The Rate of Elaboration of the Extracellular Polysaccharide, Pullulan, during Growth of Pullularia pullulans

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Summary

During the growth of the yeast-like fungus Pullularia pullulans, the pattern of glucose utilization by the cell varied from a complete inability to elaborate extracellular polysaccharide to the diversion of almost two-thirds of metabolized glucose to pullulan elaboration. The effects of carbon and nitrogen levels on this changing pattern during growth were examined. These investigations indicated that the elaboration of pullulan, and the onset of a morphological change of Pullularia from a filamentous to a yeast-like form, were related to the availability of nitrogen, and not carbon, in the growth medium.

Introduction

Pullulan, an extracellular α-glucan elaborated by the yeast-like fungus Pullularia pullulans, is predominantly α-maltotriose linked endwise through 1,6-bonds (Wallenfels, Keilich, Bechtler & Freudenberger, 1965). During an investigation into the ability of the organism to elaborate pullulan, Catley (1971b) observed that, at an early phase of growth, there appeared to be little or no production of the polysaccharide, whereas at a later stage pullulan was synthesized. These analyses were made at two arbitrarily chosen stages of growth. This communication describes changes in the organism’s patterns of glucose utilization and polysaccharide elaboration which occur throughout the course of its growth, and relates these patterns to the availability of nitrogen and carbon in the medium.

Methods

Quartermaster strain no. 3092 of Pullularia pullulans, kindly supplied by E. T. Reese of the U.S. Army Natick Laboratories, Natick, Massachusetts, U.S.A., was maintained and grown as described previously (Catley, 1971a). The carbon source of the culture medium was glucose at 35, 140 or 560 mM; nitrogen sources were 0, 4.5 or 22.5 mM-ammonium sulphate. All cultures contained 0.4% yeast extract (Difco). Rates of uptake of glucose and extracellular polysaccharide elaboration were measured as previously described (Catley, 1971b). Samples of cells, withdrawn from cultures growing under different nutritional conditions, were washed and suspended in a standard uptake medium containing: glucose, 14 mM; 0.008 μCi of uniformly labelled [14C]glucose (New England Nuclear Corp.)/ml; K₂HPO₄, 29 mM; NaCl, 17 mM; MgSO₄·7H₂O, 0.8 mM; (NH₄)₂SO₄, 4.5 mM, which was then adjusted to pH 5.0 with HCl. The incorporation of 14C, measured for 90 min, was linear for the first hour and often for the total period. Extracellular polysaccharide elaboration was determined

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at the same intervals and, when it occurred, was linear after an induction period (Catley, 1972) of about 30 min. Each uptake or elaboration rate, presented in Fig. 1 to 3(b), is therefore based on four $^{14}$C analyses. In addition, data presented in these Figures are the combination of analyses from two cultures grown under identical conditions, but with different sampling times. Periodically the pullulan content of the extracellular polysaccharide was determined by hydrolysing the elaborated polymer with pullulanase (Catley, 1971a). The action of amyloglucosidase from Aspergillus niger (Abdullah, Fleming, Taylor & Whelan, 1963) on the polymer was also examined. The chromatographically separated maltotriose, produced by pullulanase, or glucose, released by amyloglucosidase, accounted for 80% of the elaborated polymer.

Glucose was measured by the glucose-oxidase reagent (Lloyd & Whelan, 1969) and growth was recorded as the dry weight of cells (Catley, 1971a). Extracellular pullulan was measured by a technique involving precipitation with alcohol, digestion with pullulanase, and measurement of the maltotriose released (Catley, 1971a).

Cells were fractionated into yeast and hyphal forms by centrifugation on a mannitol gradient (0.12 M to 1.5 M) at 100 g for 15 min at room temperature and were then washed twice with distilled water before use.

