The Life Cycle and Growth Kinetics of an Anaerobic Rumen Fungus

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The life cycle of an anaerobic fungus isolated from the rumen of sheep was studied and was found to be similar to that of the chytrids. The organism was monocentric. At 39 °C the duration of the life cycle varied from about 26 to 32 h. The zoosporangium increased in length exponentially with a doubling time of 2·49 h.
Between 6·5 to 9·5 h after inoculation, the rate of extension of the main rhizoid declined, and no further extension occurred after 9·5 h. The main rhizoid increased in width at its base from 2·2 to 15·0 μm during the first 13 h after inoculation, indicating that intercalary wall growth occurred. Nuclei were occasionally observed in the rhizomycelium using DAPI (4,6-diamidino-2-phenylindole) staining. Zoosporangia varied in shape from spherical to columnar, and some columnar zoosporangia were observed to become spherical. The zoosporangium initially increased in volume at an exponential rate with a doubling time of 1·56 h. Between 14 to 20 h after inoculation, growth of the zoosporangium decelerated and little growth occurred after 20 h: the zoosporangium had a final volume of 2·5 × 10⁵ μm³. At about 21 h after inoculation, a septum was formed at the base of the zoosporangium, delimiting it from the rhizoidal system. The formation of this septum was correlated with the cessation of zoosporangial growth and the onset of zoosporogenesis. After zoosporogenesis, zoospores (about 88 zoospores per zoosporangium) were liberated through a pore formed in the zoosporangial wall opposite the main rhizoid. About 3 h after zoospore release the rhizoidal system became less refractile, suggesting that autolysis had occurred. Growth of the isolate was inhibited by nystatin (an inhibitor of chitin synthase), but not by amphotericin B or nystatin (antibiotics which bind to sterols).

INTRODUCTION

In 1912 Weisenberg described Callimastix cyclopis as a unicellular, polyflagellate protozoan parasite of Cyclops strenuus. Braune (1913) discovered a similar organism in the rumen of sheep and cattle which he named Callimastix frontalis. The suggestion of Weisenberg (1950) that C. cyclopis might be a zoospore of an aquatic fungus was borne out when the ultrastructure of C. cyclopis was studied by Vavra & Joyon (1966) and when a plasmodial phase was identified in the body cavity of infected hosts; these workers concluded that C. cyclopis was related to the chytrid fungi and therefore the genus Callimastix with C. cyclopis as type species, was placed in the Blastocladiales. However, since the other species of Callimastix were still thought to be flagellate

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Abbreviation: DAPI, 4,6-diamidino-2-phenylindole.
protozoa they were placed in a new genus, *Neocallimastix*, with *Neocallimastix frontalis* (formerly *C. frontalis*) as the type species.

In 1975, Orpin reported that the life cycle of *N. frontalis* consisted of two phases, a non-motile reproductive stage and a motile flagellate stage. The discovery of a non-motile stage of *N. frontalis* led to the suggestion that the 'flagellates' were zoospores of an aquatic fungus. However, unlike zoospores of other aquatic fungi, which have one or two flagella per zoospore (Sparrow, 1960), zoospores of *N. frontalis* have up to 14 flagella per zoospore. Nevertheless, *N. frontalis* had a life cycle which was very similar to that of the chytrids.

Bauchop & Mountfort (1981) isolated an anaerobic rumen fungus (PN1) which was similar to *N. frontalis*. After studies of its life cycle by light and electron microscopy, PN1 was officially described as *N. frontalis* (Heath et al., 1983). As the existing taxonomy of chytrids excluded organisms with polyflagellate zoospores (Sparrow, 1960) and because the organism was an anaerobe, Heath et al. (1983) placed *N. frontalis* in a new family, the Neocallimasticae, which was assigned to the order Spizellomycetales, in the class Chytridiomycetes. It was subsequently found that PN1 was not the only organism as the *N. frontalis* isolated by Orpin (1975) and therefore the latter organism was re-named *N. patriciarum* (Orpin & Munn, 1986).

In this paper we describe the life cycle of an anaerobic rumen fungus similar to *N. frontalis* and *N. patriciarum*, which was isolated from sheep. Development of the organism was studied in liquid and agar media and the life cycle of the present isolate is compared with *N. frontalis*, *N. patriciarum* and other chytrid fungi. The life cycle of the organism is also considered in relation to the ecological niche which it may occupy in the rumen.

