Linear growth and poly(\(\beta\)-hydroxybutyrate) synthesis in response to pulse-wise addition of the growth-limiting substrate to steady-state heterotrophic continuous cultures of *Aquaspirillum autotrophicum*

**Marco Pagni, Trello Beffa, Cédric Isch and Michel Aragno**

Laboratoire de Microbiologie de l'Université de Neuchâtel, Chantemerle 22, 2007 Neuchâtel, Switzerland

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Heterotrophic pyruvate-limited steady-state continuous cultures of the bacterium *Aquaspirillum autotrophicum* were perturbed with a pulse injection of a small volume of concentrated pyruvate solution. These cultures exhibited an instantaneous change in the growth dynamics, turning from steady state to apparently linear growth. These transient growth-responses had no lag phase and were clearly distinct from unlimited exponential growth according to the initial rates of increase of biomass and substrate disappearance kinetics. A linear accumulation with time of poly(\(\beta\)-hydroxybutyrate) was observed within the cells. Slopes of these linear responses were negatively correlated with the dilution rate. Physiological bases of linear growth are discussed in the light of the models of H. E. Kubitschek. Poly(\(\beta\)-hydroxybutyrate) synthesis in the absence of exogenous limitation may serve to protect the cells against a transient metabolic overflow.

**Introduction**

Continuous culture is most often used to study steady-state growth physiology. However, only part of the regulatory characteristics of the metabolism are then expressed. Such studies cannot account for most of the growth responses in natural environments, which are essentially a succession of transitions. Several standard transitions can be studied with continuous cultures; a typical example is represented by the instantaneous suppression of the limitation by a pulse of the limiting substrate. Such experiments have been reported by Harisson & Maitra (1969) with *Klebsiella aerogenes* and more recently by Brooke *et al.* (1989) with a thermophilic *Bacillus* strain.

The facultatively autotrophic bacterium *Aquaspirillum autotrophicum* was chosen as model organism. Apart from its ability to grow autotrophically under \(\text{H}_2/\text{O}_2/\text{CO}_2\) mixtures, it can grow heterotrophically under air with a variety of organic acids (Aragno & Schlegel, 1978a, b), as well as mixotrophically (Fasnacht, 1988). It is therefore a valuable organism to study various types of transitions with regard to energy and carbon metabolism. Moreover, it forms homogeneous suspensions in culture, and does not excrete polysaccharides. Poly(\(\beta\)-hydroxybutyrate), PHB, is synthesized as a reserve material under nitrogen limitation (Walther-Mauruschat *et al.*, 1977).

Pulse experiments similar to those described above were undertaken in two different types of continuous culture apparatus. One was designed for on-line turbidimetric measurements at short intervals (6 min), requiring relatively low biomass densities; this allowed a description of transient kinetics with satisfactory accuracy on a 1 h scale. The other apparatus was used to sample cultures for chemical analysis, thus needing higher biomass densities.

**Methods**

**Organism and culture conditions.** *Aquaspirillum autotrophicum* strain SA32 (DSM 732, ATCC 29984, Aragno & Schlegel 1978a) was precultivated in the basal mineral medium described by Schlegel *et al.* (1961) and Aragno & Schlegel (1981) with 0.5% sodium pyruvate added. The temperature was maintained at 27°C throughout the experiments.

For turbidimetric observation of growth kinetics, cells were grown in six 'home-made' fermenters running in parallel. Each fermenter had a working volume of 800 ml and was stirred at 1500 r.p.m. with a large magnetic stirrer. The feed medium consisted of the basal medium with only half the phosphate concentration and with the addition of sodium pyruvate at a final concentration of 7.5 mm. For pulse experiments, 1 or 2 ml sterile 3 M-sodium pyruvate was injected directly into the culture.
The partial pressure of dissolved oxygen and the pH were followed with immersed electrodes. Feed medium and thermostatization were common to all fermenters; these differed only in their dilution rate and in their air inflow rate. Optical density at 436 nm was measured automatically every 6 min in each fermenter, with an accuracy of more than 1%., with a PU8620 Philips spectrophotometer and a HELMA 170.004-QS flow-cell with a 2 mm optical path.