**RESULTS**

The organism's ability, as a rate/mg dry wt cells, to assimilate labelled carbon from $[^{14}]$C-glucose and to elaborate pullulan over the major period of the growth cycle is expressed in Fig. 1. The culture medium contained 140 mM-glucose and 4.5 mM-ammonium sulphate. The organism's potential for producing pullulan varied considerably during growth from a complete inability to elaborate extracellular polysaccharide at 12 h to the diversion of approximately two-thirds of metabolized glucose to pullulan production at 37 h. The rate of assimilation of glucose also varied. Increasing either the glucose or ammonium sulphate concentration fourfold or decreasing it to 1/4 did not alter the $^{14}$C-distribution pattern of the cells over the 90 min period used in the uptake analysis. In the period used for determining assimilation and elaboration rates the increase in weight of cells suspended in the uptake medium was no more than 10% when sampled at the fastest phase of growth and considerably less at others. An environment of pH 5 ensured that pullulan elaboration was expressed whenever possible (Catley, 1971b). The results obtained from these studies of the uptake medium indicate that for the short assay period the composition of the uptake medium did not alter the distribution of $^{14}$C between cell and extracellular polysaccharide.

Despite a 16-fold change in the initial glucose concentration from 35 to 560 mM, the growth characteristics of Pullularia, at least over the first 30 h, remained similar (Fig. 2a, b). The distribution pattern of glucose utilization seemed little affected by the external glucose concentration.

The change in growth rate over the first 24 h was accompanied by a morphological change. The inoculum from a 48 or 72 h culture was yeast-like, but in fresh medium rapidly became filamentous, thereafter reverting to the original single-cell form. At this transitional stage the two forms were separated by centrifugation on a mannitol gradient. The hyphal cells sedimented more rapidly and only the yeast cells produced extracellular polysaccharide.

A fivefold increase or complete removal of the ammonium ion (leaving only yeast extract as nitrogen source) gave the results shown in Fig. 3(a) and (b). Table 1 describes the cell density at which the organism, grown under different conditions of nutrition and environmental pH, reached a maximum in glucose utilization rate. From microscopic observation of the cells during growth, the rate of glucose assimilation is probably a monitor of the cell's
Growth and pullulan elaboration

Fig. 1. Growth of Pullularia pullulans in medium containing 140 mM-glucose and 4.5 mM-ammonium sulphate. □, Dry cell wt (mg/ml); ○, extracellular glucose (μmoles/ml); △, extracellular pullulan (μmoles anhydroglucose/ml); ■, rate of ¹⁴C accumulation expressed in terms of glucose assimilation (μmoles/mg cell/h); ▲, rate of extracellular polysaccharide elaboration (μmoles anhydroglucose/mg cell/h).

Table 1. Effect of pH, nitrogen and glucose levels of culture medium on the cell concentration where the rates of glucose utilization and pullulan elaboration were maximal in cultures of Pullularia

<table>
<thead>
<tr>
<th>Initial concentrations (mM) of glucose and (NH₄)₂SO₄</th>
<th>pH of cultures unadjusted</th>
<th>Initial pH adjusted to 5.1</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Cell density* at</td>
<td></td>
</tr>
<tr>
<td></td>
<td>maximal glucose</td>
<td></td>
</tr>
<tr>
<td></td>
<td>utilization rate</td>
<td></td>
</tr>
<tr>
<td>Glucose (NH₄)₂SO₄</td>
<td>35 140 560 140 140</td>
<td>4.5 4.5 4.5 0 22.5</td>
</tr>
<tr>
<td></td>
<td>0·4 0·5 0·5 0·15 0·8</td>
<td>0·5</td>
</tr>
<tr>
<td></td>
<td>Cell density* at</td>
<td></td>
</tr>
<tr>
<td></td>
<td>maximal pullulan</td>
<td></td>
</tr>
<tr>
<td></td>
<td>elaboration rate</td>
<td></td>
</tr>
<tr>
<td></td>
<td>1·7 1·4 1·6 0·4 2·7</td>
<td>1·4</td>
</tr>
</tbody>
</table>

* mg dry wt/ml.

morphological state, reflecting a faster growth for the filamentous form of Pullularia. If this change is indeed monitored by the rate of glucose uptake per unit weight of cell, the levels of nitrogen, but not carbon, determine the cell density at which this maximum utilization rate is achieved (Table 1).

