Influence of Nitrogen Sources on Glycogen Metabolism in Saccharomyces carlsbergensis

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Storage of glycogen in yeast (Saccharomyces carlsbergensis) cells was strongly suppressed by the presence of nitrogen sources. Peptone initiated glycogen breakdown within minutes. This effect could not be duplicated with ammonium ions alone nor with single amino acids or mixtures of a few amino acids, but it could be duplicated by the addition of all the amino acids in the molar ratios found in casein. If fewer amino acids were supplied, glycogen was initially synthesized but then depleted after 30–60 min at 28 °C, the time being shorter for more easily utilized amino acids. In the presence of glucose, dinitrophenol (10^{-4} M) slowed down or inhibited glycogen synthesis, but it did not initiate glycogen breakdown in the absence of glucose. Energy charge and the concentrations of adenylates and UDP-glucose were not significantly different in the presence and absence of peptone; pyruvate accumulated under the former condition. No turnover of glycogen was observed during glycogen synthesis, pointing to an effective regulation of glycogen metabolism via an unknown mechanism.

INTRODUCTION

Glycogen is a major storage product in yeast, enabling the cells to survive long starvation periods. The glycogen content of the cells changes drastically in response to physiological state. For example, during sporulation the cells first synthesize and then degrade glycogen (Fonzi et al., 1979); in continuous culture, glycogen content depends on the growth rate (Küenzi & Fiechter, 1972). Glycogen synthesis initiated during transition of a batch culture of Saccharomyces cerevisiae (Rothman & Cabib, 1969) or S. carlsbergensis (Becker et al., 1979) to growth is probably caused by nutrient depletion (Lillie & Pringle, 1980). By contrast, stationary phase cells of an inoculum rapidly degrade glycogen after transfer to media in which growth is limited by either nitrogen or carbohydrate content (Becker et al., 1979). Since in these cases depletion of glycogen takes place in the presence of a high concentration of glucose, other factors present in the medium must initiate the breakdown. In this paper, the nature of these factors is explored and some of their possible modes of action are investigated.

METHODS

Growth and preparation of yeast. The experiments were initiated with an inoculum of stationary phase cells of Saccharomyces carlsbergensis (ATCC 9080) obtained from cultures grown aerobically under carbohydrate-limited conditions for 24 h at 28 °C (Becker et al., 1979) yielding 22 ± 1 g yeast l^{-1}. The cells were harvested by centrifugation at 3000 g for 5 min, washed twice in cold basic medium (see below) and finally suspended as a 2% (w/v) suspension in cold media as specified in the experiments described. The wet weight of the cells after centrifugation serves as the basis for the cell concentration in the suspensions. The suspension was first kept in ice for 20–40 min and then incubated in a water bath at 28 °C for 10 min. The experiments were started by the addition of 100 mM-D-glucose.
Determination of glycogen and metabolites. Samples (2 ml) were withdrawn and injected into 1 ml 15% (w/v) trichloroacetic acid to stop metabolism rapidly. After 20–80 min on ice, the cells were centrifuged; glucose was determined enzymically in the supernatant (Bergmeyer, 1974), and glycogen content was assayed in a portion of the cell pellet suspended in 0.25 M Na₂CO₃ (Becker, 1978).

Metabolites were determined enzymically according to Bergmeyer (1974). UDP-glucose was assayed with UDP-glucose dehydrogenase (EC 2.7.1.12), pyruvate with lactate dehydrogenase (EC 1.1.1.27), ATP with hexokinase (EC 2.7.1.1) and glucose-6-phosphate dehydrogenase (EC 1.1.1.49), and ADP and AMP with pyruvate kinase (EC 2.7.1.40), lactate dehydrogenase and myokinase (adenylate kinase; EC 2.7.4.3). For extraction of metabolites trichloroacetic acid was replaced by perchloric acid (15%, w/v). The cells were extracted for 15 min at room temperature with occasional shaking. Pyruvate was determined immediately in the supernatant after neutralization with buffered KOH (Becker, 1978). The remaining supernatant was frozen in small samples for later enzymic analysis. Repeated freezing and thawing led to low recoveries of UDP-glucose. For glycogen determination in these cells, the perichloric acid was removed from the cell residues by two water washes prior to suspension in carbonate solution.

