The Cultivation of Sheep Rumen Oligotrich Protozoa in vitro

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SUMMARY: Oligotrich protozoa from the sheep rumen, principally Entodinium caudatum, were maintained in culture in vitro and dividing every 2 days at a density of 15,000–30,000 protozoa/ml for 18 months on a medium of rice starch, dried grass, 10% (v/v) rumen fluid (fresh or autoclaved) and 50 µg. chloramphenicol/ml. The effect on these organisms of varying each of these factors in turn and the use of different growth conditions is described. The protozoa have also been grown from an inoculum of 100–900/ml. to a density of over 50,000/ml. in 10 days.

Many unsuccessful attempts to cultivate rumen protozoa in vitro have been made in the last hundred years; the earlier attempts were summarized by Becker, Schultz & Emmerson (1929). Margolin (1930) and Westphal (1934) achieved some success in the cultivation of oligotrich protozoa, but the first real advance was made by Hungate (1942, 1943) who maintained some species of cattle oligotrich ciliates alive for over a year. Sugden (1953) was unable to maintain oligotrich protozoa from the sheep alive for more than 15 days, whereas Kandatsu & Takahashi (1955a, b, 1956) kept Entodinia spp. from goats alive for over 30 days with some multiplication. The purpose of the present paper is to describe experiments in which Entodinium spp., principally Entodinium caudatum from sheep rumen, have been maintained with frequent division for over 18 months; a preliminary communication has already appeared (Coleman, 1958).

METHODS

Media. The standard medium consisted of 30 ml. quantities of autoclaved mineral salts solution (Coleman, 1958) to which was added: Na₂S.9H₂O to a final concentration of 0.01% (w/v); c. 15 mg. dried grass (55° for 3 days) finely ground in a Lee Attrition Mill (Lee Engineering Co., Milwaukee, Wis., U.S.A.); 15 mg. rice starch, added as a suspension of known concentration in water; 3.0 ml. prepared fresh rumen fluid; chloramphenicol to a final concentration of 50 µg./ml. The final volume of complete medium was c. 35 ml. Tubes of complete medium were gassed vigorously for 3 min. with a mixture of 95% (v/v) N₂+5% (v/v) CO₂ (which had been freed from oxygen by the method of Stone & Beeson, 1936) and stoppered immediately. All media were warmed to 38° before use. Fresh rumen fluid was usually prepared and used the same day (see Coleman, 1958), except when it was inconvenient to obtain fresh rumen contents. At these times the prepared material was kept at –15° under 95% (v/v) N₂+5% (v/v) CO₂ for not longer than 4 days and warmed to 38°
before use. Autoclaved rumen fluid was prepared by autoclaving (10 lb./sq.in. for 20 min.) fresh rumen fluid in sealed McCartney bottles under 95% (v/v) N₂ + 5% (v/v) CO₂.

Cultural conditions. The procedures involved in the initial inoculation from fresh rumen contents and the subsequent manipulations that resulted in cultures of oligotrich protozoa which divided every 2–4 days have already been described (Coleman, 1958). The conditions used in the experiments described below differed initially in that the cultures were divided every 48 hr. regardless of the number of protozoa present. Later, maintenance was further simplified by omitting the preliminary centrifugation and replacement of supernatant fluid on days when the cultures were diluted with an equal volume of fresh medium (i.e. growth occurred under condition B—see below).

All experiments were carried out in 50 ml. centrifuge tubes containing c. 35 ml. medium, gassed with O₂-free 95% N₂ + 5% CO₂ gas mixture, sealed with a rubber bung and incubated at 38°. The time taken for all manipulations during which cultures had to be out of the incubator was kept to a minimum.

For the successful maintenance of the protozoa it was necessary to add every day fresh rice starch and grass with or without the other medium constituents. Cultures were diluted with an equal volume of fresh medium every 1–4 days, depending on the conditions, and the diluted cultures then divided each to make two identical cultures (this process is hereafter referred to as 'dilution' of a culture and the day on which it was carried out as a 'dilution day'). All the experiments described were carried out on one member of such a pair of tubes, the other serving as a control.

Four different growth conditions were used. The manipulations on days when the culture was diluted and on days when the volume was not changed, i.e. non-dilution days, were carried out as follows.

