CM Trisacryl M and heparin-agarose. SDS-PAGE analysis of the purified enzyme yielded a single band with a mol.wt of c. 83 000. The purified GAG depolymerase was unusual in its substrate specificity. The enzyme was initially regarded as a CS depolymerase because of its induction by CS-A. However, the GAG depolymerase exhibited greatest activity against HA, whereas the degradation rates of CS-A and CS-C were c. 8% and 2%, respectively, of the rate with HA. On this basis the enzyme could be classified as a hyaluronidase rather than a CS depolymerase. The pH optimum was around neutrality and the enzyme was unusual in having a high pl of approximately 9.3.

Introduction

The isolation of ‘Streptococcus milleri’ from human clinical specimens has become almost synonymous with the diagnosis of deep-seated sepsis [1-4] and other workers have suggested that ‘S. milleri’ bacteremia in a febrile patient should prompt a search for an internal abscess [5,6]. These bacteria are often not recognised as pathogens [7,8] primarily because of confusion over nomenclature and difficulties in routine laboratory identification. Recently, Whiley and Beighton stated that members of the ‘S. milleri’ group represented three distinct species, namely S. intermedius, S. constellatus and S. anginosus, and demonstrated simple laboratory tests for their differentiation [9].

The propensity of these organisms for causing serious pyogenic infections is well documented, but the factors mediating their pathogenicity and ability to proliferate in vivo are still unclear. One possible mechanism may be the production of extracellular enzymes that degrade host tissue components [7,8,10]. These enzymes may contribute not only to the pathogenic potential of these organisms but also fulfil an important nutritional role. Of the ‘S. milleri’ group, S. intermedius is particularly associated with brain and liver abscesses [11,12] and produces a greater range of glycoprotein- and glycosaminoglycan- (GAG) degrading enzymes, including hyaluronidase and chondroitin sulphate (CS) depolymerase, than any other species of viridans streptococci [13]. GAGs are linear polysaccharides composed of repeating disaccharide units and include chondroitin sulphate A (CS-A), chondroitin sulphate C (CS-C), dermatan sulphate, keratan sulphate, heparan sulphate, heparin and hyaluronate, important components of the extracellular matrix responsible for the structural integrity of host tissue [14].

We have demonstrated recently that the growth of S. intermedius strain UNS 35, in minimal medium supplemented with CS-A as the sole carbohydrate source, results in the production of extracellular CS depolymerase and sulphatase activities [39]. The CS depolymerase enzyme produced by S. intermedius has not been purified previously to homogeneity nor have its biochemical properties been reported. This paper describes the purification and characterisation of the CS depolymerase activity from an S. intermedius strain.
Materials and methods

Bacterial strain and culture

*S. intermedius* strain UNS 35, first isolated by Professor P. F. Unsworth from a human brain abscess, was obtained from the Central Public Health Laboratory, 61 Colindale Avenue, London, and was stored frozen on glass beads at −70°C. Bacteria were maintained by routine subculture on Fastidious Anaerobe Agar (Lab M Ltd, Lancs) containing defibrinated horse blood 5% v/v (FAA) in an anaerobic cabinet (Don Whitley, Shipley, West Yorkshire) for 48 h at 37°C.

Assay of reducing terminal N-acetylhexosamine for measurement of CS depolymerase activity

CS depolymerase activity at different stages of the enzyme purification was determined by measuring the increase in N-acetylhexosamine concentration after the method of Levvy and McAllan [15], with either CS-A (Sigma) or CS-C (Sigma) as substrate. A 20-μl sample of enzyme preparation was added to 62.5 μl of substrate (20 mg/ml) and 42.5 μl of 0.2 m sodium phosphate buffer, pH 7.5, and incubated for 18 h at 37°C. Twenty μl of 0.2 m potassium borate were added to 62.5 μl of the reaction mixture and the assay was heated at 100°C for 3 min. After cooling to ambient temperature, 375 μl of N-acetylhexosamine reagent [16] were added to each sample and these were incubated for 20 min at 37°C. Portions (200 μl) of each sample were dispensed into 96-well microtitration trays and the absorbance at 540 nm was recorded with a plate-reading spectrophotometer (Titertek, Multiscan MCC340; ICN-Flow Laboratories Ltd, Herts). The increase in reducing terminal N-acetylhexosamine was estimated by comparison with N-acetylgalactosamine standards (0–10 mM) treated in the same manner.

