The Role of Oxygen in the Regulation of Glucose Metabolism, Transport and the Tricarboxylic Acid Cycle in *Pseudomonas aeruginosa*

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The effect of dissolved oxygen concentration on the metabolism of glucose in *Pseudomonas aeruginosa* was studied with chemostat cultures using both single-step and gradual transitions from either ammonium or glucose limitation to oxygen limitation and studying transient and steady states. The pathway of glucose metabolism was regulated by the availability of oxygen. The organism responded to oxygen limitation by adjusting its metabolism of glucose from the extracellular direct oxidative pathway, which produces gluconate and 2-oxoglucunate, to the intracellular phosphorylative route. This change was a consequence of decreased activities of glucose dehydrogenase and gluconate dehydrogenase and of the transport systems for gluconate and 2-oxoglucunate, and an increased activity of glucose transport, while relatively high activities of hexokinase and glucose-6-phosphate dehydrogenase were maintained. Citrate synthase, isocitrate dehydrogenase and malate dehydrogenase activities responded to changes in dissolved oxygen concentration rather than to changes in the glucose or ammonium concentrations. The effect of oxygen limitation on the oxo-acid dehydrogenases and aconitase was probably due, wholly or in part, to repression by glucose consequent upon the increase in residual glucose concentration. Succinate dehydrogenase was repressed by an increase in ammonium concentration under an oxygen limitation.

INTRODUCTION

Studies on the effects of dissolved oxygen concentration on the enzymic activities and metabolic versatility of obligate aerobes are relatively meagre compared with those on facultative anaerobes. This is understandable in view of the diversity of fermentation patterns encountered with the latter organisms on their transition from aerobic to anaerobic conditions (for recent reviews, see Harrison, 1976; Stouthamer, 1978; Jones, 1979). However, in their natural habitat obligate aerobes encounter a wide range of oxygen concentrations. Nitrogen-fixing azotobacters exhibit the phenomenon of respiratory protection; in these organisms high oxygen concentrations are toxic, inhibiting nitrogenase (Parker, 1954; Phillips & Johnson, 1961; Dalton & Postgate, 1969). Studies on the effect of oxygen on the enzymic complement of *Azotobacter chroococcum* and *Azotobacter vinelandii* have been carried out (Drozd & Postgate, 1970; Haaker & Veeger, 1976). We have previously shown that the energy-reserve polymer poly-β-hydroxybutyrate (PHB) accumulates in *Azotobacter beijerinckii* when cultures become oxygen-limited; the reductive stage of polymer synthesis serves as an alternative electron acceptor when oxygen is no longer so readily available, and thus permits the organism to grow under such conditions (Senior et al., 1972; Senior & Dawes, 1973; Ward et al., 1977). The activities of key enzymes of PHB metabolism and of certain enzymes of the tricarboxylic acid cycle responded to changes in oxygen concentration.

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(Jackson & Dawes, 1976) but the Entner–Doudoroff enzymes were not affected (Stephenson et al., 1978; Carter & Dawes, 1979).

The observed regulation of tricarboxylic acid cycle activity by oxygen concentration in A. beijerinckii posed the question of whether this behaviour was representative of obligate aerobes in general or whether it was a manifestation of respiratory protection and therefore characteristic only of nitrogen-fixing organisms. As we have previously studied the aerobe *Pseudomonas aeruginosa*, which is a denitrifier that utilizes glucose via the Entner–Doudoroff and tricarboxylic acid cycle pathways, we chose this organism for a comparative study of oxygen effects. *Pseudomonas aeruginosa* also metabolizes glucose via an extracellular direct oxidative pathway to gluconate and 2-oxogluconate (Ng & Dawes, 1973; Midgley & Dawes, 1973; Roberts et al., 1973; Whiting et al., 1976a, b). Transport of glucose, gluconate and 2-oxogluconate occurs by independently regulated systems (Whiting et al., 1976a) and the extracellular oxidative enzymes and associated transport systems are repressed when the organism is transferred from ammonium to glucose limitation (Whiting et al., 1976b). Hunt & Phibbs (1977) observed that the extracellular route was also repressed when *P. aeruginosa* was grown anaerobically with nitrate as electron acceptor in batch culture. It was of interest to examine the effect of varying oxygen concentration on the pathways of glucose metabolism in *P. aeruginosa* under the controlled conditions of the chemostat. We have thus examined the effect of transitions from ammonium or glucose limitation to oxygen limitation on various enzymes of glucose metabolism and the tricarboxylic acid cycle, and also on the transport systems for glucose, gluconate and 2-oxoglucanlate.

