A Study of Some Motile Group D Streptococci

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SUMMARY

Thirteen strains of motile enterococci showed more similarity, in their
dphysiological reactions, to *Streptococcus faecium* than to *S. faecalis*. A
serological study of the type antigens divided the motile strains into four sets;
(1) three strains previously described as *S. faecium* serotype 29, (2) one
strain reacting as *S. faecium* serotype 38, (3) five strains showing a specific
reaction with antiserum prepared to one of them ("serotype 4725"), (4) four
untyped strains. Esterase and protein patterns from the soluble fractions of
the motile strains were examined by electrophoresis in polyacrylamide gel and
were different from those of non-motile strains of *S. faecalis*, *S. faecium* and
*S. durans*. Extracts of the three motile strains of *S. faecium* type 29 showed
a common esterase pattern, extracts from five strains of serotype 4725
showed three different esterase patterns. The motile strain of *S. faecium*
serotype 38 gave extracts with a strong esterase band which differed from
the very weak bands shown by extracts of non-motile strains. Differences in
esterase pattern could be found between motile strains whether untyped,
of different serotypes, or of the same serotype. Comparison of the 'protein
patterns' of motile strains gave some indication that major protein bands
occurred at similar positions after electrophoresis.

INTRODUCTION

A previous study by gel electrophoresis of soluble cell components (Lund, 1965)
showed that 12 strains of *Streptococcus faecalis* and its varieties 'zymogenes' and
'liquefaciens' of eight serotypes had a similar pattern of separated protein bands,
and showed strong bands with esterase activity. Strains of *S. faecium* of 19 sero-
types gave 'protein patterns' with major bands having similar mobility (amongst
these strains) but strong esterase bands were not detected. The pattern of protein
bands of *S. faecium* and *S. durans* differed from that of *S. faecalis*. Of 21 strains of
*S. faecium* and *S. durans* examined initially, one strain had a distinctive pattern of
proteins and strong bands of esterase activity; this strain was the only motile one
among those initially studied.

The purpose of the present work was to study, by gel electrophoresis, extracts of
other motile strains of enterococci from diverse sources to compare their protein
components and esterase enzymes, and to correlate this comparison with a study of
some physiological and serological properties. Physiological properties and agglutina-
tion reactions of a large number of strains of motile enterococci have previously been
studied by Graudal (1952, 1955, 1957a, b).

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METHODS

Organisms. The motile enterococci studied are listed in Table I. They were compared with strains of Streptococcus faecalis and S. faecium which have previously been described (Lund, 1965). Cultures of S. faecalis and S. faecium were stored on slopes of Hartley’s digest agar at 5°C. Because cultures of motile organisms tended to die during storage they were maintained as stab cultures in Brain Heart Infusion agar (Difco) at 5°C and were subcultured at intervals of 1 month.

Table I. Strains of motile group D streptococci studied

<table>
<thead>
<tr>
<th>Strain (species name is that used in the reference cited)</th>
<th>Source</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>S. faecium (type 29, P14/6 type strain)</td>
<td>Pig colon</td>
<td>Sharpe &amp; Fewins (1960) ; Barnes (1964)</td>
</tr>
<tr>
<td>S. faecium</td>
<td>H/2/23 Human faeces</td>
<td>Pownall (1935)</td>
</tr>
<tr>
<td>S. faecium</td>
<td>119 Piglet</td>
<td></td>
</tr>
<tr>
<td>S. faecalis</td>
<td>NCTC 4725 Probably human</td>
<td></td>
</tr>
<tr>
<td>Streptococcus sp. D 1000(b)</td>
<td>Representatives of Graudal’s collection, isolated mainly from human faeces</td>
<td>Graudal (1957a, b)</td>
</tr>
<tr>
<td>Streptococcus sp. D 1003(b) (described as pigmented strain)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Streptococcus sp. D 1006(c)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Streptococcus sp. D 1010(c) (described as pigmented strain)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Streptococcus sp. ATCC 12817</td>
<td>Oropharynx, patient undergoing treatment for pernicious anaemia</td>
<td>Hugh (1959)</td>
</tr>
<tr>
<td>Streptococcus sp. ATCC 12818</td>
<td>Mouth, patient with untreated oral cancer</td>
<td></td>
</tr>
<tr>
<td>Streptococcus sp. ATCC 12819</td>
<td>As ATCC 12817</td>
<td>Liu et al. (1955)</td>
</tr>
<tr>
<td>Streptococcus sp. ATCC 13638</td>
<td>Spinal fluid, patient with chronic meningitis</td>
<td></td>
</tr>
<tr>
<td>S. faecalis</td>
<td>ATCC 12755 Milk</td>
<td>Sherman (1937); Hugh (1959)</td>
</tr>
</tbody>
</table>