**METHODS**

**Organism.** The anaerobic fungus studied is a member of the genus *Neocallimastix* (Heath et al., 1983) which was isolated from sheep at the Animal and Grassland Research Institute, Hurley. The methods of fungal enrichment and isolation and a brief description of the fungal culture were the subject of a previous publication (Lowe et al., 1985). Since the original communication, the culture has been maintained on straw in a defined, liquid medium as described below. In order to ensure that the culture consisted of only a single fungus, culture supernatant, which contained zoospores, was serially diluted and used to inoculate roll tubes (Hugnet, 1969) of molten defined medium B agar (modified from Lowe et al., 1985; see below). The tubes were incubated at 39°C for 2 days and, after examination with a Leitz Diavert inverted microscope, a tube containing a few colonies (each of which had developed from a single zoospore) was selected, and an individual colony was picked and transferred (using a sterile dissection needle) to an anaerobic culture tube (18 x 142 mm, Belco Glass) containing 10 ml liquid defined medium B plus 25 mM-glucose. All subsequent cultures were derived from this isolate, which has been designated R1. We have decided not to name the organism at the present time because direct comparisons with other members of the genus have not been made and the evidence presented in this paper and in Lowe et al. (1987b) suggests that R1 may be a new species.

**Culture conditions.** The anaerobic techniques used were as described by Hugnet (1969), Bryant (1972) and Miller & Wolin (1974). Defined medium B is the medium B described by Lowe et al. (1985), except that coenzyme M solution, Trypticase peptone, yeast extract and antibiotics were omitted. Unless stated otherwise, the medium contained 25 mM-glucose.

Stock cultures were maintained on 0.1 g wheat-straw (milled to pass through a 1 mm dry mesh screen) in 10 ml defined medium B and were inoculated with 1 ml culture supernatant (containing zoospores); subcultures were made every 7 days. Experimental cultures were inoculated with 1 ml culture supernatant from cultures grown for 5 days on defined medium B; these were in the late-exponential phase of growth and haemocytometer counts showed that they contained about 1 x 10⁸ zoospores ml⁻¹. All cultures were transferred using anaerobic pipetting techniques and, unless stated otherwise, they were incubated at 39°C under a gas phase of 100% CO₂ without agitation. The light microscope was used routinely to ensure that cultures were free from bacterial contamination.

**Morphological observations.** Development of the fungus was followed in agar medium in roll tubes, on coverslips or membrane filters in liquid medium, and on cellophane strips in liquid medium. Roll tubes were prepared using defined medium B and 18% (w/v) agar. To ensure that only a few colonies developed in each culture, tubes containing 3 ml molten defined medium B agar were inoculated with 1 ml culture supernatant which had been diluted with defined medium B until it contained approximately 10 zoospores ml⁻¹.

Cellophane (British Cellophane) strips (3.5 x 5 cm) were washed by boiling in distilled water for 10 min and placed around the inside of anaerobic tubes (one strip per tube). After autoclaving, 10 ml defined medium B lacking glucose was added and each tube was inoculated with 1 ml zoospore suspension; cellophane was the carbon source in these cultures.
Development of the isolate in roll tubes and in 'cellophane' cultures was observed using an inverted microscope housed in a perspex incubator maintained at 37 to 39°C. Cultures were incubated on the microscope stage to allow continuous microscopic observation and photographs were taken using FP4 ASA 125 black and white film (Ilford).

Development of the isolate was also followed on glass coverslips and on membrane filters (0-22 μm pore size, Membrical Membrane Filters; Gelman Sciences) added to liquid defined medium B in anaerobic tubes. The use of coverslips to observe aquatic fungi has been described by Fuller (1962) and it proved a useful technique in studying the R1 isolate, which encysted on glass and adhered to it during subsequent growth. Circular (12 mm diameter) glass coverslips (Chance Propper) were cleaned with alcohol and autoclaved in anaerobic culture tubes, prior to the addition of 10 ml defined medium B. The tubes were inoculated with 1 ml undiluted or diluted culture supernatant; dilutions were made using defined medium B lacking glucose. Growth was stopped by addition of 1 ml 25% (v/v) glutaraldehyde solution. After 10 min at room temperature, the coverslips were mounted in 0-2 M-phosphate buffer (pH 7-2) and examined using a Reichert microscope. All microscopic measurements were made using a travelling micrometer eye piece (C. Baker).

