Electron Transport in Phosphorylating Mitochondria from *Tetrahymena pyriformis* Strain ST

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**SUMMARY**

Mitochondria isolated from *Tetrahymena pyriformis* carried out oxidative phosphorylation with succinate and 2-oxoglutarate as substrates. Electron transport was inhibited by rotenone, piericidin A, antimycin A and cyanide. Succinate-cytochrome c and ferricyanide oxidoreductases were antimycin A sensitive. NADH-cytochrome c and ferricyanide oxidoreductases were only partially inhibited by high concentrations of rotenone. Externally added NADH gave no oxygen uptake in the absence of artificial electron acceptors. The mitochondria contained haems a, c and protohaem, and difference spectra revealed the presence of cytochromes b, c, c, and a pigment with an extinction maximum at 620 nm. Steady-state and kinetic measurements of cytochrome components were made. Several kinetically distinct flavoprotein components were present. Kinetic measurements suggested that the reduced 620 nm component reacted sufficiently rapidly with molecular oxygen to have been the terminal oxidase.

**INTRODUCTION**

Mitochondria capable of oxidative phosphorylation were first isolated from *Tetrahymena pyriformis* strain GL by Kobayashi (1965). In this and in earlier studies (Ryley, 1952; Eichel, 1954) cytochrome oxidase activity was not detected when reduced mammalian cytochrome c was used as substrate for cell-free preparations; neither was the α-band of cytochrome oxidase demonstrated spectroscopically.

The present work was undertaken to obtain a better understanding of electron transport in phosphorylating mitochondria isolated from *Tetrahymena*. Strain ST has been used previously in studies of mitochondrial DNA (Suyama & Eyer, 1968), and the method used here for isolation of mitochondria is based on those of Kobayashi (1965) and Suyama & Preer (1965). A preliminary account of some of this work has previously been published (Turner, Lloyd & Chance, 1969).

**METHODS**

*Maintenance growth, and harvesting of the organism.* *Tetrahymena pyriformis* strain ST (kindly provided by Dr Y. Suyama, Department of Biology, University of Penn-

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sylvania) was maintained, grown and harvested as described previously (Lloyd, Brightwell, Venables, Roach & Turner, 1971). Volumes of culture up to 20 l. were used for the preparation of mitochondria, and organisms were harvested in late exponential phase of growth (5 to 7 x 10^6 organisms/ml).

Disruption of organisms. The organisms were washed once in preparation buffer consisting of 0.35 M-mannitol, 0.1 % (w/v) bovine serum albumin, 0.1 mM-EGTA (ethylene glycol-bis (β-amino-ethyl ether) N,N'-tetraacetic acid), 10 mM-tris-HCl, pH 7.2, and resuspended in 5 vol. of this buffer (approximately 10 mg. protein/ml.). The suspension was homogenized in a 50 ml. Kontes (Vineland, New Jersey, U.S.A.) tight fitting (type B plunger) glass hand-homogenizer in ice, until over 90 % of the cells were broken (20 to 30 strokes).

Preparation of mitochondrial fractions. All operations were carried out at 4°. The whole homogenate was transferred to 50 ml. centrifuge tubes and centrifuged at 300 g for 5 min. to remove whole organisms and pellicles. The supernatant was re-centrifuged at 7000 g for 10 min. The resulting pellet consisted of a small amount of tightly packed reserve material at the bottom of the tube, a cream-coloured layer of mitochondria, and a loosely packed white layer above this consisting of cell membranes, cilia and possibly lysosomes and peroxisomes. The supernatant (containing microsomal and 'soluble' fractions) was carefully removed and most of the upper white layer with it. Five vol. of preparation buffer were added to the residue and shaken gently to remove the mitochondrial fraction, which was resuspended and recentrifuged at 7000 g for 10 min. The separation procedure was repeated and the mitochondrial fraction retained (approx. 10 mg. protein/l. of culture).

