Further Studies on Germination of Sporangiospores of *Rhizopus arrhizus*

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**SUMMARY**

Although sporangiospores of *Rhizopus arrhizus* do not swell or produce germ-tubes in distilled water, when they are suspended in heavy water the water in the spore is exchanged with heavy water in the medium. Spores swell and some produce germ-tubes in a glucose solution. Maximum germination occurred in the presence of utilizable carbon and nitrogen sources and suitable compounds containing phosphate, sulphate, potassium and magnesium ions. Germination was accompanied by a considerable increase in oxygen uptake and by the time the germ-tube emerged, the dry weight had increased by about 500%. Respiratory inhibitors (2,4-dinitrophenol, sodium azide, potassium cyanide) inhibited germination; sodium azide inhibited both the oxygen uptake and the dry weight increase.

Electron microscope studies showed structural changes in germinating *Rhizopus arrhizus* spores similar to those reported in other *Rhizopus* species: a new inner wall layer was formed, and changes in form and probably number of mitochondria occur. The effects of either anaerobic conditions or media containing sodium azide, on fine structure of germinating *R. arrhizus* spores were similar; mitochondrial multiplication ceases, mitochondrial cristae became a disorganized collection of undulating 'plates', and the nuclear membrane became split in places, thus giving rise to small vacuoles between the two electron-dense layers of the membrane.

**INTRODUCTION**

Some aspects of the physiology of germination of sporangiospores of *Rhizopus arrhizus* Fischer have already been reported by Ekundayo & Carlile (1964), who found that initiation of germination required the presence of glucose or fructose; maximum germination required in addition a nitrogen source, PO₄³⁻ and K⁺ or Na⁺. On media containing these nutrients, the spore increased in length from 5.5 to 13.5 μ in 8 hr. Anaerobic conditions inhibited germination. Weber & Ogawa (1965) observed that only proline, in the absence of phosphate, stimulated germination of *R. arrhizus* spores among the individual amino acids, sugars, organic acids, vitamins and inorganic nitrogen sources tested in their studies. Hawker & Abbott (1963b) described changes in fine structure during germination of sporangiospores of *R. nigricans* (= *R. stolonifer*) and *R. sexualis*. Washed spores of these species also failed to germinate in distilled water (Abbott, personal communication). The present investigations extended these various lines of study.

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METHODS

General. The preparation of media and inocula and the assessment of spore germination were as previously described (Ekundayo & Carlile, 1964).

Permeability studies with heavy water (deuterium oxide) as tracer. Spores, filtered, centrifuged and dried over P₂O₅ at 0-05 torr in an Edwards High Vacuum Unit, were suspended in 1 ml. D₂O in a centrifuge tube covered with aluminium foil and incubated at 30° for 18 hr. The D₂O was decanted and distilled under reduced pressure in a micro-distillation apparatus. The density of the distillate was determined by the gradient tube method (Shaw, 1955). 50 ml. CCl₄ were put in a 100 ml. graduated cylinder, 50 ml. Kerosene were added to the top of the carbon tetrachloride, and the interface between them was stirred and allowed to stand for 24 hr in a water-bath at 20° to allow the density gradient to become established. The density of D₂O was calibrated by introducing drops of known concentrations of D₂O into the CCl₄ + Kerosene mixture and measuring the positions of the drops with a travelling microscope and vernier. Drops of the distillate were then added, their equilibrium position determined and the equivalent D₂O concentration was read from the calibration graph. The percentage exchange was taken as the difference between the percentage of D₂O in the distillate from spore suspension and from 100% D₂O control. The theoretical full exchange value was derived from the equation:

\[ \% \text{ full exchange} = \frac{x}{x+y} \times 100, \]

where \( x = \text{wt. (mg.) of water from spore sample at 100° for 24 hr; } \)

\( y = \text{wt. (mg.) of 1 ml. } D₂O \text{ at room temperature.} \)

Electron microscopy. Ungerminated Rhizopus spores were not always well fixed after 24 hr in 2% KMnO₄, glutaraldehyde or osmic tetroxide. Satisfactory fixation was achieved by shaking the spore suspension in 4% KMnO₄ in veronal buffer pH 7.4 with Ballotini glass balls in a Mickle disintegrator for 30 min. and leaving the spores in the fixative at room temperature for a further 3½ hr. The spore suspension was then centrifuged at 2000 rev./min. for 5 min. to separate the glass balls. The supernatant was decanted and centrifuged at 4000 rev./min. for 15 min. and the spore deposit washed with distilled water. Only broken or damaged spores subsequently showed internal structure; it was, therefore, assumed that the fixative penetrated through the cracks in the walls.

