Inducible and Constitutive Formation of β-Fructofuranosidase (Inulase) in Batch and Continuous Cultures of the Yeast Kluyveromyces fragilis

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Inulase production by Kluyveromyces fragilis on various fermentable and non-fermentable carbon sources was examined in carbon-limited continuous culture. Fructose and sucrose supported superior inulase yields [above 24 μmol sucrose hydrolysed min⁻¹ (mg cell dry wt)⁻¹ at pH 5.0, 50 °C], while some other carbon sources, including lactose, galactose, ethanol and lactate, did not stimulate inulase formation beyond basal levels. Thus fructose was identified as the primary physiological inducer. Isolation of a constitutive mutant also provided genetic evidence for the inducible nature of inulase in the wild-type. The mutant was generated spontaneously and selected in continuous culture. It produced high inulase activities in continuous culture irrespective of the carbon source. Inulase formation in the wild-type and mutant strain was further controlled by general carbon catabolite repression as suggested by enzyme yield patterns in batch and continuous culture.

INTRODUCTION

The non-specific β-fructofuranosidase inulase is an external glycoprotein produced by a number of yeasts and filamentous fungi, among them Kluyveromyces (formerly Saccharomyces) fragilis (Snyder & Phaff, 1960), Kluyveromyces lactis (Kovaleva & Yurkevich, 1973), Kluyveromyces marxianus (Kierstan & Bucke, 1977), Debaromyces cantarelli (Beluche et al., 1980), Candida kefyr (Negoro & Kito, 1973), Aspergillus and Penicillium spp. (Kim, 1975; Nakamura et al., 1978). The yeast K. fragilis appears to give the highest enzyme yields (Kovaleva & Yurkevich, 1973; Negoro, 1978; GrootWassink & Fleming, 1980). Inulase rapidly splits terminal β-2,1- and β-2,6-fructofuranosidic bonds, which are present in sucrose, raffinose, stachyose and oligofructosides of the inulin or levan type (Snyder & Phaff, 1960). This broad substrate spectrum renders the enzyme relatively non-specific in comparison with the other extensively studied, yeast β-fructofuranosidase, invertase. This latter enzyme acts rapidly on terminal β-fructofuranosidic bonds in disaccharides and trisaccharides but has an extremely low affinity (very high $K_m$) for saccharides with a higher degree of polymerization (GrootWassink & Fleming, 1980).

Since yeast cell membranes are impermeable to fructosides, including sucrose (Barnett, 1976; Sutton & Lampen, 1962), the primary physiological function of inulase dictates an external location. The ratio of cell wall-bound inulase to soluble inulase depends on the cultivation conditions. High enzyme levels in yeast batch cultures have been observed only if the media contained oligofructosides such as inulin (Beluche et al., 1980; Negoro & Kito, 1973; Snyder & Phaff, 1960). However, experimental limitation of carbon catabolite repression by using the continuous culture technique has shown that the rapidly metabolizable substrate sucrose can also support high yields (GrootWassink & Fleming, 1980).

In this paper we describe further efforts to identify environmental and genetic factors regulating inulase formation in K. fragilis. The range of culture conditions supporting enzyme production has been greatly expanded.
METHODS

Organisms. The homothallic diploid yeast (Van Der Walt, 1970) Kluyveromyces fragilis ATCC 12424 (wild-type), and its spontaneous mutant strain PRL Y53A selected in continuous culture (see Fig. 4), were used. Genetic stability of stock cultures was safeguarded by preserving freshly grown vegetative cells of single-colony isolates in silica gel stored at 4 °C (Fink, 1970). Inocula for experimental liquid cultures were grown by spreading a loopful of the silica gel mixture on an agar slant (10 ml of 5 g yeast extract l⁻¹, 20 g glucose l⁻¹ and 20 g Bacto agar l⁻¹ in a 25 x 95 mm vial) and incubating for 3-4 d at 28 °C.