For experiments in which samples had to be collected for chemical analyses, cells were grown in a Biostat-S fermenter with a working volume of 3.4 l stirred at 800 r.p.m. Sodium pyruvate was added to the basal mineral medium at 22.26 mm final concentration. Sterile 3 m-
sodium pyruvate (15 ml) was injected for pulse experiments. Sampling was usually performed with an automatic collector, but, during transitions, samples were collected in a large precooled glass beaker and continuously stirred on an ice bed until the temperature fell to 1 °C (less than 30 s). Optical density was measured with a Zeiss PM4 photometer at 436 nm with an optical path of 1 cm.

**PHB determination.** The PHB assay was a combination of the methods of Law & Slepecky (1961) and of Poindexter & Eley (1983): cells from a 5 ml culture sample were collected by centrifugation (8000 g, 10 min, 4 °C), and washed with 10 mm-Tris/HCl/saline (NaCl 0.9%, w/v) buffer pH 7.0. The pellet was suspended in 5 ml 0.5% (w/v) sodium hypochlorite and left for 1 h at 37 °C to allow total lysis of the cells. The PHB granules were then collected by centrifugation (4000 g, 10 min, 4 °C), washed successively in distilled water, acetone and ethanol, and finally dissolved in hot chloroform. After the chloroform-insoluble part had been discarded, the chloroform was evaporated and the residue was hydrolysed and dehydrated with concentrated sulphuric acid to obtain crotonic acid, which can be quantified by its absorbance at 235 nm, according to Zevenhuizen (1974). A complete absorbance spectrum was measured for each sample using sulphuric acid as a blank.

**Pyruvate determination.** Pyruvate concentration in the culture supernatant was measured by spectrophotometric analysis of NADH oxidation at 340 nm in the presence of lactate dehydrogenase (Czok & Lamprecht, 1974). This method permits detection of pyruvate concentrations as low as 3 μM.

**Determination of respiratory activities.** Cells were washed by centrifugation in potassium phosphate buffer, pH 7.0, and the respiratory activities were determined polarographically at 28 °C using a Clark-type oxygen electrode (Hansatech, CBH2 Box, adjustable volume) with a chart recorder. Oxygen concentration in the experimental system was calibrated using the method of Robinson & Cooper (1970) and the respiratory activities were determined as previously described (Beffa et al., 1988). The reaction mixture (1 ml final vol.) contained only the mineral basal medium for the determination of endogenous respiratory activity, and the same medium supplemented with 25 mm-pyruvate (final concentration) for the measurement of total respiratory activity. The measurements were performed with 0.2 to 0.4 mg dry wt cells ml⁻¹ and are expressed in nmol O₂ consumed (ml culture)⁻¹ min⁻¹.

**Other methods.** Total polyphosphates (acid soluble plus insoluble) were determined from 5 ml of washed cells according to Poindexter & Eley (1983) and Clark et al. (1986). Dry weight was measured after drying washed cells for 48 h at 100 °C. Total protein was estimated with the bicinchoninic acid method (P. K. Smith et al., 1985). Non-linear fitting of the models to the experimental data was performed by standard algorithms of minimization.

**Results**

The term 'exponential' will be restricted to qualify unlimited balanced growth at a constant rate. It will never be used for exogeneously limited growth at steady state. The subscript 'exp' will therefore qualify all parameters related to this kind of growth, such as the exponential growth rate μ<sub>exp</sub> (often referred to as μ<sub>max</sub>).

