Degradation and utilisation of chondroitin sulphate by Streptococcus intermedius

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Streptococcus intermedius, part of the ‘Streptococcus milleri group’, has the ability to produce glycosaminoglycan depolymerising enzymes (hyaluronidase and chondroitin sulphate depolymerase) which is unique amongst the viridans streptococci and may contribute to their virulence in brain and liver abscesses. The growth of S. intermedius strain UNS 35 was studied in basal medium supplemented with chondroitin sulphate A (CS-A, sulphated at position 4 of the N-acetylgalactosamine moiety) or chondroitin sulphate C (CS-C, sulphated at position 6 of the N-acetylgalactosamine moiety) as the major carbohydrate source. CS-A but not CS-C supported the growth of S. intermedius. Extracellular degradation of CS-A resulted in the initial accumulation of 2-acetamido-2-deoxy-3-O-(β-D-glucopyranosyluronic acid)-D-galactose (AUA GalNAc-OS), and low levels of 2-acetamido-2-deoxy-3-O-(β-D-glucopyranosyluronic acid)-4-O-sulpho-D-galactose (AUA GalNAc-4S) in the medium with GalNAc-OS being subsequently utilised during bacterial growth. Metabolic end-products included formate and ethanol but not lactate, indicating that growth was probably carbon-limited. The CS-A contained 30% CS-C, which was also depolymerised resulting in the formation of 2-acetamido-2-deoxy-3-O-(β-D-glucopyranosyluronic acid)-6-O-sulpho-D-galactose (AUA GalNAc-6S) in the culture supernate, but this unsaturated disaccharide was apparently not utilised during growth. The results indicate that S. intermedius produced CS-AC depolymerase, which was inducible and extracellular, and sulphatase activity. Experiments with authentic AUA GalNAc-4S and AUA GalNAc-6S demonstrated that AUA GalNAc-4S rather than AUA GalNAc-6S was the preferred substrate for the sulphatase. Therefore, it is suggested that the CS-AC depolymerase of S. intermedius may play a role in the destruction of CS in host tissues, facilitating bacterial spread, and also in bacterial nutrition by the liberation of nutrients at the site of infection.

Introduction

The ‘Streptococcus milleri group’ which includes S. intermedius, S. constellatus and S. anginosus [1] forms part of the normal flora of the mouth, gastrointestinal tract and genitourinary tract and is often associated with purulent infections [2-4]. The tendency of members of the ‘S. milleri group’ to be associated with abscess formation in a variety of body sites, although well documented, remains largely unexplained. Interest has focused on the production of extracellular enzymes with the ability to damage host tissues, including hyaluronidase and chondroitin sulphate (CS) depolymerising enzymes, because of the apparent association between the production of these glycosaminoglycan (GAG) depolymerising enzymes and pathogenicity of the ‘S. milleri group’ [5, 6].

GAGs are a group of linear polysaccharides composed of repeating disaccharide units. Based on differences in monosaccharide moieties and the degree and position of sulphation, five classes are recognised: chondroitin sulphate, dermatan sulphate, heparan sulphate, keratan sulphate and the non-sulphated hyaluronic acid. These macromolecules interact with other fibrous proteins to form an extracellular matrix that maintains the structural integrity of host tissues [7]. The ability to produce GAG depolymerising enzymes has been considered to be a potential virulence determinant in organisms such as S. pneumoniae, staphylococci and Clostridium perfringens [8-10]. These enzymes may play a role in the pathogenic process by acting as...
bacterial 'spreading' factors facilitating local tissue spread of the organisms. In contrast to mammalian enzymes that depolymerise GAGs in a hydrolytic fashion, bacterial GAG depolymerising enzymes cleave hexosaminy1-hexuronyl linkages of GAGs by a β-elimination reaction leading to the formation of products with an unsaturated double bond in the 4,5-position of the hexuronyl moiety [11–13].