**METHODS**

*Organism and growth. Pseudomonas aeruginosa* PAO1 was kindly provided by Professor B. W. Holloway. Routine maintenance, batch growth of the organism and harvesting procedures were as previously described (Ng & Dawes, 1973; Midgley & Dawes, 1973). For chemostat inocula, the organism was subcultured at least ten times in the appropriate medium. All cultures were grown at 37 °C.

The chemostat vessel (2.5 l volume) was built in this Department. It was fitted with automatic pH control (E.I.L. Instruments, Richmond, Surrey; with pH electrodes supplied by Activion, Kinglissie, Fife), temperature control (Fielden Electronics, Manchester) and oxygen control (Leeds & Northrup, Birmingham; Precision Products and Controls, Tulsa, U.S.A.). The oxygen electrodes were made in this Department. CO₂ was measured with an infrared analyser (Mine Safety Appliances, Glasgow) and oxygen with a paramagnetic oxygen analyser (Servomex OA 137; Servomex Controls, Crowborough, Sussex). Continuous readout of gas composition was obtained with a Kent chart recorder (George Kent, Luton, Beds). The total gas flow was 11 min⁻¹ and the required dissolved oxygen tension (d.o.t.) was secured by adjusting the proportion of oxygen in the inflowing oxygen/nitrogen gas mixture. Oxygen limitation (undetectable d.o.t.) occurred at 4.5 % (v/v) oxygen in the inflowing gas.

Medium for continuous cultivation was prepared in 40 l batches which were sterilized by filtration through a Sartorius filter (142 mm diam., pore size 0.25 μm) at 292 kPa. The medium contained (per litre): KH₂PO₄, 5.4 g; nitritrotiacetic acid, 0.286 g; (NH₄)₂SO₄, 0.9 g (for ammonium-limited growth) or 1.8 g (for glucose- and oxygen-limited growth); trace metal solutions 1 (5 ml), 2 (5-25 ml) and 3 (0-1 ml) (Ng & Dawes, 1973); glucose, 4-0 g (for carbon-limited growth) or 8-11 g (for ammonium-limited growth). For most of the single-step transitions from ammonium- or glucose-limited growth to oxygen limitation, the inflowing medium was simultaneously changed to furnish excess ammonium or glucose. Initial studies were carried out to ascertain the effect of such additions of ammonium or glucose, as noted in the Results. In the gradual transition experiments with ammonium-limited cultures, the inflowing medium was changed to provide excess ammonium as soon as ammonium ions could be detected in the medium supernatant; this occurred at approximately 9% oxygen in the inflowing gas. In similar experiments with glucose-limited cultures, glucose was detected in the supernatant at between 6 and 8% oxygen in the inflowing gas and the medium was changed at 4% oxygen to ensure an excess of glucose.

Six to ten vessel volumes were allowed to pass through the fermenter between each steady state before sampling except in experiments to study transient responses.

*Transport studies.* These were performed by the methods of Midgley & Dawes (1973) using the substrate concentrations and specific radioactivities specified by Whiting et al. (1976b). The radioactivity was assayed as described by Midgley & Dawes (1973).
Enzyme assays. Bacterial extracts were prepared by the methods of Ng & Dawes (1973). All enzyme assays were performed in cuvettes of 1 cm light-path at 37 °C with a Pye Unicam SP1800 recording spectrophotometer under previously determined optimum conditions. Specific activities are recorded as μmol h⁻¹ (mg protein)⁻¹ and are the means of at least two assays which did not differ by more than 5%.

