Cytochrome Formation, Oxygen-induced Proton Extrusion and Respiratory Activity in *Streptococcus faecalis* var. *zymogenes* Grown in the Presence of Haematin

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**INTRODUCTION**

Recent studies (Bryan-Jones & Whittenbury, 1969; Sijpesteijn, 1970; Ritchey & Seeley, 1974) have revealed that certain lactic acid bacteria, formerly thought to be incapable of cytochrome synthesis, can form a functional electron transport system when grown in media containing haematin. Among streptococci, this ability is apparently restricted to *Streptococcus faecalis* and some strains of *S. lactis* (Ritchey & Seeley, 1976) but has also been reported in *Leuconostoc mesenteroides* NCIB9917 (Sijpesteijn, 1970). The evidence for the presence of a functional, membrane-bound electron transport system in the haematin-grown bacteria includes: sensitivity of the respiration to cyanide, antimycin A and *n*-heptyl-4-hydroxyquinoline-N-oxide (HOQNO); membrane location of much of the NADH oxidase activity; ATP formation coupled to NADH oxidation by cell membrane particles; and spectroscopy of whole organisms or membranes. The published spectra (Bryan-Jones & Whittenbury, 1969; Ritchey & Seeley, 1974) indicate an *α*-band around 560 nm indicative of a *b*-type cytochrome but show no evidence of other cytochromes. Sijpesteijn (1970)
reported the presence of peaks at 423 nm, at 556 to 558 nm and at 628 to 630 nm in _S. lactis_ grown in the presence of haematin but the spectra were not published.

The present study was undertaken, firstly, to characterize more fully the spectra of the cytochromes formed in the presence of haematin and, secondly, to investigate the capacity of haematin-grown bacteria to couple respiration to proton translocation. Observations on the effect of proton-translocating ionophores on the respiration of haematin-grown cells are also reported. The bacteria were grown in oxygen-limited continuous culture using a non-fermentable substrate, lactate, as the energy source to minimize the possibility of catabolite repression and to enhance the difference between bacteria grown with and without haematin.

**METHODS**

*Organism, Streptococcus faecalis* var. *zymogenes* strain TR was kindly supplied by Professor H. W. Seeley, Cornell University, Ithaca, U.S.A. It was maintained on a lactate/trypentine/yeast extract agar of the same composition as the liquid medium described below. The maintenance medium contained 10 µg haematin ml⁻¹.

**Growth conditions.** The medium used contained (g l⁻¹): tryptone (Difco), 10; yeast extract (Difco), 6.5; K₂HPO₄, 0.5; MgSO₄.7H₂O, 0.2; MnCl₂.4H₂O, 0.05; sodium dl-lactate (70% solution, BDH), 12.8. The medium was adjusted to pH 6.7 and sterilized in 15 l batches at 121°C for 60 min. For haematin-containing media, haematin (1 mg ml⁻¹) in 0.1 M NaOH was sterilized by membrane filtration and added aseptically to give 7.5 µg ml⁻¹. The organism was grown in continuous culture in a CCI500 chemostat (LH Engineering Co., Stoke Poges, Buckinghamshire) at 30°C. The culture volume was 2.25 l and the dilution rate was 0.083 h⁻¹. Foaming was prevented by automatic addition every 30 min of 0.2 ml of a 1:4 dilution of silicone antifoam RD (Dow Corning, Barry, South Glamorgan). The pH was controlled by automatic addition of 2.5 M NaOH.

The culture was sparged with an air/N₂ mixture containing O₂ at a partial pressure of 10 mmHg (1.33 kPa) at a gas flow rate of 750 ml min⁻¹. The culture was stirred at 425 rev. min⁻¹, giving a mass transfer coefficient (kₘ) for this fermenter of 36 h⁻¹ at the gas flow rate used. No dissolved O₂ was detected by the oxygen electrode and studies at different pO₂ levels showed that, at the partial pressure used, oxygen was the growth-limiting factor. The culture was maintained for 48 h to obtain a steady state as indicated by pH and extinction measurements. Samples were examined microscopically and by plating to establish culture purity. Samples for respiration measurement, proton-pulse studies and for cytochrome and enzyme determination were collected in an ice-cold receiver. Bacteria were harvested by centrifuging (10000 g, 10 min) and washed twice in 20 mM-phosphate buffer pH 7.0.