Liquid cultures. The media generally consisted of 5 g yeast extract l⁻¹ (Anheuser Busch, St Louis) to supply nitrogen, minerals and vitamins, plus one of the following carbon sources at a concentration of 10 g l⁻¹: sucrose (Sigma, or technical grade beet sugar), glucose, fructose, mannose, galactose, lactose, inulin, glycerol, lactate, succinate, malate (all from Sigma) or ethanol (Fisher). In one experiment, the defined nutrients present in Yeast Nitrogen Base (Difco) were substituted for the complex yeast extract.

Batch cultures (50 ml) were grown in 500 ml Erlenmeyer flasks incubated at 28 °C on a rotary shaker (280 r.p.m.). Continuous cultures were grown at 30 °C in a 7.5 l bench top fermenter (Microferm; New Brunswick Scientific). Sterile medium was pumped into the fermenter from Nutrient Vessel; New Brunswick Scientific). Spent culture and air were removed from the fermenter through a fixed overflow tube. Dissolved oxygen tension in continuous cultures was maintained around 20% of saturation by automatic adjustment of the impeller speed (300-500 r.p.m.) and aeration rate (0.5-3.0 l min⁻¹), employing a dissolved oxygen controller (model DO 81-14; New Brunswick Scientific). The pH of continuous cultures was maintained at 4.2 by automatic addition of M-HCl. Culture volumes varied with agitation and aeration rates and ranged from 2800 to 3000 ml. After experiments were completed at each dilution rate, the actual volumes were measured by interrupting agitation and aeration momentarily and reading a calibration scale on the fermentation vessel. Only minor fluctuations in agitation and aeration rates occurred during steady-state growth and had a practically negligible effect on the culture volume.

All media were inoculated with water-suspended cells from agar slant cultures at a ratio of one slant l⁻¹. Continuous cultivations were begun after initial batch growth in the fermenter had just reached the stationary phase as indicated by a rapid increase in dissolved oxygen tension. Cultures were grown at a fixed or sequentially increased dilution rate.

Analyses. Sets of shake flask cultures were sampled by randomly withdrawing flasks from the shaker. Continuous cultures were sampled by collecting 25-50 ml spent culture in containers cooled in ice. Inulase activity was completely stable during sample collection and processing. To ensure that growth and enzyme biosynthesis had reached a steady-state in continuous culture, multiple sampling at 8-16 h intervals was performed after at least 3-5 culture volumes had passed through the fermenter at each dilution rate examined. For measuring inulase content, 0·1-1 ml samples of fresh whole culture samples were diluted 10-100 fold with acetate buffer (0·1 M, pH 5·0) and kept frozen until assayed. For monitoring growth, samples were Millipore filtered (pore size 0·45 μm) and the dry weight of the yeast mat was determined after freeze-drying.

Inulase was assayed by incubating appropriately diluted whole culture samples (0·1 ml) at 50 °C with 4% (w/v) sucrose (Sigma) in 0·9 ml 0·1 M-acetate buffer, pH 5·0. Periodically, 0·1 ml samples were withdrawn from the reaction mixture and analysed for an increase in total reducing sugars by the neocuproine method (Dygert et al., 1965). Thus, total inulase content could be measured since cell wall bound and soluble enzyme are both accessible to the substrate sucrose. Activity on inulin was occasionally verified but required a more elaborate assay procedure including cell disruption to minimize substrate diffusion limitations. The ratio of hydrolysis rates with 4% (w/v) sucrose and 4% (w/v) inulin was 10·5, regardless of the source of inulase. Inulase activities are expressed as units mg⁻¹, defined as μmol sucrose hydrolysed min⁻¹ (mg cell dry wt)⁻¹.

