
**Chaetomium brasiliensis** Batista & Pontual; Nutritional Requirements for Growth and Fruiting

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**SUMMARY:** *Chaetomium brasiliensis* requires aneurin for growth, and calcium (strontium or barium), aneurin and a low nutrient level for the formation of ascospores. Decreasing suboptimal doses of calcium progressively decrease the number of fertile perithecia until none is formed, but not the total number of perithecia. Above the optimum level, calcium delays formation of perithecia and ascospores and, ultimately, spore germination; it has no favourable effect on vegetative growth. The fungus appears able to synthesize aneurin slowly after growth is complete.

Lilly & Barnett (1949) found that *Chaetomium convolutum* needs external supplies of both aneurin and biotin for growth and for ascospore formation. Apart from this, no essential vitamin requirements of *Chaetomium* species are known, although *C. cochliodes* fruits better in the presence of aneurin (Hawker, 1942) and that *C. globosum* fruits more quickly with added biotin (Buston & Basu, 1948). *C. globosum* and several other *Chaetomium* species, however, require calcium for the formation of mature perithecia and ascospores (Basu, 1951). In this paper it is shown that the recently discovered *C. brasiliensis* needs calcium for fruiting, and aneurin for both growth and fruiting.

**MATERIALS AND METHODS**

The test organism (*IJMARI 117*) was undescribed when first isolated by us from rotting jute fabric. According to the Commonwealth Mycological Institute, Kew, to which a subculture was submitted, it is identical with the fungus described by Batista & Pontual (1948) as *C. brasiliensis* and by Ames (1949) as *C. velutinum*. A subculture of the latter was kindly sent us by Dr Ames and we found it to be morphologically indistinguishable from our strain.

The organism was grown on malt agar slopes at 30° and stored in the refrigerator. The basal medium used was Czapek-Dox salt solution at pH 7·0 and contained: NaNO₃ 2 g.; K₂HPO₄ 0·75 g.; KH₂PO₄ 0·25 g.; KCl 0·5 g.; MgSO₄.7H₂O 0·5 g.; water, 1 l. The source of carbon was 0·2% glucose (w/v), unless otherwise mentioned. The medium was first prepared with half the required quantity of water and 5 ml. placed in 75 ml. American-made Pyrex conical flasks, which were cleaned before each test with hot chromic-sulphuric acid. After addition of supplements, the volume of medium in each flask was made to 10 ml. with distilled water. The flasks were sterilized at 5 lb./sq.in. for 10 min. Aneurin (as hydrochloride) was added before steriliza-
tion as a freshly prepared solution. Calcium was added as CaCl₂. Flasks were inoculated with a loopful of spore suspension and incubated at 30°.

As the organism produced numerous small, densely aggregated perithecia, counting perithecia was not possible. This was of little consequence because, unlike C. globosum and the other species previously examined, the number of perithecia, when these were formed, was always of the same order; differences were noticed in fertility and size of perithecia and in their terminal hairs. Whenever practicable cultures were duplicated.

RESULTS

Effect of aneurin on growth

On malt extract medium or on basal medium plus jute extract the fungus grew and fruited well, forming a deep smoky-black mycelium closely covered with blue-green perithecia having spirally curved terminal hairs; on basal medium alone growth did not proceed much beyond spore germination (Pl. 1, fig. 1), indicating a nutritional deficiency. Aneurin, biotin, riboflavin, pyridoxin, nicotinic acid and Ca pantothenate were added at concentrations of 0·1, 0·01, 0·001 μg./flask. The fungus responded only to aneurin, growth proceeding to completion with 0·01 μg. but not with 0·001 μg. aneurin/flask. At the lower concentration some sugar remained, the mycelium was white and no perithecia were formed; at the higher concentration sugar was exhausted, the mycelium turned black and its surface was extensively covered with greyish green perithecia which, however, showed more or less straight hairs, did not form either asci or ascospores and may be described as infertile. This state of growth (see Pl. 1, figs. 2 and 3), was not altered with increased doses of aneurin or combination of aneurin with any of the other substances mentioned above (at 0·2 μg./flask). A nutritionally adequate medium gives the same type of vegetative growth, but the perithecia, in addition to being fertile and covered with spirally curved hairs (Pl. 1, fig. 4), are darker and bigger.