**Oxygen uptake measurement.** The harvested bacteria were resuspended in 100 mM-phosphate buffer pH 7.0 to a density of between 5 and 10 mg dry wt ml⁻¹. Oxygen uptake was determined in an oxygen electrode (Rank Bros, Bottisham, Cambridgeshire) at 30°C in the presence of 2.5 mM-sodium DL-lactate or 2.5 mM-glucose. The effects of uncoupling agents were examined by adding 1 or 2 µl of either carbonyl cyanide m-chlorophenylhydrazone (CCCP) or gramicidin D (both at 0.5 mg ml⁻¹ in methanol). At the concentration used, methanol had no effect on oxygen uptake.

**Determination of enzyme activities.** The harvested bacteria were disrupted in a Hughes press (Hughes, 1951) cooled to −30°C with dry ice. The homogenate was fractionated into a thoroughly washed cell wall/membrane fraction and a high speed supernatant fraction [from which particulate matter had been removed by centrifuging (90000 g, 2 h, 0°C)] according to the procedure of Gray et al. (1966).

Reaction mixtures (300 ml) for enzyme assays had the following compositions.

**NADH oxidase:** 275 µmol potassium phosphate buffer pH 7.0; 0.25 µmol NADH; 0.2 ml of either cell wall/membrane suspension (40 to 60 µg protein) or supernatant fraction (200 to 400 µg protein). The rate of decrease in extinction at 340 nm was measured.

**NADH peroxidase:** 275 µmol sodium acetate buffer pH 5.4; 0.25 µmol NADH; 5 µmol H₂O₂; 0.2 ml supernatant fraction (50 to 100 µg protein). Residual NADH oxidase activity at pH 5.4 was measured before adding the peroxide and peroxidase activity was determined from the difference in rates of NADH oxidation before and after peroxide addition.

**NADH: dichlorphenolindophenol (DCPIP) oxidoreductase:** 265 µmol Tris/HCl buffer pH 7.4; 0.25 µmol NADH; 0.1 µmol DCPIP; 0.2 ml cell wall/membrane suspension (10 to 12 µg protein). The decrease in extinction at 600 nm was measured.

Protein was determined by the method of Lowry et al. (1951).

**Cytochromes.** Cytochrome spectra of the cell wall/membrane fraction were determined at room temperature and in liquid nitrogen (−196°C); whole organism spectra were determined in liquid nitrogen. Room temperature spectra of the membrane suspension (1 to 1.5 mg protein ml⁻¹) in 20 mM-phosphate buffer
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pH 7.0 were determined in 1 ml cuvettes with a 1 cm light path in a Cary 14 spectrophotometer. Difference spectra were recorded for samples reduced with dithionite and read against samples oxidized either with air or with a few crystals of potassium ferricyanide. For CO-difference spectra, samples reduced with dithionite were gassed with CO for 60 s and spectra were determined using a dithionite-reduced sample as reference.

Spectra at −196 °C of whole organism suspensions (approx. 50 mg dry wt ml−1) or of cell wall/membrane suspensions (5 mg protein ml−1) in 0.8 M mannitol were determined using 0.2 ml cuvettes with a 2 mm light path. Difference spectra were recorded for samples reduced with lactate (about 2 μl 70 % sodium lactate solution per 0.2 ml suspension) or with a few crystals of dithionite and read against samples oxidized with air or with a few crystals of ferricyanide. Samples were immersed in liquid nitrogen and maintained at this temperature by immersion of the base of the cuvette holder in a small Dewar flask in the turbid sample compartment of a Unicam SP800 spectrophotometer.

