Some Biological Effects of Volatile Metabolites from Cultures of Saccharomyces cerevisiae Meyen ex Hansen

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SUMMARY

The gas mixtures from cultures of Saccharomyces cerevisiae inhibited growth and sporulation of Aspergillus niger and germination of seeds of Lepidium sativum. In test conditions seven volatile organic metabolites in the culture gases were identified by gas-liquid chromatography (GLC) as acetaldehyde, ethyl acetate, ethanol, n-propanol, isobutanol, and a mixture of isopentanols (1 part 2-methyl-butanol-1-ol to 2 parts 3-methyl-butanol-1-ol); changes in the concentrations of CO₂ and O₂ were also measured. Pure samples of each of these components were tested at the concentrations found in the culture gases in order to identify the inhibitory substances.

Inhibition of growth of Aspergillus niger could be produced by these culture gas concentrations of acetaldehyde and of ethanol. Inhibition by this concentration of CO₂ was just significant in these tests. The effect on sporulation could be produced by the CO₂, but not by these concentrations of any of the other identified components.

The effect of Lepidium sativum seed germination could be produced by these culture gas concentrations of ethanol and of 3-methyl-butanol-1-ol to a lesser extent. Slight effects were also observed with a lowered O₂ concentration and with a raised CO₂ concentration but not with the other constituents.

INTRODUCTION

Dick & Hutchinson (1966) reported that gases from cultures of Saccharomyces cerevisiae inhibited the growth and/or sporulation of several species of fungi, and Hutchinson (1967) reported that they inhibited spore germination and prothallial growth of Pteridium aquilinum. Lösel (1964) summarized previous work on the biological effects of volatile fungal products. Robinson & Park (1966) and Robinson, Park & Garrett (1968) identified acetaldehyde as one of the volatile products from cultures of Fusarium oxysporum and of Rhizopus stolonifer which can inhibit fungal spore germination; they also identified ethanol amongst the nine otherwise unidentified peaks which they saw in gas-liquid chromatograms of samples from these cultures, but they found that it did not inhibit spore germination in their tests. They state that none of the volatile germination inhibitors affected the growth of hyphae 'to any marked degree' in their tests. Norrman (1968) examined the effects of vapours from aqueous solutions of a variety of organic compounds on the morphology of Pestalotia rhododendri. He found that spore production was stimulated by many compounds in the order of activity alcohols > esters > acids > aldehydes > ketones, and that high vapour concentrations of many of these substances often inhibited linear growth of
the colony. We now report on the analysis of the gas mixtures produced by *S. cerevisiae* in defined conditions, and on the identification of the active components.

### METHODS

**Analysis of the gases above yeast cultures**

Cultures of *Saccharomyces cerevisiae* Meyen ex Hansen (Glasgow University Collection I, referred to as 'yeast' throughout this article) were prepared by spreading a suspension of cells over the surface of 200 ml. of 5% malt agar (50 g. Oxoid malt extract + 20 g. Oxoid agar in 1 l. of deionized water) in each of a series of 1 l. Roux bottles. The cultures were then incubated at 30° in the dark for 48 h. Five ml, 1 ml. and 0.5 ml. were then withdrawn by syringe through the cotton-wool plug for analysis of organic volatiles, O₂ and CO₂ respectively. Similar samples were also taken from the assemblies used to examine the biological effects of these gases (described below).

The samples were analysed in an Aerograph Model 204 gas-liquid chromatograph, using a flame ionisation detector for organic volatiles and a micro cross-section detector for CO₂ and O₂, under the operating conditions shown in Table 1.

**Table 1. Operating conditions used for chromatographic analysis of yeast culture gases**

<table>
<thead>
<tr>
<th>Column</th>
<th>Dimensions</th>
<th>Material</th>
<th>Packing</th>
<th>Temp. (°C)</th>
<th>Carrier gas and flow rate (ml./min.)</th>
<th>Hydrogen flow rate (ml./min.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>10 ft × ⅓ in</td>
<td>S.S.*</td>
<td>Molecular sieve</td>
<td>65</td>
<td>He 22</td>
<td>—</td>
</tr>
<tr>
<td>B</td>
<td>14 in × ⅓ in</td>
<td>Copper</td>
<td>Davison 08 grade silica gel 30/60 mesh</td>
<td>50</td>
<td>He 30</td>
<td>—</td>
</tr>
<tr>
<td>C</td>
<td>5 ft × ⅓ in</td>
<td>S.S.</td>
<td>20% Carbowax 1500</td>
<td>65</td>
<td>N₂ 25</td>
<td>25</td>
</tr>
<tr>
<td>D</td>
<td>6 ft × ⅓ in</td>
<td>S.S.</td>
<td>15% Dinonyl phthalate</td>
<td>60</td>
<td>N₂ 30</td>
<td>30</td>
</tr>
<tr>
<td>E</td>
<td>6 ft × ⅓ in</td>
<td>S.S.</td>
<td>20% 1,2,3-Tris-(2-cyanoethoxy) propane</td>
<td>50</td>
<td>N₂ 30</td>
<td>30</td>
</tr>
<tr>
<td>F</td>
<td>6 ft × ⅓ in</td>
<td>Copper</td>
<td>25% glycerol</td>
<td>60</td>
<td>N₂ 30</td>
<td>25</td>
</tr>
</tbody>
</table>