RESULTS

Continuous culture of the wild-type yeast

Time course analysis of a batch culture of K. fragilis ATCC 12424 grown on inulin had previously shown a gradual increase in specific inulase activity until a maximum of 10 units mg⁻¹ was reached at the end of the growth phase (GrootWassink & Fleming, 1980). A more sudden appearance of inulase synthesis was observed when shake flask cultures were grown on glucose, fructose or sucrose: 80% of the total of only 2 units mg⁻¹ was formed at the very end of the growth phase over a period of 2-4 h. Considering that growth in these cultures was carbon-limited, it appeared that low medium concentrations of carbon substrates, which prevailed at the end of the growth phase, were prerequisites for inulase synthesis. The higher initial concentrations of the carbon sources were probably causing catabolite repression. One way to limit this repression throughout the cultivation period is by use of the continuous culture
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Fig. 1. Inulase activities in continuous cultures of *K. fragilis* wild-type grown on various carbon substrates. (a) Sucrose (1%, w/v). (b) Glucose (1%, w/v). (c) ○, Fructose (1%, w/v); ●, mannose (1%, w/v). (d) Fructose (0·5%, w/v) plus glucose (0·5%, w/v). (e) ○, Fructose (0·5%, w/v) plus galactose (0·5%, w/v); ---, lactose, galactose, ethanol or lactate (each at 1%, w/v; results very similar).

Continuous cultures of the wild-type yeast were grown on a number of fermentable and non-fermentable carbon substrates over a range of dilution rates (Fig. 1). The most striking result from this substrate screening was that high inulase activities (24 units mg⁻¹) were produced on media containing fructose, present either bound in sucrose (Fig. 1a), alone (Fig. 1c) or in a mixture with other substrates (Fig. 1d, e). Glucose (Fig. 1b) and mannose (Fig. 1c) supported much lower activities (7 and 4 units mg⁻¹, respectively), while lactose, galactose, ethanol and lactate gave basal activities of only 0·1-0·2 units mg⁻¹ (Fig. 1e). Maximum inulase activities were strongly dependent on the dilution rate. The sucrose-grown culture (Fig. 1a) gave the highest activities at the lowest dilution rate (0·06 h⁻¹). Cultures grown on free fructose showed a distinct inulase peak between 0·11 and 0·14 h⁻¹ (Fig. 1c, d, e). These inulase profiles were reproducibly obtained under the standard operating conditions involving sequentially increased dilution rates. However, complications arose when a continuous culture, growing on fructose alone as carbon source, was passed through dilution rates in the reverse direction. The inulase yield rose dramatically to 42 units mg⁻¹ as the dilution rate was decreased from 0·23 h⁻¹ to 0·09 h⁻¹ (Fig. 2).

To test the possibility that the complex nitrogen source yeast extract (which also provides small amounts of carbon) itself had a specific and major effect on inulase production, the defined and simple nitrogen source Yeast Nitrogen Base was used as a replacement in one experiment. It was complemented with glucose as the carbon source because of its intermediate capacity to support inulase. The results (Fig. 3) were similar to those presented earlier (Fig. 1b) for the
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Fig. 2

Inulase activities as a function of sequentially decreased dilution rates in a continuous culture of *K. fragilis* wild-type grown on fructose.

Fig. 3

Inulase activities in a continuous culture of *K. fragilis* wild-type grown on minimal medium containing 0.25% (w/v) Yeast Nitrogen Base and 0.5% (w/v) glucose.

culture grown on yeast extract. This indicated that the yeast extract used did not contain any substances overriding or interfering with the unique role of the primary carbon source in the regulation of inulase formation.

