Terminal Oxidations in *Bacillus brevis*. Soluble Reduced Nicotinamide Adenine Dinucleotide Oxidase Activity in *Bacillus brevis*

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*Bacillus brevis* grown so as to produce tyrothricin possesses an active cytochrome-linked NADH oxidase system (Seddon & Fynn, 1971). Electron-transport particles, containing flavoprotein, a menaquinone and cytochromes of type \(a+a_3\), \(b\), \(c\) and \(o\) possess most of the NADH oxidase activity of this organism during exponential growth (Fynn & Seddon, 1971; Seddon & Fynn, 1971; Fynn, Thomas & Seddon, 1972). Their NADH oxidase activity is inhibited by classical inhibitors of the respiratory electron-transport chain such as cyanide, antimycin A, 2-n-heptyl-4-hydroxyquinoline-N-oxide (HQNO), amytal and rotenone showing characteristics typical of a respiratory chain electron-transport system (Seddon & Fynn, 1971). The oxidative activity of this particulate electron-transport system is also inhibited by levels of tyrothricin and tyrocidine comparable to those produced towards the latter end of the exponential growth phase of this organism and indicates the possibility that such activity may be inhibited *in vivo*. However, the soluble fraction of this organism is capable of oxidizing NADH by molecular \(O_2\) (Seddon & Fynn, 1970; Fynn & Seddon, 1971). As tyrothricin inhibits the particulate NADH oxidase system its effects on the soluble NADH oxidase are of interest since, if the soluble NADH oxidase is insensitive to tyrothricin, the possibility exists that the soluble NADH oxidase may take over a redox role should the particulate fraction be inhibited.

The present communication describes and discusses properties of the soluble NADH oxidase and the effects on it of inhibitors, including tyrocidine and tyrothricin.

**METHODS**

*Organism, growth and maintenance.* *Bacillus brevis* ATCC 10068 was maintained and grown as described by Seddon & Fynn (1971) on an asparagine-glycerol basal salts medium as used by Mach, Reich & Tatum (1963). Bacteria were harvested after exponential growth (24 to 30 h).

*Preparation of subcellular fractions.* Organisms were harvested from 1 l of media, washed, resuspended in tris-HCl buffer (0.1 M), pH 7.4, and extruded through a French pressure cell at 0 to 5 °C and 10 000 to 16 000 lb/in\(^2\). Centrifugation at 15 000 \(g\) removed whole bacteria and larger debris. The crude supernatant fraction was further separated at 105 000 \(g\) to yield a particulate fraction (electron-transport particles) and a soluble fraction. This method of fractionation has been described in detail (Seddon & Fynn, 1971).

*Assay of NADH dehydrogenase.* Ferricyanide reduction was used to measure dehydrogenase activity. The assay system contained tris-HCl buffer (pH 7.4), 300 \(\mu\) mol; potassium
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ferricyanide, 5 μmol; KCN (where added), 12 μmol; NADH, 0.33 μmol; and enzyme protein 0.1 to 5.0 mg in a total volume of 3.0 ml at 30 °C.

Assay of NADH-menadione reductase. This was assayed as described by Sanadi, Pharo & Sordahl (1967) using the following assay system: tris-HCl (pH 7.4), 300 μmol; menadione, 0.5 μmol; KCN (where present), 12 μmol; NADH, 0.33 μmol and enzyme protein 0.1 to 5.0 mg in a total volume of 3.0 ml at 30 °C.

Assay of NADH-cytochrome c reductase. Reduction of cytochrome c at 550 nm was used as described by Hatefi & Rieske (1967). The reaction mixture contained tris-HCl buffer (pH 7.4), 300 μmol; cytochrome c, 0.05 μmol; NADH, 0.33 μmol; KCN (where present), 12 μmol and enzyme protein, 0.2 to 5.0 mg in a total volume of 3.0 ml at 30 °C.

Assay of cytochrome c oxidase. The cytochrome c oxidase activity was measured using a Rank O₂ electrode. The rate of O₂ uptake was monitored upon addition of ascorbate plus cytochrome c as described by Smith & Camerino (1963). The reaction mixture contained tris-HCl buffer (pH 7.4), 300 μmol; cytochrome c, 1 μmol; sodium ascorbate, 20 μmol and enzyme protein, 1 to 20 mg. Total volume was 3.0 ml. A control without enzyme was used to correct for the autoxidation of sodium ascorbate under the conditions of incubation and assays were made at 30 °C.