The response of vegetative growth to increasing doses of aneurin was tested on basal medium which contained 3% (w/v) glucose and 0·2 M-phosphate buffer, and was adjusted to pH 6·0, the flasks being sterilized at 3 lb./sq.in. for 10 min. After growth the pH was still below 7·0. Buffer did not seem to affect growth or germination. Although glucose was still present, growth stopped in all flasks at the end of 20 days. The mycelia were washed once in distilled water, pressed lightly between filter-papers, dried at 110° for 8 hr. and, after cooling in a desiccator, weighed. Maximum growth, c. 75 mg. mycelium/flask, was obtained with 0·125 μg. aneurin HCl/flask, and half maximum growth at c. 0·04 μg. aneurin HCl/flask (see also Table 2). The mycelium was somewhat paler at low aneurin levels, darkening being absent when the amount of growth was very small. No distinct perithecia were formed. As infertile perithecia were formed on the 0·2% glucose medium, both with and without phosphate buffer, in the latter case the final pH was 8·2, it seems that the appearance of perithecia is dependent on the concentration of nutrients remaining in the medium rather than on its pH.
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Requirements for fruiting

Calcium. Production of ascospores was induced by adding Ca to the medium, the results of one experiment being summarized in Table 1. Slight precipitates were observed in flasks containing 30 p.p.m. or more of Ca, but these disappeared as growth proceeded. At the end of the incubation period (30 days), no sugar was left in any flask. Perithecia were usually spherical, sometimes oval. Infertile perithecia were recognized from their smaller size, paler colour and deficiency in coiled hairs.

Table 1. Effect of varying concentrations of calcium on perithecial development of Chaetomium brasiliensis

<table>
<thead>
<tr>
<th>Ca (p.p.m.)</th>
<th>Day of appearance of Perithecia</th>
<th>Mean length (μm)</th>
<th>Infertile (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>9</td>
<td>62</td>
<td>100</td>
</tr>
<tr>
<td>0.1</td>
<td>9</td>
<td>82</td>
<td>100</td>
</tr>
<tr>
<td>0.4</td>
<td>11</td>
<td>92</td>
<td>64</td>
</tr>
<tr>
<td>1</td>
<td>11</td>
<td>87</td>
<td>48</td>
</tr>
<tr>
<td>3</td>
<td>11</td>
<td>98</td>
<td>25</td>
</tr>
<tr>
<td>5</td>
<td>11</td>
<td>100</td>
<td>20</td>
</tr>
</tbody>
</table>

Above Ca 5 p.p.m. the days of appearance of perithecia and ascospores respectively were: at 20 p.p.m. 9 and 11; at 50 p.p.m. 12, 14 and 14, 16 (duplicates); at 1000 p.p.m. 13, 14 and 15, 16 (duplicates).

np = none produced.

Ca increased the size of the perithecia and the number of fertile perithecia, the optimum Ca concentration being 3–5 p.p.m.; higher doses not significantly affecting these characteristics. No differences were detected in the degree of fertility, i.e. the number of spores per fertile perithecium. Above the optimum Ca range, other effects were noticed, the most important of which was delay in the maturation of perithecia. At Ca 30 p.p.m. and greater, the fungus began growth as white flakes and not as the normal mat but this difference soon vanished. At Ca 1000 p.p.m., germination was delayed by 2 days and the perithecia appeared as a scattered overgrowth and were probably fewer in number. At the concentrations tried, Ca did not affect the rate of growth or the total amount of growth (mycelium dry-weight) and had no effect without aneurin in the medium. With the formation of spores the colour of the perithecia changed to a deeper shade, from grey-green to blue-green, this colour change being an accurate indication of perithecial fertility.

In previous experiments with C. globosum, Ca was found to be replaceable by Sr and Ba, although particularly with Ba the number of mature perithecia formed was fewer (Basu, 1951). These elements also induced fruiting in C. brasiliensis, the effects being equal to that of Ca, when Sr and Ba were supplied at 10 p.p.m. (as nitrate and chloride respectively).