Selection of constitutive mutants in continuous culture

It was shown above that maximum inulase yields did not vary greatly if fructose alone (Fig. 1c) was replaced by equal mixtures of fructose and glucose (Fig. 1d) or fructose and galactose (Fig. 1e) as carbon sources. This finding suggested that perhaps only a minor proportion of the carbon source was needed in the form of fructose to effect high induction of enzyme synthesis. Thus, an experiment was done to test a range of fructose/galactose concentration ratios. Galactose was chosen as the supplementary carbon source because of its inability to stimulate inulase formation. The experiment was begun by first establishing high inulase levels in a culture growing on fructose alone at a dilution rate of 0.14 h⁻¹. To achieve this, the culture was passed through a few successively higher dilution rates beginning at 0.08 h⁻¹ prior to fixing the rate at 0.14 h⁻¹. Subsequently, the feed medium was changed at appropriate intervals to contain increasingly larger fractions of galactose (Fig. 4). The data show that a less than proportional drop in inulase occurred until a fructose/galactose concentration ratio of 3:7 was reached. Further replacement of fructose by galactose had the unexpected effect of causing a rise in the inulase yield until it reached 25 units mg⁻¹ with galactose as the sole carbon source. The most attractive explanation of this phenomenon was that selection of a mutant population which no longer depended on fructose for inulase formation had taken place. To see whether this newly emerged population consisted of clones with a stably maintained constitutive phenotype, cells were plated and colonies picked for separate propagation on agar slants. The majority (80%) had markedly improved inulase productivity over the wild-type strain in shake flask culture, suggesting that they were mutants. Continual propagation during 12 serial transfers of a
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Fig. 4. Selection of an inulase-constitutive cell population in a continuous culture of *K. fragilis*. The culture was grown at a fixed dilution rate of 0.14 h⁻¹. The composition of the carbon source in the feed medium was changed stepwise from 1% (w/v) fructose to 1% (w/v) galactose. Representative isolate, designated PRL Y53A, on agar slants did not result in any loss of the enhanced inulase productivity, indicating genetic stability of the new trait.

Shake flask culture of an inulase-constitutive mutant

A comparison of inulase yields in shake flask cultures of the wild-type and mutant PRL Y53A is presented in Table 1. Cultures were grown on fermentable and non-fermentable carbon sources. Without exception, all carbon sources supported much higher inulase activities in the mutant than in the wild-type. The increase ranged from 2-fold on inulin to 30-fold on succinate. Maximum activities of 32 units mg⁻¹ were obtained on the poor carbon source glycerol. In addition to testing for this overall effect on the final yield, two carbon sources, glucose and ethanol, were tested for their effect on the kinetics of inulase formation. As in the wild-type strain, the readily metabolizable substrate glucose completely repressed inulase synthesis in the mutant during exponential growth (Fig. 5). Exhaustion of glucose and limited post-exponential growth, presumably on secondary products and some components of the yeast extract, caused a rapid rise in inulase activity. The much more slowly metabolizable and non-fermentable carbon source ethanol supported enzyme formation during the entire growth phase, with activities reaching eight times higher values in the mutant (Fig. 6). However, some inulase repression was still apparent in these ethanol-grown cultures since the specific activity increased during the course of growth.

These results indicated that inulase synthesis in the mutant, as in the wild-type, is under the control of carbon catabolite repression. However, the second control mechanism, involving induction, appeared to be inoperative in the mutant. This conclusion was based on the finding that lactose, galactose, ethanol and lactate supported enormously higher activities (50–170 times higher) in the mutant grown in shake flask cultures, as compared to the wild-type grown in continuous culture on the same substrates under non-repressive conditions.
Table 1. *Inulase yields of shake flask cultures of K. fragilis wild-type and mutant PRL Y53A*

The media contained yeast extract at 0.5% (w/v) and carbon substrate at 1.0% (w/v) except in the case of lactate, succinate and malate which were added at 0.5% (w/v) in combination with yeast extract at 0.25% (w/v). Cells grown on carbohydrates were harvested at 24 h; the remainder were harvested at 40 h.

<table>
<thead>
<tr>
<th>Carbon substrate</th>
<th>Wild-type</th>
<th>PRL Y53A</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inulin</td>
<td>10</td>
<td>19</td>
</tr>
<tr>
<td>Sucrose</td>
<td>2</td>
<td>9</td>
</tr>
<tr>
<td>Glucose</td>
<td>2</td>
<td>7</td>
</tr>
<tr>
<td>Fructose</td>
<td>2</td>
<td>6</td>
</tr>
<tr>
<td>Lactose</td>
<td>1</td>
<td>11</td>
</tr>
<tr>
<td>Galactose</td>
<td>1</td>
<td>9</td>
</tr>
<tr>
<td>Glycerol</td>
<td>3</td>
<td>32</td>
</tr>
<tr>
<td>Ethanol</td>
<td>2</td>
<td>17</td>
</tr>
<tr>
<td>Lactate</td>
<td>0.3</td>
<td>5</td>
</tr>
<tr>
<td>Succinate</td>
<td>0.3</td>
<td>10</td>
</tr>
<tr>
<td>Malate</td>
<td>0.4</td>
<td>14</td>
</tr>
</tbody>
</table>