**Condition A.** Non-dilution day: culture centrifuged, supernatant fluid removed and replaced by an equal volume of fresh medium. Dilution day: culture centrifuged, supernatant fluid removed and replaced by twice its volume of fresh medium.

**Condition B.** Non-dilution day: culture centrifuged, supernatant fluid removed and replaced by an equal volume of fresh medium. Dilution day: culture diluted with an equal volume of fresh medium.

**Condition C.** Non-dilution day: rice starch (to same concentration as on dilution day) and 15 mg. dried grass added only. Dilution day: culture centrifuged, supernatant fluid removed and replaced by twice its volume of fresh medium containing rice starch and grass.

**Condition D.** Non-dilution day: rice starch and 15 mg. dried grass added only. Dilution day: culture diluted with equal volume of fresh medium containing rice starch and grass.

All centrifugation was at 500 g for 2 min. (this did not affect the motility and viability of the protozoa). The initial concentration of all medium constituents in fresh tubes of medium was adjusted so that at the end of the manipulation the concentration of the fresh constituents was that in the medium under test, except that on dilution days under conditions B and D,
the concentration of freshly added chloramphenicol and rumen fluid (when present) was half that added on non-dilution days. The frequency of dilution of the culture under any growth condition is given in days by the number after the letter, e.g. A2.

The number of protozoa was estimated by the method of Coleman (1958); unless otherwise stated the numbers quoted always refer to the numbers of protozoa present immediately before the dilution of a culture. Where the effect of a change in growth conditions is reported, at least 8 weeks was allowed to elapse before the number of protozoa present was estimated.

RESULTS

It was shown by Coleman (1958) that for the continued growth of rumen oligotrich protozoa it was necessary to add rice starch, dried grass, chloramphenicol and fresh rumen fluid to the basal mineral salts (i.e. standard medium) and that when the supernatant fluid was not replaced every 24 hr. by fresh complete medium the protozoa soon died. In the initial stages of this work, i.e. after 40 dilutions, when the cultures were diluted (condition A) every

Table 1. Steady state population densities of Entodinia spp. under various growth conditions

<table>
<thead>
<tr>
<th>Medium</th>
<th>Growth condition</th>
<th>No. of dilutions under that growth condition</th>
<th>Population density immediately before dilution (protozoa/ml.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. R, G, FRF, CAP</td>
<td>A2</td>
<td>40</td>
<td>28,000</td>
</tr>
<tr>
<td>2. R/2, G, FRF, CAP</td>
<td>A2</td>
<td>16</td>
<td>15,000</td>
</tr>
<tr>
<td>4. R, G, FRF, CAP</td>
<td>D2</td>
<td>53</td>
<td>14,000</td>
</tr>
<tr>
<td>5. R/2, G, FRF, CAP</td>
<td>D (dil. 3 days out of 4)</td>
<td>70</td>
<td>4000 before third successive dilution</td>
</tr>
<tr>
<td>6. R (120°, 24 hr.) G (autoclaved), FRF, CAP</td>
<td>B2</td>
<td>117</td>
<td>26,000</td>
</tr>
<tr>
<td>7. R/2, G, ARF, CAP</td>
<td>A2</td>
<td>17</td>
<td>16,000–26,000</td>
</tr>
<tr>
<td>8. R/2, G, ARF, CAP</td>
<td>C2</td>
<td>171</td>
<td>24,000</td>
</tr>
<tr>
<td>9. R/2, G, 1% ARF, CAP</td>
<td>C2</td>
<td>32</td>
<td>10,000</td>
</tr>
<tr>
<td>10. R/2 (120°, 24 hr.), G (autoclaved), ARF, CAP</td>
<td>C2</td>
<td>95</td>
<td>18,000</td>
</tr>
<tr>
<td>11. R, G</td>
<td>D3</td>
<td>53</td>
<td>14,000</td>
</tr>
</tbody>
</table>

2–4 days, the predominant organisms were morphologically similar to Entodinium caudatum and E. longinucleatum as described by Bhatia (1936), but variable numbers of Metadinium medium and E. simplex were also present. It is possible that, as was pointed out by Sugden (1953), the small E. simplex could not ingest rice starch grains and was living on the other medium constituents. Latterly, however, after 150 dilutions under condition B2, over 99% of the organisms were E. caudatum. All the experiments were carried
out with cultures containing over 90% *E. caudatum* and only *E. caudatum* and *E. longinucleatum* were included in the numbers reported. The detailed effect of varying in turn each material in the standard medium under condition A2 or B2 is shown below.