Proton pulse measurement. Oxygen-induced proton pulses were determined by a procedure similar to that described by Scholes & Mitchell (1970). The harvested bacteria were thoroughly washed twice in buffer (140 mM-KCl/1 mM-glycylglycine pH 7.0). They were resuspended in this buffer to a density of about 10 mg dry wt ml−1 and carbonic anhydrase (100 μl of a 5 mg ml−1 solution) was added. The suspension was placed in a Perspex vessel maintained at 30 °C (a Rank oxygen electrode vessel with a magnetic stirrer was used for this purpose). The pH was measured with a combination glass/reference electrode (Electronic Instruments, Richmond, Surrey) inserted tightly through a silicone rubber stopper so that the suspension completely filled the chamber. Additions were made by means of a Hamilton syringe inserted between the vessel wall and the stopper. The electrode output was measured and amplified by a Vibron Electrometer (33B-2) and pH measuring unit (C33B-2) (Electronic Instruments). Approximately 30 min was allowed for the suspension to become completely anaerobic. By the end of this pre-incubation period, the pH drift had settled to a very slow rate of change and the pH was brought back to 7.0 by addition of NaOH. Sodium dl-lactate (50 μl, 0.1 M) was added and a further 5 to 10 min was allowed for the pH drift to settle again. Valinomycin was then added to give concentrations between 0.5 and 5.0 pg ml−1, followed by 50 μl KCl/glycylglycine buffer saturated with pure oxygen at 30 °C. Proton production was measured by calibrating with 10 μl additions of 10 mM-HCl. Pulse height was measured by extrapolating the pulse slope and the subsequent drift slope back to the point of intersection.

Chemicals. Haematin, valinomycin, CCCP, gramicidin D, NADH and carbonic anhydrase were obtained from Sigma. All other chemicals, except where specified, were obtained from BDH.

RESULTS

Enzyme activities

Four NADH-oxidizing enzyme systems were measured in bacteria grown with and without haematin (Table 1). Bacteria grown with haematin had a high membrane-bound NADH oxidase activity whereas membranes of the bacteria grown without added haematin had a very low NADH oxidase activity. Possibly the low activity of particulate NADH oxidase in organisms grown without haematin represents adsorbed soluble NADH oxidase activity in spite of the thorough washing of the membrane fraction (three washes in phosphate buffer) since a low level of NADH peroxidase was also present in the membrane fraction of these bacteria. However, the oxidase:peroxidase ratio was higher in the membrane fraction than in the soluble fraction from bacteria grown without haematin.

In contrast to the NADH oxidase activity, the membrane-bound NADH:DCPIP oxido-reductase activity was higher in membranes from the bacteria grown without haematin than in the haem-grown bacteria. The soluble NADH oxidase activity was also higher in bacteria grown without haematin, as found in a previous study (Ritchey & Seeley, 1974). NADH peroxidase activity was unaltered by growth in a haematin-containing medium. No catalase activity was detected in bacteria grown with or without haematin.

Cytochromes

Reduced minus oxidized difference spectra of the cell wall/membrane fraction of haematin-grown bacteria showed a Soret peak at 432 nm and α-peaks at 562 and 632 nm when measured at room temperature (Fig. 1). The 562 nm peak was superimposed on a broad absorption region between 500 and 600 nm which is probably due to adsorbed haematin, since haematin in solution also shows this broad absorbance with a λ_max at 580 nm (Fig. 1).
Table 1. Activities of NADH-oxidizing enzyme systems in *S. faecalis var. zymogenes* grown with and without haematin

Enzyme activities are expressed as μmol NADH oxidized min⁻¹ (mg protein)⁻¹.

<table>
<thead>
<tr>
<th>Enzyme system</th>
<th>Bacteria grown with haematin (7.5 μg ml⁻¹)</th>
<th>Bacteria grown without haematin</th>
</tr>
</thead>
<tbody>
<tr>
<td>Particulate NADH oxidase</td>
<td>2.31</td>
<td>0.19</td>
</tr>
<tr>
<td>(in cell wall/membrane fraction)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Soluble NADH oxidase</td>
<td>0.18</td>
<td>0.48</td>
</tr>
<tr>
<td>(in 90,000 g supernatant)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Soluble NADH peroxidase</td>
<td>0.84</td>
<td>0.89</td>
</tr>
<tr>
<td>Particulate NADH:DCPIP oxidoreductase</td>
<td>2.15</td>
<td>4.02</td>
</tr>
</tbody>
</table>

Fig. 1. Dithionite-reduced minus air-oxidized difference spectra at room temperature of the cell wall/membrane fraction (1.75 mg protein ml⁻¹) of *S. faecalis var. zymogenes* grown in the presence of haematin (--), and of a haematin solution (10 μg ml⁻¹; ---). Both the membrane suspension and the haematin solution were in 20 mM-phosphate buffer pH 7.0.