Determination of glycogen turnover. Cells were washed twice with 50 mM potassium phosphate buffer, pH 4-6, aerated for 4 h at 28 °C to decrease the glycogen concentration and centrifuged. They were resuspended in the same buffer. After 10 min aeration at 28 °C, 20 mM-[U-¹³C]glucose (21000 d.p.m. μmol⁻¹) was added and the cells were incubated for 20 min. They were spun down (3000 g, 2 min), washed in 50 mM potassium phosphate buffer, then resuspended in the original volume of the buffer and aerated. To one half of the suspended cells, 100 mM unlabelled glucose was added, while the second half was only aerated. At intervals samples were withdrawn, extracted with trichloroacetic acid and washed once with the solution. The cell residues were suspended in 0.25 M Na₂CO₃, kept in a boiling water bath for 90 min, spun down, and washed with the same solution and twice with 0.2 M sodium acetate buffer, pH 4.8. This treatment dissolves only 5–10% of the glycogen. Samples were incubated with amyloglucosidase; in blanks the enzyme was omitted. In the supernatant the concentration (enzyme assay) and radioactivity (liquid scintillation counting) of the glucose liberated were determined.

Media and chemicals. Basic medium was the semi-synthetic medium used for culturing the cells (Gosh et al., 1960) but without nitrogen sources and glucose. It contained the following (per litre): KH₂PO₄, 0.55 g; KCl, 0.42 g; MgSO₄·7H₂O, 0.125 g; CaCl₂·2H₂O, 0.125 g; FeCl₃·6H₂O, 1.25 mg; MnSO₄·H₂O, 2.50 mg; myo-inositol, 10-00 mg; calcium pantothenate, 3.50 mg; pyridoxal. HCl, 0.25 mg; thiamin. HCl, 0.25 mg; biotin, 0.03 mg. The pH was adjusted to 5.0 with 5-0 g tripotassium citrate. H₂O and titration with citric acid. Complete medium contained in addition 5 g peptone from casein and 1.88 g (NH₄)₂SO₄.

All chemicals were of analytical purity. Radioactive glucose was a product of Amersham-Buchler, Braunschweig, F.R.G. Peptone from casein was obtained from Serva Feinbiochemica, Heidelberg, F.R.G., and enzymes and cofactors were from Boehringer. For all solutions, glass-distilled water was used.

RESULTS AND DISCUSSION

Influence of nitrogen sources on glycogen metabolism

The glycogen breakdown in the cells of the inoculum after transfer to complete medium (cf. Becker et al., 1979) is likely to be initiated by one or more components of the medium. We conducted a systematic search to identify these components.

As shown in Fig. 1, cells synthesized glycogen in the basic medium. Typically, a 2% suspension of the cells accumulated glycogen at a rate of 0.5–1.5 μmol glucosyl units (g yeast⁻¹) min⁻¹ at 28 °C in the absence of all nitrogen sources. In the presence of 55 mM-ammonium sulphate, the rate of glycogen synthesis was initially lowered by a factor of about 3, an effect previously observed (Becker et al., 1979). Only after 2 h was glycogen broken down. By contrast, in the presence of peptone, whether in the absence or the presence of ammonium sulphate, glycogen breakdown was initiated within 5 min or less. Thus, we further investigated the effect of peptone. This nutrient is rather ill-defined: it consists mainly of a mixture of amino acids and small peptides, all of which could have been responsible for the effect of peptone. First, single selected amino acids were tested for their ability to induce immediate glycogen breakdown. Glutamic acid, aspartic acid, glutamine, asparagine and lysine, each at concentrations of 25, 50 and 75 mM, were added to the basic medium. In no case was glycogen breakdown initiated immediately, even if the cells were preincubated at 28 °C with the amino acid prior to the addition of glucose. Under the latter condition,
Fig. 1. Glycogen synthesis and breakdown in *Saccharomyces carlsbergensis* cells in the absence and presence of nitrogen sources. Cells were incubated in basic medium (■), basic medium with 55 mM (NH₄)₂SO₄ (□), complete medium (basic medium with 5 g peptone and 1.88 g (NH₄)₂SO₄) (○) or complete medium without (NH₄)₂SO₄ (●). The glucose consumption rate was 21 ± 3 μmol glucose min⁻¹ (g yeast)⁻¹ in all runs. Increased budding occurred after 1 h in all cases, but significant growth was observed only in the batches with nitrogen, after 3 h.

Fig. 2. Glycogen metabolism in *S. carlsbergensis* cells in the presence of an amino acid mixture (0-22 %) in the molar ratio of casein. The cells were incubated in basic medium containing all the amino acids specified in the legend to Fig. 3, either with (○) or without (●) ammonium sulphate (55 mM).