**Observation of transitions**

Similar experiments were undertaken in both fermenter installations. Each continuous culture was inoculated with an exponentially growing preculture of *A. autotrophicum* on pyruvate. After an initial exponential growth phase at rate μ<sub>exp</sub> ≈ 0.32 h⁻¹ (see below, Fig. 3), the culture became limited in pyruvate and relaxed to a pyruvate-limited steady state. Then a given volume of concentrated pyruvate solution was injected into the culture. The volume added had a negligible influence, except on the pyruvate concentration. The growth dynamics changed immediately, turning from steady state to apparently linear growth (Fig. 1). Despite the fact that the characteristic exponential curvature is only faintly marked on an hour scale, the precision of biomass measurements was sufficient to reveal the absence of any positive curvature. This typical growth response appeared in all experiments for dilution rates ranging from 0.049 to 0.246 h⁻¹, lasted for at least 1 h and was observed for up to 3 h depending on the injected volume and dilution rate. The pyruvate pulse and the reappearance of pyruvate limitation were both correlated with sudden changes in pH and oxygen signals. The aspect of linear growth termination depended on the particular experimental conditions. It was either sudden and correlated with the reappearance of pyruvate limitation for all transitions lasting less than 2 h, or progressive, before the reappearance of limitation (see below, Fig. 4). Finally, each culture relaxed back to its previously observed steady state.

The intersection of the regression lines fitting steady state and linear growth is located very close to the injection time. This was observed in all experiments and is fully correlated both with sudden changes in dissolved oxygen and in pH values. Experimental conditions do not permit the demonstration of the simultaneity of the onset of linear growth and pyruvate injection with an accuracy of more than 30 s. Changes in oxygen concentration on the other hand, can be determined in a matter of seconds.

**Modifications of cell composition**

Table 1 summarizes some relevant measurements made during a linear growth response. Only negligible PHB concentrations were detected in cultures at steady state, whereas PHB accumulated linearly during the transitions (Fig. 2). During the linear growth response, small
Table 1. Measurement of relevant physiological parameters after a pyruvate pulse

<table>
<thead>
<tr>
<th>Measurement</th>
<th>Unit</th>
<th>Time after injection (h):</th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>0</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Turbidity</td>
<td>OD_{436} (1 cm optical path)</td>
<td>3.197</td>
<td>4.017</td>
<td>4.894</td>
</tr>
<tr>
<td>Dry wt</td>
<td>mg ml^{-1}</td>
<td>0.670</td>
<td>0.845</td>
<td>1.019</td>
</tr>
<tr>
<td>Turbidity/dry wt</td>
<td>OD mg^{-1}</td>
<td>4.77</td>
<td>4.75</td>
<td>4.80</td>
</tr>
<tr>
<td>PHB</td>
<td>µg ml^{-1}</td>
<td>&lt;0.2</td>
<td>37.5</td>
<td>69.2</td>
</tr>
<tr>
<td>PHB/dry wt</td>
<td>% mg mg^{-1}</td>
<td>&lt;0.03</td>
<td>4.4</td>
<td>6.8</td>
</tr>
<tr>
<td>Protein</td>
<td>mg ml^{-1}</td>
<td>0.50</td>
<td>0.57</td>
<td>0.64</td>
</tr>
<tr>
<td>Protein/dry wt</td>
<td>% mg mg^{-1}</td>
<td>75</td>
<td>67</td>
<td>63</td>
</tr>
<tr>
<td>Polyphosphate</td>
<td>µg ml^{-1}</td>
<td>&lt;0.2</td>
<td>&lt;0.2</td>
<td>&lt;0.2</td>
</tr>
<tr>
<td>Endo. resp.</td>
<td>nmol O_{2} ml^{-1} min^{-1}</td>
<td>8.4</td>
<td>12.6</td>
<td>21.3</td>
</tr>
<tr>
<td>Tot. resp.</td>
<td>nmol O_{2} ml^{-1} min^{-1}</td>
<td>64.1</td>
<td>65.5</td>
<td>70.4</td>
</tr>
</tbody>
</table>

