Changes in Gross Chemical Components of *Trichophyton mentagrophytes* During Incubation in Increased Carbon Dioxide Tensions

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(Received 21 May 1962)

SUMMARY

Macroconidia of *Trichophyton mentagrophytes* harvested from a culture grown on a medium rich in glucose contained 33% acid-soluble carbohydrate. Paper chromatography of hydrolysates showed glucose as the main sugar in this fraction; there was also a trace of an unidentified sugar. Macroconidia from the same strain grown on the same medium but deficient in glucose contained 19% carbohydrate; hydrolysates of this fraction contained glucose, with traces of galactose and the unidentified sugar. Microconidia harvested from a second strain grown on a medium rich in glucose contained 11% acid-soluble carbohydrate; hydrolysates of this fraction also contained glucose with traces of galactose and the unidentified sugar. These differences were reflected in whole cultures induced for macroconidial formation with carbon dioxide. Strains which responded well increased in acid-soluble carbohydrate; glucose was the predominating sugar in hydrolysates of these fractions. Strains which responded poorly did not increase in carbohydrate content; glucose, and traces of galactose and an unidentified sugar, were found in hydrolysates of these fractions.

INTRODUCTION

A previous report from this laboratory (Chin & Knight, 1957) described the stimulation of macroconidial formation in *Trichophyton mentagrophytes* during incubation at increased carbon dioxide tensions. A requirement for glucose was demonstrated, since macroconidia formation was not stimulated in increased atmospheric tensions of CO$_2$ when glucose was omitted from the medium. The experiments described in the present paper attempted to analyse and compare the gross chemical components of macroconidia, microconidia and hyphae. It was our intention to find some measurable difference between macroconidia and the other morphological units of *T. mentagrophytes* with the hope that this difference might be induced in the normally non-sporulating strains when macroconidia formation was stimulated by carbon dioxide. Because of difficulties involved in separating macroconidia from microconidia and hyphae, mutant strains which differed in type of sporulation were derived from a common parent. Macroconidia are easily harvested from a sporulating mutant which does not form microconidia, and vice versa.
Mutants

All mutants were derived from Trichophyton mentagrophytes strain A-280 (from the Communicable Disease Center, Chamblee, Georgia, U.S.A.). Stock cultures were maintained by 8-monthly transfer on Sabouraud’s glucose agar (Difco) and incubation at 80° for 10–14 days followed by storage at 5°. Sabouraud’s glucose agar (4% glucose, 1% Neopeptone, 1.5% agar) was used throughout. Liquid medium was prepared by omitting the agar, and Sabouraud’s conservation medium by the omission of glucose. Cultures were incubated for 5 days before harvest. The genetic purities of parent and mutant strains were assured by single spore isolations to avoid the selection of macroconidia-forming variants from mixed clones during incubation in CO₂ (Emmons & Hollaender, 1939). A microconidium was isolated from strain A-280 by the excision and transfer to fresh medium of a germinated microconidium from ruled agar blocks of Sabouraud’s glucose agar. The clone resulting from the growth of this spore was maintained as the parent culture, strain 5-7. Spores from this clone were seeded on Petri plates containing solid media and irradiated with ultraviolet radiation. Morphological mutants differing in type of sporulation were selected from the survivors and transferred to fresh medium. In turn, clones resulting from the growth of transferred survivors were purified by the isolation of microconidia. Since strain 1-2-1 does not form microconidia, a macroconidium was isolated. The growth characteristics of all mutants have remained stable through 3 years of serial transfer. Their sporulating patterns and responses to CO₂ are recorded in Table 1.