The enzymes of glucose and 2-oxoglucunate metabolism [glucose dehydrogenase (EC 1.1.1.47), gluconate dehydrogenase (EC 1.1.99.3), glucose-6-phosphate dehydrogenase (EC 1.1.1.49), hexokinase (EC 2.7.1.1) and gluconate kinase (EC 2.7.1.12)] were assayed under the conditions described by Ng & Dawes (1973) except that glucose dehydrogenase was assayed in the presence of phenazine methosulphate (2 mM). The combined activity of 2-oxoglucunate kinase and 2-oxo-6-phosphogluconate reductase (referred to as 2-oxoglucunate enzymes) was assayed by the method of Whiting et al. (1976b). Isocitrate dehydrogenase (EC 1.1.1.42) and aconitase (EC 4.2.1.3) were assayed as described by Ng & Dawes (1973); fumarate hydratase (EC 4.2.1.2) by the method of Racker (1950); citrate synthase (EC 4.1.3.7) according to Weitzman & Dunnmore (1969); malate dehydrogenase (EC 1.1.1.37) by the method of von Tigerström & Campbell (1966) except that 5 mM-MgCl₂ was included and the pH was 8.6; pyruvate dehydrogenase (EC 1.2.4.1) and 2-oxoglutarate dehydrogenase (EC 1.2.4.2) by the method of von Tigerström & Campbell (1966) except that Mg²⁺ was omitted in the latter assay; succinate dehydrogenase (EC 1.3.99.1) by the method of Veeger et al. (1969); and NADH oxidase according to Jackson & Dawes (1976). Both pyruvate and 2-oxoglutarate dehydrogenases were assayed under anaerobic conditions in cuvettes fitted with side-arms, on account of the high NADH oxidase activities of bacterial extracts. Cuvettes were flushed with oxygen-free nitrogen for 5 min and an airtight seal was achieved with Subaseal caps. Nitrogen was flushed for a further 5 min via syringe needles penetrating the Subaseals. Cuvettes were incubated for 10 min in the spectrophotometer at 37 °C and the reaction was started by tipping the substrate from the side-arm. In control experiments for endogenous NAD reduction, water replaced the substrate. Combined Entner–Doudoroff enzymes [6-phosphogluconate dehydratase (EC 4.2.1.12) and 2-keto-3-deoxy-6-phosphogluconate aldolase (EC 4.1.2.14)] were jointly assayed by measuring the 6-phosphogluconate-dependent production of pyruvate coupled to the oxidation of NADH at 340 nm using an excess of commercial lactate dehydrogenase. Cuvettes contained (total volume 3 ml): 0.1 M-triethanolamine/HCl buffer, pH 7-8, 2 ml; 80 mM-dithiothreitol, 0.2 ml; 10 mM-NADH, 0.2 ml; 0.2 mM-6-phosphogluconate, 0.1 ml; lactate dehydrogenase (2 mg ml⁻¹), 0.2 ml; cell extract, 100 μl.