Pyruvate (15 ml; 3 M) was injected into a 3.4 l culture at steady state (D = 0.134 h^{-1}, S_{i} = 22-26 mM, Biostat fermenter). Values at 0 h correspond to the steady state.
Table 2. Determination of $q_{\text{exp}}$ at the start-up of an experiment with the Biostat fermenter and determination of $q_{\text{lin}}$ later in the same experiment after a pyruvate pulse

Pyruvate (15 ml; 3 mM) was injected into a 3-4 l culture at steady state ($D = 0.134$ h$^{-1}$, $S_f = 22.26$ mM. The fixed values were obtained from culture conditions and from biomass growth kinetics. The fitted values were computed on the pyruvate disappearance kinetics.

<table>
<thead>
<tr>
<th>Biomass kinetics:</th>
<th>Exponential</th>
<th>Linear</th>
</tr>
</thead>
<tbody>
<tr>
<td>Model used:</td>
<td>$S_{\text{exp}}(t)$ (eq. 2)</td>
<td>$S_{\text{lin}}(t)$ (eq. 6)</td>
</tr>
<tr>
<td>Fixed values:</td>
<td>$S_f = 22.26$ mM</td>
<td>$S_f = 22.26$ mM</td>
</tr>
<tr>
<td></td>
<td>$D = 0.134$ h$^{-1}$</td>
<td>$D = 0.134$ h$^{-1}$</td>
</tr>
<tr>
<td></td>
<td>$\mu_{\text{exp}} = 0.310$ h$^{-1}$</td>
<td></td>
</tr>
<tr>
<td></td>
<td>$X_0 = 0.043$ OD</td>
<td></td>
</tr>
<tr>
<td>Fitted values:</td>
<td>$S_0 = 22.02$ mM</td>
<td>$X_{\text{lin}} = 3.207$ OD</td>
</tr>
<tr>
<td></td>
<td>$q_{\text{exp}} = 1.75$ mM OD$^{-1}$ h$^{-1}$</td>
<td>$q_{\text{lin}}^0 = 1.76$ mm OD$^{-1}$ h$^{-1}$</td>
</tr>
<tr>
<td>Residual square sum:</td>
<td>RSS = 5.71</td>
<td>RSS = 3.17</td>
</tr>
<tr>
<td></td>
<td>25 data points</td>
<td>11 data points</td>
</tr>
<tr>
<td></td>
<td>Fig. 3</td>
<td>Fig. 4</td>
</tr>
</tbody>
</table>

Kinetics of substrate disappearance

During steady state, the residual pyruvate concentration was undetectable by the assay used, so the affinity constant of A. autotrophicum for pyruvate is certainly within the micromolar range. In turn, during exponential or linear growth, pyruvate concentrations in the millimolar range were recorded. As a consequence, it was postulated that the pyruvate uptake system(s) was (were) saturated during both types of growth.

During start-up exponential growth in continuous culture, the evolution of the substrate concentration $S_{\text{exp}}$ is described by the following differential equation

$$\frac{dS_{\text{exp}}}{dt} = (S_t - S_{\text{exp}})D - q_{\text{exp}} X_0 e^{(\mu_{\text{exp}} - D)t}$$

(1)

where $\mu_{\text{exp}}$ is the exponential growth rate, $D$ the dilution rate, $S_t$ the substrate concentration in the feed, $q_{\text{exp}}$ the constant substrate consumption rate per biomass unit and $X_0$ the initial biomass. This differential equation can be resolved into

$$S_{\text{exp}}(t) = S_t + e^{-D t} \left[ S_0 - S_t + \frac{X_0 q_{\text{exp}}}{\mu_{\text{exp}}} (1 - e^{\mu_{\text{exp}} t}) \right]$$

(2)

with $S_0$ as the initial substrate concentration.