Analytical methods. Polarograph measurements of oxygen uptake by mitochondrial suspensions were made with an oxygen electrode (Lloyd & Brookman, 1967). Respiratory control ratios and ADP/O ratios were calculated by the method of Chance & Williams (1956). Difference spectra were traced with a Cary model 14 split-beam spectrophotometer fitted with a 0 to 0.1 extinction slide wire. Low temperature spectra were obtained using a split-beam spectrophotometer (Yang & Legallais, 1954) with a liquid N₂ attachment (Chance, 1957). Steady-state reduction levels of electron transport components were measured using a dual wavelength spectrophotometer (Chance, 1951). Unless otherwise stated, the buffer used in all analytical experiments contained 0.35 M-mannitol, 0.1 mM-EGTA, 10 mM-KCl, 10 mM phosphate, and 0.1 % (w/v) bovine plasma albumin at pH 7.2. Rapid reaction measurements of the reoxidation of substrate-reduced electron transport components in whole mitochondria were made using the regenerative flow apparatus of Chance (1954) fitted with fluorometric and dual wavelength measuring facilities. Briefly, the method involves mixing an anaerobic mitochondrial suspension with oxygenated buffer, after which the mixture is driven rapidly through the light path of a dual wavelength spectrophotometer or fluorometer. The reoxidation of an electron transport component is thus observed a short time after initiation of the reaction, and the proportion of component oxidized after this time is recorded using a storage oscilloscope. Oxidation of the anaerobic mitochondrial suspensions was achieved by addition of 16 μM-O₂ in the form of oxygenated buffer, and reduction by respiration of the mitochondria in the presence of 10 mM-succinate and 2 μM-carbonyl-cyanide m-chlorophenylhydrazone (CCCP) until state 5 was obtained (anaerobic). The path length of the observation window was 6 mm. in all cases and the temperature 25°. The upper oscilloscope trace (see Results)
indicates the syringe velocity, and the lower trace the spectrophotometric recording of the reaction. The time taken for the flow to travel from the mixing point of suspension and buffer to the observation window is referred to as $t_{4}$. Values for $k_{1}$ are calculated after noting how far the reaction has gone towards completion once a steady flow is obtained, i.e. after time $= t_{4}$. Then

$$k_{1} = \frac{2.3 \log D_{2}}{t_{4} D_{1}}$$

where

$$\frac{D_{2}}{D_{1}} \text{ is the ratio of Concentration of reduced cytochrome at time zero}{Concentration of reduced cytochrome after t_{4}}.$$  

Pyridine haemochrome derivatives were prepared according to the method of Falk (1964). Protein was measured by the method of Lowry, Rosebrough, Farr & Randall (1951), with bovine plasma albumin as a standard.

**Enzyme assays.** NADH-cytochrome c reductase (NADH$_2$:cytochrome c oxidoreductase EC 1.6.2.1) was assayed according to the method of Mahler (1955). Succinate-cytochrome c oxidoreductase was assayed in a similar way but using 0.1 M-sodium succinate in the reaction mixture. When required, electron transport inhibitors (rotenone and antimycin A) were added as methanolic solutions.

**Electron microscopy.** Whole organisms were fixed in 2% (w/v) unbuffered potassium permanganate for 2 h. at 4°C (Luft, 1956), dehydrated via an alcohol series and embedded in Araldite. For mitochondrial preparations, the fixative contained 0.35 M-sucrose. Ultra-thin sections were cut with an LKB Ultramicrotome, and floated on to copper grids. Negative staining was done with 1% (w/v) potassium phosphotungstate, pH 6.8. The specimens were examined and photographed in an A.E.I. EM6 operating at 60 kV.

**Materials.** ADP, ATP, NAD, NADH, cytochrome c (horse heart type III), antimycin A, 2-oxoglutarate, malate, oligomycin and EGTA were obtained from Sigma Chemical Co., St Louis, Missouri, U.S.A. Bovine Albumin Powder (fraction V from bovine plasma) was obtained from Armour Pharmaceutical Co. Ltd, Eastbourne. Rotenone was a gift from Professor L. Crombie, Department of Chemistry, University College, Cardiff, and piericidin A was from Dr Karl Folkers, Stanford Research Institute, Menlo Park, California, U.S.A.