The shoulder on the cell wall/membrane spectrum at 580 nm was presumably due to this haematin absorbance.

The liquid nitrogen difference spectrum of whole organisms (Fig. 2) grown in the absence of haematin showed a trough between 460 and 480 nm presumably due to flavoprotein absorbance but no trace of any cytochrome peaks. In the haematin-grown organisms, the intense flavoprotein absorbance largely obscured the Soret peak of the cytochromes but a small peak was evident at 430 nm. Two b-type cytochrome peaks were evident in the α-band spectrum, at 558 and 562 nm. This double peak was present in the low temperature spectra of organisms of this strain of *S. faecalis* grown under a wide range of aeration conditions. It was also present in the low temperature spectrum of the cell membrane fraction, again superimposed on an adsorbed haematin absorbance (Fig. 2). Membranes obtained from bacteria grown in the absence of haematin were suspended in a haematin solution (10 μg ml⁻¹) at pH 7.0 for 2 h and then sedimented, washed and resuspended in 0.8 M-mannitol. The liquid nitrogen difference spectrum of these membranes (Fig. 2) showed the broad peak due to adsorbed haematin with a λmax at 572 nm. There was no evidence of either the cytochrome b or the cytochrome d peaks.

The CO-difference spectrum (Fig. 3) showed a peak at 417 nm characteristic of an o-type cytochrome. The CO-difference spectrum of membranes obtained from bacteria grown
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**Fig. 2.** Reduced minus oxidized difference spectra of whole organisms and of the cell wall/membrane fraction of *S. faecalis* var. *zymogenes* in 0.8 M-mannitol at -196 °C. (a) Dithionite-reduced minus air-oxidized spectrum of haematin-grown organisms (25 mg dry wt ml⁻¹). (b) Dithionite-reduced minus air-oxidized spectrum of organisms grown without haematin (50 mg dry wt ml⁻¹). (c) Dithionite-reduced minus ferricyanide-oxidized spectrum of the cell wall/membrane fraction (5 mg protein ml⁻¹) from haematin-grown organisms. (d) Dithionite-reduced minus ferricyanide-oxidized spectrum of the cell wall/membrane fraction (5 mg protein ml⁻¹) of organisms grown without haematin after the membranes had been suspended in haematin solution (10 μg ml⁻¹, pH 7.0) for 2 h, then sedimented, washed and resuspended in 0.8 M-mannitol.

**Fig. 3.** Carbon monoxide difference spectra of the cell wall/membrane fraction prepared from *S. faecalis* var. *zymogenes* grown in the presence of haematin (---) and of membranes from organisms grown without haematin to which haematin has been adsorbed from solution (---). Both suspensions had a density of 1.0 to 1.2 mg protein ml⁻¹.

without haematin and then allowed to adsorb haematin showed no such intense absorbance at 417 nm. The peaks at 535 and 570 nm may be due to haematin itself since they can be obtained in the CO-difference spectrum of a haematin solution.
Respiratory activity and sensitivity to uncoupling agents

Haematin-grown bacteria oxidized lactate 10 to 20 times faster than did bacteria grown in the absence of haematin (Table 2). The lactate respiration of haematin-grown bacteria was strongly inhibited by low concentrations of the uncoupling ionophores CCCP and gramicidin D. The low rate of lactate respiration obtained in bacteria grown without added haematin was only slightly inhibited by these uncoupling agents. In contrast to lactate respiration, glucose respiration was only about 20% higher in haematin-grown bacteria than in those grown without haematin. Furthermore, CCCP and gramicidin D stimulated glucose respiration in haematin-grown bacteria by between 16 and 27%. In the bacteria grown without haematin these uncoupling agents inhibited glucose-induced oxygen uptake by 10 to 20%. These effects of uncoupling agents on the respiration of haematin-grown bacteria were very reproducible. A second culture of haematin-grown bacteria gave the same response pattern: 70 to 75% inhibition of lactate respiration and 20 to 25% stimulation of glucose respiration by CCCP and gramicidin D.