Commercial CS-A contained CS-A (sulphated at the 4-position of the N-acetylgalactosamine residue) 70% and CS-C (sulphated at the 6-position of the N-acetylgalactosamine residue) 30%; CS-C contained CS-A 10% and CS-C 90%. The choice of substrate (CS-A or CS-C) and buffer (0.2 m Tris-HCl, pH 7.5, or 0.2 m sodium phosphate, pH 7.5) used in the assay for enzyme activity depended on the stage of enzyme purification.

Assay of 4,5-unsaturated disaccharide formation for measurement of CS depolymerase activity

For the determination of pH optima, K_m, V_max and substrate specificity, enzyme activity was assayed by a modification of the method of Saito et al. [17]. Enzyme preparation (20 μl) was added to 50 μl of the appropriate buffer and 70 μl of substrate solution and incubated at 37°C. The reactions were stopped after an appropriate incubation period by heating at 100°C for 5 min; 40 μl of the reaction mixture were added to 10 μl of enriched Tris buffer, pH 8.0, and incubated at 37°C for 30 min. Then 950 μl of 0.1 m HCl were added to each sample and, after centrifugation (13 000 rpm, 10 min), the absorbance at 232 nm was measured (Shimadzu UV 160-A recording spectrophotometer) with quartz cuvettes. Control assays contained heat-inactivated (100°C for 10 min) enzyme preparation. The concentration of unsaturated disaccharide formed in each assay was calculated from the increase in absorbance in conjunction with the use of 5.7, 5.1 and 5.5 as millimolar absorbance coefficients for ΔUA GalNAc-OS (2-acetamido-2-deoxy-3-O-(β-D-gluco-4-Δenepyranosyluronic acid)-D-galactose), ΔUA GalNAc-4S (2-acetamido-2-deoxy-3-O-(β-D-gluco-4-Δenepyranosyluronic acid)-4-O-sulpho-D-galactose) and ΔUA GalNAc-6S (2-acetamido-2-deoxy-3-O-(β-D-gluco-4-Δenepyranosyluronic acid)-6-O-sulpho-D-galactose), respectively [18].

Growth of bacteria for CS depolymerase production

Preparations of CS depolymerase for purification and characterisation were obtained from 25 L (5 × 5-L volumes) of a semi-defined medium (CasMM) [16, 19] supplemented with CS-A 5 mg/ml as the sole source of carbohydrate. Each 5 L of CS-A-supplemented CasMM was inoculated with 250 ml of a late exponential phase culture of *S. intermedius* UNS 35, grown in Brain Heart Infusion Broth (BHI; Oxoid) and incubated anaerobically at 37°C with constant stirring for 18 h. Cells were harvested by centrifugation (10 000 rpm, 20 min, 4°C) and the culture supernates were decanted for purification of the CS depolymerase.

Concentration of culture supernate and ammonium sulphate precipitation

The culture supernate (25 L) was concentrated 16-fold to c. 1.5 L by ultrafiltration (Ultrasette ultrafiltration unit, 10 kDa cut-off; Flowgen, Sittingbourne, Kent). The proteins were precipitated by the gradual addition, with gentle stirring, of ammonium sulphate to 80% saturation at 4°C. After 18 h the precipitate was pelleted by centrifugation (15 000 rpm, 1 h 4°C), dissolved in a minimal volume of ice-cold 50 mM Tris-HCl buffer, pH 7.5 (Tris-buffer) and dialysed against three changes of the same buffer at 4°C. The dialysate was divided into 10-mL volumes and stored at −20°C.