Previous studies have demonstrated the production of hyaluronidase or CS depolymerase, or both, by members of the 'S. milleri group' [5, 6, 14, 15]. However, following recent changes in the taxonomy of this group [1], Homer et al. [16], attempted to detect hyaluronidase and CS depolymerase production by representatives of each of the currently recognised species of viridans streptococci with a sensitive and quantitative spectrophotometric assay. Both S. intermedius and S. constellatus degraded hyaluronic acid but only S. intermedius produced CS depolymerase. The production of these GAG depolymerising enzymes by S. intermedius is of interest because this species, in particular, is strongly associated with brain and liver abscesses [17, 18].

Hyaluronate, when used as the sole carbohydrate supplement to minimal media, has been shown to support the growth of S. intermedius [19] but the degradation and utilisation of CS by S. intermedius has not been reported. Therefore the ability of S. intermedius to utilise CS as the sole source of carbohydrate was studied, as this might provide an insight into the association between S. intermedius and deep-seated purulent infections.

Materials and methods

Growth and maintenance of S. intermedius

S. intermedius strain UNS 35, a brain abscess isolate obtained from Dr P. Unsworth (Central Public Health Laboratory, 61 Colindale Avenue, London), was stored frozen on glass beads (Technical Service Consultants) at –70°C. Routine subculturing was performed on Fastidious Anaerobe Agar (Lab M) containing defibrinated horse blood 5% v/v (FAA) with incubation at 37°C in an anaerobic atmosphere (5% CO2, 95% N2). Samples of each culture at intervals and measuring the increase in optical density at 620 nm in flat-bottomed 96-well microtiter trays with a microtitration plate reader (Titertek Multispan MCC-340; ICN-Flow Laboratories Ltd). Bacterial doubling times were calculated as described by Meynell and Meynell [22]. Additional samples (1 ml) were removed at intervals into microcentrifuge tubes; the cells were pelleted by centrifugation (13 000 rpm, 15 min, MSE Microfuge) and culture supernate were stored at –20°C for the subsequent analysis of residual substrates and metabolic end-products.

A modification of a semi-defined medium was used in all growth studies [20]. The basal medium contained sodium acetate 30 g, casein hydrolysate (Oxoid) 25 g, anhydrous Na2HPO4 15.75 g, anhydrous NaH2PO4 10.25 g, ammonium sulphate 3 g, KH2PO4 2.2 g, K3HPO4 1.5 g, trisodium citrate 1.125 g, adenine 0.15 g, uracil 0.15 g, guanine 0.1 g, ferrous sulphate 0.05 g, manganous sulphate 0.05 g and NaCl 0.05 g in distilled water 5.0 L. All chemicals were purchased from Sigma. The medium was dispensed into 1-L volumes and sterilised by autoclaving at 121°C for 15 min. Immediately prior to use, filter-sterilised solutions (0.2-µm pore size filters) of MgSO4, cysteine-HCl, and Na2CO3 were added to final concentrations of 0.20, 0.25 and 2.2 g/L, respectively, along with 1.0 ml of a filter-sterilised vitamin solution. The vitamin solution contained nicotinamide 40 mg, pantothenate 16 mg, pyridoxamine 16 mg, riboflavin 8 mg, thiamine 8 mg, biotin 0.2 mg, folic acid 2.0 mg and p-aminobenzoate 2.0 mg in 20 ml of distilled water. The medium was dispensed into sterile screw capped containers in 10-ml volumes.

Chondroitin sulphate A (CS-A, Sigma) and chondroitin sulphate C (CS-C, Sigma) were used in the growth studies. The CS-A preparation as supplied consisted of CS-A:CS-C (70:30) and CS-C consisted of CS-A:CS-C (10:90). The two types of CS differ in that CS-A is sulphated at the 4-position and CS-C is sulphated at the 6-position on the N-acetylgalactosamine moiety. The presence of both 4- and 6-sulphated residues in the commercial preparations are due not only to the difficulty of separating the two types of CS but also to the fact that most CS molecules are hybrid structures containing both CS-A and CS-C residues [21].