Linear growth is biomass accumulation at a constant rate, because a linear function of time is the primitive of a constant function. A simple hypothesis for linear growth could involve a constant consumption rate for the whole culture and a constant growth yield. Then, the following equation would describe the substrate-disappearance kinetics during linear growth

$$\frac{dS_{\text{lin}}}{dt} = (S_t - S_{\text{lin}}) D - q_{\text{lin}}^0 X_{SS}$$

(3)

where $Q_{\text{lin}}^0$ is the constant consumption rate by volume unit of the whole culture. In fact, the physiology underlying this hypothesis is more complicated than it appears, because of the consumption rate per biomass unit, which has to decrease with time, and because of the ambiguous meaning of a constant growth yield during a growth which is known to be unbalanced. Formally, a consumption rate per biomass unit, $q_{\text{lin}}^0$, can be defined at the beginning of linear growth when the biomass is equal to the steady state biomass $X_{SS}$

$$q_{\text{lin}}^0 = \frac{Q_{\text{lin}}^0}{X_{SS}}$$

(4)

so the differential equation (3) can be rewritten

$$\frac{dS_{\text{lin}}}{dt} = (S_t - S_{\text{lin}}) D - q_{\text{lin}}^0 X_{SS}$$

(5)

and admits the solution

$$S_{\text{lin}}(t) = S_t - \frac{q_{\text{lin}}^0 X_{SS}}{D} + e^{-Dt} \left( S_0 - S_t + \frac{q_{\text{lin}}^0 X_{SS}}{D} \right)$$

(6)

$S_{\text{exp}}(t)$ and $S_{\text{lin}}(t)$ can easily be distinguished graphically, because of the negative curvature of $S_{\text{exp}}(t)$

$$\frac{d^2S_{\text{exp}}}{dt^2} = -(S_t - S_{\text{exp}}) D^2 - q_{\text{exp}} X_0 (\mu_{\text{exp}} - 2D)$$

$$e^{(\mu_{\text{exp}} - D)t} < 0 \quad \forall t, \quad t \geq 0$$

if $S_{\text{exp}}(0) < S_t + q_{\text{exp}} X_0 \frac{2D - \mu_{\text{exp}}}{D^2}$

(7)
prevailing at least at the start-up of the culture \([S_{\text{exp}}(0)] \) close to \(S_t\) and \(X_0\) small] against the slight positive curvature of \(S_{\text{lin}}(t)\)

\[
\frac{d^2S_{\text{lin}}}{dt^2} = D [Q_{\text{whole}}^{\text{calc}} - D (S_{\text{t}} - S_{\text{lin}})] > 0 \text{ as } Q_{\text{lin}}^{\text{calc}} > DS_t
\]  

Kinetic data of pyruvate disappearance were recorded during both exponential and linear growth and both models were fitted to the data. The values obtained for the parameters are shown in Table 2, and the fits of the models to experimental data are presented in Figs 3 and 4. The hypotheses underlying the two sets of substrate-consumption kinetics appeared to be satisfied for the respective growth behaviours. Cross-testing of the models did not yield acceptable fits because of the opposing curvatures. Fig. 4 shows a case where linear growth ended before pyruvate had become limiting again. Nevertheless, the hypothesis of a constant consumption rate for the whole culture still appeared valid up to the reappearance of pyruvate-limitation. The two consumption rates per biomass unit are strikingly close

\[
q_{\text{exp}} = 1.75 \text{ mmol OD}^{-1} \text{ h}^{-1} \text{ and } q_{\text{lin}} = 1.76 \text{ mmol OD}^{-1} \text{ h}^{-1}
\]

The growth at steady state seems to have only a negligible influence on the constancy of the (maximal) uptake rate of the saturated consumption system. This apparent constancy was verified by the data from the automated turbidity-measuring system. Admitting the same substrate disappearance kinetics, the solution equations, \(S_{\text{exp}}(t)\) and \(S_{\text{lin}}(t)\), were rearranged in order to relate the consumption rates with the time required for substrate disappearance. The respective consumption rate was then evaluated for each pulse experiment. The constancy of the maximal potential consumption rate of pyruvate was roughly confirmed in the range of observed dilution rates (0.049 to 0.246 h\(^{-1}\)).