Analyses. Protein was estimated by the methods of Gornall et al. (1949) and Lowry. Glucose was determined with a glucose oxidase kit (Boehringer) and 2-oxoglucunate by the method of Manning & Cohen (1951). Gluconate was determined by coupling gluconate kinase with glucose-6-phosphate dehydrogenase according to the Boehringer Handbook. Ammonia was determined by the method of Chaney et al. (1976b). Isocitrate dehydrogenase (EC 1.1.1.22), succinate dehydrogenase (EC 1.3.99.3), glucose-6-phosphate dehydrogenase (EC 1.1.1.49), hexokinase (EC 2.7.1.1) and gluconate kinase (EC 2.7.1.12) were assayed under the conditions described by Ng & Dawes (1973) except that glucose dehydrogenase was assayed in the presence of phenazine methosulphate (2 mM). The combined activity of 2-oxoglucunate kinase and 2-oxo-6-phosphogluconate reductase (referred to as 2-oxoglucunate enzymes) was assayed by the method of Whiting et al. (1976b). Isocitrate dehydrogenase (EC 1.1.1.42) and aconitase (EC 4.2.1.3) were assayed as described by Ng & Dawes (1973); fumarate hydratase (EC 4.2.1.2) by the method of Racker (1950); citrate synthase (EC 4.1.3.7) according to Weitzman & Dunnmore (1969); malate dehydrogenase (EC 1.1.1.37) by the method of von Tigerström & Campbell (1966) except that 5 mM-MgCl₂ was included and the pH was 8.6; pyruvate dehydrogenase (EC 1.2.4.1) and 2-oxoglutarate dehydrogenase (EC 1.2.4.2) by the method of von Tigerström & Campbell (1966) except that Mg²⁺ was omitted in the latter assay; succinate dehydrogenase (EC 1.3.99.1) by the method of Veeger et al. (1969); and NADH oxidase according to Jackson & Dawes (1976). Both pyruvate and 2-oxoglutarate dehydrogenases were assayed under anaerobic conditions in cuvettes fitted with side-arms, on account of the high NADH oxidase activities of bacterial extracts. Cuvettes were flushed with oxygen-free nitrogen for 5 min and an airtight seal was achieved with Subaseal caps. Nitrogen was flushed for a further 5 min via syringe needles penetrating the Subaseals. Cuvettes were incubated for 10 min in the spectrophotometer at 37 °C and the reaction was started by tipping the substrate from the side-arm. In control experiments for endogenous NAD reduction, water replaced the substrate. Combined Entner–Doudoroff enzymes [6-phosphogluconate dehydratase (EC 4.2.1.12) and 2-keto-3-deoxy-6-phosphogluconate aldolase (EC 4.1.2.14)] were jointly assayed by measuring the 6-phosphogluconate-dependent production of pyruvate coupled to the oxidation of NADH at 340 nm using an excess of commercial lactate dehydrogenase. Cuvettes contained (total volume 3 ml): 0.1 M-triethanolamine/HCl buffer, pH 7-8, 2 ml; 80 mM-dithiothreitol, 0.2 ml; 10 mM-NADH, 0.2 ml; 0.2 mM-6-phosphogluconate, 0.1 ml; lactate dehydrogenase (2 mg ml⁻¹), 0.2 ml; cell extract, 100 μl.

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Chemicals. [U-¹⁴C]Glucose and [U-¹⁴C]gluconate were obtained from Amersham. 2-Oxo[¹⁴C]gluconate was prepared by the method of Whiting et al. (1976b). The following chemicals were obtained from Sigma: DL-isocitrate, phenazine methosulphate, DL-dithiothreitol, L-malate, 2-oxoglutarate (monosodium and monopotassium), 5,5′-dithiobis-(2-nitrobenzoic acid), 6-phosphogluconate, NAD (reduced and oxidized forms), NADPH, coenzyme A, S-acetyl-coenzyme A and glucose 6-phosphate. NADP was obtained from Boehringer. L-Cysteine. HCl and oxaloacetic acid were from BDH. All other chemicals were of AnalAr standard or the highest purity available.

Enzymes. All commercial enzymes were obtained from Sigma.

RESULTS AND DISCUSSION

Two types of transition to oxygen limitation were used. The first, referred to as a single-step transition, subjected ammonium- or glucose-limited cultures to a sudden decrease from a high to an undetectable dissolved oxygen tension (d.o.t.). In the second, the d.o.t. was lowered in stages, and is termed a gradual transition.