Fig. 3. Glycogen metabolism in *S. carlsbergensis* cells in the presence of various amino acid mixtures. The cells were incubated in basic medium with the amino acids specified (concentrations in g l⁻¹). (a) Asp, 0.08; Asn, 0.1; Glu, 0.3; Gln, 0.17; Lys, 0.19. (b) Leu, 0.19; Ile, 0.12; His, 0.07; Met, 0.06; Val, 0.11. (c) Phe, 0.11; Tyr, 0.15; Trp, 0.04; Thr, 0.05; Ala, 0.07. (d) Ser, 0.07; Arg, 0.09; Pro, 0.17; Gly, 0.06. ●, Glycogen in cells; ○, glucose in medium.
50 mM-asparagine maintained a constant level of glycogen for about 30 min. Without pre-incubation and with all the other amino acids, glycogen was synthesized initially at a low rate for 30–60 min; then breakdown occurred. Cell growth was apparent by bud formation after 1–2 h and an increase in $A_{546}$.

The above experiments demonstrated that the presence of nitrogen sources supporting growth was not sufficient to induce immediate glycogen breakdown as was peptone. This could be due to two possibilities: peptone contains some substances responsible for the effect, or peptone acts immediately because it is composed of all amino acids necessary for immediate cell proliferation. While the first possibility is more difficult to prove, the second could be tested simply by adding to the cells a defined mixture of amino acids in the molar ratio found in casein (Mercier et al., 1971), from which the peptone is derived. As shown in Fig. 2, a 0.22% mixture of amino acids indeed mimicked the effects of peptone. This experiment did not rule out the possibility that a single amino acid not tested in the previous experiments may have been responsible for this result. In order to identify the amino acid(s) responsible for the induction of glycogen breakdown, amino acids were added in groups in the same concentration as in the total mixture (Fig. 3 a–d). Group (a) contained those most easily utilized as a nitrogen source, while in group (d) those serving as poor sources were combined (Suomalainen & Oura, 1971). With all four mixtures, glycogen was initially synthesized for about 30 min, followed by a degradation period. After 90 min, resynthesis of glycogen occurred. The $A_{546}$ and the number of budding cells increased after 120–180 min in all these experiments, indicating growth of the cells. With the more easily utilized nitrogen sources, the initial glycogen synthesis rate was lower, indicating that glycogen synthesis was the result of a lack of nutrients (Lillie & Pringle, 1980) at least as perceived by the cells prior to induction of enzymes enabling them to interconvert certain amino acids. These experiments showed that no single amino acid was able to trigger glycogen breakdown, but that it was initiated only if all necessary amino acids were supplied. One can assume that in the presence of only a few amino acids or ammonium ions as nitrogen sources, the enzymes necessary for synthesis of the lacking amino acids must be induced prior to growth, although this was not measured. Only after the induction period (i.e. between 30 min and 1 h) was glycogen breakdown observed.

Response of glycogen metabolism to dinitrophenol

Still, the question remains, how can cells sense within minutes the simultaneous presence of all amino acids in the nutrient broth? One possibility is that initiation of massive protein synthesis after inoculation draws heavily on the energy pool of the cells, thus initiating glycogen breakdown to make up the deficit. In this case, lowering the energy supply should also initiate glycogen breakdown.

Cells were incubated in complete medium in the presence of the uncoupler dinitrophenol (10$^{-4}$ M) with and without glucose. Controls were incubated in complete and basic medium without dinitrophenol. While glycogen breakdown was initiated by peptone, the glycogen level remained constant in the presence of dinitrophenol but in the absence of glucose. However, the uncoupler slowed glycogen synthesis in the presence of glucose, indicating that it was taken up and exerted an influence on cell metabolism. Apparently, increasing the consumption of metabolic energy does not by itself stimulate glycogen breakdown.