Motility was demonstrated by stab inoculation of the organism into Brain Heart Infusion Broth (Difco)+0.25 % (w/v) Bacto-Agar (Difco) and incubation at 25°C, 30°C or 37°C for 24 hr. The diffuse growth of motile strains could be distinguished from the more discrete growth of non-motile strains. Motility was confirmed by microscopical examination of living cells. Flagella were stained by Leifson’s technique (Leifson, 1951).

Biochemical tests. The methods used for testing haemolysis, growth in presence of 40 % bile, sensitivity to potassium tellurite, growth at 10°C, growth at 45°C, growth in presence of 6.5 % NaCl, survival at 60°C for 30 min. were those described by Barnes, Ingram & Ingram (1956). The appearance of colonies on thallous acetate-tetrazolium agar (TITg) was studied using thallous acetate agar (Baltimore Biological Laboratory) and the method described by Barnes (1956). In all the above tests incubation was at 37°C unless otherwise specified. Production of acid in carbohydrate media was studied by the method of Whittenbury (1963) utilizing a soft agar medium. The basal medium contained meat extract (Lab-Lemco), 5 g.; peptone (Evans), 5 g.; yeast extract (Difco), 5 g.; Tween 80, 0.5 ml.; agar (Davis), 1.5 g., distilled water to 1 l. As pH indicator
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5.6 ml. of a 1.6% (w/v) solution of bromocresol purple in ethanol was added to 1 l. of medium and the pH adjusted to 6.8–7.0. Carbohydrate substrates (B.D.H. Laboratory reagent grade), as 5% (w/v) solutions in distilled water, were sterilized separately at 120°C for 15 min. and added to the basal medium to give a final concentration of 0.5% (w/v). The melted medium was cooled to 45°C before inoculating and then allowed to set. Results were recorded after incubation at 37°C for 3 days. Tests for utilization of individual carbohydrates as sources of energy for growth were made by the technique of Deibel, Lake & Niven (1963).

For aerobic tests 10 ml. quantities of media in T-tubes were inoculated and shaken at 104 strokes/min. with an amplitude of 2.25 in. on a reciprocal shaker; for anaerobic tests tubes containing 10 ml. of inoculated media were placed in a desiccator under an atmosphere of 95% nitrogen + 5% carbon dioxide. After incubation for 18 hr at 30°C growth was estimated by turbidity measured in an ‘EEL’ nephelometer (Evans Electroselenium Ltd., Harlow, Essex). A positive result was taken as a turbidity, in the presence of added carbohydrate, equal to at least twice that in the basal medium.

Serological test for group D antigen. Extracts prepared as for polyacrylamide gel electrophoresis were tested for precipitin reaction with group D Streptococcus Grouping Serum (Burroughs Wellcome and Co.). Results were recorded 30 min. after testing.

Serological typing. The methods used were based on those described by Sharpe & Shattock (1952).

Type antisera. In preliminary experiments, extracts of motile strains were tested against Streptococcus faecium type antisera which had been prepared by Dr M. E. Sharpe, National Institute for Research in Dairying, Shinfield, Reading. In subsequent experiments antisera were prepared to motile strains P14/6, ATCC 13638, NCTC 4725 and the non-motile strain P17/8, using the following technique. Cultures were grown in 80 ml. of medium containing glucose, 1 g.; peptone (Evans), 1 g.; Lab-Lemco (Oxoid), 1 g.; NaCl, 0.5 g.; distilled water to 100 ml. pH 7.0–7.2 (medium A). After incubation at 37°C for 18 hr cells were separated by centrifugation, washed twice in 50 ml. of 0.85% (w/v) NaCl in water and resuspended in 20 ml. solution containing 0.85% (w/v) NaCl and 0.1% (w/v) HCHO. The suspension was kept at room temperature for 24 hr, after which tests for viable organisms were negative. The suspension was adjusted to an opacity of 7–8 using standard opacity tubes (Burroughs Wellcome and Co.) and stored at −20°C.

Two rabbits were inoculated by intravenous injections with each organism. An initial sensitizing dose of 0.4 ml. was given, and after 5 days the following series of injections was started: three injections of 0.4 ml. (week 1), three of 0.8 ml. (week 2), three of 1.0 ml. (week 3). Three days after the last injection a test bleeding showed satisfactory antibody production; 25 ml. of blood was withdrawn from each rabbit. After clot formation the separated serum was stored at −20°C without preservative.