Difference spectra. Difference spectra were obtained using a Perkin-Elmer model 356 two-wavelength double-beam spectrophotometer operated in the split beam mode. Comparison was made between an anaerobic sample (NADH reduced) and an aerobic sample (treated with O₂). Carbon monoxide-reduced minus reduced spectra were obtained by bubbling carbon monoxide for up to 10 min through the sample cuvette or by mixing the sample contents with an equal volume of carbon monoxide-saturated buffer whilst the reference solution was mixed with an equal volume of N₂-saturated buffer.

Protein determination. Protein concentrations were measured by the procedure of Lowry, Rosebrough, Farr & Randall (1951) using crystalline bovine plasma albumin (fraction V) as standard.

Chemicals. The chemicals used throughout this work were of A.R. quality when available. More specialized chemicals were obtained from the Sigma Chemical Co., St Louis, Missouri, U.S.A. except those indicated as follows: bovine plasma albumin (fraction V), Armour Pharmaceutical Co. Ltd. Eastbourne Sussex; tyrothricin, tyrocidine and gramicidin from Nutritional Biochemicals Co., Cleveland, Ohio, U.S.A.

RESULTS AND DISCUSSION

Although the specific activity of the NADH oxidase is usually lower in the soluble fraction than in the electron-transport particles the total activity present can be higher in the soluble fraction. Specific activities measured ranged from 20 to 95 ng-atoms O₂/min/mg protein with the particulate fraction and from 13 to 29 ng-atoms O₂/min/mg protein with the soluble fraction and total activities from 2.7 to 8.9 μg-atoms O₂/min and from 4.6 to 13.1 μg-atoms O₂/min with the particulate and soluble fractions respectively. Low observed NADH oxidase activity in the particulate fraction did not necessarily correspond with high NADH oxidase activity in the soluble fraction of the same preparation and vice versa. It appears that the particulate NADH oxidase varies considerably. The soluble NADH oxidase activity was found not to vary much more than twofold. As prolonged (150 min) centrifugation at 105000 g did not further decrease the soluble NADH oxidase activity it was inferred that the soluble NADH oxidase activity was substantial in amount and was not a result of contaminating particulate matter.
Table 1. Comparison of the effects of various treatments and substances on the NADH oxidase activities of the 105,000 g particulate and soluble fractions of Bacillus brevis

All measurements were made by following O₂ uptake on addition of NADH at 30 °C using a Rank O₂ electrode (Seddon & Fynn, 1971). All inhibitors except KCN and atebrin were added as small volumes (0.05 ml or less) of ethanolic solutions.

<table>
<thead>
<tr>
<th>Treatment or addition</th>
<th>Time or concentration</th>
<th>105,000 g particulateoxidase activity</th>
<th>105,000 g solubleoxidase activity</th>
</tr>
</thead>
<tbody>
<tr>
<td>Incubation at 100 °C</td>
<td>10 min</td>
<td>-100</td>
<td>-100</td>
</tr>
<tr>
<td>Storage at -16 °C</td>
<td>24 h</td>
<td>-80</td>
<td>0</td>
</tr>
<tr>
<td>KCN</td>
<td>10 mM</td>
<td>-90</td>
<td>0</td>
</tr>
<tr>
<td>Antimycin A</td>
<td>0.1 μmol/mg protein</td>
<td>-80</td>
<td>0</td>
</tr>
<tr>
<td>HQNO</td>
<td>0.5 μmol/mg protein</td>
<td>-55</td>
<td>0</td>
</tr>
<tr>
<td>Amytal</td>
<td>0.5 μmol/mg protein</td>
<td>-50</td>
<td>0</td>
</tr>
<tr>
<td>Rotenone</td>
<td>0.5 μmol/mg protein</td>
<td>-50</td>
<td>0</td>
</tr>
<tr>
<td>Atebrin</td>
<td>4 mM</td>
<td>-30</td>
<td>-70</td>
</tr>
<tr>
<td>Dialysis</td>
<td>24 h</td>
<td>-</td>
<td>-40</td>
</tr>
<tr>
<td>FMN</td>
<td>40 μM</td>
<td>0</td>
<td>+155</td>
</tr>
<tr>
<td>FAD</td>
<td>40 μM</td>
<td>0</td>
<td>+135</td>
</tr>
<tr>
<td>Dialysis + FMN</td>
<td>24 h + 40 μM</td>
<td>-</td>
<td>+155</td>
</tr>
<tr>
<td>Tyrocidine</td>
<td>2 mM or 0.1 mM</td>
<td>-100</td>
<td>0</td>
</tr>
<tr>
<td>Tyrothricin</td>
<td>2 mM or 0.1 mM</td>
<td>-100</td>
<td>0</td>
</tr>
<tr>
<td>Gramicidin S</td>
<td>2 mM or 0.1 mM</td>
<td>-100</td>
<td>0</td>
</tr>
<tr>
<td>Gramicidin</td>
<td>2 mM or 0.1 mM</td>
<td>-100</td>
<td>0</td>
</tr>
</tbody>
</table>