Chromatography with diethylaminoethyl (DEAE) trisacryl M

A column (1.5 cm × 24 cm) of DEAE Trisacryl M (Sepracor SA, Villeneuve la Garenne Cedex, France) was prepared and equilibrated with 80 ml of Tris-buffer. The dialysed ammonium sulphate fraction (10 mL volumes) was applied to the column at the rate of 40 ml/h. After washing the column with 80 ml of Tris-buffer, a step gradient (80 ml each of 0.1 M, 0.2 M,
GLYCOSAMINOGLYCAN DEPOLYMERASE FROM S. INTERMEDIUS

0.3 M and 0.4 M NaCl in Tris-buffer) was used to elute the proteins from the column. Fractions (8 ml) were collected and enzyme activity was monitored by the N-acetylhexosamine assay with CS-A as substrate. The column was regenerated by washing with 40 ml of 1 M NaCl in Tris-buffer, followed by 40 ml of 0.5 M Tris-HCl, pH 7.5, and then re-equilibrated with 80 ml of Tris-buffer, before the application of further samples.

Enzyme activity was detected in the effluent from the wash buffer and each of the step gradient buffers used for elution. The fractions with the greatest activity (15-21) eluted at the 0.1 M NaCl step (Fig. 1) were pooled, desalted (in Tris-buffer) and concentrated with a stirred cell concentrator (10 000 Da cut-off, Flowgen). The fractions containing minor levels of depolymerase activity were not investigated further.

Chromatography with carboxymethyl (CM) trisacryl M

A CM trisacryl M (Sepracor) column (1.5 cm × 24 cm) was prepared and equilibrated with 80 ml of Tris-buffer. The partially purified and concentrated enzyme fraction (11 ml) from the DEAE trisacryl M column was applied to the CM trisacryl M column, washed with 80 ml of the Tris-buffer and eluted with a linear gradient of 0-0.5 M NaCl in Tris-buffer (total volume 200 ml) at a flow rate of 40 ml/h. The fractions (8 ml) were monitored for enzyme activity by the N-acetylhexosamine assay with CS-A or CS-C as substrate. Active fractions (19-28) were pooled, desalted with exchange of buffer to 50 mM potassium phosphate, pH 7.5, and then concentrated as described previously to c. 750 μl (Fig. 2).

Affinity chromatography with heparin-agarose

Heparin-agarose affinity chromatography was carried out with modifications of a previously described method [20]. The enzyme preparation from the CM trisacryl M column was applied to a heparin-agarose column (Type 1, Sigma; 1.5 cm × 8.5 cm), equilibrated with 5 column-volumes of 50 mM potassium phosphate buffer, pH 7.5. Preliminary experiments indicated that the depolymerase bound only weakly to the heparin-agarose column, so to avoid co-elution of the enzyme with non-adsorbed material, the sample was divided and applied to the column in two separate batches. The samples were applied to the column at a flow rate of 17 ml/h and, after washing with 30 ml of equilibration buffer, the protein was eluted with a gradient of 0-0.5 M NaCl in 50 mM potassium phosphate, pH 7.5, in 5 column-volumes. The flow rate was 30 ml/h and 4-ml fractions were collected.

The column was regenerated between sample applications by washing with up to 20 column-volumes of 0.1 M Tris-HCl buffer containing 0.5 M NaCl, pH 7.5, followed by an equal volume of 0.1 M sodium acetate containing 0.5 M NaCl, pH 5.0, and finally re-equilibrated by washing with 5 column volumes of 50 mM potassium phosphate, pH 7.5. The N-acetylhexosamine assay was used to monitor enzyme activity in the fractions, with CS-C as the substrate. The active fractions (15-19) were pooled, divided in 5-ml volumes and stored at −20°C before further characterisation (Fig. 3).
**Fig. 2.** Separation of GAG depolymerase on a CM trisacryl M column (1.5 × 24 cm). Fractions of the second peak (shown in Fig. 1) from DEAE trisacryl M column were applied to CM trisacryl M. Proteins were eluted with 50 mM Tris-HCl, pH 7.5 (fractions 1–9) followed by application of a continuous salt gradient indicated by the arrow (0–0.5 M NaCl in 50 mM Tris-HCl, pH 7.5). Fractions (8 ml) were collected and the flow rate was 40 ml/h. N-acetylhexosamine assay with CS-A and CS-C as substrates was used to determine enzyme activity and protein was determined by the Pierce protein assay: ⊙, protein; ■, enzyme activity.