<table>
<thead>
<tr>
<th>Mutant</th>
<th>Normal air</th>
<th>16% (v/v) CO₂</th>
</tr>
</thead>
<tbody>
<tr>
<td>5-7</td>
<td>rare</td>
<td>profuse</td>
</tr>
<tr>
<td>1-1-2</td>
<td>none</td>
<td>poor</td>
</tr>
<tr>
<td>1-2-1</td>
<td>abundant</td>
<td>profuse</td>
</tr>
<tr>
<td>1-8-1</td>
<td>none</td>
<td>poor</td>
</tr>
</tbody>
</table>

Inoculum and growth

Inoculum for solid media was prepared by dispersing the growth on a stock culture slope with sterile water and pipette. Cultures for chemical analysis were grown as surface mats on liquid media. A floating inoculum was prepared by pipetting 1 ml medium on to the surface of a stock culture slope. The culture tube was rolled gently to dislodge spores and hyphal fragments, the majority of which floated on the surface of the medium. In inoculation, this preparation was poured into a flask containing liquid medium. Flasks which were incubated with increased atmospheric tensions of CO₂ (0–16% v/v) were fitted with rubber stoppers and glass tubing outlets. The gassing procedure was as previously described (Chin & Knight, 1957). After incubation the floating cultures were washed by pouring off the media and refloating the culture mats with water 4 times. The mats were then removed from the flasks and suspended in cold acetone (Umbreit, Burris & Stauffer, 1957). The organism was separated from the acetone by centrifugation and the supernatant acetone drawn off through a capillary pipette connected to an aspirator pump. Samples of organism were dried in vacuo overnight.
Spores

Spores were harvested from cultures grown on solid media in Roux bottles; macroconidia were harvested from strain 1-2-1, microconidia from strain 5-7. After 5 days of incubation, 40 ml. water were added to each bottle, the contents vigorously shaken and strained through gauze. Although many spores were retained with mycelium in the filter, the effluent usually contained a high percentage of the spores in the original harvest. Spores were separated from the effluent by centrifugation. The spores were washed 4 times, after which they were suspended in cold acetone and dried as described above.

Spore walls

Macroconidia were harvested and broken, prior to acetone treatment, in a Serval Omni-mix with water and glass microbeads for 30 min. The proportions of the mixture were spores+glass beads+water (1+2+8, by vol.). Microscopic examination of samples stained with lactophenol cotton blue revealed that 95% of the macroconidia were broken in all compartments. The spore walls were washed with water in excess of twenty times, suspended in cold acetone, and dried.

Extraction procedure

Weighed samples were transferred to test tubes and sequentially extracted in a boiling water bath with water for 15 min., with 5% (w/v) trichloroacetic acid (TCA) for 30 min., with NaOH for 80 min. The tubes were capped with glass marbles to prevent evaporation. After each extraction, the contents of the tubes were separated by centrifugation. Samples of extracts were removed for analysis with a pipetting bulb; the rest of the extract was discarded through a capillary pipette connected to an aspirator pump. The cell residues remaining after extraction with hot NaOH were washed with water four times by repeated centrifugation and transferred to tared aluminum cups. The residues were dried overnight at 80° and the weights determined by difference.

Chemical analyses

Carbohydrate was measured with the anthrone reagent, with glucose as the standard (Loewus, 1952). Samples from TCA extracts were hydrolysed with 0.6 N-HCl in a boiling water bath before paper chromatography. The solvent systems used for paper chromatography were isopropanol + water (8+2, by vol.) and amyl acetate + pyridine + water (3+8+1, by vol.). Paper chromatograms were developed with AgNO3 reagent (Smith, 1958).

Protein was measured with the Folin phenol reagent method based on the modification of Lowry, Rosenbrough, Farr & Randall (1951).

Total nucleic acids (NA) were estimated by the method of Logan, Mannell & Rossiter (1952) at 268.5 mμ in a Beckman model DU spectrophotometer with 80 ml. silica cuvettes after extraction by the method of Schneider (1945). Samples were read against a blank of 5% TCA. Herring sperm deoxyribonucleic acid (DNA Nutritional Biochemical Co.), similarly hydrolysed, was used as the standard.

DNA content was determined by the method of Kech (1958). Ribonucleic acid (RNA) was estimated by the difference between total NA and DNA.
RESULTS

The gross chemical compositions of macroconidia, microconidia, and whole culture of *Trichophyton mentagrophytes* strain 5-7, which produces microconidia but no macroconidia on Sabouraud's glucose agar, are described in Table 2. The

Table 2. Gross chemical components of macroconidia, microconidia, and whole culture of strain 5-7 of *Trichophyton mentagrophytes*

