Development of Candida albicans Hyphae in Different Growth Media – Variations in Growth Rates, Cell Dimensions and Timing of Morphogenetic Events

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In six liquid culture media, all of which stimulated Candida albicans to grow in the hyphal form, the rates of hyphal extension and increase in cellular ATP concentration, hyphal diameters, times of evagination of hyphae, times of septum formation and positions of septa in the hyphae appeared to vary independently. There were no discernible associations between properties such as length or volume of hyphal compartments at the time of septation and temporal parameters of hyphal growth. The results suggest that growth environment influences many of the processes contributing to hyphal development, but that these processes are not necessarily interrelated.

INTRODUCTION

Considerable information has been published on the cell biological and molecular factors apparently involved in the morphological development of Candida albicans. Stationary-phase yeast-form cells of the fungus can be induced to form hyphal germ tubes in a variety of growth media (Odds, 1985). In developing germ tubes, new cell wall material is synthesized predominantly at the hyphal apex with minimal lateral expansion, whereas in yeast-form cells there is an additional component of generalized wall synthesis that leads to lateral expansion of the cells into an ovoid shape (Staebell & Soll, 1985). Intermediate forms between parallel-sided hyphae and swollen yeast cells indicate that regulation of lateral wall synthesis is a critical determinant of cell shape (Soll et al., 1985).

Hypha-form cell walls have a higher chitin content than those of yeast-form walls (Chattaway et al., 1968) and plasmalemma-bound chitin synthase activity has been found to be higher in developing C. albicans germ tubes than in yeast-form cells (Braun & Calderone, 1978; Chiew et al., 1980). In hypha-form cells the volume of the cytoplasmic contents appears to remain unchanged from that of the parent yeast cells and the cytoplasm stays in proximity with the developing hyphal apex, so that the hyphae consist of nucleated but vacuolated and collapsible cell compartments behind their growing tips (Gow & Gooday, 1984; Brawner & Cutler, 1985).

It is possible that regulation of wall structure and hence cell shape is a phenotypic property rather than the direct result of differential gene transcription (Soll, 1984). Certainly, germ tube development proceeds in the presence of actinomycin D (Oliver et al., 1982) even though this compound blocks induction of enzyme pathways (Bhattacharya et al., 1974a, b), and it proceeds despite a progressively lowered rate of protein synthesis (Torosantucci et al., 1984).

Soll and his coworkers have accounted for developmental events in C. albicans in terms of temporal and spatial controls of processes such as emergence of new cell wall material, septum formation and cell wall synthesis (Mitchell & Soll, 1979a; Bedell et al., 1980; Soll et al., 1981 b, 1985; Soll, 1984; Staebell & Soll, 1985).

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Abbreviations: AAS, amino acids/salts medium; EMEM, Eagle's minimal essential medium; MAM, methionine assay medium; MSAB, modified Sabouraud broth; NAG, buffered N-acetylglucosamine.
This brief summary appears to present an integrated account of *C. albicans* morphogenesis, but it is entirely eclectic in one important respect: the information comes from experiments with the fungus grown in or on more than five different media. Investigators have rarely studied the consistency of their observations in different hypha-stimulating environments, so it is possible that some of the observations and models in the literature may be based on data related to specific growth environments and are not generally applicable to the development of the fungus. We have therefore undertaken a study of the basic morphogenetic development of *C. albicans* hyphae in six culture media, with emphasis on events during the first 3 h of growth – the period that has received most previous attention.

**METHODS**

**Fungus and preparation of inocula.** *C. albicans* isolate 73/055 was used in all experiments. It was maintained on Sabouraud glucose agar (Oxoid Mycological Peptone 10 g l⁻¹, glucose 40 g l⁻¹, agar 20 g l⁻¹). For preparation of inocula, yeast cells were grown in M ycological Peptone 10 g l⁻¹, glucose 40 g l⁻¹, shaken at 160 r.p.m. at 37 °C for 24 h. The yeasts were harvested by centrifugation at 2500 g for 10 min, washed twice with 0.1 M citrate buffer pH 5.0 and resuspended in this citrate buffer overnight at 27 °C to starve the cells. Of several starvation procedures tested, citrate starvation gave the most homogeneous suspensions of single yeast cells for use as inoculum, in our hands.

The starved yeasts were washed twice with water, resuspended in water, and added to culture media to give an initial concentration of 10⁶ cells ml⁻¹.