**RESULTS**

**Electron Microscopy.** Isolated mitochondria (Pl. I, fig. 1) suffered little apparent structural damage when compared with the mitochondria within the intact organism (Pl. I, fig. 2). In sections the inner and outer membranes are clearly distinguishable. The extent of the tubular cristae is readily seen in negatively stained preparations of isolated mitochondria (Pl. I, fig. 3) as the phosphotungstate penetrates the intermembrane space and fills the cristae tubules, making them electron-dense.

**Oxidation of substrate by isolated mitochondria.** Table 1 shows typical oxidation rates obtained. Succinate, 2-oxoglutarate and DL-lactate were the only substrates tested which gave high oxygen uptake rates. Oxidation rates of isocitrate and of glutarate were increased on adding low concentrations of malate. NADH oxidation was undetectable in both intact and sonicated mitochondria. Respiratory control could be demonstrated for 2-oxoglutarate and succinate but not for other substrates.
Fig. 1. Respiratory control in isolated mitochondria. The oxygen electrode traces are examples of those obtained with 10 mM-succinate and 10 mM-2-oxoglutarate as substrates (each ADP addition was 80 nmoles). Temperature of incubation 25°. The range of ADP/O values and respiratory control (RC) ratios obtained in four experiments were as follows:

<table>
<thead>
<tr>
<th>Substrate</th>
<th>ADP/O</th>
<th>R.C.</th>
</tr>
</thead>
<tbody>
<tr>
<td>2-Oxoglutarate</td>
<td>1.3 to 2.3</td>
<td>1.2 to 2.2</td>
</tr>
<tr>
<td>Succinate</td>
<td>0.8 to 1.5</td>
<td>1.1 to 1.6</td>
</tr>
</tbody>
</table>

Table 1. Oxidation of substrates by mitochondria isolated from *Tetrahymena pyriformis*

Oxygen uptake was followed polarographically at 25° with a mitochondrial suspension containing 0.5 mg./ml. protein. Rates of O₂ uptake shown are in the presence of substrate and 80 nmoles of ADP (state 3) after subtraction of endogenous respiration rate (no added substrate).

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Concentration (mm)</th>
<th>Specific activity (nmoles/min./mg. protein)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tris-2-oxoglutarate</td>
<td>9</td>
<td>43.0</td>
</tr>
<tr>
<td>Sodium succinate</td>
<td>9</td>
<td>93.0</td>
</tr>
<tr>
<td>Sodium DL-lactate</td>
<td>9</td>
<td>19.0</td>
</tr>
<tr>
<td>Sodium citrate</td>
<td>9</td>
<td>8.0</td>
</tr>
<tr>
<td>Sodium isocitrate</td>
<td>9</td>
<td>4.5</td>
</tr>
<tr>
<td>+ sodium malate</td>
<td>0.9</td>
<td>16.5</td>
</tr>
<tr>
<td>Sodium aspartate</td>
<td>9</td>
<td>1.8</td>
</tr>
<tr>
<td>+ sodium malate</td>
<td>0.9</td>
<td>8.3</td>
</tr>
<tr>
<td>Sodium malate</td>
<td>0.9</td>
<td>9.0</td>
</tr>
<tr>
<td>+ sodium malate</td>
<td>9</td>
<td>12.6</td>
</tr>
<tr>
<td>sarcosine</td>
<td>0.9</td>
<td>9.0</td>
</tr>
<tr>
<td>Sodium α-glycerophosphate</td>
<td>9</td>
<td>9.0</td>
</tr>
<tr>
<td>Ethanol</td>
<td>5</td>
<td>6.5</td>
</tr>
<tr>
<td>Sodium glutamate</td>
<td>2</td>
<td>1.8</td>
</tr>
<tr>
<td>+ sodium malate</td>
<td>0.9</td>
<td>18.1</td>
</tr>
<tr>
<td>NADH</td>
<td>0.1</td>
<td>2.0</td>
</tr>
</tbody>
</table>
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tested. Typical traces are shown in Fig. 1 together with a range of ADP/O and respiratory control values calculated from four experiments. Phosphorylation cycles could be repeated until the oxygen in the reaction vessel was exhausted.