* Defined as μmol sucrose hydrolysed min⁻¹ (mg cell dry wt)⁻¹.

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Fig. 5. Time course of shake flask cultures of a constitutive mutant of *K. fragilis* grown on glucose. ---, Mutant PRL Y53A; ----, wild-type; ○, cell density; □, glucose concentration.

Fig. 6. Time course of shake flask cultures of a constitutive mutant of *K. fragilis* grown on ethanol. ---, Mutant PRL Y53A; ----, wild-type.
Continuous culture of an inulase-constitutive mutant

If indeed the hyperproducing mutant PRL Y53A was constitutive, it would be expected to produce high levels of inulase in continuous culture on all utilizable carbon sources. Special interest was focused on those substrates that had previously failed to give more than basal levels in continuous culture of the wild-type. The results in Table 2 show that enzyme yields of the mutant cultures grown on lactose, galactose, ethanol and lactate were similar to those obtained in wild-type cultures grown on the inducing substrates sucrose and fructose. The inulase levels were inversely related to the dilution rate, as exemplified in Fig. 7 for a mutant culture grown on galactose. The maximum activities observed were as high as 42 units mg⁻¹ in a culture grown on glucose at the relatively low dilution rate of 0·08 h⁻¹ (Table 2).

Table 2. Inulase yields of continuous cultures of K. fragilis wild-type and mutant PRL Y53A

<table>
<thead>
<tr>
<th>Carbon substrate</th>
<th>Dilution rate (h⁻¹)</th>
<th>Dry wt (mg ml⁻¹)</th>
<th>Inulase activity (units mg⁻¹) *</th>
<th>Wild-type</th>
<th>PRL Y53A</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sucrose</td>
<td>0·11</td>
<td>6·0</td>
<td>18</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Fructose</td>
<td>0·11</td>
<td>5·0</td>
<td>20</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Glucose</td>
<td>0·08</td>
<td>5·0</td>
<td>7</td>
<td>42</td>
<td></td>
</tr>
<tr>
<td>Glucose</td>
<td>0·11</td>
<td>5·5</td>
<td>6</td>
<td>26</td>
<td></td>
</tr>
<tr>
<td>Lactose†</td>
<td>0·17</td>
<td>5·6</td>
<td>0·1</td>
<td>22</td>
<td></td>
</tr>
<tr>
<td>Galactose</td>
<td>0·11</td>
<td>5·4</td>
<td>0·2</td>
<td>26</td>
<td></td>
</tr>
<tr>
<td>Ethanol</td>
<td>0·08</td>
<td>5·5</td>
<td>0·1</td>
<td>18</td>
<td></td>
</tr>
<tr>
<td>Lactate</td>
<td>0·11</td>
<td>4·2</td>
<td>0·1</td>
<td>22</td>
<td></td>
</tr>
</tbody>
</table>

* Defined as μmol sucrose hydrolysed min⁻¹ (mg cell dry wt)⁻¹.
† Cells grown on lactose below a dilution rate of 0·14 h⁻¹ showed an unstable constitutive phenotype.

Fig. 7. Inulase production in a continuous culture of a constitutive mutant of K. fragilis grown on galactose. ——, Mutant PRL Y53A; ——, wild-type.
The results clearly confirmed that the *K. fragilis* mutant does not require an inducer for inulase synthesis and thus is constitutive.