NADH-reduced minus oxidized difference spectra of the soluble fraction showed that, although the 500 to 600 nm region of the spectrum was not distinct enough to detect any α-peaks of the cytochromes, the Soret region has only one absorbance band with a peak around 420 nm (possibly a cytochrome of type c) and suggests the absence of type α+α₂ cytochrome oxidase. In agreement with this finding CO-NADH-reduced minus NADH-reduced difference spectra showed nothing in the 400 to 450 nm region and thus failed to indicate the participation of a carbon monoxide reactive cytochrome. The soluble fraction showed component enzyme activities of 0.812, 1.145 and 0.856 μmol substrate oxidized or reduced/min/mg protein at 30 °C for NADH-cytochrome c reductase, NADH-menadione reductase and NADH dehydrogenase respectively. Cytochrome c oxidase activity could not be detected. The absence of cytochrome c oxidase activity again indicates the lack of cytochromes of type α+α₂. The overall NADH oxidase activity (13 to 29 ng-atoms O/min/mg protein) is much less than any of the three reductase activities assayed and suggests a rate-limiting constraint on the oxidase activity.

NADH-reduced minus oxidized difference spectra also showed a large trough at 450 to 460 nm, indicating the involvement of flavin in the oxidation of NADH by the soluble fraction. The flavin involved with the soluble NADH oxidase is about 70% of the total flavin content (estimated by dithionite reduction). Dialysis of the soluble fraction at 0 to 5 °C against 0.1 M-tris-HCl buffer, pH 7.4, led to a loss of 40% of the soluble NADH oxidase activity. Addition of FMN or FAD restored the activity as shown in Table 1. The slightly higher activity with FMN may indicate FMN as the flavin component of the NADH oxidase.

Table 1 shows that the soluble NADH oxidase, in contrast to the particulate NADH oxidase, is not inhibited by cyanide, antimycin A, HQNO, amytal or rotenone. It was, how-
ever, inhibited by atebrin again implicating flavin. Tyrothricin, tyrocidine, gramicidin S and gramicidin inhibit the particulate NADH oxidase (Table 1) but not the soluble NADH oxidase. Whether the electron-transport chain NADH oxidase remains sensitive to tyrothricin production in the intact organism is not yet known, but if it does then the soluble NADH oxidase could provide an uninhibited route for reducing equivalents, though it is unlikely that this would lead to efficient ATP formation since such systems do not normally possess sites of phosphorylation (Asano & Brodie, 1965).

Soluble NADH oxidase is found in a wide variety of micro-organisms (Dolin, 1961) and the particular function such a system may play is not known. At the end of active growth, sporulating strains of bacilli require an active tricarboxylic acid cycle, presumably for the synthesis of materials necessary for spore formation. During sporulation, although increase in bacterial numbers has stopped, there must be a complete redirection of synthesis geared to the production of the mature spore. A functional tricarboxylic acid cycle could play a central role in the degradation of vegetative material and the synthesis of material essential for the development of the mature spore. Commitment to sporulation appears to be an irreversible process (Gould & Hurst, 1969) and the possible inhibition of particulate NADH oxidase by tyrothricin production would also be expected to be irreversible since such antibiotics interfere with the integrity of membranes (Hunter & Schwartz, 1967). The soluble NADH-oxidizing system would ensure the reoxidation of NADH such that the NAD+/NADH redox couple participation in the tricarboxylic acid cycle could operate even under conditions when the electron-transport NADH oxidase system may be lost or absent.

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REFERENCES