Oxygen-induced proton pulse production

When an anaerobic suspension of washed, haematin-grown bacteria suspended in very weak buffer (1 mM-glycylglycine/140 mM-KCl) in the presence of lactate and valinomycin was injected with a 50 nl pulse of oxygen-saturated buffer, a rapid acidification of the medium occurred (Fig. 4). Maximum acidification was obtained in the presence of 1 mM-lactate and 0.5 µg valinomycin ml⁻¹; higher concentrations of either of these did not increase the size of the proton pulse. With increasing incubation time there was a slight decrease in pulse size; this was probably due to leakage of compounds from the cells which increase the buffering capacity of the suspension medium since pulses produced by injection of standard acid were correspondingly decreased. The proton pulses were eliminated by addition of CCCP. Calculation of the →H⁺/O stoichiometry from 22 pulses gave a value of 1.37 ± 0.27. Bacteria grown without added haematin normally showed no proton pulses on addition of oxygen-saturated buffer. One batch of bacteria grown in the absence of haematin at a very low pₒ₂ (4 mmHg) did produce small and rather sluggish proton pulses. This particular batch of cells had a higher rate of lactate respiration [11 nmol min⁻¹ (mg dry wt)⁻¹] than those used in the present study.

DISCUSSION

Our results confirm earlier findings (Bryan-Jones & Whittenbury, 1969; Ritchey & Seeley, 1974) that some strains of S. faecalis can synthesize a functional, membrane-bound cytochrome system when grown in the presence of added haematin. In addition to the b-type cytochrome there are two possible oxidase components, a cytochrome d₅₃₀ and the CO-binding cytochrome. The production of proton pulses by such cells when supplied with oxygen indicates that the cytochrome system is vectorially oriented in the membrane, as in other cytochrome-containing aerobes, so that electron transport to oxygen may be chemiosmotically coupled to phosphorylation or to transport processes. The magnitude of the proton pulse produced in response to a given amount of oxygen (the →H⁺/O ratio) has been used to estimate the P/O ratio of bacterial cells (Lawford & Haddock, 1973; Jones et al., 1975) on the assumption that two H⁺ are involved in the production of one ATP via the proton-translocating ATPase system. The average →H⁺/O ratio obtained with the haematin-grown bacteria in the present study was 1.4 which corresponds to a P/O ratio of 0.7. From a comparison of the molar growth yield on mannitol and glucose, Ritchey & Seeley (1974) calculated a P/O ratio of 0.75 for this same strain of S. faecalis. Direct measurement of the P/O ratio using phosphorylating membrane particles yielded lower values of between 0.3 and 0.4 (Bryan-Jones & Whittenbury, 1969; Ritchey & Seeley, 1974) but such low values are commonly obtained from in vitro measurements with bacterial membranes (Gel'man, Lukoyanova & Ostrovskii, 1967). Calculation of the P/O ratio from the →H⁺/O value does
Table 2. Effect of uncoupling agents on lactate and glucose oxidation by S. faecalis var. zymogenes grown with and without haematin

Oxygen uptake rates of washed suspensions of bacteria grown in a lactate/tryptone/yeast extract medium (QO2 values) are expressed as nmol O2 min⁻¹ (mg dry wt)⁻¹.

<table>
<thead>
<tr>
<th>Oxidation substrate and treatment</th>
<th>Bacteria grown with haematin (7.5 µg ml⁻¹)</th>
<th>Bacteria grown without haematin</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na DL-lactate (2.5 mM)</td>
<td>106.2</td>
<td>43.1</td>
</tr>
<tr>
<td>Lactate+CCCP (0.5 µg ml⁻¹)</td>
<td>51.0</td>
<td>37.0</td>
</tr>
<tr>
<td>Lactate+CCCP (1.0 µg ml⁻¹)</td>
<td>24.2</td>
<td>3.2</td>
</tr>
<tr>
<td>Lactate+gramicidin D (0.5 µg ml⁻¹)</td>
<td>26.7</td>
<td>3.4</td>
</tr>
<tr>
<td>Glucose (2.5 mM)</td>
<td>103.5</td>
<td>80.0</td>
</tr>
<tr>
<td>Glucose+CCCP (0.5 µg ml⁻¹)</td>
<td>131.0</td>
<td>ND</td>
</tr>
<tr>
<td>Glucose+CCCP (1.0 µg ml⁻¹)</td>
<td>131.0</td>
<td>60.4</td>
</tr>
<tr>
<td>Glucose+gramicidin D (0.5 µg ml⁻¹)</td>
<td>126.5</td>
<td>71.1</td>
</tr>
</tbody>
</table>

ND, Not determined.