**Fig. 3.** Separation of GAG depolymerase on a heparin-agarose column (1.5 × 8.5 cm). Fractions 1–7 were eluted with 50 mM potassium phosphate, pH 7.5, and later fractions were eluted with a continuous salt gradient indicated by the arrow (0–0.5 M NaCl in 50 mM potassium phosphate, pH 7.5). Fractions (4 ml) were collected and the flow rate was 30 ml/h. CS-C was used as substrate in the N-acetylhexosamine assay and protein was measured with the Pierce protein assay ⊙, protein; ■, enzyme activity.

**Protein estimation**

The protein concentration of the enzyme preparations at various stages of purification was determined routinely by a Coomassie Blue dye-binding assay (Sigma) in comparison with a standard curve of bovine serum albumin (0–500 μg/ml). The protein content of column fractions was monitored by a more sensitive Coomassie Blue dye-binding assay (Pierce) with bovine serum albumin (0–20 μg/ml) as standard.

**Mol. wt determination of purified CS depolymerase**

Before mol. wt determination by SDS-PAGE, a 3-ml sample of the enzyme preparation from the heparin-agarose column was desalted and concentrated (Microsep Microconcentrator, 10 kDa cut-off; Filtron Technology Corporation, MA, USA). To 5 μl of the concentrated sample (protein concentration 82.3 μg/ml), 10 μl of sample buffer (10 mM Tris-HCl, pH 8.0,
1 mM EDTA, SDS 2.5%, β-mercaptoethanol 5%) was added and heated at 100°C for 5 min. SDS-PAGE was performed with pre-formed homogeneous polyacrylamide 12.5% gels on the Phast system (Pharmacia Biotech, Uppsala, Sweden) with 1-μl samples applied to the gels. A set of low mol. wt protein markers (Pharmacia) was used for mol. wt determination.

To separate the proteins, the Phast system was programmed with an initial low output (100 V, 1.0 mA, 1.0 W, 4 Vh) to draw samples through the stacking gel, followed by an increased output phase (250 V, 3.0 mA, 3.0 W, 66 Vh) as described by Slayne et al. [21]. On completion of the electrophoretic separation the gel was transferred to the development chamber and stained by the silver staining protocol (Pharmacia).

The protein bands were visualised by high performance CCD video camera (COHU) and the image was captured with Bioscan Snapshot soft-ware (Datacell). The relative positions of the protein standards and the purified CS depolymerase were determined and the loglo mol.wt of the standard proteins were plotted against migration distance to calculate the mol. wt of the purified protein.

Determination of the isoelectric point (pI)

The pI of the CS depolymerase was determined using the Phast System (Pharmacia). A pH gradient was generated on a Phast Gel IEF 3-9 (Pharmacia) during a pre-focusing step (2000 V, 2.5 mA, 3.5 W, 75 Vh). Following this, the enzyme (1 μl each at both the anode and cathode positions) and 1 μl of pI calibration kit proteins (broad range pI kit, Pharmacia) were applied. The voltage was reduced to 200 V during the application step. After the focusing step (2000 V, 2.5 mA, 3.5 W, 410 Vh), the proteins were visualised by silver staining as described by the manufacturer (Pharmacia). The pI of the depolymerase was determined by comparison with the relative positions of the calibration proteins, in the pI range 3–10.

Determination of the pH optimum of the CS depolymerase

The enzyme activity at various pH values was measured under the standard assay conditions by the A$_{232}$ method with 0.2 M MES (2-[N-morpholino]ethane sulphonic acid) buffer (pH 5.5, 6.0, 6.5), 0.2 M sodium phosphate buffer (pH 6.5, 7.0, 7.5) and 0.2 M Tris-HCl buffer (pH 7.5, 8.0, 8.5, 8.9). The substrates used in the assays were CS-A and CS-C 10 mg/ml and hyaluronic acid (HA; Sigma; 5 mg/ml).

Determination of $K_m$ and $V_{max}$ values

The $K_m$ and $V_{max}$ values were determined for the purified protein with CS-A, CS-C and HA as substrates from double reciprocal (Lineweaver-Burk) plots. The initial reaction velocities were measured at different substrate concentrations with a fixed enzyme concentration. A 1 in 50 dilution of the enzyme preparation was used for determining the $K_m$ and $V_{max}$ values for HA. The rates of product formation were measured by the A$_{232}$ assay and all assays were performed at the pH optimum of the enzyme.