60/80 mesh Chromosorb W was used as support in columns C-F.

* Stainless steel.

The constituents of the culture gases were identified by comparison with retention times of authentic pure samples and by syringe reactions (Hoff & Feit, 1964). The concentration of each identified metabolite was estimated by comparing the height of the peaks in representative samples with those of known concentration of authentic pure samples in parallel tests.

**Examination of the effects of culture gases on Aspergillus niger**

A suspension of conidia of *Aspergillus niger* van Tieghem (Glasgow University Collection 1) was made in 2% malt agar (20 g. Oxoid malt extract + 20 g. Oxoid agar in 1 l. deionized water), at approximately 45°, poured immediately into a Petri dish and incubated for 18 h. at 24°.

A series of 1 l. Roux bottles each containing 200 ml. of 2% malt agar were then each inoculated centrally with a 4 × 2 mm. disc cut from this suspension, the same
Volatile metabolites from yeast

suspension being used for all cultures in any one experiment. This inoculation method produced a single symmetrical colony in each vessel. All cultures were incubated at 24° in a low light intensity. The effects of complete yeast culture gases were examined by joining *Aspergillus niger* cultures and yeast cultures by means of a glass T-piece passing through a rubber bung in the neck of each Roux bottle. Gas samples were taken by a syringe through a serum cap closing the lateral arm of the T-piece. Control cultures were paired with Roux bottles containing uninoculated 5% malt agar. The effects of identified organic components of the yeast-culture gas mixture were examined by placing samples of authentic pure compounds (purity verified by GLC) on filter-paper strips in the necks of the Roux bottles, and replacing the cotton-wool plug by a serum cap. The weight of each sample was that required to give an initial vapour pressure of the metabolite in the sealed vessel equal to the highest found at any stage during the incubation of yeast cultures with *A. niger*, or with cress seedlings. Untreated filter papers and serum caps were added to the control cultures at the same time.

Changes in O₂ and in CO₂ content during each experiment in the sealed assemblies were measured using a micro cross-section detector and column A (O₂) or column B (CO₂). Atmospheres with comparable O₂ and CO₂ concentrations to the final ones were made by partially evacuating freshly inoculated Roux bottles and replacing the air by N₂ (for reduced O₂) or by authentic CO₂. Obviously by this method the N₂ content was increased and the initial CO₂ content was decreased in proportion to changes in the O₂ content. It is thought that these changes can be disregarded at this stage, particularly as the reduction of CO₂ concentration would be counteracted very quickly by respiration.

After 8 days incubation, linear growth of a colony was assessed by taking the mean of two diameters at right angles to each other, and any gross effects on sporulation were estimated by eye.

*Examination of the effects of culture gases on germination of cress seeds and on seedling growth*

Seeds of *Lepidium sativum* L. orient (cultivar Dobbies 'white curled' 1964, referred to as cress throughout this article) were sterilized by shaking in aqueous HgCl₂ (0.01%, w/v) for 1 min, followed by three changes of deionized water. Twenty sterilized seeds were placed in each of a series of 1 l. Roux bottles containing 200 ml. 0.5% Oxoid agar in 1 l. deionized water. These were then paired with yeast cultures, control bottles, or received authentic single substances by the methods described above. All cultures were held at 22–26° under continuous illumination from G.E.C. 'Warm White' fluorescent tubes.

After 5 days incubation a seed was recorded as germinated when it had produced a radicle as long as the diameter of the seed. Gross differences in the appearance of seedlings were noted by eye.

*R E S U L T S*

The only metabolites found in samples in sufficient concentration for identification were CO₂, acetaldehyde, ethyl acetate, ethanol, n-propanol, isobutanol, and isopen- tanols. Oxygen concentrations below atmospheric were also noted. They were all found in samples both from pure yeast cultures and from the paired assemblies during biological activity tests.
Figure 1 shows the chromatogram of a typical sample taken from a pure yeast culture after 2 days growth.