**Growth media.** The six media and incubation conditions used to examine their effects on hyphal development of *C. albicans* were as follows: (1) amino acids/salts medium (AAS) (Lee et al., 1975); (2) buffered N-acetyl-D-glucosamine (NAG) (Shepherd et al., 1980); (3) horse serum (Difco); (4) methionine assay medium (MAM), comprising Difco methionine assay medium at 10·5 g l⁻¹, 10 ml; 0·2 M-HEPES pH 7·0, 10 ml; 0·5 M-NaHCO₃, 5 ml; water, 75 ml; (5) modified Sabouraud broth (MSAB) (Evans et al., 1975); and (6) Eagle’s minimal essential medium (EMEM) (prepared according to Odds et al., 1985). The first three media were incubated at 37 °C in air, shaken at 160 r.p.m. MSAB was shaken in air at 40 °C, and EMEM and MAM were incubated statically at 37 °C under 5% (v/v) CO₂ in air.

**Treatment of culture samples.** Samples were removed from the culture at zero time, then every 15–30 min from 1 h to 3 h, and every 30–60 min up to 7 h. ATP concentrations were measured immediately (Odds, 1982) to determine viable fungal biomass. The remainder of each sample was mixed with 0·1 vol. formaldehyde/sodium dodecyl sulphate preservative solution (Odds et al., 1985) and stored at room temperature for subsequent microscopic examination. Calcofluor was added to samples to a concentration of 200 µg l⁻¹ to facilitate microscopic visualization of septa.

**Measurement of cell dimensions and other properties.** For all samples from 1 h to 3 h the following were determined with the aid of a UV/phase contrast microscope equipped with a calibrated eyepiece graticule: the percentage of yeast cells bearing a structure recognizable as a hyphal evagination, the percentage of hyphae with discernible septa, the length and diameter of parent yeast cells, the length and diameter of hyphal outgrowths, and the distance between the septum and the parent yeast/hypha cell junction. For each sample, the percentages were determined by counting of at least 100 yeast/yeast-plus-hypha cell units and the dimensions were calculated from measurements of at least 10 cells. All experiments were repeated twice, and results presented are pooled data from the duplicate experiments.

**Calculation of morphogenetic parameters.** Growth rates of the fungi were calculated by least-squares regression analysis of plots of log(ATP concentration) vs time, hyphal extension rates by regression analysis of plots of hyphal length vs time. Hyphal volumes were calculated from the formula π(d/2)²(l − d/2) + (2π/3)(d/2)³, where *d* is hyphal diameter and *l* is hyphal length. Yeast cell volumes were calculated from the formula for a prolate spheroid: (πlw²)/6, where *l* is yeast cell length and *w* is yeast cell width.

**RESULTS**

The overall microscopic appearance of *C. albicans* germ tubes/hyphae varied little between the different media, although the lengths of hyphal outgrowths were not always the same at the same culture ages (Fig. 1). In NAG the germ tubes were more slender than germ tubes in the other five media, but in all other respects the hyphae looked similar – phase-dark, parallel-sided structures with a slight constriction at the junction with the parent yeast cell. In later samples, septa were particularly conspicuous in hyphae grown in NAG, but they were visible only with great difficulty in hyphae grown in AAS medium, even with the aid of Calcofluor and UV microscopy.
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Fig. 1. *C. albicans* hyphal development after 1.5 h in (a) AAS medium, (b) EMEM, (c) MSAB, (d) NAG, (e) serum, and after 3 h in (f) MAM. Bar, 5 μm.

![Hyphal development images](image)

Fig. 2. Evagination of hyphae from parent yeast cells inoculated into AAS medium (●), EMEM (□), MAM (▲), MSAB (▲), NAG (○) and serum (■).

![Evagination graph](image)

**Growth parameters of *C. albicans* in six liquid media**

*C. albicans* hyphae developed fairly synchronously in four of the culture media – EMEM, MAM, NAG and serum – but the emergence of hyphal evaginations in AAS and MSAB occurred over a relatively protracted period (Fig. 2).

Fig. 3 shows the exponential changes in fungal ATP concentration with time in the six growth media tested. Linear regression analysis of these plots between 1 h and 6 h showed viable biomass growth rates, in terms of ATP increase, ranging from 0.00 h⁻¹ in NAG to 0.80 h⁻¹ in MAM (Table 1). Lag phases in the six media were never more than 1 h (Fig. 3).

In all six media, *C. albicans* hyphae extended at a linear rate (Fig. 4). The lowest extension rate was 10.9 μm h⁻¹ in NAG, and the highest was 21.9 μm h⁻¹ in MAM (Table 1). There was thus some correlation between biomass growth rate and hyphal extension rate (Table 1).

The diameters of *C. albicans* hyphae were similar (approximately 2 μm) in five of the six media, but substantially less in NAG (Table 1). Mean diameters of the hyphae grown in MAM and NAG appeared to increase by 0.1–0.2 μm between 1.5 and 3 h of growth; measurements for hyphae in the other four media were consistent throughout this period of growth.