Effects of oxygen concentration: single-step transitions

The transition from ammonium limitation to oxygen limitation resulted in decreases in the specific activities of glucose, gluconate and glucose-6-phosphate dehydrogenases, gluconate kinase and the 2-oxoglucunate enzymes, while hexokinase increased in activity (Table 1). With the exception of gluconate kinase and the 2-oxoglucunate enzymes, new steady-state activities were reached within 40 h of the transition, as were the residual nutrient concentrations. The activity of glucose dehydrogenase decreased 20 to 30% faster than that of gluconate dehydrogenase.
Table 1. Effect of single-step transition to a different limitation on steady-state activities of enzymes of glucose metabolism and the tricarboxylic acid cycle and on residual concentrations of nutrients

In the transition from ammonium to glucose limitation, 15 vessel volumes were allowed to pass through the fermenter before sampling. Ammonium- and glucose-limited cultures were grown at 80% of air saturation (23% oxygen in the inflowing gas) while oxygen limitation (undetectable d.0.t.) was secured with 2.5% oxygen in the inflowing gas; the total gas flow rate (oxygen/nitrogen) was 11 min -1. In the transition from ammonium or glucose limitation to oxygen limitation, the ammonium or glucose concentration in the inflowing medium was increased as described in Methods to ensure that only oxygen was growth-limiting. The values for oxygen limitation refer to 120 h after the transition unless otherwise indicated. Each value in the table represents the average of duplicate assays, which did not vary by more than 5%, from one particular steady state.

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Ammonium limitation (D = 0-36 h)</th>
<th>Glucose limitation (D = 0-36 h)</th>
<th>Ammonium limitation (D = 0-35 h)</th>
<th>Oxygen limitation (D = 0-35 h)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pyruvate dehydrogenase</td>
<td>3.8</td>
<td>9.8</td>
<td>11.9</td>
<td>18.8</td>
</tr>
<tr>
<td>Citrate synthase</td>
<td>4.1</td>
<td>9.4</td>
<td>12.0</td>
<td>19.2</td>
</tr>
<tr>
<td>Aconitase</td>
<td>15.9</td>
<td>30.3</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Isocitrate dehydrogenase</td>
<td>77.1</td>
<td>66.7</td>
<td>60.2</td>
<td>49.5</td>
</tr>
<tr>
<td>2-Oxoglutarate dehydrogenase</td>
<td>10.4</td>
<td>22.9</td>
<td>6.8</td>
<td>32.7</td>
</tr>
<tr>
<td>Succinate dehydrogenase</td>
<td>14.1</td>
<td>14.2</td>
<td>18.9</td>
<td>11.0</td>
</tr>
<tr>
<td>Fumarate hydratase</td>
<td>142</td>
<td>92</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>Malate dehydrogenase</td>
<td>14/2</td>
<td>14/1</td>
<td>22.1</td>
<td>15.5</td>
</tr>
<tr>
<td>Glucose dehydrogenase</td>
<td>12/1</td>
<td>15/4</td>
<td>21.9</td>
<td>15.2</td>
</tr>
<tr>
<td>Gluconate dehydrogenase</td>
<td>—</td>
<td>—</td>
<td>16.0</td>
<td>2.9</td>
</tr>
<tr>
<td>2-Oxogluconate enzymes</td>
<td>—</td>
<td>—</td>
<td>100</td>
<td>13.5</td>
</tr>
<tr>
<td>Gluconate kinase</td>
<td>—</td>
<td>—</td>
<td>2.0</td>
<td>0.1</td>
</tr>
<tr>
<td>Hexokinase</td>
<td>—</td>
<td>—</td>
<td>4.5</td>
<td>0.25</td>
</tr>
<tr>
<td>Glucose-6-phosphate dehydrogenase</td>
<td>—</td>
<td>—</td>
<td>32.3</td>
<td>30.0</td>
</tr>
<tr>
<td>Entner-Doudoroff enzymes</td>
<td>—</td>
<td>—</td>
<td>51.7</td>
<td>66.3</td>
</tr>
<tr>
<td>Bacterial dry weight (mg ml)</td>
<td>—</td>
<td>—</td>
<td>36.0</td>
<td>43.0</td>
</tr>
<tr>
<td>Residual ammonium (mM)</td>
<td>—</td>
<td>—</td>
<td>0.8</td>
<td>2.16</td>
</tr>
<tr>
<td>Residual glucose (mM)</td>
<td>—</td>
<td>—</td>
<td>21.1</td>
<td>5.6</td>
</tr>
<tr>
<td>Residual gluconate (mM)</td>
<td>—</td>
<td>—</td>
<td>24.8</td>
<td>35.8</td>
</tr>
<tr>
<td>Residual 2-oxogluconate (mM)</td>
<td>—</td>
<td>—</td>
<td>2.3</td>
<td>ND</td>
</tr>
<tr>
<td></td>
<td>—</td>
<td>—</td>
<td>3.1</td>
<td>ND</td>
</tr>
</tbody>
</table>