<table>
<thead>
<tr>
<th>Extract</th>
<th>Component</th>
<th>Macroconidia</th>
<th>Microconidia</th>
<th>Strain 5-7 (% of dry wt.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>H$_2$O</td>
<td>CHO*</td>
<td>11.8</td>
<td>13.2</td>
<td>7.6</td>
</tr>
<tr>
<td></td>
<td>PRO</td>
<td>11.5</td>
<td>7.3</td>
<td>12.3</td>
</tr>
<tr>
<td>TCA</td>
<td>CHO</td>
<td>33.3</td>
<td>11.1</td>
<td>11.9</td>
</tr>
<tr>
<td></td>
<td>PRO</td>
<td>5.5</td>
<td>5.4</td>
<td>5.8</td>
</tr>
<tr>
<td></td>
<td>RNA</td>
<td>2.6</td>
<td>3.5</td>
<td>1.4</td>
</tr>
<tr>
<td></td>
<td>DNA</td>
<td>0.2</td>
<td>0.3</td>
<td>0.1</td>
</tr>
<tr>
<td>NaOH</td>
<td>PRO</td>
<td>15.6</td>
<td>16.0</td>
<td>18.4</td>
</tr>
<tr>
<td></td>
<td>CR</td>
<td>10.6</td>
<td>12.0</td>
<td>30.1</td>
</tr>
</tbody>
</table>

*CHO*: carbohydrate; PRO, protein; RNA, ribonucleic acid; DNA, deoxyribonucleic acid; CR, cell residue.

Fig. 1. Changes in gross chemical composition, in percent of dry weight, of *Trichophyton mentagrophytes* strain 5-7 after incubation in increased carbon dioxide tensions. CHO-H$_2$O, hot-water-extractable carbohydrate; PRO-H$_2$O, hot-water-extractable protein; CHO-TCA, hot-TCA-extractable carbohydrate; PRO-TCA, hot-TCA-extractable protein; RNA, ribonucleic acid; DNA, deoxyribonucleic acid; PRO-NaOH, hot-NaOH-extractable protein; CR, cell residue.

Fig. 2. Changes in acid-soluble carbohydrate, in percent of dry weight, in mutants which vary in type and degree of sporulation during incubation in increasing carbon dioxide tensions. All mutants were harvested as surface cultures on liquid Sabouraud's medium, strain 5-7 was grown on medium with and without glucose (S GLU).
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Macroconidia contained 38% hot-water-insoluble, hot-acid-soluble carbohydrate as compared with 11% in microconidia and 12% in strain 5-7. These differences became more striking when paper chromatograms of hydrolysates of these fractions were examined. Glucose and trace amounts of another, at present, unidentified sugar were found in the hydrolysate of the TCA extract from macroconidia; in microconidia and whole organisms of strain 5-7, glucose and trace amounts of galactose and the same unidentified sugar were found. The unidentified sugar is suspected to be fructose, rhamnose or sorbose. Macroconidia harvested from strain 1-2-1 grown on Sabouraud's conservation agar (prepared by the omission of glucose) contained 19% acid-soluble carbohydrate. Again, paper chromatography showed that glucose was the predominant sugar in hydrolysates of this fraction; only traces of galactose and the unidentified sugar were found. Walls of macroconidia from strain 1-2-1 grown on Sabouraud's glucose agar contained 9% acid-soluble carbohydrate.

After pre-treatment with acetone, the cytoplasm of macroconidia, microconidia and hyphae stain red-brown with Lugol's iodine solution. Hyphal and spore walls do not take the stain.