**Rice starch.** The optimum concentration of rice starch was 0.5 mg./ml. and with growth occurring under condition A2, 28,000 organisms/ml were present (Table 1). When the amount of starch was halved there were only 18,000 protozoa/ml.; doubling the starch concentration produced an initial increase in numbers which was followed by a steady decrease, probably associated with an increase in the number of bacteria. Omission of the rice starch resulted in death of the protozoa within 3 days. The rice starch was not replaceable by soluble starch or autoclaved rice starch, although in the presence of the latter a steadily decreasing number of organisms persisted for 46 days (100 protozoa/ml. after 15 dilutions).

The effect of heating the rice starch was investigated as it was hoped to be able to grow the protozoa in axenic culture by using sterile starch. When the dry starch was heated in an oven to 160° for 1 hr. before use, the protozoa soon died. Heating the starch at 140° for 1 hr. did not affect the protozoa when the numbers exceeded 15,000–20,000/ml. but resulted in their death when there were less than 5000–10,000/ml. The method eventually found to have least effect on the protozoa was to heat the dry starch in an oven at 120° for 24 hr.

**Dried grass.** Omission of the dried grass was followed by the death of the organisms within ten days when the culture was diluted normally; when the culture was not diluted small numbers of protozoa (i.e. 5000 protozoa/ml.) occasionally persisted for over 50 days. Solka Floc (Johnsen, Jørgensen & Wettre, Ltd., 26 Farringdon Street, London, E.C. 4) completely replaced the dried grass for 4 days, but thereafter the number of protozoa declined as in the absence of grass. The dried grass was not replaceable by 0.5 ml. of the supernatant fluid obtained after autoclaving 1 g. dried grass in 20 ml. water. The grass could be sterilized by autoclaving in the 80 ml. quantities of basal salts or in McCartney bottles after damping with water. The use of grass sterilized in either manner produced an initial depression in the growth rate of the protozoa, but this slowly improved to that of the original culture. After a 6-month period of adaptation cultures were obtained which grew almost as well on grass + rice starch sterilized as described above as on unsterile rice starch + grass (Table 1). The use of rice starch and dried grass dry-heated at 140° for 1 hr. before use produced erratic results with no or only slow growth, sometimes followed by death of the protozoa.

**Chloramphenicol.** The optimum concentration of chloramphenicol was 50 µg./ml.; an increase to 150 µg./ml. killed the protozoa. When the antibiotic was omitted, the organisms grew normally for 3–4 days but then, even when the culture was not diluted, the numbers decreased and the protozoa died or were present in small numbers. Under these conditions the bacterial growth was much heavier than in the presence of chloramphenicol and this may have been the cause of the failure of the protozoa to grow.
Rumen fluid. When the fresh rumen fluid was omitted, the protozoa grew normally for 3–4 days and then, in the absence of further dilution of the culture, the numbers steadily decreased. When fresh rumen fluid was replaced by an equal volume of autoclaved rumen fluid, the culture could still be maintained under condition A2 with 16,000–26,000 protozoa/ml. in the presence of only 0.25 mg. rice starch/ml. which was the optimum concentration under these conditions (Table 1). An increase in rice starch to 0.5 mg./ml. increased the bacterial growth without improving the protozoal growth and this heavier bacterial growth was occasionally associated with sudden death of the protozoa. Reproducible growth in the presence of 10% (v/v) autoclaved rumen fluid under condition A2 was difficult to attain and the cultures often contained many dead protozoa; with condition C2 consistent growth with 24,000 protozoa/ml. was obtained. Condition C2 was hereafter adopted for all experiments with autoclaved rumen fluid. Decrease of the autoclaved rumen fluid concentration from 10% (v/v) to 1% decreased the number of protozoa to 10,000/ml.