Germinated spores were fixed with 2% KMnO₄ in veronal buffer pH 7.4 (Luft, 1956) for 4 hr at room temperature.

After fixation the spores were dehydrated with acetone, embedded in vestopal W, sectioned, and observed with either an Akashi TRS-50 or a Siemens Elmiskop I electron microscope.

RESULTS

Permeability of Rhizopus arrhizus spores

Ekundayo & Carlile (1964) showed that R. arrhizus spores did not swell or produce germ-tubes in sterile distilled water; maximum swelling and germ-tube production by all spores required the presence of utilizable carbon and nitrogen sources and compounds containing phosphate, potassium or sodium ions.
Germination of sporangiospores of Rhizopus arrhizus

In the present studies, wetting the spores with Gemex Z-11, Manoxol OT, Nonidet P-42 or Teepol either before or during their incubation in water did not make them swell or produce germ-tubes in water.

Experiments in which spores were suspended in solutions of the standard medium containing high concentrations of (NH₄)₂SO₄ showed that saturated solutions of (NH₄)₂SO₄ did not plasmolyse ungerminated Rhizopus arrhizus spores. Spores which had been incubated for about 3 hr on the standard medium were, however, plasmolysed by standard medium containing 3.0 M (NH₄)₂SO₄.

Heavy water was used as a tracer to find whether ungerminated Rhizopus arrhizus spores could exchange the water in the spore with heavy water in the medium. The results are given in Table 1. Experimental values were higher than theoretical figures obtained by the use of the equation given under Methods; full exchange values were always obtained. This suggests that R. arrhizus spores are permeable to water even though they do not swell in it in the absence of suitable nutrients.

Vacuum-dried spores did not swell or produce germ-tubes in distilled water; but all the spores germinated in the standard medium. The ability of the spores to exchange their water with the heavy water in the medium, demonstrated above, was therefore not due to an effect of drying on the permeability of the spore wall.

Table 1. The estimation of heavy water (deuterium oxide) in permeability studies with Rhizopus arrhizus spores

<table>
<thead>
<tr>
<th>Wt. (mg.) of vacuum-dried spores</th>
<th>Wt. (mg.) of oven-dried spores</th>
<th>Wt. (mg.) of water in spores</th>
<th>Theoretical full-exchange value (%)</th>
<th>Experimental exchange value (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>389-3</td>
<td>282-0</td>
<td>107-4</td>
<td>9.7</td>
<td>15.4</td>
</tr>
<tr>
<td>296-9</td>
<td>167-1</td>
<td>129-8</td>
<td>11.5</td>
<td>17.8</td>
</tr>
</tbody>
</table>

\( M = \text{wt. (mg.) of oven-dried spores} \)

Table 2. The effect of respiratory inhibitors on germination of Rhizopus arrhizus on standard medium at 30°C

<table>
<thead>
<tr>
<th>Chemical inhibitor in medium</th>
<th>Least molar concentration inhibiting germination</th>
<th>Average spore diameter (μ) at time 0</th>
<th>Average spore diameter (μ) at 8 hr</th>
<th>Spores (%) with germ-tubes at 8 hr</th>
</tr>
</thead>
<tbody>
<tr>
<td>2,4-Dinitrophenol</td>
<td>( 5 \times 10^{-3} )</td>
<td>5-5</td>
<td>6-0</td>
<td>15</td>
</tr>
<tr>
<td>Sodium azide</td>
<td>( 10^{-3} )</td>
<td>5-5</td>
<td>5-5</td>
<td>5</td>
</tr>
<tr>
<td>Potassium cyanide</td>
<td>( 5 \times 10^{-3} )</td>
<td>5-5</td>
<td>8-0</td>
<td>30</td>
</tr>
</tbody>
</table>

Effect of respiratory inhibitors on spore germination

Mandels & Darby (1958) reported the inhibition by sodium azide of spore swelling in Myrothecium verrucaria in the basal nutrient medium.

The effect of 2,4-dinitrophenol, sodium azide and potassium cyanide on the germination of Rhizopus arrhizus spores was, therefore, investigated. A decimolar solution of each inhibitor in liquid standard medium was made and samples of it were withdrawn into different Petri dishes. Melted standard medium agar was added to each Petri dish to a total volume of 20 ml. and allowed to solidify. Four sterile cellophane strips were placed on the solidified medium in each Petri dish.
and two drops of spore suspension were spread on each strip. After 8 hr the least concentration of each inhibitor causing incomplete swelling and/or less than 50% germ-tube production was determined. 2,4-Dinitrophenol, sodium azide and potassium cyanide concentrations up to about 10^{-3}M inhibited both spore swelling and germ-tube emergence in R. arrhizus (Table 2).