Measurements of the dimensions of the parent yeast cells showed these to have a constant volume of 50–70 μm³ throughout the first 3 h of growth (Table 1).

**Times of evagination and septum formation in different growth media**

Table 1 shows various measurements of developmental events calculated from experimental observations. The times at which 50% of the yeast cells in the culture bore hyphal outgrowths were calculated by regression analysis of the data for percentage hyphae vs time. They ranged from 40 to 72 min after inoculation. Regression analysis was also used to calculate the times at...
Fig. 3. ATP concentrations in *C. albicans* hyphae grown in AAS medium (●●●●●), EMEM (□□□□□), MAM (▲▲▲▲▲), MSAB (△△△△△), NAG (○○○○○) and serum (■■■■■). Data are means from duplicate experiments, normalized to an initial ATP concentration of 120 nM, with regression lines drawn.

Fig. 4. Hyphal extension of *C. albicans* grown in AAS medium (●●●●●), EMEM (□□□□□), MAM (▲▲▲▲▲), MSAB (△△△△△), NAG (○○○○○) and serum (■■■■■). Data are means ± SEM for at least 20 replicates. Regression lines are drawn.

**Table 1. Cell dimensions, growth rates and critical times for *C. albicans* hyphae grown in six different media**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>AAS</th>
<th>EMEM</th>
<th>MAM</th>
<th>MSAB</th>
<th>NAG</th>
<th>Serum</th>
</tr>
</thead>
<tbody>
<tr>
<td>Growth rate (h⁻¹) by ATP measurement</td>
<td>0-60</td>
<td>0-67</td>
<td>0-80</td>
<td>0-67</td>
<td>0-00</td>
<td>0-33</td>
</tr>
<tr>
<td>Hyphal extension rate (µm h⁻¹)</td>
<td>15-5</td>
<td>18-9</td>
<td>21-9</td>
<td>20-1</td>
<td>10-9</td>
<td>20-4</td>
</tr>
<tr>
<td>Hyphal diameter (µm ± SD, at 180 min)</td>
<td>2.0 ± 0.2</td>
<td>1.9 ± 0.1</td>
<td>1.9 ± 0.1</td>
<td>2.0 ± 0.3</td>
<td>1.2 ± 0.2</td>
<td>1.9 ± 0.1</td>
</tr>
<tr>
<td>Time 50% cells evacuated (T50, min)</td>
<td>73</td>
<td>40</td>
<td>50</td>
<td>66</td>
<td>72</td>
<td>63</td>
</tr>
<tr>
<td>Time of hyphal initiation*</td>
<td>34</td>
<td>36</td>
<td>44</td>
<td>42</td>
<td>37</td>
<td>47</td>
</tr>
<tr>
<td>Time 50% hyphae septated (S50; min)</td>
<td>195</td>
<td>173</td>
<td>180</td>
<td>158</td>
<td>189</td>
<td>142</td>
</tr>
<tr>
<td>Length of hyphae at T50 (µm)†</td>
<td>8-1</td>
<td>1-2</td>
<td>2-0</td>
<td>10-8</td>
<td>6-4</td>
<td>5-6</td>
</tr>
<tr>
<td>Length of hyphae at S50 (µm)†</td>
<td>41-6</td>
<td>43-1</td>
<td>49-6</td>
<td>41-6</td>
<td>27-6</td>
<td>32-5</td>
</tr>
<tr>
<td>Volume of hyphae at S50 (µm³)</td>
<td>130</td>
<td>121</td>
<td>140</td>
<td>130</td>
<td>31</td>
<td>91</td>
</tr>
<tr>
<td>Volume of parent yeast cells (µm³)</td>
<td>47-67</td>
<td>51-69</td>
<td>51-66</td>
<td>48-69</td>
<td>50-60</td>
<td>46-68</td>
</tr>
<tr>
<td>Distance between parent cell and septum</td>
<td>3-4 ± 2.4</td>
<td>5-6 ± 2.3</td>
<td>4-3 ± 1.5</td>
<td>6-3 ± 1.9</td>
<td>5-4 ± 2.3</td>
<td>6-0 ± 2.0</td>
</tr>
</tbody>
</table>

*Calculated by extrapolation of data in Fig. 4 to a hyphal length of 0.
† Determined from data in Fig. 4.

which 50% of the hyphae contained septa. This time ranged from 142 min in serum to 195 min in NAG (Table 1). Extrapolation of the regression curves for the data in Fig. 4 was used to calculate the mean time of initial emergence of hyphae (the time at which mean hyphal length was 0 µm). This time varied from 34 min in AAS medium to 47 min in serum.