—, Not determined; ND, not detected.
* Oscillations; no steady state achieved in oxygen limitation.
† Damped oscillations for 180 h before attaining steady state.
‡ No change in activity until 120 h; steady state not attained in oxygen limitation.
Effect of oxygen on glucose metabolism

Fig. 1. Effect of single-step transition to oxygen limitation (undetectable d.o.t.) of an ammonium-limited culture on the activities of gluconate kinase (O) and the 2-oxoglucconate enzymes (●). At the time indicated by the arrow, the oxygen supply rate in the oxygen/nitrogen mixture was decreased from 230 to 25 ml min⁻¹ (total gas flow 1 l min⁻¹) and the (NH₄)₂SO₄ concentration in the inflowing medium was increased from 0.9 to 1.8 g l⁻¹: D = 0.35 h⁻¹. Enzyme specific activities are expressed as μmol h⁻¹ (mg protein)⁻¹.

Table 2. Effect of a single-step transition to oxygen limitation of an ammonium-limited culture on the steady-state activities of the transport systems for glucose, gluconate and 2-oxoglucconate

The conditions were as described in Fig. 1. The activities for oxygen limitation are steady-state values obtained 110 h after the transition; activities oscillated during the initial 100 h period.

<table>
<thead>
<tr>
<th>Transport system</th>
<th>Ammonium limitation</th>
<th>Oxygen limitation</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glucose</td>
<td>1.5</td>
<td>5.5</td>
</tr>
<tr>
<td>Gluconate</td>
<td>5.5</td>
<td>3.0</td>
</tr>
<tr>
<td>2-Oxoglucconate</td>
<td>105</td>
<td>43</td>
</tr>
</tbody>
</table>

A similar response for hexokinase and glucose-6-phosphate dehydrogenase was observed in the transition from glucose limitation to oxygen limitation (Table 1). The Entner-Doudoroff enzymes were unchanged in transitions from ammonium or glucose limitation to oxygen limitation. In the latter transition, gluconate dehydrogenase was unchanged while glucose dehydrogenase decreased slightly in activity (Table 1). Gluconate, which is known to induce gluconate dehydrogenase (Whiting et al., 1976a), was not detected in the medium during the transition which probably accounts for the invariance of this enzyme. This interpretation is supported by the anaerobic batch culture studies of Hunt & Phibbs (1977).

In transitions from ammonium to oxygen limitation the 2-oxoglucconate enzymes increased markedly in activity up to 80 h, before falling to a new steady-state level (Fig. 1). A similar response was observed for an initially glucose-limited culture. The reasons for this peak are not apparent but it may reflect transient changes in intracellular metabolite concentrations as a consequence of abrupt oxygen limitation. This response was observed on each occasion the experiment was performed, although the peak height did vary.

Transition from ammonium to oxygen limitation also resulted in an increase in the initial rate of glucose transport and decreases in those of gluconate and 2-oxoglucconate (Table 2). In contrast, transition from glucose to oxygen limitation gave no change in these initial rates. This correlates with the virtual absence from the culture of gluconate and 2-oxoglucconate (Table 1), compounds which induce their respective transport systems, while gluconate
additionally represses the activity of the glucose transport system (Whiting et al., 1976a). The decrease in the residual concentrations of these compounds probably explains the changes observed in transport rate in the transition from ammonium to oxygen limitation. The transport systems showed oscillations in the transition from ammonium to oxygen limitation, a new steady state being reached after 100 h (Table 2). The reasons for this response are unclear. No oscillations in the concentrations of glucose, gluconate or 2-oxoglucuronate were observed. Oscillations in the concentration of some intracellular modulator or metabolite may have given rise to this response, as may also be the case for the 2-oxoglucuronate enzymes. However, further study is required to clarify this behaviour.