**Effect of 10^{-3} M-NaN_3 on germinating Rhizopus arrhizus spores**

Experiments in which ungerminated spores were placed in contact with the inhibitor for different periods, after which they were transferred to fresh media not containing the inhibitor, showed that R. arrhizus spores on the standard medium containing 5 \times 10^{-3} M-NaN_3 remained unswollen and without germ-tubes for at least 24 hr, but if then transferred to the standard medium not containing the inhibitor, all spores swelled and produced germ-tubes after a further 8 hr. The effect of the inhibitor on the ungerminated spore was, therefore, not permanent.

The following experiment was accordingly made to investigate the effect of 10^{-3} M-NaN_3 on spores which had been incubated for 8 hr on the standard medium. Such partially swollen spores were transferred either to the standard medium containing 10^{-3} M-NaN_3 or to water agar (with or without 10^{-3} M-NaN_3) for 5 hr. The results are given in Table 3. Spores transferred to the standard medium containing 10^{-3} M-NaN_3 did not swell further or produce germ-tubes, but spores transferred to water agar, with or without the inhibitor, produced germ-tubes but stopped swelling. This suggested that NaN_3 was taken into the spores only when utilisable exogenous nutrients were present.

**Table 3. Effect of 10^{-3} M-NaN_3 on germinating Rhizopus arrhizus spores at 30°**

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Condition at time of transfer (8 hr)</th>
<th>Condition at 8 hr</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Average spore diam. (μ)</td>
<td>Spores (%) with germ-tubes</td>
</tr>
<tr>
<td>3 hr on standard medium, then:</td>
<td></td>
<td></td>
</tr>
<tr>
<td>5 hr on medium + 10^{-3}M-NaN_3</td>
<td>7.5</td>
<td>0</td>
</tr>
<tr>
<td>5 hr on water agar + 10^{-3}M-NaN_3</td>
<td>7.5</td>
<td>0</td>
</tr>
<tr>
<td>5 hr on water agar</td>
<td>7.5</td>
<td>0</td>
</tr>
<tr>
<td>5 hr on fresh standard medium</td>
<td>7.5</td>
<td>0</td>
</tr>
<tr>
<td>All 8 hr on standard medium</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>All 8 hr on plain agar</td>
<td>—</td>
<td>—</td>
</tr>
</tbody>
</table>

**Dry weight changes during germination**

Dry weight was determined at intervals during the germination of Rhizopus arrhizus spores in the standard medium with or without sodium azide. 25 ml. samples of spore suspension in water were measured into 150 ml. Erlenmeyer flasks containing either 25 ml. of double-strength standard medium with or without 10^{-3} M-NaN_3 (to maintain the usual standard medium nutrient level on dilution with spore suspension), or 25 ml. of water. Four replicates of each treatment were set up. Air, filtered by passing through cotton wool plugs, was bubbled through the suspensions which were maintained at 30 ± 1°. After 8 hr, the contents of
similarly treated flasks were pooled and filtered through a sterile sintered crucible of fine porosity lined with sterile glass paper of known weight. Spores were washed with distilled water and then acetone, and dried at 100° for 6 hr. A considerable increase (about 500%) in dry weight accompanied the germination of R. arrhizus spores in the standard medium, but in the presence of 10^-3 M-NaN₃, no increase in dry weight occurred (Table 4).

![Graph](image)

**Table 4. Dry weight changes during germination of R. arrhizus spores**

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Weight (mg.) of spores</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spores at time 0</td>
<td>1.6</td>
</tr>
<tr>
<td>Spores in sterile distilled water for 8 hr</td>
<td>1.5</td>
</tr>
<tr>
<td>Spores in standard medium for 8 hr</td>
<td>8.1</td>
</tr>
<tr>
<td>Spores in standard medium + 10^-3 M-NaN₃ for 8 hr</td>
<td>1.8</td>
</tr>
<tr>
<td>Spores on standard medium for 8 hr</td>
<td>8.7</td>
</tr>
<tr>
<td>Spores in standard medium far 3 hr, then 5 hr in</td>
<td>3.5</td>
</tr>
<tr>
<td>medium containing 10^-4 M-NaN₃</td>
<td></td>
</tr>
</tbody>
</table>