Coverslip cultures were also used to examine the effects of antibiotics on growth. Nikkomycin (kindly supplied by Professor G. Gooday), amphotericin B (dissolved in 35% (w/v) sodium deoxycholate) or nystatin were included in the medium prior to inoculation, or were added to growing cultures 12 h after inoculation. An appropriate concentration of sodium deoxycholate was included in the medium of cultures used as controls for the amphotericin B treatments.

**DAPI staining.** Nuclei fluoresce when stained with the fluorochrome DAPI (4',6-diamidino-2-phenylindole) and illuminated with UV light (Hooley et al., 1982). The stain solution contained 0-3 g DAPI (Sigma) per litre of 0-05 M-Tris/HCl buffer (pH 7-2), containing 100 mM-NaCl and 10 mM-EDTA. Coverslips removed from growing cultures were placed on a slide with a drop of the stain and left at room temperature in the dark for 10 min before being examined. A Zeiss transmitted-light photomicroscope III fitted with an exciter filter (G 365), a barrier filter (LP 418) and Neofluar objectives were used to observe the nuclei. Photographs were taken using FP4 ASA 1000 film.

**Cryo-scanning electron microscopy (cryo-SEM).** Cultures growing on membrane filters were used. Frozen, hydrated material was prepared for cryo-SEM in a similar manner to that described by Webb & Jackson (1986). Membranes (cut to 3 mm square) were removed from the culture medium, rinsed quickly in distilled water and blotted almost dry on filter paper. These were then attached to the specimen stub with carbon cement, frozen rapidly in nitrogen slush and transferred, under vacuum, to the cooled prechamber of a Hxland CT1000 cryotransfer system which was interphased to a Philips 305 scanning electron microscope. Excess water was sublimed away by heating to about -90°C and the specimen was cooled to about -160°C and sputter-coated with gold. Coated specimens were viewed by SEM at about -180°C.

**RESULTS**

**The zoospores**

The uninucleate zoospores were usually spherical (Fig. 1a) to ovoid and measured 10.71 ± 1.79 μm × 9.64 ± 1.88 μm (mean ± standard error, n = 30). Between 8 and 17 flagella were present on each zoospore and each flagellum was several times longer than the zoospore. During motility, the flagella united to form a single locomotory organelle. Zoospores moved in an erratic manner, sometimes swimming in a circular motion or moving with a rapid darting motion, interrupted by frequent pauses and changes in direction. Zoospores were occasionally observed to move in an amoeboid manner and during these periods the flagella sometimes separated from one another and spherical bodies were occasionally observed on their tips (Fig. 1b). Some zoospores remained motile for a number of hours but others encysted within a matter of minutes of their release from zoosporangia. Encystment of a zoospore was rapidly followed by the formation of a main (germination) rhizoid. Flagella remained attached to the zoospore after encystment but occasionally young plants (rhizoids plus immature zoosporangium) were observed which lacked flagella.

**Rhizoid and zoosporangium development**

Studies were made of the development of a single plant in a roll tube (Figs 2, 3 and 4a) and of a population of plants on coverslips in liquid culture (Figs 3 and 4b). At the outset the encysted zoospore in the roll tube culture was 12.6 μm long and 10.4 μm wide (Fig. 4a). During the first 6-5 h after inoculation there was a rapid development of an extensive rhizoidal system (Figs 2
and 3); during this period the main rhizoid increased in length exponentially with a doubling time of 2·49 h (specific growth rate = 0·28 h⁻¹). Between 6·5 to 9·5 h after inoculation, there was a deceleration in the rate of extension of the main rhizoid, and no further extension occurred after 9·5 h, at which time the rhizoid was 734 μm long. Growth of the main rhizoid provided a convenient indicator of the growth of the whole rhizoidal system (Fig. 2a–d).

The zoosporangium initially increased in volume at an exponential rate (Fig. 4a) with a doubling time of 1·56 h (specific growth rate = 0·44 h⁻¹). Between 14 to 20 h after inoculation, growth of the zoosporangium decelerated and little growth occurred after 20 h. Cessation of zoosporangial growth was correlated with the formation of a septum at the base of the zoosporangium; observations of a number of plants showed that septum formation occurred 21·3 ± 1·00 h after inoculation. The zoosporangium in Fig. 4(a) had a final volume of 2·5 × 10⁶ μm³.