The effect of both transitions on various tricarboxylic acid cycle enzymes is recorded in Table 1. Most showed a decrease in activity under oxygen limitation, while isocitrate dehydrogenase activity was unchanged in the transition from ammonium limitation but increased in activity in the transition from glucose limitation. Steady-state enzyme activities in organisms that had been changed from ammonium to oxygen limitation were much less stable than in those that had initially been glucose-limited. Pyruvate dehydrogenase did not reach a steady state within 240 h, while citrate synthase showed damped oscillations, reaching a steady state after 180 h. The residual nutrient concentration and bacterial dry weight reached new steady-state values after 40 h. These oscillations may be due to changes in intracellular modulator concentrations, as a consequence of the severe oxygen limitation. Why they were not observed in the transition from glucose to oxygen limitation is not apparent and further study is needed.

To differentiate between glucose rather than oxygen acting as a repressive agent in these experiments, the effect of glucose limitation on an ammonium-limited culture was examined (Table 1). Isocitrate, succinate and malate dehydrogenases were unaffected by glucose limitation, while the two oxo-acid dehydrogenases and aconitase increased in activity. The effect of oxygen on the last three enzymes may thus be due, wholly or in part, to glucose repression. Citrate synthase and fumarate hydratase decreased in activity in glucose limitation. The response observed in oxygen limitation suggests that these enzymes may respond to low d.o.t. Experiments with ammonium-limited, oxygen-excess cultures, in which the residual steady-state glucose concentration was raised from 25 to 36 mM, showed that the enzyme and transport activities remained constant. Thus, the changes observed when an ammonium-limited culture was oxygen-limited are not due to the increase in the residual glucose concentration.

A transient peak was observed for NADH oxidase when glucose- or ammonium-limited cultures were oxygen-limited, the maximum activity occurring after 40 h; a new steady state was reached after 120 h. The new steady-state activity was usually slightly higher. This
Effects of oxygen on glucose metabolism

Fig. 3. Effect of gradual transition to oxygen limitation of an ammonium-limited culture on (a) steady-state activities of glucose dehydrogenase (○) and gluconate dehydrogenase (●), (b) steady-state activities of the transport systems for glucose (□), gluconate (■) and 2-oxoglucconate (▲), and (c) residual medium concentrations of glucose (○), gluconate (▲), 2-oxoglucconate (▲) and ammonium (×). The conditions were those described for Fig. 2: all values refer to steady states. Enzyme specific activities are expressed as μmol h⁻¹ (mg protein)⁻¹. Transport activities are expressed as μmol min⁻¹ (g dry wt)⁻¹.

behaviour is quite different from the response of A. beijerinckii in which the enzyme is induced at high d.o.t. (Jackson & Dawes, 1976). Further investigation is required to elucidate the reasons for this transient peak.

Effects of oxygen concentration: gradual transitions

In the transition from ammonium limitation to oxygen limitation, the d.o.t. showed a linear relationship to the partial pressure of oxygen in the inflowing gas (Fig. 2). The cell yield displayed a peak at 20% (v/v) oxygen, possibly because oxygen was toxic at higher d.o.t. values. Glucose dehydrogenase activity decreased with d.o.t., accompanied by an increase in the residual concentration of glucose and a decrease in those of gluconate and 2-oxoglucconate (Fig. 3a, c). Gluconate dehydrogenase did not decrease significantly until the glucose
C. G. MITCHELL AND E. A. DAWES