**DISCUSSION**

In micro-organisms, many catabolic enzymes with hydrolytic activity are subject to simultaneous induction and repression by a single growth substrate (Demain, 1971; Reese, 1972). Maximum enzyme expression depends on the optimum balance between the contribution of the two regulatory mechanisms. Both systems are strongly concentration dependent. Full induction of a gene requires saturation of a DNA-binding protein with a special inducer molecule, while minimum repression requires the lowest possible intracellular concentrations of certain substrate catabolites. A distinction must be made between a physiological inducer and a molecular inducer acting at the gene level. A physiological inducer might be the growth substrate itself or an enzymic break-down product thereof, as is the case with polymeric substrates. A molecular inducer is often a metabolic derivative of the physiological inducer. Non-metabolizable synthetic analogues may also be functional as molecular inducers. Only in very few instances have the natural molecular inducers been identified, the classical example being allolactose derived from lactose for β-galactosidase induction in *Escherichia coli* (Barkley *et al.*, 1975). Cellulase induction in the mould *Trichoderma reesei* is believed to be mediated by sophorose derived from cellobiose which in turn is a product of cellulose hydrolysis (Sternberg & Mandels, 1979).

The object of this work was to gain insight into the regulation of inulase formation by carbon substrates. The continuous culture technique was relied upon to provide much of the information. Profiles of several bacterial enzymes developed by determining the yields at different dilution rates had been interpreted by other workers to reflect the involvement of induction and/or repression mechanisms (Clarke *et al.*, 1968; Dean, 1972; Bull, 1972). Similarly, in our study the peaking of inulase levels at intermediate dilution rates in cultures grown on media containing fructose could imply a balance between induction and repression (Fig. 1). Thus, a lack of inducer molecules predominated at the lower dilution rates and an excess of repressors at the higher rates. However, this interpretation became somewhat ambiguous after finding that inulase activities only peaked if the order of dilution rates was from low to high. In the converse order a totally different profile was observed, with the highest inulase yields produced at the lowest dilution rate (Fig. 2). Also contrary to expectations, when a culture which had passed through dilution rates from low to high was returned to a low rate, very high inulase activities (~40 units mg⁻¹) were observed. This difference between the trajectory of ascent and descent is referred to as hysteresis and has previously been found to occur in continuous culture (Tanner, 1978). Its origin might lie in phenotypic and genotypic adaptations that vary with culture history. It is well known that continuous cultivation imposes tremendous selective pressures.

The rate of displacement of a cell population by a better adapted population is a function of the complex interaction between the rate of cellular changes (mutational and/or physiological) and increased specific growth rate. Every microbial continuous culture system has its unique selection characteristics, and no generalizations as to the time required for adaptation can be made. Although several genotypic adaptations in continuous cultures of bacteria are known (Harder *et al.*, 1977), to date only a few such adaptations of yeasts have been reported. Downie & Garland (1973) observed the emergence of a variant population of cells in about 48 generations (7 d) in a copper-limited continuous culture of *Candida utilis*. The variants showed constitutive formation of a terminal oxidase alternative to cytochrome oxidase. In an ongoing study of the evolution of metabolic pathways, P. E. Hansche and co-workers are focusing on the genotypic adaptations of *Saccharomyces cerevisiae* in phosphate-limited continuous cultures. Mutations involving amino acid substitutions that effected a shift in the pH optimum and an increase in substrate affinity of acid phosphatase were observed after 180 generations (35 d) and 400 generations (77 d), respectively (Francis & Hansche, 1972). Mutants with increased acid phosphatase activities were isolated in another experiment after 400 generations, and were shown to result from gene duplication and transposition (Hansche, 1975; Hansche *et al.*, 1978).
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After 1000 generations a cell population emerged with a mutation in a repressor gene, resulting in constitutive formation of acid phosphatase (Lange & Hansche, 1980). These examples show that it is difficult to affirm whether our cultures of *K. fragilis* on fructose lasted long enough (30–40 generations) to allow genotypic adaptation. However, considering the clear-cut mutant selection that took place during only approximately 40 generations of growth on the mixed carbon substrates fructose and galactose (Fig. 4, discussed below), it cannot be ruled out that genotypic adaptation was also at least partially responsible for the observed hysteresis effect in the fructose-grown continuous cultures.