Filter-sterilised stock solutions of CS-A and CS-C, each at 40 mg/ml, and glucose at 20 mM were prepared. The stock solutions were added to the basal medium to give a final volume of 20 ml and the following concentrations of carbohydrate sources: CS-A 20 mg/ml; CS-C 20 mg/ml; 10 mM glucose, CS-A 10 mg/ml and 5 mM glucose; CS-C 10 mg/ml and 5 mM glucose. Media were pre-reduced prior to inoculation with 1 ml of a late exponential phase culture of S. intermedius grown in Brain Heart Infusion Broth (BHI; Oxoid) for 3–4 h. The cultures were placed on a rotating drum (26 rotations/min) and incubated at 37°C in an anaerobic atmosphere. Growth was monitored over a 16 h period by removing 200-µl samples of each culture at intervals and measuring the increase in optical density at 620 nm in flat-bottomed 96-well microtiter trays with a microtitration plate reader (Titertek Multispan MCC-340; ICN-Flow Laboratories Ltd). Bacterial doubling times were calculated as described by Meynell and Meynell [22]. Additional samples (1 ml) were removed at intervals into microcentrifuge tubes; the cells were pelleted by centrifugation (13 000 rpm, 15 min, MSE Microfuge) and culture supernate were stored at –20°C for the subsequent analysis of residual substrates and metabolic end-products.

All cultures were set up in duplicate and all assays (except the 232-nm spectrophotometric assay method, see below) for media constituents or metabolic end-products were performed in duplicate for each culture, with the data being presented as the mean of the four determinations.
Estimation of glucose and metabolic end-products

Glucose concentrations were estimated with Sigma kit No. 510. The A_440 was determined and glucose concentrations were calculated by comparison with a standard curve obtained with glucose in distilled water (linear up to 1 mM). Lactate was determined with Sigma kit No. 826. The increase in absorbance at 340 nm was measured and the A_340 at the end-point of the reaction was used to determine lactate concentrations with reference to standard lactate solutions prepared in distilled water (linear up to 10 mM). Ethanol was determined with a commercial diagnostic alcohol reagent kit (Sigma); the increase in absorbance at 340 nm was directly proportional to ethanol concentration up to an absorbance of 2.0. Formic acid was determined with a commercial kit (Boehringer Mannheim); the increase in absorbance at 340 nm was measured and formate concentration was calculated according to the manufacturer's instructions.

Determination of residual glycosaminoglycan

Residual CS was determined by the Stains-all method of Homcr et al. [23]. The Stains-all solution contained 1-ethyl-2-[3-(1-ethylthiazol-2-ylidene)-2-methylpropenyl]-naphtho-[1,2-d] thiazolium bromide (Sigma) and was prepared after the method of Benchetrit et al. [24]. The dye was dissolved to a final concentration of 0.1 mM in water: 1,4-dioxan (50:50 v:v) containing 1 mM acetic acid and 0.5 mM ascorbic acid. 1,4-Dioxan (AnalaR, BDH Ltd, Poole, Dorset) contained 2,6-di-tert-butyl-4-methyl phenol c. 25 ppm as a stabilising agent. Residual CS concentrations in culture supernates were estimated by mixing 20 μl of sample (appropriately diluted with distilled water) with 180 μl of the Stains-all solution and 100 μl of distilled water in microtitration trays and determining A_440 within 5 min. CS solutions, in distilled water gave a linear response over the concentration range up to 200 μg/ml when treated in the same manner and were included as standards.

Determination of N-acetylgalactosamine equivalents

The supernatant concentrations of N-acetylgalactosamine (GalNAc) equivalents, GalNAc moieties at the reducing terminal of saccharide species generated by the action of CS depolymerase, were estimated by a modification of the method of Levy and McAllan [25]. The reagent consisted of concentrated HCl 11 ml, distilled water 1.5 ml, glacial acetic acid 87.5 ml and 4-(N,N-dimethylamino)-benzaldehyde (Sigma) 10 g. Immediately prior to use, 10 ml of the reagent were diluted to 100 ml with glacial acetic acid. Samples (125 μl) were added to 25 μl of 0.2 M dipotassium tetraborate in a microcentrifuge tube and placed in a boiling water bath for 3 min. Samples were cooled to room temperature, 750 μl of the diluted reagent were added and the samples were incubated at 37°C for 20 min. The treated samples were cooled to room temperature, 200-μl volumes of each were dispensed into a microtitration tray and A_450 was determined. The concentration of reducing terminal GalNAc in culture supernate was determined by comparison with a series of standard GalNAc concentrations up to 10 mM, prepared in distilled water.