Substrate specificity of the CS depolymerase

The substrates CS-A, CS-C, HA, heparin, heparan sulphate (Sigma), dermatan sulphate and keratan sulphate (Fluka Chemicals Ltd, Gillingham, Dorset) (10 mg/ml) were used to determine the substrate specificity of the purified enzyme. The assays were set up with 70 μl of substrate, 20 μl of the enzyme preparation and 50 μl of 0.2 M sodium phosphate buffer at pH 7.0. The reaction mixtures were incubated at 37°C for 2 h and the rate of substrate degradation was determined by monitoring the increase in A$_{232}$.

Results

Enzyme purification

The CS depolymerase of S. intermedius strain UNS 35 was purified to homogeneity by a combination of ammonium sulphate precipitation, ion exchange and affinity chromatography. Table 1 summarises the total activity, specific activity and recovery of the enzyme at each step of purification. An overall purification of 6.4-fold and a yield of 3.2% were obtained.

Earlier studies have indicated the inherent difficulties encountered in purifying bacterial CS depolymerases to homogeneity. Preliminary studies with crude

<table>
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<th>Stage of purification</th>
<th>Protein (mg)</th>
<th>Activity (μmol/min)</th>
<th>Yield (%)</th>
<th>Specific activity (μmol/min/mg)</th>
<th>Purification (fold)</th>
</tr>
</thead>
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<tr>
<td>Concentrated supernate</td>
<td>24.9</td>
<td>96.6</td>
<td>100</td>
<td>3.8</td>
<td>1.0</td>
</tr>
<tr>
<td>(NH$_4$SO$_4$ precipitate)</td>
<td>13.7</td>
<td>54.7</td>
<td>56.0</td>
<td>4.0</td>
<td>1.0</td>
</tr>
<tr>
<td>DEAE column</td>
<td>1.528</td>
<td>24.4</td>
<td>25.3</td>
<td>16.0</td>
<td>4.2</td>
</tr>
<tr>
<td>CM column</td>
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<td>13.4</td>
<td>13.9</td>
<td>19.4</td>
<td>5.1</td>
</tr>
<tr>
<td>Heparin column</td>
<td>0.125</td>
<td>3.1</td>
<td>3.2</td>
<td>24.5</td>
<td>6.4</td>
</tr>
</tbody>
</table>

Table 1. Purification of GAG depolymerase from S. intermedius
enzyme preparations obtained from the culture supernate of *S. intermedius* grown in minimal medium supplemented with CS-A indicated the presence of a sulphatase activity (with AUA GalNAc-4S rather than AUA GalNAc-6S as its preferred substrate) in addition to a CS-AC depolymerase (chondroitinase AC) activity [39]. When these crude preparations were incubated with commercial CS-A (CS-A 70%, CS-C 30%) there was a progressive increase in the concentrations of AUA GalNAc-6S and AUA GalNAc-OS, whereas that of AUA GalNAc-4S remained low during the incubation period due to the rapid desulphation of the latter disaccharide. The chemical assay for reducing N-acetylhexosamine gives a positive reaction with AUA GalNAc-6S (derived from CS-C) and AUA GalNAc-OS but not with AUA GalNAc-4S (derived from CS-A) [18]. Therefore, when this assay is used to determine enzyme activity during the purification process with CS-A as the substrate, a strong positive reaction is indicative of the combined action of the depolymerase and the sulphatase. This assay was used to demonstrate indirectly the presence of the sulphatase and its subsequent removal during the protein purification process.