Strains 5-7 and 1-2-1 responded markedly to incubation in increased atmospheric tensions of CO, and macroconidia formation was stimulated greatly. Strains 1-1-2
and 1-3-1 did not respond well to CO$_2$ and macroconidia formation was poorly stimulated. The changes in gross chemical components of the parent strain, 5-7, grown in increasing amounts of CO$_2$ are shown in Fig. 1. The greatest fluctuations occur in three fractions: acid soluble carbohydrate (CHO-TCA), NaOH-extractable protein (PRO-NaOH), and cell residue (CR). The changes in these fractions, for all mutants, in response to CO$_2$ are described in Fig. 2 (CHO-TCA), Fig. 3 (PRO-NaOH), and Fig. 4 (CR). In strains 5-7 and 1-2-1, increases in carbohydrate and protein, and decreases in cell residue, accompanied increased macroconidia formation as it was induced by increasing CO$_2$ tensions. Strains 1-1-2 and 1-3-1 were poorly stimulated to form macroconidia and did not show increases in carbohydrate. Under increased atmospheric tensions of CO$_2$, strain 1-1-2 increased in protein and decreased in cell residue; strain 1-3-1 did not increase in protein or decrease in cell residue. Paper chromatograms of TCA extract hydrolysates from all cultures showed that glucose was the predominant sugar in these fractions; traces of galactose and the unidentified sugar were also present. When strain 5-7 was grown on Sabouraud's conservation medium under increased tensions of CO$_2$, macroconidia formation was poorly stimulated. Analyses of gross chemical components during incubation with CO$_2$ revealed no increase in carbohydrate, an increase in protein, and no decrease in cell residue.

DISCUSSION

The formation of macroconidia and microconidia by *Trichophyton mentagrophytes* implies that differences in structure and function, as well as size, exist between the two spore forms. Unfortunately, information about functional differences between macroconidia and microconidia is not available. One difference in function has been implied (Wilhelm, 1947). Since macroconidia are multinucleate and microconidia are uninucleate (Emmons, 1934), a heterokaryotic state may exist in the multinuclear compartment of a macroconidium and provide a survival advantage by allowing the macroconidium to carry a recessive gene.

The stimulation of macroconidia formation by incubation in increased CO$_2$ tensions is difficult to investigate, since very little is known about the physiology of the organism. If any existing differences in the gross chemical components of morphological structures were known, it should be possible to investigate the physiological changes in mycelia during the process of sporulation and the effect of CO$_2$ on these changes. The experiments described in this paper were designed to this end.

It was previously demonstrated that stimulation of macroconidia formation by incubation in increased CO$_2$ tensions did not occur when glucose was omitted from the medium. This requirement for glucose may be a direct reflection of the higher concentration of carbohydrate in macroconidia than in microconidia or hyphae. Perhaps the formation of macroconidia requires higher concentrations of glucose in the medium for a structural precursor of the spore itself. The carbohydrate appears to reside in the cytoplasm.

It is our experience that macroconidia germinate more rapidly than microconidia on Sabouraud's glucose agar. Wheeler, Cabaniss & Cawley (1958) showed that macroconidia of *Microsporum fulvum*, another dermatophyte, were capable of germinating in distilled water in 4-5-10 hr. (confirmed in our laboratory). Macroconidia of
Trichophyton mentagrophytes germinate in distilled water in 8 hr. when they are aerated; microconidia will not germinate under the same conditions. We have postulated that the higher concentration of carbohydrate in macroconidia may be more available as an energy source for germination than the lower quantity in microconidia. This is not the only factor to be considered, however, since microconidia suspended in 4% glucose solution will not germinate in 48 hr.

Other changes in gross chemical components of cultures also occur during incubation in increased CO₂ tensions, particularly, in protein and cell residue fractions. However, investigations into these two fractions have been deferred because the separation of strong macroconidia-forming mutants and weak macroconidia-forming mutants is not as clear as in the carbohydrate fraction. These fractions have not, by any means, been dismissed from further consideration since the process of macroconidia formation must be complex. Physiological and structural changes during sporulation will rely heavily upon the formation of new and more protein. Generalizations about the sporulating processes of fungi are difficult to make. The factors which influence spore formation are many and varied (Cochrane, 1958); an agent which stimulates sporulation in one organism may inhibit it in another. Comparisons of sporulating processes will have to await more studies similar to those of Wright & Anderson (1958) on Dictyostelium discoideum and Cantino (1956) on Blastocladiella emersonii. It is of interest that D. discoideum accumulates cellulose during one phase of sporulation and that CO₂ induces the formation of resistant sporangia in B. emersonii. Cantino & Horenstein (1956) implicated a succinate-ketoglutarate-isocitrate cycle in the sporulation of Blastocladiella. Whether these aspects of carbohydrate metabolism are related to the accumulation of glucose in Trichophyton mentagrophytes remains to be seen.

This work was supported by Grant E-1201 from the National Institutes of Health, U.S.A.

REFERENCES