**Effects of dilution rate**

The steady state optical density, \(X_{SS}\), and the slope, \(\Delta X/\Delta t\), of the linear response were recorded at different dilution rates in the parallel fermenter installation in order to compute the initial linear growth rate

\[
\mu_{\text{lin}} = \frac{1}{X_{SS}} \frac{\Delta X}{\Delta t} + D
\]  

This initial linear growth rate was taken into account instead of the slope alone, because while the potential maximal consumption rate appeared constant, the steady state biomass varied according to the dependence of growth yield on dilution rate (Pirt 1965; Esener et al., 1983). Fig. 5 shows the \(\mu_{\text{lin}}\) distribution against the dilution rate. A negative regression correlates both parameters satisfactorily

\[
\mu_{\text{lin}} = 0.55 - 0.61 D
\]  

showing that the initial linear growth rate is always greater than the exponential growth rate \((\mu_{\text{exp}} = 0.32 \text{ h}^{-1})\) at any observed dilution rate. Biomass linear over-production is proportional to the difference between the conditioning dilution rate and the maximal growth rate, or, from another point of view, to the sudden over-consumption of substrate. Moreover, the intercept of the regression line with \(\mu_{\text{exp}}\) is close to the washout dilution
rate; this implies that no linear response would be expected if extra substrate were injected into an already (non-limited) exponentially growing culture. Therefore, in these cells PHB accumulation is not to be expected in significant amounts, just as was observed.

Discussion

Pyruvate as sole limiting factor

It was assumed that pyruvate was the sole limiting factor at steady state according to the following experimental evidence. (a) Steady-state pyruvate concentration is not detectable for any tested feed concentration. (b) Schlegel's medium allows *A. autotrophicum* to grow up to 2.5 mg dry wt ml⁻¹, twice the highest biomass recorded here; ammonium then becomes limiting, provided carbon and energy are in excess (Fasnacht 1988, and personal observations). (c) After several successive pulse experiments, the pyruvate feed concentration was increased; the culture then grew up to a new steady state with a biomass concentration higher than recorded during the pulse experiments. (d) The characteristic linear response was observed under various dissolved oxygen partial pressures ranging from 16 to 60% air saturation, depending on air inflow rates, while oxygen limitation was observed only below 3% air saturation in other experiments. (e) The characteristic transitions were similar in both basal and half-phosphate concentration media.

Linear growth kinetics

The growth kinetics following a pyruvate pulse are clearly non-exponential: the initial transient growth rate is distinctly higher than the exponential one (Fig. 5). This growth is unbalanced, as revealed by the accumulation of PHB and by the decrease in the protein fraction of the biomass (Table 1). PHB accumulation is related to an increase in endogenous respiration. However, PHB does not account for the total increase in biomass: in 2 h, the dry biomass increased by 349 μg ml⁻¹, whereas only 69-2 μg PHB ml⁻¹ accumulated. Simultaneously, the protein content of the culture increased by 140 μg ml⁻¹. Clearly, both phenomena (PHB and protein synthesis) participate in the biomass increase.

Despite the complexity of such unbalanced growth, a simple linear growth model for total biomass suffices to describe the observed kinetics of growth and substrate disappearance. It should therefore be considered as a useful local description. Its adequacy is certainly restricted to the accuracy of the measurements which is probably not the case for true exponential growth. The effect of the dilution rate, which should give a slight negative curvature to the kinetics of biomass accumulation at a constant rate, is actually negligible.

