An Expanded Concept for the Glucose Effect in the Yeast *Saccharomyces uvarum*: Involvement of Short- and Long-term Regulation

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When *Saccharomyces uvarum* was cultivated in continuous culture, it exhibited the typical growth behaviour of a glucose-sensitive yeast. Metabolic changes related to glucose-repressed growth were assessed by an analysis of overall culture parameters (biomass formation, ethanol and acetate production and gas exchange rates) and by measuring the mitochondrial cytochrome content. These functions were mainly affected by the glucose effect; the steady state values of these variables were first established in the chemostat as a function of dilution rate.

The short- and long-term regulation taking place when the cells were submitted to repression was assessed by administering glucose pulses and by shifts in the dilution rate. The primary response of the cells to the initiation of repressed growth was the formation of ethanol and acetate. Since there was no repression of oxygen uptake rate or cytochrome content prior to this response, it was concluded that ethanol and acetate formation was not the consequence of repression of respiratory activity, but resulted from the regulation of pyruvate dehydrogenase and pyruvate decarboxylase activities. Long-term adaptation of the cells occurred within 24 to 48 h of the initiation of repressed growth as manifested by a decrease of mitochondrial cytochrome content to the steady state value corresponding to that of repressed growth.

**INTRODUCTION**

Glucose metabolism in glucose-sensitive yeasts is known to have some remarkable features. Respiration, the formation of gluconeogenetic enzymes, as well as the synthesis of enzymes and transport systems involved in disaccharide utilization, are repressed in the presence of glucose (Linnane et al., 1972; Perlman & Mahler, 1974; Witt et al., 1966; Polakis & Bartley, 1965; Duntze et al., 1969; van Wijk et al., 1969; Zimmermann & Eaton, 1974). Numerous biochemical studies concerning the so-called glucose effect in yeast have been published, but the mechanisms involved are still unknown. Genetic information is also limited. All the regulatory mutants obtained so far affect only certain segments of the whole complex of carbon catabolite repression (Montene court et al., 1973; Zimmermann & Eaton, 1974; Schamhaart et al., 1975; Rytka et al., 1976; Ciriacy, 1978; Entian, 1980), or lead to a rather diffuse pattern of regulation (Ciriacy & Breitenbach, 1979). In no case have the genetic elements and physiological effectors been identified which regulate repression and derepression of mitochondrial functions and enzymes.

The slow progress in the elucidation of mechanisms involved in glucose repression seems to be caused by a general absence of comprehensive cultivation techniques used in these studies. For instance, in order to establish in batch culture glucose-repressed and -derepressed states, substrate changes are required which generally involve centrifugation procedures. Furthermore, few data have been reported on the kinetics and the sequence of events observed that underly a classical repression/derepression description of the glucose effect. In our studies we took
advantage of the continuous culture technique for establishing glucose-repressed and -derepressed cells.

By pulsing glucose into the culture and by applying dilution rate shift techniques in the chemostat, it was possible to demonstrate that a short-term (on/off) regulation of ethanol formation and a long-term adaptation of the mitochondrial cytochrome content occurs when the cells are submitted to repressed growth.

METHODS

Organism. Saccharomyces uvarum H2055 (from this Department), a glucose-sensitive yeast, was used in all experiments.

Medium. The synthetic medium had the following composition [mg (g glucose)\(^{-1}\)]: (NH\(_4\))\(_2\)SO\(_4\), 200; (NH\(_4\))\(_2\)HPO\(_4\), 64; KCl, 30; MgSO\(_4\) \(\cdot\) 7H\(_2\)O, 15; CaCl\(_2\) \(\cdot\) 2H\(_2\)O, 10; FeCl\(_3\) \(\cdot\) 6H\(_2\)O, 0.5; ZnSO\(_4\) \(\cdot\) 7H\(_2\)O, 0.3; MnSO\(_4\) \(\cdot\) 2H\(_2\)O, 0.6; CuSO\(_4\) \(\cdot\) 5H\(_2\)O, 0.08; mezo-inositol, 2; calcium pantothenate, 1; nicotinic acid, 0.2; pyridoxine. HCl, 0.5; biotin, 0.001. This medium was carbon-limited as judged by an X\(^{-}\)S\(_0\) diagram and was used throughout this work. Glucose was used at concentrations of 30 and 60 g l\(^{-1}\). The medium was heat-sterilized at 121°C, 20 min) after reducing the pH to 2.5-3.0 with concentrated hydrochloric acid. For continuous cultivation, the sterilization was carried out in 300 l batches.