Spectrophotometric assay for determination of 4,5-unsaturated disaccharides

The presence of 4,5-unsaturated disaccharides was determined by a modification of the method of Saito et al. [26]. The unsaturated disaccharides have an absorbance maximum in the ultraviolet range (232 nm) at pH 2. Enriched Tris Buffer (ETB, pH 8.0) was prepared by dissolving tris (hydroxymethyl)amino methane 3 g, sodium acetate 2.4 g, NaCl 1.46 g and crystalline bovine serum albumin (Sigma) 50 mg in 100 ml of 0.13 M HCl. Samples of culture supernate (40 μl) were added to 10 μl of ETB in a microcentrifuge tube and incubated at 37°C for 30 min. To this reaction mixture, 950 μl of 0.01 M HCl were added and the absorbance of each treated sample was measured in a quartz cuvette at 232 nm with a Shimadzu UV 160-A recording spectrophotometer. The units for this assay were ΔA_232 relative to the control assay.

Preparation of enzyme extracts

A portion (25 ml) of a late exponential phase culture of S. intermedius grown in BHI was used to inoculate 500 ml of basal medium supplemented with either glucose (final concentration 10 mM) or CS-A (final concentration 5 mg/ml). The cultures were incubated anaerobically at 37°C with constant stirring for 18 h. Cells were harvested by centrifugation (10,000 rpm, 20 min, 4°C) and supernates were decanted. Ammonium sulphate (80% saturation) was added to each of the culture supernates with gentle stirring at 4°C. After 18 h at 4°C, the solutions were centrifuged at 15,000 rpm for 1 h at 4°C. The precipitates were dissolved in a minimal volume of ice-cold 50 mM sodium phosphate buffer, pH 7.0, and dialysed against three changes of the same buffer at 4°C. The dialysates were divided into small volumes and stored at −20°C.

The cell pellets were washed twice by centrifugation at 4°C in 50 mM sodium phosphate buffer, pH 7.0, and disrupted by shaking with glass beads (Ballaotini No. 12, Jencons) for 15 min at 4°C in a tissue disintegrator (Mickle Engineering Co.). Cell debris was removed by centrifugation (13,000 rpm, 5 min, 4°C) and the cell-free extracts were stored at −20°C.

Determination of protein concentrations

The protein content of concentrated culture supernates and cell extracts was determined by the Coomassie
Blue dye-binding assay (Sigma) and by comparison with a standard curve of bovine serum albumin (0–500 µg/ml).

**HPAEC-PAD quantification of 4,5-unsaturated disaccharides**

The concentrations of 2-acetamido-2-deoxy-3-O-(β-D-glucopyranosyluronic acid)-4-O-sulphido-D-galactose (AUA GalNAc-4S), 2-acetamido-2-deoxy-3-O-(β-D-glucopyranosyluronic acid)-6-O-sulphido-D-galactose (AUA GalNAc-6S) and 2-acetamido-2-deoxy-3-O-(β-D-glucopyranosyluronic acid)-D-galactose (AUA GalNAc-OS) in culture supernates were measured by high pH anion exchange chromatography with pulsed amperometric detection (HPAEC-PAD) by a modification of the method of Shibata et al. [27]. Prior to analysis the unsaturated disaccharides in culture supernates were stabilised to alkali conditions by borohydride reduction. Briefly, samples of culture supernates were centrifuged (13 000 rpm for 90 min) through a Micron 3 microconcentrator with a 10 kDa cut-off (Amicon Ltd). Sodium borohydride solution (100 µl of 1 M NaBH₄ in 100 mM NaOH) was added to 500 µl of ultrafiltrate to give final concentration of 200 mM NaBH₄. The borohydride reduction reaction was performed at room temperature for 1 h and was terminated by the addition of 180 µl of 1 M acetic acid to the sample placed on ice. A 100-µl sample of each reduced sample was diluted with 400 µl of 0.1 M NaOH and this was subjected to HPAEC-PAD analysis for the determination of concentrations of individual disaccharides.