Zoospores were liberated from the zoosporangium 27 h after inoculation, but instead of swimming away, they encysted in the agar medium at the site of the parent zoosporangium (Fig. 2e). Each encysted zoospore rapidly developed its own rhizoidal system and these rhizoids grew radially away from the central area of developing plants (Fig. 2f). Extension of rhizoids of the second generation of plants continued, and 3 h after the zoospores had been released, the rhizoidal system of the parent plant became less refractile. Development of the second generation of zoospores into mature plants was followed by the liberation of a third generation of zoospores and these swam away from the parent zoosporangia to encyst and germinate at the outer edge of the region of rhizoidal growth.

Development of the R1 isolate on coverstips in liquid cultures was also studied. Growth of zoosporangia which developed on coverstips in liquid culture was similar to that observed for the zoosporangium which developed in the agar roll tube (Fig. 4a, b). The most significant change in size of the population of zoosporangia occurred 12 to 26 h after inoculation during the latter part of the life cycle. During this period the developing zoosporangia became 'ovoid' and 26 h after inoculation they had a mean size of 103·3 ± 5·70 × 81·5 ± 6·01 μm (n = 18) (Fig. 4b). These mean measurements suggest that the zoosporangia were all ovoid in shape. However, the population contained both spherical (120 × 120 μm) and columnar (140 × 62 μm) zoosporangia. Direct observations of growing cultures showed that the shape of a zoosporangium sometimes changed during its development. Most zoosporangia were initially spherical, but subsequent development sometimes resulted in the formation of columnar zoosporangia. Some columnar zoosporangia were also observed to change in shape, becoming spherical. When this happened the length of the zoosporangium remained approximately constant but its width increased. Cryo-scanning electron micrographs of individual plants of the R1 isolate, which were observed
Fig. 2. Development of a single plant of R1 in a roll tube (a) 1·5 h, (b) 5·0 h, (c) 9·0 h and (d) 24 h after inoculation, and (e) 0·5 h and (f) 2·0 h after the onset of liberation of zoospores from the zoosporangium. Zoospores were liberated 27 h after inoculation. Bars, 100 μm.
on membrane filters after 48 h incubation in defined medium B, are shown in Fig. 5. The main rhizoid and range of shapes of the zoosporangia were typical morphological characteristics of the R1 isolate (Fig. 5).

Although the length of the main rhizoid of the population of plants developing on coverslips was not determined, the width of the rhizoid immediately beneath the base of the zoosporangium was measured (Fig. 3). Four hours after inoculation the main rhizoid had a mean width of $2.2 \pm 0.11 \mu m$ ($n = 18$). Subsequent growth resulted in an increase in width and 14 h after inoculation the main rhizoid had a mean width of $15.0 \pm 0.61 \mu m$ ($n = 18$). No further increase occurred during the rest of the life cycle.

Young zoosporangia initially contained one nucleus per zoosporangium (Fig. 6a, b), but mature zoosporangia contained very many nuclei (Fig. 6c, d). The presence of DAPI-staining bodies (presumably nuclei) within the main rhizoid was observed on a number of occasions and similar DAPI-staining structures were also observed in rhizoids distant from the zoosporangium.

The number of plants formed on coverslips recovered from liquid medium was determined to give an estimate of the length of the life cycle and the number of zoospores formed per
zoosporangium (Fig. 7). In cultures with a high concentration of zoospores the number of plants formed per coverslip increased steadily with time. Two hours after inoculation a mean of 48 plants were attached to each coverslip, and after 26 h this had increased to 413 plants. By 29 h after inoculation there were too many plants present to count. These observations suggest that the mean duration of the life cycle under these conditions was between 26 and 29 h.

Cultures inoculated with a dilute zoospore suspension had counts of 5 to 16 plants per coverslip during the first 29 h after inoculation (Fig. 7). The number of plants per coverslip did not increase appreciably until 32 h after inoculation, by which time 37 plants were present per coverslip. At 47 h after inoculation the mean number of plants per coverslip (782) indicated that zoospore release had occurred. From these data, it was calculated that about 88 zoospores were released from each zoosporangium. Fig. 7 suggests that the duration of the life cycle may be influenced by zoospore concentration and that low concentrations of zoospores may develop more synchronously than high concentrations of zoospores.