Culture conditions. The organisms were grown in a 10 l bioreactor (Chemap AG, Mändorf, Switzerland) equipped with a flat store turbine stirrer. The working volume was kept constant at 3 l by a syphon. A pH of 5-5 was automatically maintained with 10 mol NaOH l\(^{-1}\). The reactor was aerated at a flow rate of 300-480 l h\(^{-1}\) corresponding to 1.7-2.7 vol. air per vol. liquid min\(^{-1}\). The applied aeration rates yielded oxygen partial pressures above 30% of air saturation. The temperature was constant at 30°C and for agitation, a stirring speed of 900 r.p.m. was used. For a constant medium flow during continuous cultivation, Watson-Marlow MHRE 100 flow inducers (Falmouth, U.K.) were used.

Pulse and shift technique. The volume of the glucose solution pulsed into the fermenter (no other medium components were added) was between 3 and 5% of the total working volume. Addition of the pulse was carried out under sterile conditions to cells growing in steady state. Shifts in dilution rate were achieved by increasing the flow rate of the inflowing medium.

Sampling. In order to minimize the sample volume, samples were withdrawn with a syringe whose needle was inserted into the reactor, through a membrane, into the culture liquid.

determination of biomass, glucose, ethanol and acetate. For measuring the biomass concentration, 10 ml culture liquid was centrifuged at 2000 g for 5 min in previously tared centrifuge tubes. The sediment was washed twice with distilled water and then dried to constant weight at 105°C.

Glucose was determined in a glucose anaalysier (YSI, Yellow Springs, Ohio, U.S.A.) with a detection limit of 100 mg l\(^{-1}\). Ethanol and acetate were analysed by gas chromatography (Hewlett Packard gas chromatograph 5830A, Avondale, Pa., U.S.A.). Prior to injection, the cells were removed by centrifugation and known amounts of methanol and propionic acid were added as internal standards. The columns were as follows: for ethanol, Propack QS, mesh 100-120, stainless steel, 6 ft (1.8 m); for acetate, 10% SP-1200, 1% H\(_3\)PO\(_4\) on 80/100 Chromosorb WAW, glass column, 6 ft. For both columns the temperature was 150°C.

Gas exchange rates. The specific oxygen uptake rate (Q\(_{O_2}\)) and carbon dioxide production rate (Q\(_{CO_2}\)) were calculated from the data provided by a gas analyser (Hartmann & Braun, F.R.G.) connected to the air outlet of the reactor, and were based on an inert gas balance.

determination of cytochrome content. Cytochromes were determined according to Gmünder et al. (1981).

RESULTS AND DISCUSSION

Growth and respiratory activity of S. uvarum in continuous culture

The yeast S. uvarum showed the typical growth pattern of a glucose-sensitive yeast (Fiechter et al., 1981). Up to a dilution rate of 0.16 h\(^{-1}\) the breakdown of glucose was purely oxidative (Fig. 1). Biomass and CO\(_2\) were the only products formed.

The R.Q. was slightly below unity, indicating a stochiometric oxidation of glucose. The oxygen uptake rate and carbon dioxide production rate of the culture increased linearly with specific growth rate to a maximum value of approximately 5.5 mmol g\(^{-1}\) h\(^{-1}\) at a dilution rate of 0.16 h\(^{-1}\). The yield of biomass on glucose was 0.47 at low dilution rates and increased to 0.53 at 0.16 h\(^{-1}\). The glucose uptake rate at a dilution rate of 0.16 h\(^{-1}\) was 2 mmol g\(^{-1}\) h\(^{-1}\).