When Ca 10 p.p.m. was added, after completion of growth, to flasks initially without Ca and containing 1 μg. aneurin, there followed quickly a deepening
of colour and formation of spores, the final result being the same as when Ca was added before inoculation. Thus Ca was concerned only with the process of fruiting and not with growth. The strain from Ames was also tested; the responses were the same as with the usual test organism.

Addition of a hot-water extract of jute to the basal medium caused good growth and fruiting. Mn, Fe, Co, Ni, Sn, and Pb, found in the ash of jute extract by Basu (1951), were tested at 1 and 10 p.p.m. in the medium but no effect on growth or fruiting was observed; Co and Ni were inhibitory to growth.

Table 2. Effect of aneurin on fruiting of Chaetomium brasiliensis
(Ca 10 p.p.m. present throughout.)

<table>
<thead>
<tr>
<th>Aneurin HCl (µg./flask)</th>
<th>Day of appearance of</th>
<th>Perithecia</th>
<th>Ascospores</th>
</tr>
</thead>
<tbody>
<tr>
<td>0.001</td>
<td>np</td>
<td>np</td>
<td></td>
</tr>
<tr>
<td>0.003</td>
<td>np</td>
<td>np</td>
<td></td>
</tr>
<tr>
<td>0.006</td>
<td>np</td>
<td>np</td>
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</tr>
<tr>
<td>0.010</td>
<td>26</td>
<td>30</td>
<td></td>
</tr>
<tr>
<td>0.020</td>
<td>20</td>
<td>23</td>
<td></td>
</tr>
<tr>
<td>0.030</td>
<td>18</td>
<td>21</td>
<td></td>
</tr>
<tr>
<td>0.040</td>
<td>11</td>
<td>17</td>
<td></td>
</tr>
<tr>
<td>0.050</td>
<td>9</td>
<td>11</td>
<td></td>
</tr>
<tr>
<td>0.075</td>
<td>9</td>
<td>11</td>
<td></td>
</tr>
<tr>
<td>0.100</td>
<td>9</td>
<td>11</td>
<td></td>
</tr>
</tbody>
</table>

Glucose remaining after 30 days' incubation were 9.1, 6.1 and <2.0 mg./flask with 0.001, 0.003 and 0.006 µg. (or higher) aneurin HCl/flask. np = none produced.

When Whatman no. 5 filter-paper was used as the source of carbon instead of glucose and aneurin was supplied, *C. brasiliensis*, like *C. globosum*, formed many fertile perithecia in 7 days without added Ca. Fewer fertile perithecia formed by *C. brasiliensis* with Whatman no. 30 and no. 44 filter-papers, which have progressively smaller ash contents, only a few spores being formed with no. 44, although they appeared equally quickly with each filter-paper tried. That *C. globosum* fruited equally well on all these filter-papers is probably related to the fact that, unlike *C. brasiliensis*, it sporulates to some extent on a medium with a very low initial glucose content. The reaction of *C. brasiliensis* to the different filter-papers was reproduced when equivalent quantities of their ash were added to the usual glucose medium. Ordinary macro-chemical analysis showed Ca, but not Sr or Ba, in the ash of no. 5 filter-paper. Cellulose decomposition of filter-paper, determined by weighing the residue of no. 5 filter-paper left after 18 days' growth, was the same with both species, with or without Ca 10 p.p.m. added to the medium, which is consistent with the observation that Ca does not affect vegetative growth.

Aneurin. In Table 2 are given the results of an experiment designed to see whether aneurin was necessary for fruiting, apart from growth. Germination and rate of growth of mycelium did not appear to vary with the dosage of aneurin. Fruiting was progressively delayed below 0.05 µg./flask and at 0.006 µg./flask no perithecia were formed in 30 days, although growth had
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reached completion. It seems that a much higher concentration of aneurin is needed for fruiting than for growth and that the fungus can slowly synthesize the extra amount after growth is complete, when a little more aneurin than needed to reach this stage is originally present. The average size of perithecia and the percentage of infertile perithecia at 45 days did not differ significantly between flasks and were of the same order as obtained under optimum conditions (Table 1). The same applies to the total number of perithecia, except that at 0.01 μg. aneurin/flask the number was smaller.