Separation and quantification of CS-derived disaccharides was performed with a Dionex DX-500 chromatography system (Dionex Corporation, Sunnyvale, CA, USA) equipped with a Dionex Carbotrap PA 100 column (10 µm, 250 mm × 4 mm i.d.) fitted with a guard column (Dionex, PA100). Samples (500 µl) were transferred to sample vials (Dionex) fitted with 20-µm filters and injected with an autosampler. The column was equilibrated in 0.1 M sodium acetate in 0.1 M NaOH at a flow rate of 1.0 ml/min for 10 min before injection of sample (25 µl sample loop). Separation of the disaccharides (ΔUA GalNAc-4S, ΔUA GalNAc-6S and ΔUA GalNAc-OS) was achieved with a linear gradient of sodium acetate (0.1–1.0 M) in 0.1 M NaOH over 90 min, followed by isocratic elution with 1.0 M sodium acetate in 0.1 M NaOH for 10 min. It was necessary to re-equilibrate the column for 10 min under the starting conditions before application of a subsequent sample.

Eluted disaccharides were detected by a pulsed amperometric detector (PAD, Dionex) fitted with a gold working electrode. Peak areas were calculated with the DX-500 data handling system and concentrations of individual disaccharides were calculated by comparison with standard curves produced with authentic samples of ΔUA GalNAc-4S, ΔUA GalNAc-6S and ΔUA GalNAc-OS (Dextra Laboratories), over the concentration range 0–200 µg/ml, reduced and treated as described above.

**Measurement of sulphatase activity**

The ability of the concentrated supernate preparations to desulphate the authentic unsaturated disaccharides, ΔUA GalNAc-4S and ΔUA GalNAc-6S derived, respectively from CS-A and CS-C, was determined. Reaction mixtures contained concentrated culture supernate (20 µl), 0.2 M sodium phosphate buffer, pH 7.0 (30 µl) and ΔUA GalNAc-4S or ΔUA GalNAc-6S 2 mg/ml (50 µl). Separate samples were prepared for times 0, 2 and 6 h and incubated at 37°C. Assays were heated for 10 min in a boiling water bath to inactivate the enzyme, ultrafiltered and reduced as described previously and the concentration of the residual disaccharides was determined by HPAEC-PAD.

**Results**

**Growth of S. intermedius UNS 35**

Growth of *S. intermedius* in the presence of glucose was rapid with a doubling time of 1.3 h (Fig. 1a); glucose was completely utilised by 5 h. In minimal medium containing CS-A as the sole carbohydrate source, bacterial growth was not detected until the concentration of reducing terminal GalNAc was increased to c. 2 mM, probably indicative of CS-A depolymerisation (Fig. 1b). The doubling time of the organism on CS-A in the period of exponential growth was 3.6 h. The bacterial growth with the combination of glucose and CS-A is shown in Fig. 1c; glucose utilisation was rapid with a concomitant increase in absorbance and the doubling time was 1.2 h. Following the utilisation of glucose the absorbance of the culture continued to increase slowly which is in contrast to the decrease in absorbance observed in the cultures with glucose alone upon the complete utilisation of glucose from the medium. The CS-A concentration was 10 mg/ml and the utilisation of this no doubt contributed to the continued growth of the cells in the absence of glucose. No growth occurred when *S. intermedius* was grown in medium containing CS-C but in the medium supplemented with CS-C and glucose the strain grew with a doubling time of 1.4 h (data not shown).