The region of growth rates above 0.16 h\(^{-1}\) was characterized by a decrease in biomass yield (yield = approximately 0.1) and the accumulation of glucose and ethanol in the medium. The
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Fig. 1. Influence of the dilution rate on steady state biomass (■), glucose (▲) and ethanol (●) concentration, the specific oxygen uptake rate (QO₂) (○) and carbon dioxide production rate (QCO₂) (□) as well as the respiratory quotient (R.Q.) (▼) in a chemostat culture of *S. uvarum* with a reservoir glucose concentration *S₀* = 55 g l⁻¹.

increased CO₂ production rate led to a R.Q. value above 2, indicating that glucose was only partly oxidized. The oxygen uptake rate remained at the level reached at the dilution rate of 0·16 h⁻¹. This is in accordance with the observations made by Barford & Hall (1979) who found no repression of respiration in *Saccharomyces cerevisiae* grown under glucose excess conditions. Previous data on *S. cerevisiae* as summarized by Fiechter *et al.* (1981) are currently being re-examined. Differences in the metabolic behaviour of the cells seem to be possible and are mainly due to the composition of the medium used by the different authors (O. Käppeli, unpublished observations). Steady states were established up to a dilution rate of 0·25 h⁻¹, indicating a critical dilution rate of approximately 0·26 h⁻¹ for *S. uvarum*.

**Steady state cytochrome content**

Cytochrome c increased sharply with increasing dilution rate under glucose-depressed growth conditions (Fig. 2). Its steady state content was approximately doubled when the dilution rate changed from 0·04 to 0·15 h⁻¹. Cytochrome b also increased with increasing dilution rate up to values where glucose repression became effective. Cytochrome a showed no uniform correlation with the dilution rate. From the data of the oxygen uptake rate and the cytochrome content, no direct relationship can be established between the two parameters. The oxygen uptake rate increased linearly from 1·5 mmol g⁻¹ h⁻¹ at *D* = 0·04 h⁻¹ to 5·5 mmol g⁻¹ h⁻¹ at *D* = 0·16 h⁻¹, whereas the content of any cytochrome was maximally doubled. Hence, it is not possible to attribute a particular cytochrome content to a certain oxygen uptake rate.

When the dilution rate was increased above 0·16 h⁻¹, the steady state content of all cytochromes dropped considerably. It is, however, remarkable that the content of cytochromes b and c under glucose-repressed conditions corresponded roughly to that at low dilution rates (below 0·1 h⁻¹) in the glucose-derepressed state. Cytochrome a was the only cytochrome whose lowest level was clearly under glucose repression. Again, the change in cytochrome content is not reflected in the oxygen uptake rate. The oxygen uptake rate did not decrease above the dilution of 0·16 h⁻¹, but instead remained approximately constant.
Kinetics of glucose repression

Both the short-term and long-term kinetics of glucose repression were investigated. The short-term responses were determined after administering a glucose pulse to glucose-derepressed cells. The long-term kinetics were assessed by an upward shift in dilution rate from the glucose-derepressed region into the glucose-repressed one. The response of the culture to these changes was recorded by monitoring the specific gas exchange rates, ethanol, acetate and biomass concentrations, as well as the mitochondrial cytochrome contents.

Glucose pulse

The response of the overall culture parameters on a glucose pulse is shown in Fig. 3. After the pulse, ethanol and acetate were formed immediately. Acetate always occurred when repressed growth was initiated, but disappeared later on and was never observed under steady state conditions (see Fig. 5). According to this partly incomplete oxidation of glucose, the R.Q. value rose above unity and increased steadily as long as glucose was present. The biomass concentration did not change significantly over the whole period. The course of the oxygen uptake rate clearly indicated that there was definitively no repression of overall respiration. The oxygen uptake rate increased as was the case for the carbon dioxide production rate, however to a lesser extent. Of the cytochrome contents, only that of cytochrome c slightly decreased (Fig. 4). For all the other cytochromes, no change occurred.