Liberation of the zoospores

Liberation of zoospores from a zoosporangium formed on cellophane in liquid medium is shown in Fig. 8. The release of zoospores from zoosporangia has been observed on several occasions and the position of the exit point was always the same, viz. a pore in the zoosporangial wall formed opposite the main rhizoid. Some gyratory movement of the zoospores occurred
Fig. 6. (a) An immature zoosporangium of R1; (b) the same zoosporangium under fluorescence illumination showing a single DAPI-staining nucleus. (c) A mature zoosporangium of R1; (d) the same zoosporangium under fluorescence illumination showing numerous DAPI-staining nuclei. Bars, 50 μm.

Fig. 7. Increase in the number of plants of R1 on coverslips in liquid cultures. Coverslip-containing anaerobic tubes were inoculated with either 1 ml culture supernatant (△), or 1 ml culture supernatant diluted 1:49 (▲). Five replicate cultures were sampled at each time interval. The error bars represent SE of the mean.
within the zoosporangium prior to their release, and this motion increased until eventually the zoospores were liberated. At this time some zoospores were abruptly shot out of the zoosporangium (Fig. 8b) and these began swimming in the medium. The remaining zoospores in the zoosporangium swam out through the pore and 3.5 min after the onset of dispersal, the zoosporangium contained only a few zoospores (Fig. 8f). Once empty, the 'ghost' zoosporangium remained and eventually the rhizoidal system autolysed.
Fig. 9. Effect of nikkomycin on the development of zoospores and immature plants of R1. Nikkomycin was either present at the time of inoculation with zoospores (△), or was added to cultures (containing immature plants) 12 h after inoculation (▲). The numbers of zoospores and immature plants which developed in the presence of nikkomycin were recorded 13 h and 30 h after inoculation respectively. The results are expressed as a percentage of the number of zoospores and immature plants which developed in cultures lacking antibiotics.

Effect of antibiotics

The effect of nikkomycin, amphotericin B and nystatin on development of zoospores and immature plants was examined. Nikkomycin is an inhibitor of chitin synthase (Muller et al., 1981) and was used to assess the importance of chitin synthase in development of the R1 isolate. Amphotericin B and nystatin bind to membrane sterols and inhibit the growth of many fungi (Medoff & Kobayashi, 1980). The effect of nikkomycin on growth of R1 is shown in Fig. 9. At 0·1 μg ml⁻¹ (final concentration) the development of immature plants was not affected by nikkomycin; however, at this concentration of antibiotic, only 65% of the zoospores developed into plants. Zoospores and immature plants incubated in the presence of 3 μg ml⁻¹ and 5 μg ml⁻¹ of nikkomycin respectively, did not develop into mature plants. Amphotericin B and nystatin were included in defined medium B at concentrations up to 10 μg ml⁻¹ but had no effect on the growth and development of zoospores and immature plants of R1.

DISCUSSION

The morphology and life cycle of the R1 isolate resembled that of eucarpic chytrids, and consisted of a reproductive body, the zoosporangium, supported by a filamentous, vegetative structure of branching, tapering rhizoids. The formation of a single zoosporangium is indicative of monocentric chytrids. The morphology of the R1 isolate resembles that of the chytrid *Chytridium*. In *Chytridium* the zoospore encysts and produces a main rhizoid which penetrates the substratum. The rhizoids continue to elongate and transport nutrients back to the encysted zoospore. Development of *Chytridium* is described as epibiotic because the zoosporangium is present on the surface of the substratum and the rhizoidal system grows within the plant material. During the later stages of development the zoosporangium becomes separated from the rhizoidal system by the formation of a septum at its base (Sparrow, 1960).

According to Waterhouse (1962) "the swimming motion of zoospores may be extremely jerky and erratic, and interspersed with periods of dormancy". The zoospores of the R1 isolate have been observed to behave in a similar manner, and during the pauses some zoospores underwent
amoeboid changes in shape. This type of behaviour is characteristic of chytrid zoospores (Sparrow, 1960) and may be a prelude to encystment.

Observations of the life cycle of the R1 isolate showed that, after zoospore encystment, there was a period of rapid development of the rhizoidal system, with only a small increase in the size of the developing zoosporangium. This type of growth has also been observed by Bauchop (1979a, b, 1980, 1981) in his studies on fungal colonization of feed material in the rumen. He observed extensive development of the rhizoidal system during the first 12 h of growth of the fungi on material in the rumen, and then the zoosporangium increased in size until the life cycle was completed. Septum formation in the R1 isolate occurred about 21 h after inoculation, towards the latter stages of the life cycle. Heath et al. (1983) also found that the onset of zoosporogenesis was correlated with septation. No pore was observed in the septum, which therefore formed a barrier to cytoplasmic flow from the rhizoidal system to the zoosporangium. After zoospore release, the zoosporangium and the rhizoidal system autolysed; this phenomenon has been reported for other chytrids (Sparrow, 1960).