No systematic attempts to estimate the number or types of bacteria present were made, but when serial tenfold dilutions of the complete culture plus protozoa grown on 10% (v/v) autoclaved rumen fluid under condition C2 were made in the starch yeast extract tryptose medium of McPherson (1953) and the cultures incubated aerobically, over 10^6 bacteria/ml. were found.

In an attempt to determine whether it was the supernatant fluid or the bacterial bodies which was the essential part of the autoclaved rumen fluid, fresh rumen fluid was fractionated as follows. Fresh rumen fluid prepared in the normal way was centrifuged at 30,000 g for 50 min. and the supernatant fluid removed. The deposit (bacterial bodies fraction) was washed twice on the centrifuge with basal salts solution and made up with basal salts solution to the same volume as the original rumen fluid. The two fractions were autoclaved in sealed McCartney bottles under a gas mixture of 95% (v/v) N₂ + 5% (v/v) CO₂ and tested at a concentration of 10% (v/v) under condition C2. The autoclaved rumen fluid was replaceable by the supernatant fluid fraction for a period exceeding 80 days, although the number of organisms was 10–40% lower. The bacterial bodies fraction replaced the autoclaved rumen fluid completely for 15 days, but thereafter the number of protozoa decreased steadily until there were only 300 protozoa/ml. after 30 days.

In the presence of 10% (v/v) autoclaved rumen fluid the use of sterile grass and rice starch instead of nonsterile material decreased the number of protozoa to 18,000/ml. (cf. 24,000/ml.).

Rumen fluid and chloramphenicol. When 15 mg. rice starch + 15 mg. dried grass was added each day for a week to a standard culture after dilution, containing e.g. 16,000 protozoa/ml., without replacing the supernatant fluid, the number of protozoa increased to 36,000/ml. and could then be maintained by treatment under condition D3 (Table 1). Under these conditions the bacterial growth was much heavier than in the presence of fresh rumen fluid and chloramphenicol. Attempts to increase the dilution rate of the culture to every 2 days were unsuccessful as the culture eventually died out.
Temperature. The usual growth temperature was 38°; incubation at 41° resulted in the formation of distorted protozoa which died within 14 days, whether or not the culture was diluted. When incubated at 38°, the protozoa died within 4 days.

Basal mineral salts. Replacement of the potassium in the basal mineral salts solution by an equimolar concentration of sodium while growth was occurring under condition B2 produced a temporary 50% decrease in the number of protozoa, but after 30 days the number had returned to normal. However, when the basal mineral salts solution was replaced by the high sodium chloride medium of Sugden (1953), the numbers decreased by over 95% in 2 days and the culture died. To determine whether the Sugden salt solution would support the protozoa, a fresh protozoal suspension from the rumen (prepared by the method of Coleman, 1958) was inoculated into Sugden salts with sulphide + rice starch + dried grass + fresh rumen fluid + chloramphenicol, using the general cultural methods described above and treated under condition A without dilution. Some protozoa were still alive after 5 weeks (when the experiment was discontinued) as compared with none after 15 days in Sugden's experiments.

Attempts to stimulate growth under condition A2

The addition of 1 mg. Difco yeast extract/ml. produced an initial increase in the number of protozoa, but this was followed by heavy bacterial growth and death of the protozoa: 0.01 mg. yeast extract/ml. had no obvious effect. In view of the work of Kandatsu & Takahashi (1956) who found that vitamin B12 stimulated the growth of Entodinia spp. from goat rumen, this compound was tested at 0.3 μg./ml. but had no effect on the growth of the protozoa. A mixture of β-alanine, p-aminobenzoic acid, biotin (at 0.001 μg./ml.), calcium pantothenate, folic acid, nicotinamide, pyridoxin, riboflavin and thiamine (each at 0.01 μg./ml.), tested with or without 0.3 μg. vitamin B12/ml. had no effect on the growth rate. The addition of maize starch, potato starch or wholemeal flour at 0.25 mg./ml. increased the small number of Metadiniun medium present but had no stimulatory effect on the growth of Entodinium caudatum.

Further attempts to stimulate growth were made by increasing the dilution rate of the cultures. When the culture was diluted 2 days out of three under condition A, there were 5000–10,000 protozoa/ml. on the second dilution day; only 4000/ml. were present when the culture was diluted 3 days out of 4 under condition D, but these numbers were constant over 90 days.