Absorbed antisera. For the initial experiments the absorbed type antisera to a range of serotypes of Streptococcus faecium had been prepared by Dr M. E. Sharpe. Further absorbed antisera were prepared by the following technique. The organism used for the absorption was grown in 200 ml. medium A at 30°C for 18 hr, the cells harvested by centrifuging, washed and resuspended in 10 ml. 0.85% (w/v) NaCl + 0.1% (w/v) HCHO. The suspensions were then heated at 70°C for 30 min. and washed on the centrifuge in saline-formaldehyde solution. The packed cells were resuspended in four
times their volume of antiserum and incubated at 37° for 2 hr, stirring frequently. After storage at +1° overnight the cells were sedimented by centrifugation, the antiserum removed and tested against an extract of the homologous organism.

*Extraction of type antigens.* Cultures in 20 ml. of medium A were incubated at 25° for 24 hr. Hydrochloric acid extracts were prepared by a technique based on Lancefield's (1933) method. The sediment from 20 ml. of growth medium was resuspended in 0.5 ml. of HCl (0.05 N or 0.01 N) in 0.85% (w/v) NaCl, heated in a boiling-water bath for 10 min., cooled, centrifuged and the clear supernatant neutralized by the addition of 1N- or 0.2N-NaOH to an end-point with phenol red; 1 drop of 1% (w/v) solution of Thiomersal (British Drug Houses Ltd.) was added as preservative.

*Removal of flagella before extraction of type antigens.* Cultures in 20 ml. of medium A were incubated at 25° for 24 hr. Cells were separated by centrifuging at 1000 g for 20 min. and resuspended in 0.85% (w/v) saline. A sample stained by Leifson's (1951) technique showed the presence of flagella, many of which had become detached. In order to detach remaining flagella 10 ml. of the bacterial suspension was treated in a homogenizer (Measuring and Scientific Equipment Ltd.) using a blending speed of approximately 12,000 rev./min. for 2 min. After centrifuging at 10,000 g for 15 min. detached flagella tended to remain in the supernatant, the sedimented cells were resuspended in 10 ml. of fresh 0.85% (w/v) saline. This centrifuging and resuspension in fresh saline was repeated 4 times, giving a final sediment which consisted mainly of intact, deflagellated organisms. A hydrochloric acid extract was prepared as previously described, using for comparison a cell sediment prepared without removal of flagella.

*Precipitin tests.* Strong precipitin reactions were observed by the capillary tube technique of Swift, Wilson & Lancefield (1943). For weaker reactions it was necessary to use the precipitin ring technique (Sharpe & Shattock, 1952). In each case extracts prepared with 0.05 N-HCl in 0.85% (w/v) NaCl (pH 1-6) gave a stronger precipitin reaction than those prepared with 0.01 N-HCl in 0.85% (w/v) NaCl (pH 2-1); this differed from the results obtained by Maxted & Fraser (1966) with *Streptococcus faecalis* strains.

*Gel diffusion tests.* These were made on microscope slides 3 in. × 1 in. covered with 2 ml. of the following medium: barbitone-acetate buffer (Oxoid), 8.25 g., thiomersal (B.D.H.), 0-1 g., Ionagar no. 2 (Oxoid), 10 g., distilled water to 1 l. Wells were cut in the agar, filled with antiserum or antigen extract and the slides placed at 0-5° for 24 hr in a moist atmosphere for precipitin lines to develop.

*Immuno-electrophoresis.* Lantern cover glasses 3½ in. × 3½ in. were covered with 12 ml. of the medium described for gel diffusion tests. Wells were cut and filled with antigen extract. For electrophoresis the tank buffer contained barbitone-acetate buffer (Oxoid) 8.25 g.; distilled water to 2 l.; a potential difference of 8–10 V. per cm. was applied for 1-5 hr. Trenches were made, filled with antiserum, and the preparation was placed at 0-5° in a moist atmosphere for 24 hr to allow development of precipitin lines.

*Effect of periodate on type antigens.* Attempts were made to detect sensitivity of the antigens to periodate, using a method similar to that of Sharpe (1964). Neutralized hydrochloric acid extract 0.5 ml., or the acetone-precipitated fraction of this extract, was treated with an equal volume of 0.4M-acetate buffer pH 4 containing 0.08M-sodium periodate. After incubation at 37° for 16 hr 0.05 ml. of 5% (v/v) ethylene glycol was added to destroy the periodate, the pH was adjusted to 7 with N-NaOH.
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and the extract was tested against the homologous antiserum. Controls were included in which antigen extract was replaced by water.