**Respiration of Rhizopus arrhizus spores**

As it had been found that respiratory inhibitors (2,4-dinitrophenol, sodium azide, and potassium cyanide) inhibited germination of R. arrhizus spores, the following experiments were made to investigate the dependence of spore germination on respiration. Measurements of oxygen uptake were made with Warburg
respirometers (Umbreit, Burris & Stauffer, 1957). The results are summarized in Fig. 1. Oxygen uptake by *R. arrhizus* spores in the standard medium was very high; in water, oxygen uptake was low and could be estimated only by using thick spore suspensions. $10^{-3}$m-$\text{Na}_2\text{S}_2\text{O}_3$ inhibited oxygen uptake. Weber & Ogawa (1965) observed that $10^{-3}$m-$\text{Na}_2\text{S}_2\text{O}_3$ and $10^{-3}$m-$\text{NaF}$ completely inhibited oxygen uptake, whereas 2,4-dinitrophenol ($10^{-3}$m) stimulated respiration of *R. arrhizus* spores in proline-phosphate mixtures.

Electron microscope studies

Hawker & Abbott (1963b) studied the maturation and germination of sporangiospores of *Rhizopus nigricans* (*R. stolonifer*) and *R. sexualis*. *R. arrhizus* spores have been similarly studied; in Plate 1, fig. 1, can be seen changes in fine structure of germinating sporangiospores (on standard medium) similar to those observed by Hawker & Abbott (1963b) in *R. stolonifer*. A new inner wall (IW) has been formed, and the large mitochondria (M) have occasional constrictions along the longer axis. Spores germinating on glucose agar contained similar structures, but no changes in spore fine structure were observed in spores on water agar.

*Effect of sodium azide and anaerobic conditions on germinating spores.* *Rhizopus arrhizus* spores which had been incubated on the standard medium for 3 hr and then transferred for 5 hr either to standard medium in a sealed system filled with nitrogen or to standard medium containing $10^{-3}$m-$\text{Na}_2\text{S}_2\text{O}_3$ usually stopped swelling and producing germ-tubes. Fine structure of the spores so treated, was, therefore, studied and it was found that mitochondrial division ceased, as shown by the large size of these organelles (Pls. 1, 2; figs. 2-4). Cristae which appeared as nearly parallel ‘plates’ in control spores became a disorderly collection of undulating plates (Pl. 2, figs. 8, 4). The nuclear membrane became split in places, thus giving rise to small vacuoles between the two electron-dense layers of the membrane. Gale & McLain (1964) observed similar effects on spores of *Candida albicans* exposed to thiobenzoate; and Hirano & Lindegren (1961, 1968) observed that mitochondria of *Saccharomyces cerevisiae* developed some abnormalities in partially anaerobic conditions, the mitochondrial cristae becoming replaced by a complex system of inner membranes. The finding that sodium azide and anaerobic conditions have similar effects on fine structure of *R. arrhizus* spores is interesting. If NaN$_3$ inhibits spore germination by its inhibitory action on the cytochrome oxidase-cytochrome c oxygen transfer system thus causing anaerobic conditions to occur (Stannard & Horecker, 1948) it might be expected that anaerobic conditions would have the same effect as that of NaN$_3$ on spore structure.

*The microfibrillar structure of *R. arrhizus* cell-walls.* Hawker & Abbott (1963b) had observed that a thin inner wall consisting of tangentially arranged elements, resembling the wall of the vegetative hypha (Hawker & Abbott, 1963a), but differing from the original spore wall, was laid down during germination of *Rhizopus stolonifer* and *R. sexualis* spores.

A similar wall-layer was seen in germinating *Rhizopus arrhizus* spores. A comparative study of shadowed preparations of isolated cell walls of ungerminated and germinating spores and of young hyphae (about 16 hr old) by Dawson’s (1949) method showed that the microfibrils of *R. arrhizus* cell walls were oriented in a
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rather irregular and random fashion and no clear differences in microfibrillar structure of original spore walls, new inner spore walls and hyphal walls was detected.

DISCUSSION

The initiation of germination in Rhizopus arrhizus spores has been shown to require a utilisable carbon compound (Ekundayo & Carlile, 1964). Earlier, it was thought that the metabolism of the carbon compound, probably at the spore surface, brought about induced permeability of the spore walls to mineral ions and water. Heavy water exchange experiments have since shown that R. arrhizus spores can exchange the water in the spore with heavy water in the medium in the absence of carbon compounds. This suggests that the spore is already permeable to water, and, probably, to mineral ions. Spore swelling, however, involves an increase in spore size and requires the spore wall to be extensible. Since swelling occurs on media containing utilisable carbon compounds (e.g. glucose) without any mineral salts, glucose is sufficient to induce the walls of R. arrhizus spores to become more extensible.