The effect of peptone on metabolites

The above conclusion was substantiated by direct measurement of the adenylate pool and other key intermediates of glycolysis in basic or complete medium. Since the concentration of the intermediates in the cells was too low for exact determination in a 2% cell suspension, the concentration of the cells was increased to 10% (w/v). At this high cell concentration, peptone inhibited glycogen synthesis, but glycogen breakdown was observed only initially. Probably, important amino acids were quickly consumed by the concentrated cell
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Fig. 4. Glycogen synthesis and metabolite concentrations in S. carlsbergensis in the absence (a) and presence (b) of nitrogen sources. A 10% suspension of cells was prepared in basic (a) or complete (b) medium. Measurements were made of glucose concentration in the medium and of the concentrations of glycogen, pyruvate, UDP-glucose and adenine nucleotides in the cells before and after the addition of glucose (arrows).

suspension, resulting in an insufficient supply of amino acids. Therefore, glycogen breakdown occurred for only a short time. Nevertheless, the suppression of glycogen synthesis was very pronounced in the presence of peptone. One can assume that the change of intermediates in response to peptone should be the same in a less concentrated cell suspension.

A representative experiment is shown in Fig. 4, illustrating the effect of peptone on glycogen synthesis, glucose consumption, the cell contents of pyruvate and UDP-glucose, and adenylate concentration. While glycogen synthesis differed considerably in the presence and absence of peptone, there was little effect of peptone on the levels of the metabolites, all of which followed similar patterns in the absence and presence of peptone, though in the latter case, pyruvate concentration tended to be elevated. This points to an activation of glycolysis, which is also reflected by the higher glucose consumption rate (Fig. 4b, top). The behaviour of UDP-glucose was puzzling because as the substrate for glycogen synthase, its concentration should have been lowered during glycogen synthesis, which was not the case. Its concentration varied similarly in the presence and absence of peptone. However,
UDP-glucose serves as the precursor for the synthesis of several molecules, all of which govern its steady-state concentration. It is difficult to assume that any of the key metabolites assayed can serve as a regulatory signal, since their concentrations in the cell were not greatly different in the presence of peptone from those in its absence. The adenine nucleotides showed similar behaviour, with an energy charge (Atkinson, 1970) between 0.59 and 0.69 under both conditions. The sum of the adenylates appeared slightly higher in the cells without nitrogen sources and this may exert some regulatory effect (Knowles, 1977).

**Absence of glycogen turnover**

Because glycogen metabolism in *S. carlsbergensis* cells reacted much faster to changes of environmental factors than that in *S. cerevisiae* (strain a-S-288 C) when tested under the same conditions, it was possible that glycogen metabolism was not regulated. Thus, flow into and out of glycogen storage may simply follow the law of mass action, resulting in a turnover of glycogen by generation of a futile substrate cycle: glycogen would still be degraded while net synthesis took place. This phenomenon should be especially apparent in yeast cells suspended in phosphate buffer: such cells rapidly accumulate glycogen in the presence of glucose but degrade it as soon as glucose is removed (cf. Becker, 1978; Becker et al., 1979).

For detection of the turnover, the cells were initially supplied with radioactive glucose to label the glycogen. The radioactive glucose was subsequently removed and the cells were washed and suspended in the presence or absence of unlabelled glucose. Glycogen concentration and radioactive label incorporated in glycogen were measured (Fig. 5). The results showed clearly that glycogen breakdown was completely stopped in the presence of external glucose: i.e. no turnover of glycogen took place, as already demonstrated by Rothman & Cabib (1969) in *S. cerevisiae* under different conditions. In the control, radioactively labelled glucose was broken down, which proved that the loss of radioactive glucose could be measured under the experimental conditions.

This result points to a highly effective regulation of glycogen breakdown, although at present no obvious mechanism by which it was achieved can be suggested. Yeast phosphorylase is known to be regulated by phosphorylation and dephosphorylation (Fosset et al., 1971; Becker et al., 1981). However, the specific molecular activity of phosphorylase was only correlated with the growth phase of the batch culture, while extracts from cells...
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Degradation or synthesizing glycogen showed identical phosphorylase activities (Becker, 1982). The very similar pattern of metabolites in the presence and absence of peptone did not yield information for a simple regulatory mechanism mediated by these effectors via allosteric interaction with regulatory enzymes.

As in the case of glycogen synthesis, where the cells start to accumulate the polysaccharide prior to depletion of glucose (Rothman & Cabib, 1969) or other nutrients (Becker et al., 1979; Lillie & Pringle, 1980), the signal for glycogen breakdown, which must relay complex information concerning the nutrient level, is not known. The behavior of the cells resembles the stringent response of bacteria to starvation (Bridger & Paranchych, 1979), which is mediated by pleiotropic nucleotides and which also affects glycogen metabolism (Dietzler & Leckie, 1977). Such a mechanism, however, has not yet been shown in eukaryotes and will be difficult to prove unless suitable mutants are found.

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