Effect of trypsin on type antigens. Crystalline trypsin was added to the neutralized hydrochloric acid extract to give a concentration of 2 mg./ml. After incubation at 37° for 3 hr the extract was tested against the homologous antiserum.

Preparation of cell extracts for electrophoresis. Cultures were grown in 800 ml. volumes of Brain Heart Infusion Broth (Difco Laboratories) in static culture for 16 hr at 30°. The bacteria were harvested by centrifugation, washed, and the wet pellet of cells resuspended in half its weight of tris-citrate buffer pH 8.7 (see below). The suspension was frozen at −20°, disrupted in an X-press (AB Biox, Box 235, Nacka 2, Sweden), melted and clarified by centrifugation as previously described (Lund, 1965). The final extracts, with a protein content of 30–50 mg./ml., as estimated by the method of Lowry, Rosebrough, Farr & Randall (1951) were stored at −20°.

Electrophoresis in polyacrylamide gel. Conditions for electrophoresis, staining for protein and for esterase enzymes (hydrolysing α-naphthyl acetate) and for photography were as previously described (Lund, 1965). The gel was prepared in buffer containing tris, 38 mM; citric acid 2.5 mM (pH 8.7 at 25°C); the buffer for electrode vessels contained boric acid, 7.22 g./l., borax, 15.75 g./l. (0.28 M in terms of borate) pH 8.8 at 25°.

RESULTS

Biochemical properties of motile strains

Table 2 summarizes the properties of motile strains of enterococci and includes a comparison of the motile and non-motile strains in serotype 38.

The term 'enterococci' [introduced by Sherman (1937) to include streptococci giving positive results in the 'Sherman tests' (Deibel, 1964)], excludes Streptococcus bovis and S. equinus. In general the motile strains gave positive results in the Sherman tests except for 1 strain, ATCC 12755 which failed to grow at 45°. The nature of haemolysis, growth in presence of 0.04% (w/v) potassium tellurite, production of acid from sorbitol and L(+)-arabinose, and appearance of colonies on thallous acetate-tetrazolium agar are well-established tests used to differentiate S. faecalis from S. faecium and S. durans (Deibel, 1964). Sensitivity of the motile strains to potassium tellurite was intermediate between that of S. faecalis and that of S. faecium and S. durans, in agreement with the observation of Graudal (1957b). Like S. faecium, motile strains grew well in the presence of L(+)-arabinose, and marked acid production occurred in 24 hr. In contrast with S. faecium, many of the motile strains showed some production of acid in the presence of sorbitol, but generally the change in colour of the indicator was slower and less well-marked than in the case of S. faecalis. Strain NCTC 4725 was received as a strain of S. faecalis. It differed from typical S. faecalis in (i) appearance on thallous acetate-tetrazolium agar, (ii) fermentation of L(+)-arabinose and not of sorbitol, (iii) failure to utilize citrate or glycerol + fumarate as energy sources. The results in Table 2 show that this strain differed in several respects from typical S. faecalis and in common with other motile strains showed more similarity to S. faecium than to S. faecalis in these tests.
### Table 2. Biochemical reaction of motile enterococci compared with type strains of S. faecalis and S. faecium

(Reactions of non-motile S. faecium type 38 are included for comparison with the motile type 38 strain.)