Nickerson's (1968) work on yeast suggested an explanation of how glucose might bring about changes in the Rhizopus spore wall and thereby make it become extensible. Plasticity of the yeast wall was thought to result from a reduction of disulphide linkages in the protein component of the wall to sulphydryl groups, the ultimate reducing agent being NADH\(_2\) generated from glucose. If this theory applies to Rhizopus, a disulphide-containing protein complex must be present in the spore wall and the spore must contain enzyme systems capable of reducing disulphide linkages. Results of investigations by other workers support such a possibility: Crook & Johnston (1962) demonstrated the presence of protein-carbohydrate complexes in a wide range of fungal cell-walls. Bartnicki-Garcia & Reyes (1964) demonstrated the presence of such complexes in Mucor rouxii which, like Rhizopus, is a member of the Mucoraceae. Hatch & Turner (1960) showed that in the presence of systems to provide NADH\(_2\) or NAPH\(_2\) pea-seed protein disulphide reductases reduced protein disulphide linkages. Robson & Stockley (1962), by using autoradiographic methods detected the presence of sulphydryl groups in the cell walls of many fungi.

If the above hypothesis is correct, swelling in Rhizopus spores would result from glucose-induced increased plasticity of the spore wall. Once wall plasticity is achieved, the wall stretches in response to water intake caused by the high osmotic pressure in the spores. The observations (1) that Rhizopus spores can exchange the water in the spore with heavy water in the medium and (2) that Rhizopus spores which have been incubated on the standard medium for about 8 hr, with probable dilution of their contents, become plasmolysed when placed in hypertonic salt solutions, support the view that these spores take in water by osmosis. It must be emphasised, however, that these observations prove only that water can be extruded from Rhizopus spores by osmotic methods. Systems are known in which water intake and extrusion take place by different processes, e.g. in amoeba in which water enters the organism passively but its excretion through the contractile vacuole requires respiratory energy.

Spore swelling in Rhizopus is accompanied by a considerable increase in oxygen
uptake and an increase of about 500% in dry weight of the spores. It is inhibited by respiratory inhibitors (Table 2, and Weber & Ogawa, 1965) and anaerobic conditions. It is thus likely that synthetic processes occur during spore swelling. Indeed, the formation of new internal structures can be demonstrated by electron microscopy during swelling. If the spores are low in nutrient content, such synthesis may account for the dependence of Rhizopus spores on exogenous nutrients for spore swelling. Anaerobic conditions and respiratory inhibitors may prevent swelling by blocking the metabolic processes necessary for the initiation of germination.

The above discussion suggests that glucose is needed for the germination of Rhizopus spores for two reasons:

(i) To generate NADH₂, to reduce disulphide linkages in the spore wall and permit wall extensibility and hence osmotic swelling.

(ii) As a source of carbon and energy for the synthesis of the new inner wall of the spore which subsequently becomes the germ-tube wall.

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REFERENCES


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EXPLANATION OF PLATES

Electron micrographs of Rhizopus arrhizus.

SW = spore wall, IW = Inner wall, N = nucleus, NM = nuclear membrane, M = mitochondria, MC = mitochondrial cristae, P = plasmalemma, TM = tonoplast, V = vacuole, IV = intramembranous vacuole.

PLATE 1

Fig. 1. Approximately transverse section through sporangiospore incubated on standard medium for 3 hr; showing typical nuclei, development of inner wall and mitochondria (e.g. MI) with constrictions along the longer axis.

Fig. 2. Section through a spore incubated on standard medium for 3 hr followed by 5 hr on standard medium containing 10-4 M-Na+. Note presence of disorganized mitochondrial cristae, vacuoles have appeared in cytoplasm.

PLATE 2

Fig. 3. Section through a spore incubated on standard medium for 3 hr followed by 5 hr in N2-saturated standard medium. Note mitochondrial cristae have become disorganized and nuclear membrane split in places thus giving rise to small vacuoles between the two electron-dense layers of the membrane. Features are essentially similar to those observed in spores placed under anaerobic conditions (Fig. 2).

Fig. 4. Two mitochondria from Fig. 3, enlarged. Note outer electron-dense layer of mitochondrial membrane has disappeared in some places and mitochondrial cristae have become a collection of undulating 'plates'.