Cells grown in the presence of glucose produced only lactate as the major end-product of metabolism, whereas cells grown in the presence of glucose and CS-A produced a mixture of lactate, formate and ethanol. The formation of acetate could not be demonstrated due to the formulation of the basal medium. *S. intermedius* grown with CS-A produced ethanol and formate, but not lactate as the major end-products (Table 1). After growth for 16 h, a reduction in the concentration of CS-A was demonstrable (by
Fig. 1. Growth characteristics of *S. intermedius* UNS 35 in semi-defined media supplemented with: a 10 mM glucose
b, CS-A 20 mg/ml; c, 5 mM glucose and CS-A 10 mg/ml. The cultures were incubated anaerobically at 37°C and
samples were taken at the times indicated for determination of optical density (■), glucose utilisation (▲), production
of lactate (▼) and N-acetylamino sugar (●).
the Stains-all assay) while no change in the concentration of CS-C was observed in either of the CS-C supplemented cultures. Similarly, after 16 h there was an increase in the concentration of reducing terminal GalNAc and in the concentration of 4,5-unsaturated disaccharides (determined by the 232-nm assay) in the cultures supplemented with CS-A, but these were present only at low levels in the CS-C supplemented cultures (Table 1).

**Breakdown products of CS-A by S. intermediusUNS 35**

The chemical assays used in this study are sensitive but not necessarily specific: the 232-nm assay does not distinguish between ΔUA GalNAc-4S (derived from CS-A), ΔUA GalNAc-6S (derived from CS-C) and ΔUA GalNAc-0S, and the N-acetylamino sugar assay gives a positive reaction with ΔUA GalNAc-6S and ΔUA GalNAc-0S but not with ΔUA GalNAc-4S [26]. As S. intermedius UNS 35 grew with CS-A but not with CS-C and the N-acetylamino sugar assay results paralleled the increase in cell numbers, HPAEC-PAD analysis was used to characterise and quantify the individual unsaturated disaccharides present in culture supernatants during the growth of S. intermedius in the presence of CS-A.

The separation of ΔUA GalNAc-0S, ΔUA GalNAc-4S and ΔUA GalNAc-6S was readily achieved (Fig. 2a) and the detector response was linear for each disaccharide (Fig. 2b). The changes in the supernate concentration of the individual unsaturated disaccharides over a 16-h incubation period are shown in Fig. 3. The concentration of ΔUA GalNAc-0S increased in a linear fashion to a peak at c. 8 h and then, presumably as a consequence of bacterial growth, was gradually removed from the medium. ΔUA GalNAc-6S was detectable at 4 h and its concentration rose progressively, paralleling cell production, reaching a maximum at 14 h after which the concentration remained level. The changes in the concentrations of ΔUA GalNAc-0S and ΔUA GalNAc-6S are in contrast to the behaviour of ΔUA GalNAc-4S, the major breakdown product of CS-A, which accumulated only slowly in the medium (Fig. 3). The initial rise of GalNAc (Fig. 1b) was due to the accumulation of ΔUA GalNAc-0S, and after 8 h ΔUA GalNAc-6S, derived from CS-C, was the predominant unsaturated disaccharide, which suggests that bacterial growth was a consequence of the utilisation of ΔUA GalNAc-0S.

The presence of both sulphated (ΔUA GalNAc-6S, ΔUA GalNAc-4S) and non-sulphated (ΔUA GalNAc-0S) unsaturated disaccharides in these cultures indicates the production, by S. intermedius, of CS-AC depolymerase and sulphatase activities.

**Location and induction of CS depolymerase**

The CS depolymerase activity associated with bacteria and in the culture supernate was measured after growing bacteria in minimal medium supplemented with either glucose or CS-A. Culture supernates were concentrated before performing these assays as the level of activity in untreated culture supernates was too low to be measured reliably. CS depolymerase activity was detected in the culture supernate of minimal medium with glucose (1.24 mg of CS-A degraded/h/mg of protein) but activity was much higher in the supernate with CS-A (34.20 mg of CS-A degraded/h/mg of protein). CS depolymerase activity was associated with cells grown in media containing CS-A but this was only 0.3% of the supernate activity, which was similar to the proportion (0.4%) present in the cells grown with glucose.