**Sugar concentration.** Apart from adequate amounts of Ca and aneurin, another requirement for fruiting is that the concentration of free sugar in the medium must be very low; this was indicated by the quicker fruiting of the fungus on filter-paper. With increasing amounts of glucose in the medium fruiting was correspondingly delayed, at the following initial concentrations of glucose (%) the days of appearance of perithecia and ascospores respectively were: 0.2%, 8, 11; 0.3%, 10, 14; 0.3%, 13–15, 15–18; 0.5%, 15–18, 20–21; 0.6%, 21, 23. In every case less than 2 mg. glucose were left when ascospores were first seen. In another experiment, with 2% glucose, 1 μg. aneurin, and 20, 50 or 100 p.p.m. Ca in the medium fruiting did not take place even when only 18 mg. sugar were left after growth. Even immature perithecia are not formed, either with or without Ca in the medium, unless the sugar content approaches exhaustion.

**DISCUSSION**

A low nutrient level favours fruiting in several other species of *Chaetomium* (Basu & Bose, 1950), *C. convolutum* (Lilly & Barnett, 1949) and in some other Ascomycetes (Asthana & Hawker, 1936; Hawker & Chaudhuri, 1946). In general, environmental conditions which restrict growth favour reproduction, particularly the formation of sexual spores.

In its aneurin requirement for growth, *C. brasiliensis* resembles *C. convolutum* which is the only other Chaetomium species shown to have a specific vitamin requirement. Lilly & Barnett (1949), examining the effect of aneurin on *C. convolutum*, found that at a given nutrient concentration, increasing doses of aneurin successively initiated vegetative growth, formation of perithecia and of ascospores; within certain limits, the number of mature perithecia and ascospores increased. With *C. brasiliensis* more aneurin is needed for the formation of perithecia than for complete vegetative growth, and when enough calcium is present fruiting always follows the appearance of perithecia. The only other noticeable effect of aneurin on this species was in the acceleration of the appearance of perithecia and ascospores. *C. brasiliensis* does not require an external supply of biotin, which has the same effect as aneurin on *C. convolutum*.

Instances of the stimulation of spore formation by calcium are rare. Brian & Hemming (1950) found that conidia formation by the imperfect fungus *Trichoderma viride* was stimulated by 0.01 to 0.1% calcium chloride. Stimulation of sexual spore formation by calcium, strontium or barium is known only in other species of *Chaetomium*, in particular *C. globosum* (Basu, 1951).
replaceability of calcium by strontium or barium is even more marked in
C. brasiliensis than in C. globosum, but manganese, which engendered a few
fertile perithecia in the latter, had no effect on the former. Similarity with
C. globosum is also observed in that calcium has no effect on vegetative growth
and both species react to calcium even when added after complete growth.
However, from a consideration of the two strains of C. brasiliensis and the
three strains of C. globosum tested, a rather fundamental point of difference
is noted between the two species, that while in C. brasiliensis calcium induces
only the formation of ascospores in the large number of infertile perithecia
which develop even without the metal, in C. globosum the transition is from
a few or no perithecia, infertile or fertile, to a large number of fertile perithecia.

I wish to thank the Research Director, Indian Jute Mills Association Research
Institute, for permission to publish these results.

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EXPLANATION OF PLATE

Fig. 1. Culture of C. brasiliensis after 14 days on the basal medium. \( \times 1.4 \).

Fig. 2. Culture of C. brasiliensis after 14 days on the basal medium plus 1 \( \mu g \) aneurin,
showing densely aggregated infertile perithecia. \( \times 1.4 \).

Fig. 3. Crushed perithecium of C. brasiliensis after 20 days on basal medium plus 1 \( \mu g \)
aneurin, showing absence of asci, ascospores or curved terminal hairs. \( \times 250 \).

Fig. 4. Crushed perithecium of C. brasiliensis after 14 days on basal medium plus 1 \( \mu g \)
aneurin and 5 p.p.m. calcium, showing cylindrical asci, ascospores, and curved terminal
hairs. \( \times 250 \).

(Received 21 May 1951)
S. N. Basu—Nutrition of Chaetomium brasiliensis. Plate 1