Other results which suggest that fructose may be the primary physiological inducer are the superior inulase yields on this carbon source in continuous culture. The fact that fructose is the breakdown product of inulase action on the natural oligomeric substrate inulin (Snyder & Phaff, 1962) makes this monomer the obvious candidate. Thus, our experimental findings agree with this putative regulatory function of fructose. The next logical question is whether fructose itself is also the natural molecular inducer acting at the gene level. No direct answers may be expected from a physiological study like this. However, the observation that glucose and mannose also stimulate inulase formation could be explained in terms of a common glycolytic intermediate of the three monosaccharides, i.e. the actual gene inducer molecule. The marked differences between the inducing capacities of these sugars suggest that the pool size of the common intermediate is of critical importance. Mannose, which is catabolized like fructose except for a single isomerization step, gave only one-seventh as much inulase as fructose. Furthermore, galactose and lactose totally lacked induction capability, even though lactose contains a glucose moiety and both sugars presumably enter the glycolytic pathway after initial epimerization of the galactose.

The most convincing evidence for the broader conclusion that inulase is an inducible enzyme in wild-type *K. fragilis* comes from the isolation of a regulatory mutant which produced inulase constitutively. As long as catabolite repression was restricted, this mutant formed high levels of inulase regardless of the carbon source.

It is not obvious which forces were responsible for the selection of inulase constitutive mutants in the continuous culture in which fructose was gradually replaced by galactose as carbon substrate (Fig. 4). Development of mutant populations constitutive for substrate-capturing enzymes has been noted in continuous cultures (Harder *et al.*, 1977; Lange & Hansche, 1980), but since inulase was not thought to have a fructose-capturing function, the present selection was totally unexpected. If inulase indeed has an affinity for fructose, mutant cells that contain large amounts of enzyme in the cell wall may have an advantage due to the higher fructose concentration in the micro-environment just outside the cell membrane. However, binding of fructose would probably cause competitive inhibition of enzymic activity in vitro. This has neither been reported nor observed in our studies. Another explanation for selection could be that enzymes or transport proteins involved in fructose metabolism are controlled by the same regulatory mechanism as inulase. Similar co-regulation of enzymes of galactose and melibiose utilization has been observed in *Saccharomyces carlsbergensis* (Kew & Douglas, 1976). Thus, mutants with enhanced fructose metabolism would show gratuitous hyperproduction of inulase.

Finally, it is tempting to compare the regulation of inulase formation to that of the very closely related and extensively studied yeast β-fructofuranosidase, invertase. At present, this enzyme is widely considered to be non-inducible and only controlled by carbon catabolite repression (Lampen, 1971; Zimmerman & Scheel, 1977; Hackel & Khan, 1978). However, conclusive biochemical and genetic evidence seems to be scanty. The conclusion is based mainly on the repressive effects of glucose and sucrose observed in continuous culture experiments using *K. fragilis* and *Saccharomyces cerevisiae* (Davies, 1956; Gascon & Ottolenghi, 1972). Similar effects were observed on addition of several other carbon compounds to resting-cell cultures (Dodyk & Rothstein, 1964). In the light of the present study on inulase, the data in these reports do not unequivocally exclude the involvement of an induction mechanism in the control of invertase formation. Simultaneous induction and repression by metabolic derivatives of the growth substrates, or lack of induction capability by the added compounds in the respective studies,
could account equally well for the results. Induction of invertase has been recognized by Toda (1976) in a more recent publication on invertase formation in continuous cultures of *S. carlsbergensis*. Also, Zimmerman & Scheel (1977), in a genetic study of *S. cerevisiae* mutants resistant to carbon catabolite repression, could not explain all their results without noting that perhaps repression is not the only mechanism controlling invertase formation.

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