<table>
<thead>
<tr>
<th>Motile strains</th>
<th>Serotype</th>
<th>'Sherman' test*</th>
<th>Haemolysis on horse blood agar</th>
<th>Growth on blood agar + 0.04% K₂ tellurite†</th>
<th>Appearance of colonies on thallous acetate-terreillium agar</th>
<th>Production of acid in medium containing</th>
<th>Utilization of substrates as source of energy</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>White to pale pink colonies†</td>
<td>Sorbitol</td>
<td>Arabinose</td>
</tr>
<tr>
<td>P 14/6</td>
<td>29</td>
<td>+ o or α</td>
<td>±</td>
<td></td>
<td></td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>H/2/23</td>
<td>29</td>
<td>+ α</td>
<td>+</td>
<td></td>
<td></td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>NCTC 4725</td>
<td>'4725'</td>
<td>+ a</td>
<td>+</td>
<td></td>
<td></td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>ATCC 12819</td>
<td>'4725'</td>
<td>+ a</td>
<td>+</td>
<td></td>
<td></td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>D 1000(b)</td>
<td>'4725'</td>
<td>+ a</td>
<td>+</td>
<td></td>
<td></td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>D 1006(c)</td>
<td>'4725'</td>
<td>+ a</td>
<td>+</td>
<td></td>
<td></td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>ATCC 12817</td>
<td>Untyped</td>
<td>+ a</td>
<td>+</td>
<td></td>
<td></td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>ATCC 12818</td>
<td>Untyped</td>
<td>+ a</td>
<td>+</td>
<td></td>
<td></td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>ATCC 12755</td>
<td>Untyped</td>
<td>§ a</td>
<td>+</td>
<td></td>
<td></td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>D 1003(b)</td>
<td>Untyped</td>
<td>+ a</td>
<td>+</td>
<td></td>
<td></td>
<td>+</td>
<td>+</td>
</tr>
<tr>
<td>ATCC 13638</td>
<td>38</td>
<td>+ a</td>
<td>-</td>
<td></td>
<td></td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Non-motile strains</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P 17/8</td>
<td>38</td>
<td>+ α</td>
<td>-</td>
<td></td>
<td></td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>30/66</td>
<td>38</td>
<td>nt α</td>
<td>-</td>
<td></td>
<td></td>
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</tr>
</tbody>
</table>

Reactions of typical strains of:

<table>
<thead>
<tr>
<th>S. faecalis</th>
<th>+ o or β</th>
<th>Red colonies</th>
<th>+</th>
<th>1/2+</th>
<th>1/2+</th>
<th>1/2+</th>
<th>1/2+</th>
<th>1/2+</th>
<th>1/2+</th>
<th>1/2+</th>
<th>1/2+</th>
</tr>
</thead>
<tbody>
<tr>
<td>S. faecium</td>
<td>+ o α</td>
<td>White to pale pink colonies</td>
<td>+</td>
<td>1/2+</td>
<td>1/2+</td>
<td>1/2+</td>
<td>1/2+</td>
<td>1/2+</td>
<td>1/2+</td>
<td>1/2+</td>
<td>1/2+</td>
</tr>
<tr>
<td>S. durans</td>
<td>+ β</td>
<td>+</td>
<td>+</td>
<td>1/2+</td>
<td>1/2+</td>
<td>1/2+</td>
<td>1/2+</td>
<td>1/2+</td>
<td>1/2+</td>
<td>1/2+</td>
<td>1/2+</td>
</tr>
</tbody>
</table>

* The term 'Sherman tests' is used to include the following: growth in presence of 40% bile; growth at 10°; growth at 45°; growth in presence of 6.5% NaCl; survival at 60° for 30 min.
† ± denotes production of small, greyish colonies, in contrast with good growth of black colonies of S. faecalis.
‡ Strain P14/6 produced variant, red colonies.
§ Failed to grow at 45°.
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Serological typing

Of the 13 motile strains, three belonged to *Streptococcus faecium* serotype 29 of which one, P14/6, is the type strain. Extracts of the remaining 10 strains were tested against antisera which were available to 17 serological types of *S. faecium*; only one strain gave a reaction, ATCC 13638, which belonged to serotype 38 (type strain non-motile P17/8). To confirm these results antisera were prepared to type strain P14/6 (type 29, motile), type strain P17/8 (type 38, non-motile), ATCC 13638 and also to the untyped strain, NCTC 4725. Precipitin reactions with these antisera are shown in Table 3.

Table 3. Precipitin reactions of extracts of motile enterococci and of a non-motile type 38 strain of *S. faecium*

<table>
<thead>
<tr>
<th>Antiserum to</th>
<th>P14/6</th>
<th>P17/8 (non-motile)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>S. faecium</em> type 29, type strain</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td><em>S. faecium</em> type 38, type strain</td>
<td>+</td>
<td>-</td>
</tr>
<tr>
<td>ATCC 13638</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>ATCC 12819</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>ATCC 12817</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>ATCC 12818</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>ATCC 12755</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>NCTC 4725</td>
<td>-</td>
<td>+</td>
</tr>
<tr>
<td>D1000(b)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>D1006(c)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>D1010(b)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>D1010(c)</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

In each case absorption of the antiserum with a heterologous reacting organism abolished the precipitin reaction with the homologous organism. Gel diffusion tests showed reactions of identity between precipitin lines of homologous and heterologous extracts reacting with antiserum to NCTC 4725. The term 'type 4725' will be used to describe organisms reacting with antiserum to NCTC 4725, but not with antiserum to P14/6.