These are mostly common respiratory metabolites and their production is likely to be affected by many environmental factors. It was therefore deliberately decided to restrict the investigation to the study of their production and effects in the sealed assemblies used for the biological activity tests.

Table 2. Volatile organic compounds identified in gases above Saccharomyces cerevisiae cultures

<table>
<thead>
<tr>
<th>Constituent</th>
<th>Range of maximum concentration (μg/l. of air)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Acetaldehyde</td>
<td>260–320</td>
</tr>
<tr>
<td>Ethyl acetate</td>
<td>13–15</td>
</tr>
<tr>
<td>Ethanol</td>
<td>800–1000</td>
</tr>
<tr>
<td>n-Propanol</td>
<td>6–8</td>
</tr>
<tr>
<td>Isobutanol</td>
<td>12–16</td>
</tr>
<tr>
<td>2-Methyl-butan-1-ol</td>
<td>4–6</td>
</tr>
<tr>
<td>3-Methyl-butan-1-ol</td>
<td>8–10</td>
</tr>
</tbody>
</table>
Table 2 shows the range of maximum concentrations of each metabolite measured in these tests. Figure 2 shows the changes in concentration of each measured in a representative test with *Aspergillus niger*. The changes followed this pattern consistently within the different ranges of concentration of each metabolite found in the three experiments in which this was followed.

![Figure 2](image1.png)

**Fig. 2.** Changes in concentration of identified volatile metabolites in test assemblies of *Aspergillus niger* and yeast, expressed as % of concentration of each, present immediately after the cultures were paired. Acetaldehyde, ○; ethyl acetate △; ethanol, ●; n-propanol ▲; isobutanol, ×; 2-methyl-butan-1-ol + 3-methyl-butan-1-ol, +.

**Fig. 3.** Changes in concentration of authentic pure compounds in tests with *Aspergillus niger*, expressed as percentage of concentration present at the start of the tests. The initial concentration of each was equal to the highest found at any stage in the other activity tests. Symbols on graph as in Fig. 2, with oxygen, □. Concentrations in µg./l. of air: acetaldehyde, 320; ethyl acetate, 15; ethanol, 1000; n-propanol, 8; isobutanol, 16; 2-methyl-butan-1-ol + 3-methyl-butan-1-ol, 16.

The O₂ content of the sealed assemblies containing yeast and *Aspergillus niger* decreased from approximately 13 % (v/v) immediately after pairing to approximately 8 % (v/v), and the CO₂ content increased from approximately 15 % (v/v) to approximately 25 % (v/v).

Figure 3 shows the initial concentration of authentic pure compounds in the synthetic gas/air mixtures, and the changes in this concentration during a representative test of the effect of the mixture on *Aspergillus niger*. Figure 4 compares the effects of the total culture gases and of the synthetic air/gas mixtures on the size of the diameter of *A. niger* colonies after 8 days growth. Each experiment involved the measurement and comparison of at least three cultures in each treatment with at least three controls and in each case at least three replicate experiments have given similar results.
The diameters are reduced in the presence of complete culture gases and in the assemblies with an initial concentration of 320 μg/L acetaldehyde, or of 1000 μg/L of ethanol, or of 25% (v/v) CO₂. These concentrations are the maximum found in any tests of yeast culture gases; since acetaldehyde concentration falls off rapidly while that of ethanol varies relatively slightly (Fig. 3), then it appears that both can contribute to the effects of complete culture gases; it seems likely that CO₂ may also contribute to the effects of total culture gases in some conditions, although in these tests the concentration found only slightly exceeded the minimum inhibitory one (between 20% (v/v) and 25% (v/v)). Sporulation, assessed by eye, was consistently reduced by about half by the presence of complete culture gases, or of 25% (v/v) initial CO₂ concentration, and by about one-third in the presence of 20% initial CO₂ concentration. It seems likely therefore that CO₂ may contribute to the effects. No other identified compounds have been shown to have any significant biological effect on Aspergillus niger in the concentrations tested. The possibility that ethyl acetate and/or n-propanol can contribute to the effects of the total culture gases has not been eliminated, however, as their concentrations fall off rapidly in the tests of authentic pure compounds. Our reasons for deliberately deciding not to investigate this possibility further are given in the discussion below.

The O₂ content of the sealed assemblies containing yeast and cress seeds decreased from approximately 17% (v/v) immediately after pairing to approximately 10% (v/v), and the CO₂ content increased from approximately 7% (v/v) to approximately 20% (v/v). The range of concentrations of other identified metabolites found in any test is shown in Table 2, and the changes in concentration measured during a representative test are shown in Fig. 5. Figure 6 shows the initial concentration of pure compound in the synthetic gas–air mixture and the changes in this concentration during a representative test. Figure 7 shows the effects of total culture gases and of pure compounds on seed...
germination. Each experiment involved the measurement and comparison of at least four cultures in each treatment with at least four controls, and at least three replicate experiments have given similar results.