From the data of the glucose pulse (Figs 3 and 4), it can be concluded that the change from completely to partly oxidative glucose metabolism in \textit{S. uvarum} is not the consequence of repression of respiration, but begins before any decrease in the overall respiration or cytochrome content of the cells takes place. Therefore, we postulate that ethanol and acetate formation, respectively, are primarily governed by a regulation of the two enzymes pyruvate decarboxylase and pyruvate dehydrogenase, which stand at the beginning of two competitive pathways proceeding from pyruvate. The former leads to ethanol formation and the latter yields acetyl-CoA, the link to complete oxidation. The change to ethanol formation seems to be the consequence of a limited respiratory capacity of the cells (maximum oxygen uptake rate of approximately 5-5 mmol g\(^{-1}\) h\(^{-1}\)), leading to an increased pyruvate pool concentration and the subsequent degradation of pyruvate via the pyruvate decarboxylase pathway.
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Fig. 3. Short-term response of S. uvarum to excess glucose as initiated by a glucose pulse to derepressed-growing cells. ■, Biomass; ▲, glucose; ●, ethanol; Δ, acetate; ○, O2, □, CO2, ▼, R.Q. The dilution rate of the chemostat was 0.1 h⁻¹ and the initial glucose concentration $S_0 = 30$ g l⁻¹. At zero time 25 g glucose l⁻¹ was added.

Fig. 4. Short-term response of the mitochondrial cytochrome content of S. uvarum to excess glucose as initiated by a glucose pulse to derepressed-growing cells. The conditions of the chemostat culture were the same as in Fig. 3.

Shift in dilution rate

In order to elucidate the long-term adaptation of S. uvarum to repressed growth conditions, a shift in dilution rate from 0.14 to 0.21 h⁻¹, i.e. from derepressed to repressed growth, was carried out (Fig. 5). The biomass dropped within 45 h to the level of repressed growth. Ethanol production started immediately after the dilution rate shift and again acetate accumulated at the beginning of the initiation of repressed growth. The R.Q. increased continuously from unity to approximately three after 45 h. Again, the oxygen uptake rate showed no repression after the shift and remained at the level of the corresponding steady state rate (see Fig. 1). The change in R.Q. was mainly due to the increased carbon dioxide production rate which increased from 5 to about 15 mmol g⁻¹ h⁻¹ in the new steady state.
Fig. 5. Long-term adaptation of *S. uvarum* to repressed growth as initiated by a shift in dilution rate from derepressed to repressed growth conditions. ■, Biomass; ▲, glucose; ●, ethanol; △, acetate; ○, \( Q_0 \); □, \( Q_{CO_2} \); ▼, R.Q. The initial glucose concentration \( S_0 = 55 \) g l\(^{-1}\).

Fig. 6. Long-term adaptation of the mitochondrial cytochrome content of *S. uvarum* to repressed growth as initiated by a shift in dilution rate from derepressed to repressed growth conditions.

The content of all measured cytochromes dropped to a lower level (Fig. 6). It is, however, remarkable that the decrease of mitochondrial cytochrome content started at approximately 2 h after the shift in dilution rate. This substantiates the evidence from the glucose pulse that the formation of ethanol is not the consequence of a repression of mitochondrial functions, but
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...occurs before any change in respiration is observed, as measured by the oxygen uptake rate or by the cytochrome content.

There is, however, a long-term adaptation of the cells to repressed growth which takes place within 24 to 48 h after its introduction. A slow decrease of the mitochondrial cytochrome content is characteristic for this adaptation process. Considering the relatively long time period required for the mitochondrial cytochrome content to reach the level corresponding to repressed growth, it is difficult to attribute this adaptation to a complete cessation of mitochondrial cytochrome synthesis. Such a repression would lead to a faster disappearance of the cytochromes at the dilution rate of 0.21 h⁻¹. However, this behaviour could originate from a gradually diminishing synthesis of mitochondrial cytochromes.

The data presented above show that a proper analysis of the growth of a glucose-sensitive yeast under defined conditions yields a remarkable insight into the phenomenon generally referred to as glucose repression, a term that certainly needs to be re-evaluated. This type of approach would be of great importance to genetic work in the area. The conclusions often drawn from growth experiments done in complex media where many factors besides glucose may affect enzyme regulation are questionable. In view of our results, it becomes evident that regulation of overall metabolism based exclusively on a repression/derepression pattern inevitably leads to more confusion than clarity, as has been demonstrated by Ciriacy & Breitenbach (1979). More comprehensive work which takes into account all possibilities for metabolic regulation needs to be done in order to elucidate the particular behaviour of glucose-sensitive yeasts.

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REFERENCES