The extracts reacting with antiserum to NCTC 4725 included those of the *Streptococcus faecium* type 29 strains P14/6, H/2/23 and 119. Antiserum to P14/6, which produced a strong precipitin reaction with the homologous extract, gave no reaction with an extract of NCTC 4725 (Table 3). The precipitin line produced by extract of P14/6 and antiserum to NCTC 4725 did not give a reaction of identity with the line produced by P14/6 and the homologous antiserum (Fig. 1). This indicates that two antigenic groups were present in acid extracts of strain P14/6, one of which reacted with antiserum to NCTC 4725 but failed to induce detectable antibody formation. The two antigens could be distinguished by immuno-electrophoresis of an acid extract of strain P14/6. The antigen revealed by the homologous antiserum showed a lower
mobility than that revealed by antiserum to NCTC 4725 (Fig. 2). Immuno-electrophoresis of extracts of 'type 4725' strains revealed a single precipitin with a similar mobility in each extract under these conditions.

Gel diffusion tests showed that the precipitin line produced by *Streptococcus faecium* type 38 strain P17/8, and by ATCC 13638 against antiserum to either of these strains gave a reaction of identity; by immuno-electrophoresis a single antigen with the same mobility was detected in extracts of each strain.

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Fig. 1. Diagram of gel diffusion test of extract of strain P14/6 (S. faecium type 29, type strain, motile) against homologous antisera and against antiserum to strain NCTC 4725. Centre well contained 0.05N-HCl extract of strain P14/6. Outer wells A and B contained antisera, from two different rabbits, to strain P14/6; C contained antiserum to strain NCTC 4725.

Fig. 2. Diagram of immuno-electrophoresis of extract of strain P14/6. An 0.05N-HCl extract of strain P14/6 was placed in the three wells. After electrophoresis trench A was filled with antiserum to strain NCTC 4725, trench B was filled with antiserum to strain P14/6.

The nature of the type antigens studied. The experiments relating to the chemical nature of the antigens were inconclusive. Treatment with 0.06 M-sodium periodate for 16 hr at 30° failed to destroy the antigens, although precipitin reactions of periodate-treated extracts tended to be weaker than those of control extracts. Treatment with 2 mg./ml. of trypsin for 3 hr at 37° failed to inhibit precipitin reactions.

The immunization procedure for serological typing of enterococci involves the use of formalin-killed suspensions. In the case of motile organisms this procedure may involve production of antibodies to flagella. During the production of acid extracts by the Lancefield method it is not known whether the serological reaction of the flagellar protein would be destroyed. The failure of trypsin to abolish the precipitin
reaction was inconclusive, since the flagellar protein of *Salmonella typhimurium* has been reported to be insensitive to trypsin and to other proteolytic enzymes (Stocker & Campbell, 1959). Cultures of strain P14/6 (type 29) and of strain NCTC 4725 were harvested and the flagella removed before preparation of acid extracts for typing. The resulting extract of P14/6 and a control extract of untreated organisms gave equally strong precipitin reactions with homologous antisera and with antiserum to NCTC 4725; in the case of strain NCTC 4725, an extract prepared after removal of flagella and a control extract also gave equally strong precipitin reactions. It is concluded that the antigens studied in acid extracts of these motile streptococci were not derived from the flagella.

*Electrophoresis of soluble components of disrupted organisms*

(i) *Esterases*

Preliminary experiments (Lund, 1965) showed the presence of esterase bands in extracts of motile strain P14/6. The relative intensity of these bands was affected by the presence of buffer during cell disruption. In experiments described below organisms were suspended in tris-citrate buffer before disruption.

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![Fig. 3](image1)

Fig. 3. Diagram of esterases of motile, type 29 strains of *S. faecium* separated by electrophoresis on polyacrylamide gel. (a) = strain P14/6 (*S. faecium*, type 29, type strain); (b) = strain N/2/23; (c) = strain 119.

![Fig. 4](image2)

Fig. 4. Diagram of esterases of a non-motile (type 38) strain of *S. faecium* and a motile (type 38) strain of streptococcus, studied by electrophoresis on polyacrylamide gel. (a) = strain P17/8 (non-motile); (b) = strain ATCC 13638 (motile).
Type 29 strains. P14/6, H/2/23, 119, all motile, showed similar esterase patterns (Fig. 3).

Type 38 strains. Esterase bands of the motile strain ATCC 13638, and non-motile type 38 strain P17/8 (type strain), are illustrated in Fig. 4. The motile strain showed strong esterase bands, the non-motile strain showed weak bands with esterase activity and with different mobilities from those of the motile strain.