Germination is significantly reduced in the presence of complete culture gases, in assemblies with 10% (v/v) initial $O_2$, and in those with an initial concentration of 1000 $\mu g./l.$ ethanol or of 10 $\mu g./l.$ 3-methyl-butan-1-ol. Since this is the maximum concentration found in any tests of yeast culture gases, and since the ethanol and 2-methyl-butan-1-ol concentrations vary only slightly (Figure 6), then it appears that both these can contribute to the effects of the complete gases. The significant reduction of germination in experiments with acetaldehyde at 500 $\mu g./l.$ of air (i.e. $\times 1.5$ the highest concentration above yeast cultures) and in those using isobutanol at 30 $\mu g./l.$ of air (i.e. twice the yeast culture concentration) suggest that these might have an effect in some conditions.

The evidence that the concentration of acetaldehyde fell off rapidly during all these tests suggests, however, that it does not contribute to the effect in these conditions.
The possibility that ethyl acetate can contribute to the effect of the total culture gases has not been entirely eliminated, as its concentration falls off by 40% during the test of the authentic pure compound. Our reasons for deliberately deciding not to investigate this further are given in the discussion below.

The few seedlings which grew in cultures with complete culture gases or with ethanol, and those which grew with CO₂ (20%, v/v) were severely stunted and chlorotic; in most of these cultures growth was restricted to formation of a radicle a few millimetres long. They were less stunted and less chlorotic in cultures with acetaldehyde. The few seedlings which grew in the presence of 3-methyl-butan-1-ol were slightly stunted but similar in colour to those in the controls and to those in cultures with ethyl acetate, n-propanol and isobutanol.

**Fig. 7.** Effects of yeast culture gases on germination of cress seeds. At least three replicates were used in each experiment. The vertical axis represents the mean germination of test cultures expressed as a percentage of the mean germination of the controls. ↔ = least significant difference at 5% level. * = Differences clearly significant by empirical observation. Each section on the horizontal axis represents a group of experiments using, from left to right, complete culture gases, 10% O₂, 20% CO₂, acetaldehyde, ethyl acetate, ethanol, n-propanol, isobutanol, 2-methyl-butan-1-ol + 3-methyl-butan-1-ol, 2-methyl-butan-1-ol, and 3-methyl-butan-1-ol.

**DISCUSSION**

No unusual metabolites have been found in the yeast culture gases; their biological effects can be due to the production of a number of common respiratory metabolites which may be effective singly or as parts of a mixture. The results contrast with those of the study of *Fomes annosus* (Glen, Hutchinson & McCorkindale, 1966), which showed that the biological activity of this fungus is due to the production of a single unusual metabolite (hexa-1,3,5-triyne); the effects on hyphal growth of *A. niger* extend the work of Robinson *et al.* (1966, 1968), who report that acetaldehyde and ethanol had no such effects on the species which they examined in these conditions. Only a limited comparison can be made with the results of Norrman (1968), owing to differences in concentrations and conditions tested. The differences in response to ethanol may be related to the fact that the maximum concentrations he tested were less than one-fifth of those found in the yeast/Aspergillus assemblies and used in our tests of
authentic pure ethanol. In his tests with *Pestalotia rhododendri* acetaldehyde, *n*-propanol, and isobutanol stimulated sporulation, isobutanol and *n*-propanol inhibited linear growth, all at the concentrations similar to those at which they showed no appreciable effect on *Aspergillus niger* colonies in our tests of authentic pure compounds. He found that 0.1 % ‘iso-amyl alcohol’ inhibited sporulation; it is not, however, possible to compare this result with our tests, as he does not specify whether this is a mixture of 2-methyl-butan-1-ol and 3-methyl-butan-1-ol or pure 3-methyl-butan-1-ol. Neither *P. rhododendri* nor *A. niger* were apparently affected by similar concentrations of ethyl acetate.

The composition of the gas mixtures of these common metabolites from yeast will vary greatly in different cultural conditions and hence any detailed analysis of one condition would have very limited general biological application. At the present time it does not seem justifiable to continue the analysis of such obviously complex possible interactions and of the possible patterns in the cumulative effects of mixtures. The report is therefore published as an illustration of a complex interaction of a mixture of common respiratory metabolites which may have obvious ecological importance in some conditions—for example, in the ecology of soil micro-organisms.

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REFERENCES