**Distances between yeast cells and hyphal septa**

The distribution of distances between parent yeast cells and hyphal septa was sometimes slightly skewed. Table 1 shows the arithmetic means, modes and median values for this measurement. The data indicate that septa were generally positioned closer to the parent cells in AAS and MAM than in the other media, although the modal septal distance in NAG was also lower than for hyphae in serum, MSAB and EMEM.
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Table 1 lists the lengths and volumes of hyphal outgrowths calculated at the times when 50% of the cells had visible hyphal outgrowths and when 50% of the cells contained septa. For four of the media tested, MSAB, AAS, MAM and EMEM, the hyphal lengths and volumes at the time 50% of the hyphae had formed septa were similar (41–50 μm and 120–140 μm³ respectively), but the lengths and volumes of the hyphae at the 50% septation time in serum and NAG were less (Table 1).

DISCUSSION

The literature on morphogenesis in C. albicans contains many discrepancies and contradictions concerning dimensions and extension rates of hyphae. Mitchell & Soll (1979b) reported that hyphae in AAS had mean dimensions of 11.2 x 1.5 μm at the time of septation, much smaller than the 35.7 x 2.6 μm for hyphae grown on serum agar and measured at the same stage by Gow & Gooday (1982). The recorded hyphal extension rates in serum- or plasma-based media are 18.7 μm h⁻¹ (Gow & Gooday, 1982), 22.2 μm h⁻¹ (Wain et al., 1975) and 39.8 μm h⁻¹ (Davies & Denning, 1972). These figures are all higher than the 4.8 μm h⁻¹ found in AAS medium by Soll et al. (1981b).

The data in the present study agree fairly closely with most published data for C. albicans hyphae grown in serum: we found an extension rate of 20.4 μm h⁻¹ and a hyphal dimension of 32.5 x 1.9 μm at the time of septation (Table 1). For AAS medium, our figures do not agree at all with those in the literature: we measured the extension rate as 15.5 μm h⁻¹, and the dimensions at the time of 50% septum formation as 41.6 x 2.0 μm. It is difficult to believe that differences of this order of magnitude can be explained simply by strain differences between the isolates tested, and even differences of technique should not lead to such discrepancies with relatively simple microscopic measurements.

One possible explanation is that the nutrient status of the C. albicans yeast cells used in our experiments differed from that of the yeasts used by Soll et al. (1981b). Soll (1985) has emphasized the significance of zinc starvation of the yeast cells on subsequent developmental events. Whereas in zinc-starved cells there is a delay of approximately 135 min before 50% of the cells grow a discernible hyphal evagination, in cells unstarved of zinc this time is reduced to 70–80 min (Soll et al., 1981a) – a time of precisely the same order as the 73 min recorded in the present study (Table 1). Data concerning septation times and cell dimensions for hyphae grown from a non-zinc-starved inoculum have not been published: however, since our starvation protocol for the yeast cells was not designed to ensure a zinc-depleted inoculum it seems likely that this factor accounts for some, at least, of the differences between our data and those published from Soll's laboratory. The water used in our experiments was distilled, not deionized, and we have experienced extreme difficulty in establishing reproducible conditions for zinc starvation in this laboratory.

Since we observed a large difference in hyphal diameter between cells grown in NAG medium as compared with cells from the other five media tested, we conclude that C. albicans is able to alter its hyphal diameter in response to environmental pressures: specifically in response to adverse nutrient conditions, since NAG supported hyphal development but not an increase in viable biomass. It is therefore possible that hyphae grown from inocula which are starved of zinc, like hyphae grown in NAG, attain a diameter less than the approximately 2 μm seen in all the nutrient-rich media. This would account for the differences between the hyphal diameter in AAS measured by Mitchell & Soll (1979b) and the larger diameters measured by ourselves (Table 1) and by Gow & Gooday (1982).

We could find no relation between hyphal length or volume and the time at which 50% of hyphae contained septa, nor was there a constant time between hyphal initiation (whether determined as the time at which hyphal length was 0 or $T_{50}$ – see Table 1) and the time at which 50% of hyphae contained septa. The distances between septa and parent yeast cells appeared to be independent of the other parameters measured or calculated. In common with Mitchell & Soll (1979b) we found an uneven distribution of septum location distances from parent yeast cells.
Our mean septal distance of 3.8 μm (Table 1) is greater than the mean of 2.2 μm reported by Mitchell & Soll (1979b). More importantly, the distances to the septa in cells grown in AAS were markedly smaller than the corresponding distances in hyphae grown in the other five media.

Overall, the rates of cell development, the dimensions of hyphae, the positions of septa and the timings of hyphal emergence and septum formation appear to be independent of each other, but they are all influenced by the growth environment. The mechanisms by which these different processes are coordinated and controlled remain to be elucidated.

REFERENCES