Fig. 4. Effect of gradual transition to oxygen limitation of a glucose-limited culture on the steady-state activities of hexokinase (■), gluconate kinase (■) and glucose-6-phosphate dehydrogenase (○). When the oxygen supply in the inflowing gas had been decreased to 4%, the glucose concentration in the inflowing medium was increased from 4.0 to 8.11 g l⁻¹ to ensure that glucose was in excess: $D = 0.36$ h⁻¹. Enzyme specific activities are expressed as μmol h⁻¹ (mg protein)⁻¹.

dehydrogenase activity had fallen by two-thirds and the gluconate concentration by one-half. As in the single-step experiments, glucose dehydrogenase decreased in activity faster than gluconate dehydrogenase, supporting its role as a central control point in glucose metabolism. In the transition from glucose limitation, gluconate dehydrogenase was unchanged in activity, while glucose dehydrogenase showed a further decrease when the culture approached oxygen limitation, confirming the single-step transition result.

In the transition from ammonium limitation, glucose transport activity remained constant until the d.o.t. reached 18 kPa, when it started to increase (Fig. 3b). This rise was associated with a decrease in residual gluconate concentration. Gluconate transport activity initially increased slightly as the d.o.t. was decreased, but both gluconate and 2-oxogluconate transport activities declined markedly below 10-65 kPa d.o.t. No change in activity for these systems was observed in the transition from glucose limitation, confirming the single-step studies.

The transition from glucose to oxygen limitation resulted in an initial increase in activity of glucose-6-phosphate dehydrogenase and hexokinase between 7.0 and 3.5 kPa d.o.t., prior to the decline observed in oxygen limitation (Fig. 4). Gluconate kinase did not respond to the change in d.o.t. while the 2-oxogluconate enzymes were completely repressed in glucose limitation. Although the Entner–Doudoroff enzymes displayed approximately the same specific activity [45 to 50 μmol h⁻¹ (mg protein)⁻¹] under both glucose and oxygen limitation, a sharp rise in activity occurred below 7 kPa d.o.t., reaching a peak of 120 μmol h⁻¹ (mg protein)⁻¹ at 5.3 kPa and then declining as the culture became oxygen-limited.

In a similar transition from glucose limitation, all the tricarboxylic acid cycle enzymes decreased in activity as the d.o.t. was lowered, except isocitrate dehydrogenase which increased slightly in activity (Fig. 5). Pyruvate dehydrogenase and aconitase activities did not decrease until glucose appeared in the culture. To establish the effect of d.o.t. on these catabolite-sensitive enzymes, further investigations with a different carbon source are necessary. NADH oxidase remained relatively constant in activity, except for a peak at 7.0 kPa d.o.t.

In both types of transition experiment, the effect of oxygen was shown to be reversible, eliminating the possibility of variant selection being responsible.
Our experiments clearly demonstrate that oxygen limitation of *P. aeruginosa* causes a diversion of glucose metabolism from the extracellular oxidative pathway to the intracellular phosphorylative route. This pattern of behaviour was previously observed when a glucose limitation was imposed on an ammonium-limited culture (Whiting et al., 1976b) and when the organism was grown anaerobically with nitrate in batch culture (Hunt & Phibbs, 1977). The enzymes and transport systems of the extracellular route are all decreased under oxygen limitation and it is apparent from the gradual transition experiments that they start to be affected when the d.o.t. falls below air saturation. The possibility that the observed effects might be attributed, at least in part, to nitrogen catabolite repression or to changes in the availability of ammonia was investigated by comparing the results recorded in Table 1 with those of experiments in which no ammonium addition was made when the oxygen limitation was imposed. In the latter case the residual ammonium concentration was about 6 mM compared with about 21 mM in the former experiments. With the exception of succinate...
dehydrogenase, which did not decrease in activity compared with a maximum fall of some 70% when ammonium was added during the transition (Table 1), there was no evidence for significant repression of these enzymes by ammonium.

Our experiments also show that certain tricarboxylic acid cycle enzymes, such as citrate synthase, malate dehydrogenase and isocitrate dehydrogenase, respond to changes in d.o.t. which suggests that such behaviour is not a unique manifestation of nitrogen-fixing organisms displaying respiratory protection.

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