Fig. 4. Changes of pH in anaerobic suspensions of S. faecalis var. zymogenes in response to oxygen pulses: (a) haematin-grown bacteria; (b) bacteria grown without haematin. Bacteria were suspended (10 mg dry wt ml⁻¹) in 5 ml of 1 mM-glycylglycine/140 mM-KCl buffer pH 7.0. Sodium DL-lactate (5 µmol) and valinomycin (4.5 nmol) were added prior to pulsing with oxygen-saturated buffer. Carbonyl cyanide m-chlorophenylhydrazone (CCCP; 12.25 nmol) was added at the time indicated.

not take into account any oxygen uptake mediated by the soluble, flavoprotein electron transport system in S. faecalis (Dolin, 1955). The error due to neglecting this component would be very small since there is apparently a very low rate of lactate respiration by this system alone (Table 2). Bacteria grown without added haematin do not usually show any oxygen-induced proton pulses although the low activity of membrane-bound NADH oxidase and the small CCCP-sensitive component of lactate respiration may indicate that even these bacteria have traces of a cytochrome system due possibly to haem compounds present in the yeast extract used in the medium. This could account for the occasional finding of small and sluggish proton pulses in non-haem-grown bacteria at high cell densities.

The stimulation of the rate of oxygen uptake by CCCP and gramicidin during glucose oxidation is further evidence of respiratory control in haematin-grown bacteria. The oxidation of glucose by such bacteria is presumably mediated in part by the membrane-bound NADH oxidase system and in part by the soluble, flavoprotein oxidase. Glucose respiration by non-haem-grown bacteria, which must be mediated almost solely by the soluble oxidase system, was inhibited 10 to 20 % by the uncoupling agents. If this same inhibition affects the soluble oxidase component of glucose respiration in the haematin-grown bacteria then the
stimulation of the membrane-bound component by uncouplers may be even greater than the 20 to 25% observed.

The high rate of lactate oxidation by haematin-grown bacteria and its sensitivity to inhibition by uncoupling agents markedly contrasts with the situation in bacteria grown without haematin. Harold & Levin (1974) have shown that lactate accumulation by S. faecalis is a function of the pH difference between the cell interior and the medium and is inhibited by ionophores which dissipate the pH gradient. The increased rate of lactate respiration in haematin-grown bacteria is probably due to an enhanced pH gradient due to the oxygen-stimulated proton extrusion. The inhibition of uptake by uncoupling agents would greatly outweigh any stimulatory effects on subsequent oxidation by the membrane-bound oxidase system.

One unresolved problem concerns the mechanism of lactate oxidation in the haematin-grown bacteria. No evidence has been obtained from the present study that lactate can be oxidized directly by the membrane-bound oxidase system. Membrane suspensions showed no oxygen uptake with added lactate (in contrast to a very active uptake with NADH) nor was there any detectable cytochrome reduction by lactate in membrane suspensions although lactate fully reduced the cytochromes in whole cells. There was no lactate DCPIP oxidoreductase activity in either the membrane fraction or the soluble fraction of haem-grown cells (even in the presence of phenazine methosulphate) in contrast to the high NADH: DCPIP oxidoreductase activity in the membrane fraction. Therefore, either the lactate oxidase system is very labile or lactate oxidation in whole cells is due to the combined action of a soluble, NAD-dependent lactate dehydrogenase and the NADH oxidase system. The NAD-dependent lactate dehydrogenase of S. faecalis has an obligatory requirement for fructose-1,6-diphosphate for activity (Wittenberger & Angelo, 1970). The concentration of this activator is unlikely to be very high in bacteria grown in the absence of glucose so the enzymic mechanism responsible for the high rate of lactate oxidation found in haematin-grown bacteria is in need of further investigation.

REFERENCES