The enzyme of interest was prepared from 25 L of strain UNS 35 culture supernate; 56% of enzymically-active protein was precipitated from the concentrated culture filtrate by ammonium sulphate 80% precipitation. After dialysis, the enzyme preparation was fractionated by chromatography on a DEAE trisacryl M column by step elution with increasing concentrations of NaCl. A typical elution pattern is shown in Fig. 1. Fractions from the second peak, which eluted at 0.1 M NaCl, showed the highest enzymatic activity. The strong positive results from the N-acetylhexosamine assay described previously indicated that the enzyme preparation at this point still contained the sulphatase activity. These fractions were collected and subjected to further purification on a CM trisacryl M column. The enzymically active fractions eluted from the column as a single distinct protein peak (Fig. 2). Tests for enzyme activity with CS-A as substrate resulted in weak positive reactions with the N-acetylhexosamine assay. When CS-C was substituted for CS-A, a three-fold increase in absorbance values at A540 was demonstrated, indicating that the enzyme preparation was free of contaminating sulphatase activity. However, SDS-PAGE analysis of the enzyme preparation at this stage demonstrated several contaminating protein bands (data not shown).

After ion-exchange chromatography the enzyme preparation was further purified with a heparin-agarose column. Fig. 3 shows the enzymically active fractions eluting as a single broad peak which corresponded with a distinct protein peak. This preparation was subsequently shown to be homogeneous by SDS-PAGE.

**Determination of protein purity, mol. wt and pl**

SDS-PAGE analysis of the purified enzyme yielded a single band demonstrated by silver staining. The mol. wt of the enzyme was c. 83080, as determined by comparison with SDS-PAGE protein mol. wt markers (Fig. 4). The purified protein was subjected to isoelectric focusing with proteins over the pl range 3.5-9.3 as standards. The purified CS depolymerase co-migrated with trypsinogen, indicating a pl of c. 9.3 (data not shown).

**pH optima of the CS depolymerase**

The purified enzyme was active against both CS-A and CS-C over a wide pH range (6.0–8.0). Maximum activity against both substrates occurred at pH 7.0, but

![Fig. 4. SDS-PAGE (12.5%) of the purified enzyme for *S. intermedius* (lane A) compared with the following marker proteins of known mol. wt (B): phosphorylase b (94 kDa); bovine serum albumin (67 kDa); ovalbumin (43 kDa); carbonic anhydrase (30 kDa); soybean trypsin inhibitor (20.1 kDa); lactalbumin (14.4 kDa). The gel was silver stained according to Pharmacia Phastgel silver kit protocol.](image-url)
with equal substrate concentrations the activity against CS-C was half that obtained with CS-A. The enzyme was highly active against HA over a pH range 6.5–8.0, with optimum activity at pH 7.5 (Fig. 5).

**Km and Vmax values**

A study of the activity of the CS depolymerase against CS-A, CS-C and HA indicated that substrate degradation followed typical Michaelis-Menten kinetics. Lineweaver-Burk analysis of these data indicated that the Km values for CS-A, CS-C and HA were 26.6, 12.7 and 1.1 mg/ml respectively. The estimated Vmax for CS-A, CS-C and HA were 1.6, 0.3 and 19.5 μM/min/mg of protein, respectively. The specificity constants (Vmax/Km) for the three substrates were 0.06 (CS-A), 0.03 (CS-C) and 17.0 (HA).

**Fig. 5.** pH optima for the degradation of various GAGs by the purified enzyme. The enzyme activity was measured in the standard reaction mixture with 0.2 M MES (pH 5.5, 6.0 and 6.5; ○); 0.2 M sodium phosphate buffer (pH 6.5, 7.0, 7.5 and 8.0 for hyaluronic acid; □); and 0.2 M Tris-HCl (pH 7.5, 8.0, 8.5 and 8.9; △): ······, CS-A; --- ---, CS-C; ———, HA.

**Table 2.** Comparative activities of the purified enzyme against glycosaminoglycans

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Relative activity (%)</th>
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<tbody>
<tr>
<td>Hyaluronic acid</td>
<td>100</td>
</tr>
<tr>
<td>CS-A</td>
<td>7.7</td>
</tr>
<tr>
<td>CS-C</td>
<td>5.6</td>
</tr>
<tr>
<td>Dermatan sulphate</td>
<td>1.4</td>
</tr>
<tr>
<td>Keratan sulphate</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Heparan sulphate</td>
<td>&lt;0.01</td>
</tr>
<tr>
<td>Heparin</td>
<td>&lt;0.01</td>
</tr>
</tbody>
</table>

**Substrate specificity**

The enzyme was most active against HA and exhibited less activity towards CS-A, CS-C and dermatan sulphate (Table 2). Activity against keratan sulphate, heparan sulphate and heparin could not be detected.