**Measurement of sulphatase activity**

Preliminary experiments were performed with authentic ΔUA GalNAc-4S and ΔUA GalNAc-6S to determine which of these was the preferred substrate for the S. intermedius sulphatase activity to identify the origin of the ΔUA GalNAc-0S detected in the culture supernate of bacteria grown with CS-A. The rates of loss of ΔUA GalNAc-4S and ΔUA GalNAc-6S were >370 and 70 μmol/h/mg of protein, respectively, indicating that the majority of ΔUA GalNAc-0S was derived from the desulphation of ΔUA GalNAc-4S, the 4,5-unsaturated disaccharide produced as a result of CS-A depolymerisation by S. intermedius.
**Discussion**

*S. intermedius* is unique amongst the viridans streptococci in its ability to produce CS depolymerase activity. Previous investigations into the substrate specificity of this enzyme activity have tended to suggest that *S. intermedius* has the ability to degrade CS-C [14, 28], whereas in a study of 10 strains of *S. intermedius*, CS-A but not CS-C was depolymerised [16]. In these studies a rich, complex medium (BHI) containing glucose and supplemented with the appropriate form of CS was used. The data presented in this study provide an explanation for the apparent discrepancies between these previous reports.

In the present study, CS-A, but not CS-C, sustained the growth of *S. intermedius* UNS 35. In these experiments the source of CS-A also contained 30% CS-C and, as a consequence of growth in CS-A-supplemented media, it was clear that *S. intermedius* depolymerised both CS-A and CS-C. This was apparent from the indirect chemical assays and from the direct measurement of 4,5-unsaturated disaccharides in culture supernates by HPAEC-PAD. The formation of the individual 4,5-unsaturated disaccharides in the culture supernates provided information on the sequence of metabolic events involved in the depolymerisation and utilisation of CS-A. Although ΔUA GalNAc-4S is the product derived initially from CS-A depolymerisation, the

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**Fig. 2.** HPAEC-PAD identification and quantitation of unsaturated disaccharides derived from CS. a, Separation of ΔUA GalNAc-6S (1), ΔUA GalNAc-4S (2), ΔUA GalNAc-6S (3) with Carbopac PA100. Samples (25 µl) of authentic disaccharides at a final concentration of 100 µg/ml after sodium borohydride reduction were injected and eluted with a sodium acetate gradient as described in Materials and methods. b, Linearity of PAD response with varying disaccharide concentrations.
Fig. 3. Quantification of unsaturated disaccharides released from the breakdown of CS-A in the culture supernate of *S. intermedius* UNS 35 grown in a semi-defined medium supplemented with CS-A as the major fermentable carbohydrate. Samples of culture supernate were taken at the times indicated (corresponding to data illustrated in Fig. 1) and ΔUA GalNAc-0S (■■■), ΔUA GalNAc-4S (■■■■), ΔUA GalNAc-6S (■■■■■) concentrations were determined by HPAEC-PAD analysis.

concentration of ΔUA GalNAc-4S remained low throughout the growth period. However, there was a significant rise in the concentration of ΔUA GalNAc-0S in the culture supernate, while the concentration of ΔUA GalNAc-6S increased throughout the incubation period, indicating that the ΔUA GalNAc-4S, rather than ΔUA GalNAc-6S, was the preferred substrate for the sulphatase activity of *S. intermedius* and this was confirmed in experiments with authentic unsaturated disaccharides. Enzymes with sulphatase activity against CS-derived disaccharides have been demonstrated in other bacteria [29,30]. The chondrosulphatases of *Proteus vulgaris* have been shown to have a high degree of specificity for the 4- and 6-sulphate esters, respectively [30].

In streptococci, the formation of lactate as the major or sole acid end-product is indicative of carbon excess as reported here in the glucose-supplemented cultures. The formation of mixed fermentation products, by heterofermentative metabolism, as was found in the CS-A-supplemented cultures, is indicative of carbon-limited growth in streptococci [31,32]. The mechanism underlying the apparent carbon-limited growth in CS-A supplemented cultures is unknown.