Attempts to facilitate the maintenance of protozoa

Initial experiments showed that to maintain the protozoa it was necessary to replace the supernatant fluid every 24 hr. by fresh complete medium. Later, the addition of grass and rice starch each day was found to maintain the organisms for periods of at least a week and that a culture on standard medium under condition A2 could be diluted without the preliminary centrifugation and replacement of supernatant fluid, i.e. changed to condition B2, with
slight improvement in the growth rate. Further simplification in the procedure by changing to condition D2 produced a fall in the number of protozoa from 32,000/ml. to 14,000/ml. When no additions were made to the medium on non-dilution days under condition D2, the protozoa grew normally for 4 days but were dead after 8 days.

**Growth from a small inoculum**

In the experiments described above the minimum number of protozoa was usually greater than 10,000/ml. and it was of interest to determine whether or not the protozoa would survive and multiply when less than 1000 protozoa/ml. were present. Figure 1 shows that when 900 protozoa/ml. were inoculated (as
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85\% (Fig. 1). Similar results, but with slower growth rate and lower maximum population density, were obtained by inoculation of standard medium or standard medium without rumen fluid and chloramphenicol followed by treatment under condition C without dilution (Fig. 1). Cultures were regularly maintained under these three conditions by inoculating fresh medium each week with a 1.5\% (v/v) inoculum grown under the same conditions as that to be used for the inoculated tube; the average number of protozoa present on the seventh day is shown in Table 2. After maintenance for 25 weeks under either condition without daily replacement of supernatant fluid, 50–80\% of the protozoa were *Entodinium longinucleatum*; most of the remainder were *E. caudatum*.

The slower growth under condition C without dilution compared with condition A without dilution is probably associated with the heavier bacterial growth under the former condition.

Table 2. *Growth of Entodinia spp. from a small inoculum*

<table>
<thead>
<tr>
<th>Culture</th>
<th>No. of weekly transfers</th>
<th>1 week average number of protozoa/ml.</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>40</td>
<td>38,000</td>
</tr>
<tr>
<td>2</td>
<td>41</td>
<td>18,000</td>
</tr>
<tr>
<td>3</td>
<td>27</td>
<td>10,000</td>
</tr>
</tbody>
</table>

**DISCUSSION**

As far as the author is aware, the present studies report the first successful attempts to maintain and grow *Entodinia* spp. from sheep rumen *in vitro*. The medium differed from that used unsuccessfully by Sugden (1953) in that:

(a) the mineral salt solution was well buffered and rich in potassium instead of sodium; (b) the starch and dried grass were not heated at 140° for 1 hr. before addition to the medium. The successful result of inoculating fresh protozoa from the rumen into Sugden’s salt solution suggests that, although the experimental conditions were not exactly comparable with those of Sugden, the salts plus rice starch, grass, rumen fluid and chloramphenicol would maintain the protozoa. It seems not improbable that the reason for Sugden’s failure was the use of heated starch and hay. Heating the rice starch and grass at 140° for 1 hr. produced a slight charring of both reagents and caused the dry powdery starch to form lumps. It is possible that the products of this charring reaction were either toxic to the protozoa or their essential bacteria or that they encouraged the growth of unfavourable bacteria. This latter possibility is supported by the observation that when heated rice starch and grass are
added in the standard medium, the contents of the tube form a thick slimy deposit in 2 or 3 days.

The success of the present studies may be also attributed to the presence of chloramphenicol in the medium. This antibiotic partially suppressed the growth of the bacteria in the fresh rumen fluid but allowed some bacterial growth. When the fresh rumen fluid was replaced by autoclaved rumen fluid bacteria were still present, and it is likely that these were chloramphenicol resistant. In view of the finding of Gutierrez & Davis (1959) that Entodinia spp. ingest *Streptococcus bovis* and possibly require that bacterium for growth, it is possible that the chloramphenicol in the present experiments suppressed bacteria harmful to the protozoa while allowing favourable bacteria to survive. Heavy bacterial growth was always associated with slower protozoal growth and decrease in the yield of organisms/ml.

Experiments are now in progress to improve the maintenance of the protozoa and to grow the organisms in axenic culture.

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REFERENCES


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