In his work on growth kinetics of single cells or synchronized cultures of *Escherichia coli*, Kubitschek (1990) has proposed a linear growth model as a general model for dry weight increase during one cell cycle. The macromolecular (cytoplasmic) fraction accumulates exponentially (Ecker & Kokaïs, 1969), but as shown by Kubitschek & Pai (1988) the precursor-pool sizes vary with a maximum at mid-cycle and a minimum at division time, so the total weight increase, macromolecules plus precursors, remains linear. This linear growth pattern is explained by the constancy of uptake during the cell cycle, the proteins responsible for consumption (periplasmic binding-proteins and membrane-bound proteins) being synthesized only near division time (reviewed in Kubitschek, 1990). The linear biomass increase and the constant substrate consumption-rate for the whole culture are consistent with our observations during the pulse experiments. Therefore, pulsing the culture with the limiting factor appears to induce a coordinated behaviour of the cell population. In *A. autotrophicum*, cell numbers are poorly correlated to biomass parameters during transient growth, which is not surprising for an elongated organism subject to variation in cell length. Thus we could not introduce with sufficient accuracy the cell numbers as a parameter for a short (2 h) period. Even if our results cannot be used as experimental evidence for Kubitschek's model, this model does allow prediction of biomass behaviour consistent with our observations.

Role of PHB

The role traditionally attributed to PHB is that of a reserve material, which is synthesized during growth limited by a non-carbon compound (reviewed in Ander-
son & Dawes, 1990). Senior & Dawes (1971) have also proposed that PHB can be viewed as a sink of reducing power. However, it has to be pointed out that during our pulse experiments, PHB was shown to be synthesized, although in small amounts, under a complete lack of exogenous limitation. This accumulation was apparently linear (Fig. 2). Such behaviour has to be considered in the light of other similar dilimination experiments. I. W. Smith et al. (1954) showed that Klebsiella aerogenes transferred from a phosphate-limiting to a non-limiting medium transiently synthesized polyphosphate inclusions. With a glucose-limited continuous culture of the same organism, Harisson & Maitra (1969) showed that a pulse of glucose causes a transient excretion of pyruvate and another unidentified substance. Moreover, one of their figures presents a transient linear excess production of CO₂ and slightly positively curved glucose disappearance kinetics; a linear growth pattern can be suspected from these data. A methanol pulse on a methanol-limited continuous culture of a thermophilic Bacillus strain resulted in a rapid excretion of formaldehyde into the medium (Brooke et al., 1989); despite the subsequent autopoisoning of the cells, formaldehyde accumulation appeared strikingly linear. Excretion or accumulation of ‘byproducts’ (polyphosphate, pyruvate, formaldehyde or PHB) during unbalanced growth is certainly of major importance for actual survival, because the cells are temporarily incapable of metabolizing all the consumed substrate. Importance of overflow metabolism for cell survival can be compared with the need for efficient uptake-regulation in yeasts: with a mutant of Saccharomyces cerevisiae defective for maltose uptake-regulation, Entian & Loureiro-Dias (1990) showed that a maltose pulse results in cell auto-poisoning caused by intracellular glucose accumulation. The fact that exponential growth metabolism cannot be achieved instantaneously by a previously slow-growing culture is underlined here by the governing role of dilution rate in culture conditioning (Fig. 5). Many different hypotheses can be formulated in order to identify the nature of the bottleneck involved, ribosome concentration or nitrogen-uptake regulation, for example, admitting such a bottleneck really does exist. But more generally, as the macromolecular composition depends on the growth rate (discussed by Esener et al., 1982, and Wanner & Egli, 1990), a certain lag-time is undoubtedly necessary to adapt the macromolecular composition to the new environmental and growth conditions.

A. autotrophicum temporarily ‘wastes’ the over-consumed substrate by synthesizing PHB, which could be considered as a sink protecting the cell against a transient metabolic overflow. The classical role of PHB as a 'strategic' reserve for future survival remains of first ecological significance, but it is secondary for actual cell survival.

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References