'Type 4725' strains. Esterase patterns are shown in Figs 5 and 6, inserts (e) to (j). Strains reacting with antiserum to NCTC 4725 gave a range of different esterase patterns, only strain D1010 (c) gave a pattern identical to that of NCTC 4725.

Untyped strains gave esterase patterns shown in Fig. 5 and Pl. 1, fig. 1, inserts (a) to (d). Strains D1003 (b) and ATCC 12755 gave esterase bands at similar positions.

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Fig. 5. Diagram of esterases of a range of motile group D streptococci, studied by electrophoresis on polyacrylamide gel. Inserts (a)–(d) contained extracts of untyped strains; inserts (e)–(j) contained extracts of strains reacting with antiserum to NCTC 4725.

(a) = strain ATCC 12817; (b) = strain ATCC 12818; (c) = strain D1003 (b); (d) = strain ATCC 12755; (e) = strain NCTC 4725; (f) = strain D1010 (c); (g) = strain ATCC 12819; (h) = strain D1000 (b); (j) = strain D1006 (c).

(ii) Protein bands

Initial experiments showed that the pattern of protein bands from a motile strain (P14/i) differed from that of other strains of Streptococcus faecalis and S. faecium (Lund, 1965). The major differences were most clearly seen in the case of bands with $E_r$ values (Fowler, Coble, Cranmer & Brown, 1963) between 25 and 60 (approx.), the region indicated by the bracket in Pl. 1, fig. 2. No difference was found between the protein patterns of extracts prepared in water and those prepared in tris-citrate buffer.
Motile group D streptococci

Protein patterns of other motile strains are shown in Pl. 1, fig. 2. In general the motile strains resembled the original strain (P14/6) in showing five clear bands with $E_r$ values between 25 and 60. In one case (strain ATCC 12755) the forward band was relatively weak, in other cases there were slight differences in the position of these bands and some strains showed a sixth band in this region.

Type 38 strains. There was a clear difference between the protein patterns of motile and non-motile strains within this serotype; strain ATCC 13638 gave a pattern similar to that of the remaining motile strains, P17/8 gave a pattern resembling that of other non-motile strains of Streptococcus faecium (Lund, 1965).

The effect of presence of flagella. It did not seem likely that subunits of flagellar protein contributed to the protein pattern since flagella tended to be broken off and separated from the cells during harvesting and washing. Deliberate removal of flagella, by a method similar to that described in the section on serological methods, before preparation of extracts of strain P14/6 in the X-press, had no observable effect on the protein pattern obtained.

DISCUSSION

Although streptococci are generally considered to be non-motile, many workers have studied motile strains belonging to group D, isolated from a wide variety of sources (e.g. Pownall, 1935; Auerbach & Felsenfeld, 1948; Ødegaard & Gardborg, 1953; Liu, Lindberg & Mason, 1955; Hugh, 1959; Langston, Gutierrez & Bouma, 1960; Courtieu, Le Tellier, Guillermet, Imbert & Longeray, 1964; de Saint Aubert, Dubouclard, Roumiantszeff & Vaugon, 1964). Cowan & Steel (1965) stated ‘we confirmed Pownall’s (1935) observation that many group D streptococci are motile’. The most extensive studies of motile enterococci appear to have been made by Graudal [1952, 1955, 1957a, b, Graudal & Birch-Anderson, 1958] who examined 129 strains, mainly isolated from human faeces, and concluded (1957b) that motile organisms constituted a special subgroup distinct from Streptococcus glycerinaceus (S. faecalis), S. faecium and S. durans. He further subdivided the motile organisms into a yellow pigmented group and a non-pigmented group. (In the case of strains of the pigmented group studied in this present work, slight pigment formation was observed in colonies grown on horse blood agar at $30^\circ$ for several days.) Hugh (1959) studied six motile strains, including four of the ATCC strains used in this present work, and concluded that they were a variety of S. faecalis, without referring to the distinction between S. faecalis and S. faecium. He reported that the organisms fermented arabinose and not sorbitol, reactions generally characteristic of S. faecium.

In the present work several motile strains were found to produce acid when growing in the presence of sorbitol. The change in colour of the indicator was usually less marked than in the case of typical Streptococcus faecalis and possibly it would not be observed in slightly different reaction conditions. The results in Table 2 show that the motile strains differed in several respects from typical S. faecalis and showed a greater similarity to S. faecium in these tests.