The present observations provide an explanation for the apparent anomalies between the reports of Osano *et al.* [14] and Hibi *et al.* [28] and our own [16]. We reported that *S. intermedius* strains degraded CS-A in BHI broth but the studies of Osano *et al.* [14] indicated that *S. intermedius* produced CS-C depolymerase when grown on BHI agar supplemented with CS-C; enzyme activity was detected as described by Smith and Willet [33]. We have used this assay technique and found that, when *S. intermedius* UNS 35 was stabbed into the medium, with CS-A as the substrate, the zones of depolymerisation were 6 mm in diameter, whereas with CS-C the zones were smaller (4 mm) and less distinct (data not shown). Hibi *et al.* [28] also demonstrated that a partially purified enzyme produced from a culture of *S. intermedius* grown on BHI supplemented with CS-C apparently had only a CS-C activity after incubation for 48 h. However, the present results demonstrate that *S. intermedius* depolymerised both CS-A and CS-C, but the resulting 4- and 6-sulphated disaccharides were subsequently desulphated at significantly different rates. As a result, prolonged incubation of a mixture of CS-A and CS-C with a partially purified enzyme preparation of *S. intermedius* is likely to result in the formation of ΔUA GalNAc-6S and ΔUA GalNAc-4S and the latter will be desulphated rapidly as reported here. The desulphated disaccharide (ΔUA GalNAc-0S) may be further metabolised by an intracellular 'glucuronidase'-like activity [11,12] released during supernate preparation. This sequence of events could explain why only ΔUA GalNAc-6S was detected by Hibi *et al.* [28]. Therefore, *S. intermedius* like many other bacterial species, produces a CS-AC depolymerase and not a specific CS-C depolymerase. Specific CS-C depolymerases are produced by only a few bacterial species [34].

The production of CS depolymerase by *S. intermedius* UNS 35 was inducible; little activity was produced by cells growing in the presence of glucose. This finding is consistent with the low level of production of polysaccharide-degrading enzymes by other bacteria when a more readily catabolisable substrate is available [35,36]. Also, a previous report has shown that exoglycosidic enzymes, (neuraminidase, β-galactosidase, β-N-acetylgalactosaminidase and β-N-acetyl-D-glucosaminidase) with the ability to degrade the oligosaccharide of human glycoproteins, are repressed in *S. intermedius* when glucose is provided as a carbohydrate source [37]. It is energetically unfavourable for micro-organisms to produce enzymes to liberate nutrients from larger molecules when a more readily available nutrient source is provided and this may explain the inducible nature of the CS depolymerase of *S. intermedius*.

In gram-negative bacteria, including *Bacteroides thetaiotaomicron*, *P. vulgaris* and *Flavobacterium heparinum*, CS depolymerase is apparently located in the periplasmic space [30,38]. However, we found that >99% of the CS depolymerase activity of *S. intermedius* was in the culture supernate, with <1% being cell-associated. These observations are in accord
with those of Hibi et al. [28] who found CS-C depolymerase activity to be predominantly in the culture supernate.

CS is widely distributed throughout connective tissues and in most cases has a hybrid structure with both 4- and 6-sulphate groups present in the same molecule [39]. The present study demonstrated that S. intermedii strain UNS35 produces CS depolymerase activity with the ability to depolymerise both CS-A and CS-C. The organism was better able to desulphate ΔUA GalNac-4S, derived from CS-A, than ΔUA GalNac-6S, derived from CS-C to yield ΔUA GalNac-0S, which may be slowly metabolised to yield cell mass. Further investigations are required to characterise the CS depolymerase and sulphatase activities of S. intermedii and to understand the mechanism of transport and intracellular metabolism of ΔUA GalNac-0S. These findings indicate that CS depolymerase production by S. intermedii may play a role not only in the destruction of host connective tissue but also in bacterial nutrition by the release of nutrients, so that the organisms can thrive and persist in an otherwise nutritionally compromised infected site.

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References