**Discussion**

Bacterial GAG depolymerases differ from the mammalian equivalents in that they cleave GAGs by an elimination reaction leading to the production of 4,5-unsaturated disaccharides [18, 22, 23]. The chondroitinase ABC produced by Proteus vulgaris [18] and Bacteroides thetaiotaomicron [20] degrades primarily CS-A, CS-B (dermatan) and CS-C, with little activity against HA (10% of chondroitinase activity). Flavobacterium heparinum produces three distinct enzymes, each with CS depolymerising activity: chondroitinase AC which is equally active against CS-A and CS-C but also has some activity against HA (20% of the chondroitinase activity) [18]; chondroitinase C which shows equal activity against CS-C and HA [24]; and chondroitinase B which acts specifically on CS-B [25]. Except for the chondroitinase AC of F. heparinum which is constitutive [18], the other reported chondroitinases are induced by growing the organisms in the presence of the appropriate CS. A chondroitinase C from S. intermedium has been partially purified but its activity on HA was not reported [26].

The GAG depolymerases of S. intermedium UNS 35 exhibited an unusual substrate specificity. It depolymerised HA rapidly and acted on CS-A and CS-C at about 8% and 2%, respectively, of the rate with HA. Although the Km for the CS-A was twice that observed for CS-C, comparison of Vmax and specificity constants (Vmax/Km) confirmed that, of the two sulphated GAGs, CS-A is the preferred substrate (the Vmax value for CS-A was approximately five-fold greater than that for CS-C). However, the specificity constant for HA is c. 280-fold greater than that for CS-A and on the basis of this unusual substrate specificity the depolymerase activity would best be classified as a hyaluronidase rather than a CS depolymerase. Of the bacterial chondroitinases previously purified, the chondroitinase C of F. heparinum has similar properties to the depolymerase reported here in that it is induced by CS but has slightly greater activity against HA than CS-C, based on the relative amount of disaccharides produced. It was suggested that this enzyme chondroitinase C, could be classified as a hyaluronidase type II C following the nomenclature proposed by Gibian [27].

GAG depolymerases that act mainly on HA have been purified from streptococci [28-30], pneumococci [31], Staphylococcus aureus, [32], clostridia [33], Escherichia freundii [34] and peptostreptococci [35]. The bacterial hyaluronidases have mol. wts of 55 000–
Multiple molecular forms of GAG depolymerases have been described for Staph. aureus [32], B. thetaiotaomicron [20] and F. heparinum [18, 24, 25]. S. intermedius UNS 35 grown in medium supplemented with CS-A has been shown to produce both hyaluronidase and CS depolymerase activities [36, 37] but whether the GAG depolymerase reported here is responsible for both activities or whether they are distinct enzymes has not been ascertained.

In mammalian tissues, CS-C is less susceptible to hyaluronidase degradation than CS-A; this is a poorer substrate for hyaluronidase than HA. Despite these observations, degrading activities of CS in mammalian tissues have been attributed to the combined actions of hyaluronidase and exo-enzymes including glycosidases and O-sulphatases [38]. Bacterial chondrosulphatases differ from mammalian O-sulphatases by their activity on sulphated 4,5-ununsaturated disaccharides and do not act on longer oligosaccharides [18]. S. intermedius UNS 35 produces a sulphatase with AUA GalNAc-4S (derived from CS-A) as its preferred substrate [39], forming the desulphated disaccharide, AUA GalNAc-OS; this is subsequently catabolised by S. intermedius grown in batch culture. Although the GAG depolymerase produced by S. intermedius has greater hyaluronidase activity than CS depolymerase activity, on the basis of the mammalian model of CS degradation, it seems reasonable to propose that it is its ability to produce a sulphatase activity that enables S. intermedius to degrade CS-A and utilise the resulting desulphated disaccharide (ΔUA GalNAc-OS) as a source of nutrient. This would explain our in vitro observations that the organism can grow in minimal medium supplemented with CS-A as the sole carbohydrate source.


Reference added in proof: