**Dictyostelium dimigraformum,**

*Dictyostelium laterosorum* and *Acytostelium ellipticum:*

New Acrasieae from the American Tropics

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**SUMMARY**

Two new species of *Dictyostelium*, *D. dimigraformum* and *D. laterosorum,* and a new species of *Acytostelium*, *A. ellipticum,* are described. All three species were first isolated from the surface humus layer of tropical forest soils on the island of Trinidad, W.I. *D. dimigraformum* is distinguished from other species of the genus by its ability to form both stalkless and stalked migrating pseudoplasmodia. *D. laterosorum* is a member of the crampon-based Dictyostelia. It differs from other members of this group in bearing lateral, sessile sori along the terminal half of the sorophore. *A. ellipticum* is the second species in the genus *Acytostelium* to be described. It is distinguished from *A. leptosomum* by its elliptical spores, pattern of aggregation and development, and smaller size.

**INTRODUCTION**

As a part of my continuing investigation of forest soils for Acrasieae, I surveyed, during the summer of 1968, tropical forest soils in Puerto Rico, Trinidad and Tobago, West Indies, and Guyana, South America. Previous investigations indicated that a centre of diversity for Acrasieae exists in the tropics of Central and South America (Cavender & Raper, 1968; Cavender, 1969), as reflected in the several new Acrasieae which have been isolated from this area: *Dictyostelium rosarium, D. deminutivum, D. rhizopodium, D. lavandulum, D. coeruleo-stipes, D. vinaceo-fuscum and D. mucoroides* var. *stoloniferum.* Distribution studies have focused on the lowland seasonal evergreen forest which is the optimum forest habitat for Acrasieae in the tropics (Cavender & Raper, 1968).

On the island of Puerto Rico little such forest remains, consequently most collecting was done in the lower montane forest type prevalent in the Loquillo Experimental Forest. Only seven species and variants were isolated (Table 1). Two isolates, one resembling *Polysphondylium violaceum* in form but lacking pigment, and the other a member of the *Dictyostelium mucoroides* complex which produces exceptionally numerous sorocarps, are worthy of further study but have not yet been investigated. Possibly the less complex nature of the forests of Puerto Rico compared with those of the South American continent limits the number of Acrasieae species that survive there, or perhaps the island is simply too remote from the continent.

Floristically Trinidad resembles north-eastern South America, a consequence of long union with the continent. The forest vegetation has been studied extensively by J. S. Beard (e.g. 1946). Most of the forests sampled are seasonal evergreen; annual
rainfall 200 to 250 cm. with a dry season from January to May. Mora and crappoguatarea (Carapa–Eschweilera) are the most widespread forest types. I collected from eight sites in the northern part of the island in the vicinity of, and on the lower slopes of, the Northern Range where elevations reach 500 m. Trinidad is an excellent area for Acrasieae. Fourteen species were isolated (Table 1), including the three species described here, which represent approximately 75% of the known soil Acrasieae. The adjacent island of Tobago supports fewer species of plants because of its greater isolation and smaller size. Less extensive collecting was done there. Only six species were found.

Table 1. Species of Acrasieae isolated from tropical forest soils of Puerto Rico, Trinidad, Tobago and Guyana

<table>
<thead>
<tr>
<th>Species</th>
<th>Puerto Rico</th>
<th>Trinidad</th>
<th>Tobago</th>
<th>Guyana</th>
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<tr>
<td>Dictyostelium mucoroides</td>
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<td>D. mucoroides variant I²</td>
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<td>D. mucoroides variant II³</td>
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<td>D. purpureum</td>
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<td>D. polycephalum</td>
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<td>D. rhizopodiurn</td>
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<td>D. vinaceo-fuscum</td>
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<td>D. aureum³</td>
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<td>D. minutum³</td>
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<td>D. laterosorurn</td>
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<td>D. dimigraformum</td>
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<td>Polysphondylium pallidum</td>
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<td>P. pallidum variantI³</td>
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<td>P. violaceum</td>
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<tr>
<td>Acytostelium leptosomum</td>
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<tr>
<td>A. ellipticum</td>
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1 A delicate, hyaline-spored acrasian common in the tropics.
2 Similar to the above but producing exceptionally numerous sorocarps.
3 Golden-yellow sorocarps similar to D. aureum (E. Olive).
4 In the tropics there are a variety of small hyaline-spored acrasians that resemble D. minutum.
5 Unpigmented but resembling P. violaceum in form.

A limited area in Guyana was sampled—the forests located west of Bartica on the Essequibo River and along Moraballi Creek which are on either red or white sand. They are either mixed forests or consociations with mora, greenheart, morabukea and wallaba as the principal dominants (Davis & Richards, 1934). Ten species of Acrasieae were isolated (Table 1).

Two of the three Acrasieae described are considered new species of the genus Dictyostelium because they have unbranched cellular stalks. The third slime mould has an acellular stalk, consequently it is assigned to the genus Acytostelium.

METHODS

Soils were collected in small glass vials which hold about 30 g. Leaf mould, humus and some surface mineral soil were taken since these parts of the forest soil profile harbour the greatest number and kinds of Acrasieae (Cavender & Raper, 1965b). When I returned to the laboratory the samples were placed at 4°. The isolating technique of Cavender & Raper (1965a) was used. Developing Acrasieae were transferred
to dilute hay infusion agar media cross-streaked with *Escherichia coli* by touching a sorus with a needle and implanting the spores at the intersection of the bacterial streaks (see Raper, 1951). Further studies were carried out either by using this method of cultivation or by ‘seeding’ the surface of a dilute nutrient agar medium with a suspension of bacteria and spores which was then spread uniformly over the entire agar surface or in a broad band. This is a more favourable method for studying spore germination, aggregation and pseudoplasmodium formation (Raper, 1951). Other agar media used in this study, in addition to the hay infusion, were low nutrient media containing either lactose or dextrose and peptone (LP, DP) in varying amounts and non-nutrient agar supplemented with a pre-grown bacterial food supply.

The amount of growth per day of the slime mould and production of typical fruiting bodies were used as a basis for determining optimum conditions for cultivation. Temperature studies were carried out at 16, 20, 22, 25 and 30°. Five different bacteria, *Escherichia coli*, *Aerobacter aerogenes*, *Pseudomonas fluorescens*, *Serratia marcescens* and *Sarcina lutea* were tested as food for the Acrasieae using two types of media: (1) non-nutrient agar streaked with pre-grown bacterial samples, and (2) LP or DP agar media streaked with bacterial suspensions. Spores were implanted at the ends of the streaks. Most cellular slime moulds respond best to either *E. coli* or *A. aerogenes* and this was found to be true for the Acrasieae described here.

**Dictyostelium dimigraformum** Cavender, sp. nov.

Sorocarpi erecti, vel proni, non ramosi, magnitudine proportioneque varii; in nigritie pseudoplasmodia vaga, typice sine caulibus, in uniducto lumine phototropica, cum caulibus vel sine caulibus; sorophora incolora usque ad leviter flava, usque ad plerumque 3 ad 8 mm. in longitudinem; sori subglobosi usque ad citroformes, plerumque 200 ad 400 μm. in diametro, lactei usque ad citrini: spori elliptici usque ad reniformes, magnitudine maxime varii, plerumque 7·0 ad 12·0 x 2·5 ad 3·5 μm. sed interdum usque 26 x 5 μm.

Habitat: In foliari humo et summo solo, tropica humida silva, Trinidad, W.I.

Typica cultura: AR-5b.

Sorocarps erect or inclined, unbranched, variable in size and proportions, in the dark pseudoplasmodia wandering, typically without producing a stalk, in one-sided light phototropic, migration occurring with or without stalk production; sorophores unpigmented to yellowish, mostly 3 to 8 mm. in length; sori subglobose to citroform, mostly 200 to 400 μm. in diameter, white to lemon-yellow; spores elliptical to reniform, extremely variable in size, mostly 7·0 to 12·0 x 2·5 to 2·5 μm. but sometimes as large as 26 x 5 μm.

Habitat: In leaf mould and surface soil, tropical moist forest, Trinidad, W.I.

Type culture: AR-5b

**Isolation and Cultivation**

The only isolate of *Dictyostelium dimigraformum* was obtained from the surface soil of a forest reserve in Trinidad located just above the Asa Wright Nature Centre on the south slope of the Northern Range along the road between Arima and Blanchisseuse. One clone of the organism appeared on the isolation plate.

*Dictyostelium dimigraformum* was characterized by robust growth but less so than *D. discoideum*, *D. purpureum* or some of the larger strains of *D. mucoroides*. It grew

well on a variety of media of low nutrient content. Those most often used were 0.1% LP and thin hay infusion agar (15 g. of dried, leached bluegrass per litre). Optimum growth on these media occurred with *Escherichia coli* although good growth was obtained with *Aerobacter aerogenes* and *Serratia marcescens*. The pigment of *S. marcescens*, prodigiosin, was not digested by the myxamoebae, consequently the slugs and sorocarps were pink to red in colour. Optimum temperature for growth and development was between 20 and 25°C. At 16°C growth and development were slow but normal, although much slime was left behind by the migrating slugs. No growth occurred at 30°C but growth and differentiation proceeded normally at 28°C.

**Growth and morphogenesis**

*Spore germination.* Germination took place in 4 to 5 h. when spores were spread on 0.1% LP medium. The germination process was similar to that described for *Dictyostelium discoideum* by Cotter & Raper (1968). The swelling spore (Pl. 1, fig. 1) in some cases approached spherical proportions before pressure from within caused the cellulose wall to split and allowed the myxamoeba to emerge (Pl. 1, fig. 2). The slime surrounding spores removed from a sorus dried very quickly making them difficult to spread on the agar surface without leaving clumps. Some such clumps of several hundred or more spores germinated *en masse* (Pl. 1, fig. 6). Aggregation and culmination followed without an intervening growth phase. However, this refruiting habit did not occur to the extent described for *D. mucoroides* var. *stoloniferum* (Cavender & Raper, 1968). An interesting characteristic found in *D. dimigraformum* was the range in spore size. Spores are elliptical in shape, occasionally reniform. Most spores are 7 to 12 × 2.5 to 3.5 μm. with a mean of 8.0 × 3.0 μm. Spores 15 to 18 μm in length were not uncommon (Pl. 1, fig. 4); larger spores were rare but one measured 26 × 5 μm. (Pl. 1, fig. 5). Spores of different size ranges are known for *D. discoideum* and for the crampon-based Dictyostelia (Raper & Fennell, 1967), but this is the first report of such extreme variation in spore size in Acrasiaceae. Sussman & Sussman (1962) found a relationship between ploidy and spore size in *D. discoideum*.

*Vegetative growth.* The myxamoebae appeared similar to those of *Dictyostelium discoideum* or *D. mucoroides* (Pl. 1, fig. 3). Variation in size was apparent from spore dimensions. Shape changed during the life-cycle: while feeding, myxamoebae were circular in outline with diameters averaging about 11 μm.; while moving, they elongated to about 20 × 10 μm. This elongation was even more exaggerated when the myxamoebae entered an aggregation stream.

*Aggregation.* In a culture prepared by inoculating the surface of 0.1% LP medium with a mixture of *Escherichia coli* and slime mould spores, aggregation began after available bacteria had been consumed. At 25°C this period was about 40 h. for *Dictyostelium dimigraformum*. Centres were spaced at an average density of 6/cm² on dark-grown plates. In the light the density was greater. Almost 100% of the myxamoebae entered the aggregates. Streams appeared before actual centres could be detected and were often flat and sheet-like at this stage. Wheel-like aggregations soon developed, each with a definite centre or hub and well-defined streams or spokes (Pl. 1, fig. 7). A later stage had fewer but larger streams (Pl. 2, fig. 9). Usually one pseudoplasmodium or slug was formed from each centre. Where myxamoebae were very dense and large centres were formed, several slugs might emerge.

*Pseudoplasmodia.* When a culture plate was illuminated from above, sorocarps were
constructed vertically from the point of aggregation (Pl. 2, fig. 14). If grown with one-sided illumination, some slugs moved away from the point of aggregation and a prostrate stalk was formed as the slug migrated (Pl. 1, fig. 8). The occasional production of a stalkless slug (Pl. 2, fig. 11) distinguished this species from all those with stalk-forming slugs. The stalkless-slug left behind as a trail only the collapsed slime case that is continually secreted. This slug, identical in appearance and behaviour to its counterpart in the life-cycle of Dictyostelium discoideum, was especially abundant on 0.1% LP medium and much less abundant on buffered hay infusion agar. In absolute darkness most slugs were stalkless. They might wander over the agar surface for a week or more before sorocarp formation occurred, but sorocarp formation could be initiated dramatically by exposing the culture to overhead light.* Slugs then stopped moving and within 1 to 2 h. became orientated on a vertical axis, and somewhat flattened to a pear shape (Pl. 2, fig. 12) as the internally formed stalk made contact with the agar surface (Pl. 2, fig. 10). As the stalk lengthened, the mass of cells left the substratum, perhaps partially through the individual efforts of the myxamoebae but principally through the force exerted by the lengthening stalk (Raper & Fennell, 1952).

A dark-grown slug might form a length of stalk and then cease stalk formation while continuing to migrate. The environmental factors which ‘trigger’ stalk formation in the dark are not known. Sorocarps that were formed were generally on the sides of the Petri dish, indicating that a decreasing relative humidity might act as a trigger as it does for other acrasians such as Dictyostelium discoideum.

Migrating slugs produced well-defined slime tracks often (especially at temperatures below 20°) leaving behind clumps of cells which occasionally formed small slugs or fruiting bodies (Pl. 1, fig. 8).

Sorocarps. The formation of sorocarps was not studied cytologically but appeared to follow closely the pattern described for Dictyostelium discoideum by Raper & Fennell (1952) except that a basal disc was not formed. Sorocarps varied greatly in size and orientation depending upon the density of myxamoebae and the availability and direction of light. Commonly, when produced on LP medium after a period of migration, they resembled the upright portions of sorocarps of D. purpureum or larger members of the D. mucoroides complex. The unbranched, tapering, upright stalk, 3 to 8 mm. high, usually had a short supporting horizontal portion surrounded by slime (Pl. 2, fig. 13). The terminal sori were globular or citriform, their diameters (usually proportional to the stalk length) most commonly 200 to 400 μm. The sori often developed an intense lemon-yellow colour derived from the pigmentation of the spores, but this did not always develop, and a gradation in colour from cream-white to lemon-yellow was found in most cultures.

Dictyostelium laterosorum Cavender, sp.nov.

Sorocarpi typice erecti vel proni, interdum procidui, non saepe ramosi, magnitudine proportioneque variis; sorophora caesia usque leviter violacea, saepe subbruna, plerumque 5 ad 10 mm. in longitudinem sed ex 1 mm. usque 20 mm. varia, sorophorum basi exiliter digitatae et instar claviculae; sori globosi, caesii usque leviter violacei, saepe subbruni, et ad laterales et terminales in positionibus, laterales sori plerumque minores, sessiles, 80 ad 150 μm. in diametro, 1 ad 10 in numero, terminales sori

* Light induction of sorocarp formation was recently discovered for D. discoideum by Newell, Telser & Sussman (1969).
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130 ad 200 μm. in diametro; spori vari, elliptici, reniformes vel recurvi, 6.0 ad 13.0 × 2.5 ad 4.0 μm.

Habitat: In foliari humo et summo solo, tropica humida silva, Trinidad et Tobago, W.I.; Colombia, S.A.

Typica cultura: TB11-1

Sorocarps typically erect or inclined, sometimes prostrate, infrequently branched, variable in size and proportions, sorophores bluish grey to light violet, often with a brownish cast, usually 5 to 10 mm. in length but varying from 1 to 20 mm.; sorophore bases weakly digitate and crampon-like; sori globose, bluish grey to light violet often with a brownish cast, both lateral and terminal in position, lateral sori smaller, sessile 80 to 150 μm. in diameter, 1 to 10 in number, terminal sori 130 to 200 μm. in diameter; spores variable in shape and size, elliptical, reniform or recurved, 6.0 to 13.0 μm. × 2.5 to 4.0 μm.

Habitat: Leaf mould and surface soil, tropical moist forest, Trinidad and Tobago, W.I.; Colombia, S.A.

Type culture: TB11-1

Isolation and cultivation

Initially two isolations of this species were made, strain TB11-1 from semi-deciduous forest on the north slope of the Main Ridge, Tobago, along the road from Roxborough to Charloville, and strain AE-4 from seasonal evergreen forest near Arena, Trinidad. Three other isolations have subsequently been made from Colombia. Excellent growth and development of Dictyostelium laterosorum was obtained at 22 to 25° on 0.1% LP or hay infusion agar media in association with Escherichia coli or on non-nutrient agar streaked with pre-grown bacteria.

Growth and morphogenesis

Spores and myxamoebae. Spores gave rise to myxamoebae after 4 to 5 h. at 25° on LP medium. A large percentage failed to germinate. Spores of the two strains TB11-1 and AE-4 were originally different in shape and size; strain TB11-1 produced predominantly elliptical spores 6 to 9 × 2.5 to 4 μm. (Pl. 3, fig. 17); strain AE-4 produced larger spores, 9 to 13 × 3.5 to 4.5 μm., mostly reniform or recurved in shape (Pl. 3, fig. 16). After about eight months of laboratory cultivation diploidization apparently occurred. Strain TB11-1 began producing spores similar in size and shape to those of AE-4 (Pl. 3, fig. 18); other differences between the two strains remained constant.

The myxamoebae were distinguishable from those of Dictyostelium dimigraformum by their elongated triangular shape while moving (Pl. 3, fig. 19), the relatively broad front advancing, the tapering apex trailing behind. This posterior apex might be as long as the body of the myxamoeba, so narrow that it was just visible when magnified × 600, and occasionally it was forked. Up to five peripheral nucleoli were conspicuous in the nucleus. Feeding myxamoebae were circular in outline, their diameters mostly 10 to 15 μm.

On LP medium aggregation began after the bacteria had been consumed by the myxamoebal population. At 25° this period was c. 42 h. for strain AE-4 and c. 45 h. for strain TB11-1. The two strains differed somewhat in their patterns of aggregation. Myxamoebae of strain AE-4 tended to develop strong wheel-like centres without delay (Pl. 3, fig. 23), whereas those of strain TB11-1 did not form definite centres immediately
New Acrasieae

but first formed small clumps or mounds (Pl. 3, fig. 20) which gradually increased in size (Pl. 3, fig. 21). Some of these early aggregations developed short streams. After 72 h. certain centres had become dominant and attracted myxamoebae from most of the early aggregations (Pl. 3, fig. 22), though the radiate pattern exhibited by AE-4 did not often develop (Pl. 4, fig. 24) and the streams often broke up into numerous smaller centres, especially when aggregation occurred in the light.

*Pseudoplasmodia and sorocarp formation.* Dictyostelium laterosorum typically produced pseudoplasmodia that constructed stalks vertically or at an angle from the aggregation centre although the somewhat larger Colombian strains were prone to wander over the agar surface. The phototropic response was not particularly strong although one-sided illumination induced migration toward the light. Light is not essential for completion of the life-cycle.

The rising sorogen resembled that of *Dictyostelium mucoroides* during the early stage, but as the terminal half of the stalk was formed masses of myxamoebae were periodically abstricted from the posterior portion (Pl. 4, fig. 28). These myxamoebae differentiated into spore cells without concomitant formation of lateral stalks as in *Polysphondylium* (Pl. 4, fig. 26). The result was a series of sessile globular sori which gave the sorocarp a beaded appearance (Pl. 4, fig. 29). This method of increasing the efficiency of spore distribution was first discovered in the slime mould *Dictyostelium rosarium* (Raper & Cavender, 1968). *D. laterosorum* closely resembled this species in the nature and disposition of the sori. Occasionally a lateral branch bearing a sorus at the tip (Pl. 4, fig. 30) was seen. The masses of abstricted myxamoebae occasionally retained the capacity to construct a stalk. Sorocarps may have up to ten or more lateral sori, each 80 to 150 μm. in diameter, though the terminal sorus is slightly larger (130 to 200 μm.).

The pigmentation of the sorocarps approximated that of *Dictyostelium lavandulum* (Raper & Fennell, 1967), i.e. bluish grey deepening with age to light violet or lavender, often with a brownish cast.

The sorocarps were supported by a crampon base enveloped by a layer of supporting slime material (Pl. 4, fig. 27). The ramifications of the crampon, much less developed than in *Dictyostelium rhizopodium*, were short cellular extensions, often knob-like, protruding from the expanded base of the stalk (Pl. 4, fig. 25). Of other Acrasieae these crampons resembled most closely the expanded base type of *D. coeruleo-stipes* (Raper & Fennell, 1967).

Sorocarps have a broad size range. Lengths of mature stalks varied from 1 to 20 mm. but most were 5 to 10 mm. Strain AE-4 produced more large, rangy sorocarps with a greater proportion of stalk to spore mass than strain TB11-1. Some of the Colombian strains also had this habit but produced more lateral sori than AE-4.

**Acytostelium ellipticum** Cavender, sp.nov.

Sorocarpi maxime delicati, solitarii vel gregarii, 200 ad 1000 μm. in altitudinem; sorophora incolora, acellularia, 1 ad 2 μm. in diametro attenuata usque < 10 μm. in extrema parte; surgens sorogen ventricosa-rostrata; sori globosi, 18 ad 40 μm. in diametro; spori elliptici, 5–5 ad 8–0 μm. × 2–0 ad 3–0 μm.

Habitat: In foliari humo et summo solo, tropica humida silva, Trinidad, W.I.; Guyana, Colombia, S.A.

Typica cultura: AE-2
Sorocarps extremely delicate, solitary or gregarious, 200 to 1000 μm. in height; sorophores unpigmented, acellular, 1 to 2 μm. in diameter tapering to less than 1 μm. at the tip; rising sorogen ventricose-rostrate; sori globose, 18 to 40 μm. in diameter; spores elliptical, 5-5 to 8-0 μm × 2-0 to 3-0 μm.

Habitat: Leaf mould and surface soil, tropical moist forest, Trinidad, W.I.; Guyanas, Colombia, S.A.

Type culture: AE-2

Isolation and cultivation

*Acytostelium ellipticum* was isolated from the Trinidad Government Forest Reserves at Melajo and Arena, from the lowland forest along Moraballi Creek in Guyana, and more recently from the Amazon Basin of Colombia. The isolates were very small and delicate, barely visible on the isolation plates at ×30 magnification, and so considerably smaller than those of *A. leptosomum* on the same isolation plates. Cultivation was originally attempted on 0.1% LP medium streaked with *Escherichia coli*. Germination and growth occurred with no further morphogenesis. Weak hay infusion medium (8 g. dried leached bluegrass/l.) with *E. coli* gave good growth and development. After several months cultivation the slime mould was able to complete its life-cycle on 0.1% LP, but optimum growth and development occurred on a weak glucose-peptone medium (0-05% glucose and 0-025% peptone) at 20 to 25° when a suspension of *E. coli* and slime mould spores was spread in a broad band on the agar. *E. coli* and *Aerobacter aerogenes* were the best food sources. No development took place at 30°.

Growth and development

The elliptical spores measured 5.5 to 8.0 × 2.0 to 3.0 μm. (Pl. 5, fig. 31). After germination the myxamoebae formed a densely packed feeding front (Pl. 5, fig. 32). On DP medium inoculated with a suspension of spores and *Escherichia coli* the bacteria were consumed after 40 h. at 25°. The myxamoebae then clumped together (Pl. 5, fig. 32) in increasing numbers without streaming, to form small mounds (Pl. 5, fig. 35). Crude streams were observed only when a culture was cooled to 4° or on a medium of higher nutrient content (0-25% DP) (Pl. 5, fig. 34) on which subsequent development was abnormal. The aggregation pattern therefore differs from *Acytostelium leptosomum* (Pl. 5, fig. 40), where radiate streaming occurs. Each mound-like aggregation produces one to several sorocarps (Pl. 5, fig. 38) depending upon its size: the highly gregarious fruiting of *A. leptosomum* (Pl. 5, fig. 40) does not occur. The developing sorogen is long and tapered (Pl. 5, fig. 33), and as the stalk lengthens it develops a bulbous posterior portion (Pl. 5, fig. 36); in contrast, the developing sorogens of *A. leptosomum* are naviculate (Pl. 5, fig. 40). The cells in the narrow anterior portion of the *A. ellipticum* sorogen probably secrete the acellular stalk tube: Raper & Quinlan (1958) found that in the sorogen of *A. leptosomum* the myxamoebae in the anterior end were arranged perpendicular to the stalk, indicating a secretory rather than a locomotive function. Following aggregation, secretion of the entire stalk (200 to 1000 μm. in length) took 2 to 3 h. Its delicate nature in respect to the mass of spores which it supported (Pl. 5, fig. 39) was an engineering wonder. A globular unpigmented sorus (18 to 40 μm. across) developed at the terminus of the stalk (Pl. 5, fig. 37). The average dimensions of both stalk length and sorus were less than for *A. leptosomum*. 
DISCUSSION

Of the three new Acrasieae described here, perhaps *Dictyostelium dimigriformum* is of greatest interest because it possesses characteristics intermediate between those of *D. mucoroides* and *D. discoideum*. Its similarity to *D. discoideum* is most apparent in the migrating pseudoplasmodial phase of development when stalk formation does not occur. This is a more efficient means of migration than we find in *D. mucoroides* since cells are not needlessly expended. My observations concerning the response of the pseudoplasmodia to light essentially duplicate those of Newell et al. (1969) working with *D. discoideum*. The myxamoebae show a physiological similarity to those of *D. discoideum* since they are unable to digest the pigment prodigiosin. Myxamoebae of *D. mucoroides* digest this pigment. The sorocarps themselves as well as the stalked migrating pseudoplasmodia resemble those produced by *D. mucoroides* or *D. purpureum*. The organism provides good experimental material for the study of environmental factors which initiate stalk formation. The 17-fold variation in size of the spores of *D. dimigriformum* is of interest, for it has been argued that genetic heterogeneity brought about through para-sexual mechanisms is basic to cell differentiation in the Acrasieae (Ashworth & Sackin, 1969). The possible role which these large cells play in development is worthy of investigation.

*Dictyostelium laterosorum* has the beaded sorocarps of *D. rosarium* but differs from this species in pigmentation, spore shape, crampon base, and developmental pattern (see Raper & Cavender, 1968). *D. laterosorum* also resembles *D. lavandulum* in pigmentation of the sorocarps and the presence of a crampon base. The crampon-based Acrasieae are tropical in distribution. Hopefully, *Coenonia*, a crampon-based genus which is more highly differentiated than any cellular slime mould now in culture, will someday turn up in tropical soils.

*Acytostelium ellipticum* is one of the smallest Acrasieae. The sorocarps are more delicate than those of *Dictyostelium deminutivum* but the myxamoebae are larger. The pattern of aggregation and sorocarp distribution is closer to this species than to *A. leptosomum*.

Strains AR-5b, TB11-1 and AE-2 have been deposited with the American Type Culture Collection, Washington, D.C.

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REFERENCES


EXPLANATION OF PLATES

**PLATE 1**

*Dictyostelium dimigraformum*

Fig. 1. A germinating spore just before the splitting of the spore wall. ×1140.

Fig. 2. The same germinating spore. A myxamoeba is emerging from the split spore case. ×1140.

Fig. 3. A moving myxamoeba. ×1140.

Fig. 4. Spores, including a large one 17 μm. in length. ×1140.

Fig. 5. A very large spore, 26 μm. in length. ×900.

Fig. 6. A pseudoplasmodium forming from a clump of spores which have germinated *en masse*. Some of the myxamoebae are wandering away. ×70.

Fig. 7. A typical wheel-like aggregation. ×29.

Fig. 8. A migrating pseudoplasmodium which is producing a stalk. Note the partially dried sorogen which developed from myxamoebae cast off in the slime track of a stalkless migrating slug. ×90.

**PLATE 2**

*Dictyostelium dimigraformum*

Fig. 9. An aggregation at a slightly later stage in development from that in Pl. 1, fig. 7. ×35.

Fig. 10. A pseudoplasmodium which has stopped migration after exposure to light and which is producing a stalk, visible in the interior. ×58.

Fig. 11. A stalkless migrating pseudoplasmodium or slug. ×58.

Fig. 12. A pseudoplasmodium which has ceased migration and become orientated almost vertically. The pre-spore mass is about to be lifted off the agar surface. ×100.

Fig. 13. A mature sorocarp produced from a stalkless migrating pseudoplasmodium. Note the short horizontal portion surrounded by slime. ×44.

Fig. 14. A small rising sorogen. The stalk is composed of a single row of cells. ×35.

Fig. 15. Mature sorocarps, which have developed without migration, collapsed on the agar surface. ×5.
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PLATE 3

Dictyostelium laterosorum

Fig. 16. Spores of strain AE-4. Note the granules at each end. ×1140.
Fig. 17. Spores of strain TB11-1 before size change. ×1140.
Fig. 18. Spores of strain TB11-1 after size change. ×1140.
Fig. 19. The central myxamoeba is moving toward the lower right. Note the triangular shape and long tail. ×450.
Fig. 20. TB11-1. An early stage in aggregation. The cells have gathered into small clumps. ×340.
Fig. 21. TB11-1. A slightly later stage. The clumps have increased in size. ×290.
Fig. 22. TB11-1. An intermediate stage in aggregation. Dominant centres are forming which attract the myxamoebae from surrounding clumps. ×115.
Fig. 23. Wheel-like aggregates typical of strain AE-4. ×29.

PLATE 4

Dictyostelium laterosorum

Fig. 24. TB11-1. A late stage in aggregation. Strong centres have developed which attract myxamoebae from a relatively large area. ×9.
Fig. 25. A typical crampon-like base which supports the sorocarp. ×450.
Fig. 26. Terminal portion of the sorocarp showing the disposition of the sori. ×46.
Fig. 27. The expanded disc of slime material which surrounds the base of the stalk. ×115.
Fig. 28. A developing sorogen which has just abstricted a mass of myxamoebae from its posterior portion. The myxamoebae will differentiate into spores forming together a lateral sorus. ×53.
Fig. 29. Mature sorocarps showing the typical beaded appearance. ×23.
Fig. 30. A sorocarp on which a lateral branch has developed. ×23.

PLATE 5

Acytostelium ellipticum

Fig. 31. Spores showing the characteristic shape for the species. ×1140.
Fig. 32. Myxamoebae densely packed in a feeding front at the left. At the right aggregation is beginning to occur. ×290.
Fig. 33. Characteristic shape for the species of a young rising sorogen. ×230.
Fig. 34. Rudimentary stream formation by strain AE-2 on 0·25% dextrose-0·25% peptone agar. Subsequent development was abnormal. ×29.
Fig. 35. Typical aggregates formed on 0·25% dextrose-0·05% peptone agar. ×29.
Fig. 36. Rising sorogens. At this stage the sorogen has a relatively long narrow anterior portion and a more bulbous posterior. ×92.
Fig. 37. Sorocarps at various stages of development. ×46.
Fig. 38. Aggregates from which 1 to 3 sorocarps are developing. ×92.
Fig. 39. Terminal portion of a mature sorocarp collapsed on the agar surface. Note the dimension of the stalk tube in relation to the spore mass it supported. ×290.
Fig. 40. A typical aggregate of A. leptosorum showing stream formation, numerous clustered sorocarps and naviculate sorogens. ×23.