Serology. Three of the motile strains had previously been described as Streptococcus faecium type 29 (Sharpe & Fewins, 1960; Barnes 1964). The remaining motile strains were tested against available antisera to 7 S. faecium types. Only one further strain gave a reaction (ATCC 13638, type 38); antisera were prepared to confirm these results, and antiserum to strain NCTC 4725 was prepared in order to elucidate
further the relationships between serotype and esterase pattern in this group of organisms.

In relating the serological studies of motile organisms to those of non-motile enterococci the nature of the antigens detected in motile bacteria by this typing technique should be considered. Elliott (1960) showed that type-specific antigens of three strains of *Streptococcus faecalis* and one of *S. durans* were located in the cell wall and were probably carbohydrate. The chemical nature of the carbohydrate type antigen of these and other strains of *S. faecalis* has been the subject of later publications (Bleiweis & Krause, 1965; Willers & Michel, 1966). Sharpe (1964) compared the serological type strains of *S. faecalis* from several workers and confirmed that the type antigen was derived from the cell wall. The inactivation of these antigens by periodate was taken as evidence of their carbohydrate nature. This also appeared to be true for the type antigens of many strains of *S. faecium* (Barnes, 1964), in most cases these antigens were destroyed by treatment with 0.01 M-potassium periodate. The antigens of motile streptococci studied by these techniques showed no evidence of inactivation by periodate, treatment of extracts with sodium periodate at a higher concentration and for a longer time than used by Sharpe (1964) or Barnes (1964) had very little effect on the subsequent precipitin reaction. Among the range of *S. faecium* strains studied by Barnes, several were reported to give type antigens resistant to periodate, these included strains of types 29 and 38, two of the three 'serotypes' studied in the present work.

The location of the antigen extracted from motile enterococci in these experiments has not been determined. Preliminary results indicate that it occurs at the cell surface since whole cells were used for immunization and for absorption of antisera. It is not flagellar since removal of flagella before acid extraction of strains P14/6 and NCTC 4725 gave no detectable decrease in precipitin reaction. An aim of the present work was to relate the study of esterase patterns to the most widely used method of serotyping the enterococci; no study has been made of the H antigens. The work of Graudal (1957a) seems to indicate a surprising lack of variation in flagellar antigens of motile, enterococci. Using agglutination reactions to study 129 strains he reported a great number of serological O-types, but 128 strains constituted one H-type and 1 strain another H-type.

**Esterases.** The finding that motile strains show diverse patterns of esterase enzymes indicates that this technique may be a useful aid to characterization of these strains. The results with strains of type 38 illustrate the difference which may occur between motile and non-motile strains within the same serotype, the former (ATCC 13638) having strong esterase bands, the latter (strain P17/8) having a different, much fainter esterase pattern. Motile strains within the same serotype may show different esterase patterns ('serotype 4725'); this contrasts with the results found for *Streptococcus faecalis* (Lund, 1965), where a similar esterase pattern was found in strains of *S. faecalis* and its varieties 'zymogenes' and 'liquefaciens', and in strains from five serotypes.

**Protein patterns.** The protein patterns, in contrast to those of the esterases, indicated some features which seem to be common to these motile strains. Despite the diverse esterase patterns observed it is possible that much of the enzyme protein of these organisms is very similar. The motile strain of serotype 38 showed a different protein pattern from the non-motile strain of this serotype.
Motile group D streptococci

Relationship of motile strains to Streptococcus faecalis and S. faecium. The protein patterns indicate that these motile strains form a group distinct from S. faecalis and S. faecium. The esterase patterns show a distinction from S. faecalis, strains of which give esterase patterns different from those of any of the motile strains, and from S. faecium, strains of which show much fainter esterase bands.

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REFERENCES


EXPLANATION OF PLATE
(Studies of soluble components of disintegrates of motile group D streptococci by electrophoresis on polyacrylamide gel.)

Fig. 1. Esterases of untyped motile streptococci (inserts a-d) and of strains reacting with antiserum to NCTC 4725 (inserts e-j). After electrophoresis the gel was incubated with α-naphthyl acetate and Fast Blue BB to detect esterase enzymes. Photograph of gel shown diagrammatically in Fig. 5 (approximately actual size).

Fig. 2. Protein bands of untyped motile streptococci (inserts a-d) and of strains reacting with antiserum to NCTC 4725 (inserts e-j). After electrophoresis the gel was stained with naphthalene black to detect protein bands. Inserts contained the same series of extracts as in Fig. 5 and Pl. 1, fig. 1 (approximately actual size).