The observation of DAPI-staining bodies (nuclei) within the rhizoids was unexpected. It is generally accepted that members of the Chytridiales produce a limited rhizoidal system, and the rhizoids possess cytoplasm but no nuclei. During development of the main rhizoid, intercalary wall growth occurred at the point where it joined the zoosporangium. It is not known if growth of the zoosporangium is autocatalytic or, as is more likely, supported at least in part by growth in the rhizoidal system. The observation that the zoosporangium had a faster specific growth rate (0.44 h⁻¹) than the main rhizoid (0.28 h⁻¹) supports the latter hypothesis.

Zoosporangia of the R1 isolate varied in shape from spherical to columnar. Similar observations have been made by Orpin (1975) and Orpin & Munn (1986) in their studies of *N. patriciarum*. Variation in the shape and size of zoosporangia of other chytrids also occurs (Sparrow, 1960) and this is particularly true of *Endochytrium operculatum* (Karling, 1937), which forms both spherical and pyriform zoosporangia. Chytrids either form a single pore through which zoospores are discharged or several pores; the manner of zoospore release is characteristic for a particular species and all the zoospores released develop and release their zoospores in the same manner (Sparrow, 1960). Release of zoospores by the R1 isolate always occurred from a point in the zoosporangium wall opposite the main rhizoid. In this respect the R1 isolate differed from *N. frontalis* (Heath et al., 1983) in which zoospore release occurred by 'nonlocalized dissolution of the entire zoosporangium wall'. Examination of photographs of zoospore release from a zoosporangium of *N. patriciarum* (Orpin, 1975) suggests that zoospores of this fungus were released from a number of pores in the zoosporangial wall. These differences in the methods of zoospore release between R1, *N. frontalis* and *N. patriciarum* suggest that they are different species.

Movement of zoospores within the zoosporangium prior to their release may act as a trigger for zoospore discharge (Sparrow, 1960). Our observations of zoospores leaving zoosporangia are typical of descriptions given by Sparrow for other chytrids in which the first zoospores to leave are ejected by force and the remainder leave the zoosporangium either by swimming or by amoeboid movement. The zoospores of the R1 isolate sometimes swam for several hours prior to encystment; Munn et al. (1981) have detected glycogen in zoospores of *N. patriciarum* and this could supply the source of energy needed for prolonged periods of swimming and for subsequent encystment and germination.

On average, about 88 zoospores were released per zoosporangium, but small zoosporangia have been observed to release fewer zoospores. Observations by Joblin (1981) on zoospore release from rumen fungi showed that between 10 and 120 viable zoospores were liberated per zoosporangium. With *N. patriciarum* between 2 and 38 zoospores were formed within zoosporangia (Orpin, 1975). Autolysis of the rhizoidal system after zoospore release suggests that each plant has a finite life span.

The effect of nikkomycin on growth of R1 suggests that chitin synthase is an important enzyme in this organism and that its walls contain chitin. Orpin (1977) has reported the presence of chitin in *N. frontalis*, *Pirononas communis* and *Sphaeromonas communis*, and Brownlee (1986) has recently demonstrated the presence of chitin synthase in anaerobic fungi. Analysis of *N.*
*frontalis* (Body & Bauchop, 1985) suggested that 12% of its lipids consisted of sterols. However, Kemp et al. (1984) failed to detect sterols in *P. communis*. Oxygen is essential for sterol synthesis (Goldfine & Bloch, 1963), but anaerobic organisms may obtain these compounds from their environment. However, sterols do not seem to be essential for the growth of R1, as nystatin and amphotericin B, antibiotics that bind to membrane sterols and cause cell leakage, did not affect growth.

Rhizoids facilitate the attachment of rumen fungi to plant material, and such attachment would increase their residence time in the rumen. The main function of rhizoids is to absorb nutrients from the substrate, and their ability to penetrate and degrade plant tissues (Bauchop, 1980; Lowe et al., 1987b) may allow these fungi to reach growth substrates not accessible to bacteria or protozoa. The production of extracellular cellulases and xylanase by rumen fungi (Mountfort & Asher, 1985; Pearce & Bauchop, 1985; Lowe et al., 1987a) enables them to utilize cellulose and hemicellulose as substrates for growth.

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