Pullulan was produced with the onset of a decrease in culture growth rate in each of the five nutritional conditions presented (Fig. 1 to 3). However, pullulan production declined at a later stage of growth. The nitrogen level of the medium seemed to determine pullulan
Fig. 2. Growth of *Pullularia pullulans* in medium containing initial glucose concentrations of (a) 35 mM; (b) 560 mM. For key see Fig. 1.

Table 2. Utilization of pullulan by *Pullularia* growing in conditions of glucose limitation

<table>
<thead>
<tr>
<th>Sample time (h)</th>
<th>Cell density†</th>
<th>Pullulan (c.p.m./ml)</th>
<th>Cells (c.p.m./ml)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>1660</td>
<td>0</td>
</tr>
<tr>
<td>19</td>
<td>0·8</td>
<td>1522</td>
<td>17</td>
</tr>
<tr>
<td>42</td>
<td>2·0</td>
<td>1530</td>
<td>16</td>
</tr>
<tr>
<td>63</td>
<td>1·8</td>
<td>1350</td>
<td>19</td>
</tr>
<tr>
<td>87</td>
<td>1·9</td>
<td>1390</td>
<td>16</td>
</tr>
</tbody>
</table>

* Medium containing glucose (10 mM) and [14C]pullulan (10 mM-anhydroglucose).
† mg dry wt/ml.

formation since the maximum pullulan-elaboration rate for cultures with high initial levels of nitrogen occurred at a cell density some 6·7 times greater than those relying solely on yeast extract for nitrogen (Table 1).

The distribution of 14C incorporation into the cells and pullulan was examined with cultures grown in medium with an initial pH of 5·1 instead of 7·4. Although the pH fell to 2·9 during the 3 days of growth there were essentially no differences from those cells commencing growth at the higher pH, and the cell densities at which maximum glucose utilization and pullulan elaboration rates occurred were the same (Table 1).
Growth and pullulan elaboration

Fig. 3. Growth of *Pullularia pullulans* in medium containing initially (a) 22.5 mM-ammonium sulphate or (b) no ammonium sulphate. For key see Fig. 1.

The utilization of pullulan by *Pullularia* growing in conditions of glucose limitation was studied. In medium with initially 10 mM-glucose (in contrast to the normal 140 mM) and containing \(^{14}\text{C}\)pullulan, there was a negligible uptake of radioactive carbon by the cells and little disappearance of label from the medium (Table 2).

**DISCUSSION**

The appearance of extracellular pullulan does not parallel the increase in cell mass of growing *Pullularia* cultures (Catley, 1971a, b); the accumulation of extracellular pullulan is linear with time during a phase of the culture where there is rapid multiplication. If ability to elaborate pullulan, once initiated, remained constant, extracellular polysaccharide should accumulate at an increasing rate over this period. Elaboration of pullulan during the major growth period might vary or pullulan might be both produced and reutilized by the organism, and an imbalance of these two processes would lead to the accumulation of the polysaccharide observed. *Pullularia* does not normally metabolize pullulan (Catley, 1970; Table 2). Thus in conditions of glucose limitation the cells seemed unable to utilize their own extracellular product; and this alternative rationale for the observed rate of pullulan elaboration must be dismissed.

Pullulan was not elaborated at all stages of growth, and it is clear from the results presented here and earlier (Catley, 1971b) that the level of nitrogen present in the medium is
critical to glucose utilization patterns. The change to the yeast form indicates the point at which the ability to elaborate pullulan appears. The inhibition of polysaccharide elaboration observed at a later phase of growth is again related to nitrogen availability. This decrease contrasts with observations on other micro-organisms. Thus in *Escherichia coli* (Sigal, Cattaneo & Segel, 1964) limitation in both sulphur and nitrogen sources leads to an accumulation of internal polysaccharide reserves.

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REFERENCES