**Inhibition of oxidation by antimycin A, rotenone, piericidin A and cyanide.** Inhibition curves for antimycin A, rotenone and piericidin A are shown in Fig. 2. Tetrahymena mitochondria were a 1000-fold less sensitive to antimycin A and rotenone inhibition than rat liver mitochondria (Slater, 1967) but only threefold less sensitive to piericidin A (Hall et al. 1966). Cyanide (1 mM) gave 99% inhibition of state 3 respiration with both 2-oxoglutarate and succinate.

![Fig. 2. Inhibition of electron transport by (a) antimycin A, (b) rotenone and (c) piericidin A in isolated mitochondria. Inhibitors were added as methanolic solution to suspensions of mitochondria respiring in the presence of 9 mm-substrate and 0.1 mm-ADP at 25°. For inhibitors acting at or near site I (rotenone and piericidin A) the substrate was 2-oxoglutarate, and for site II inhibitor (antimycin A), succinate. Methanol controls showed no inhibitory effect with either substrate.](image)

**Table 2. Reduction of mammalian cytochrome c and ferricyanide by NADH and by succinate in the presence of mitochondria isolated from Tetrahymena pyriformis**

The assay mixture (1 ml.) contained 0.3 mg. cytochrome c or 3 mM-ferricyanide, 10 mM-KCN and mitochondria (0.1 to 0.3 mg. protein) in preparation buffer. The reactions were started by adding 10 mM-succinate or 0.1 mM-NADH, blanks contained no substrates, cytochrome c reduction was followed at 550 nm., ferricyanide reduction at 400 nm. Inhibitors were added as methanolic solutions. Figures in parenthesis indicate extent of inhibition by an equal volume of methanol. Results typical of those obtained with four different mitochondrial suspensions. Specific activities expressed as cytochrome c or ferricyanide reduced. Temperature of incubations 18°.

<table>
<thead>
<tr>
<th>Enzyme system</th>
<th>Specific activity (nmoles/min./mg. protein)</th>
<th>Inhibition (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Succinate-cytochrome c oxidoreductase + antimycin A</td>
<td>12.8 (1460 nmoles/mg. protein)</td>
<td>94 (18)</td>
</tr>
<tr>
<td>Succinate-ferricyanide oxidoreductase + antimycin A</td>
<td>240.0 (1460 nmoles/mg. protein)</td>
<td>84 (15)</td>
</tr>
<tr>
<td>NADH-cytochrome c oxidoreductase + rotenone</td>
<td>27.8 (500 nmoles/mg. protein)</td>
<td>38 (7)</td>
</tr>
<tr>
<td>NADH-ferricyanide oxidoreductase + rotenone</td>
<td>520.0 (500 nmoles/mg. protein)</td>
<td>19 (5)</td>
</tr>
</tbody>
</table>
Reduction of artificial electron acceptors. Externally added mammalian cytochrome c and ferricyanide were both reduced by isolated mitochondria in the presence of either NADH or succinate. Table 2 shows activities measured as saturating substrate concentrations, together with the inhibitory effects of rotenone and antimycin A. NADH-cytochrome c and NADH-ferricyanide oxidoreductases were only partially inhibited by rotenone in 'intact' mitochondria at a concentration which completely inhibited 2-oxoglutarate oxidation, suggesting that these oxidoreductions did not involve the respiratory chain which is presumably located in the inner mitochondrial

![Difference spectra (reduced minus oxidized) of Tetrahymena mitochondria. The suspension of mitochondria contained 3.0 mg. protein/ml. (path length 10 mm.). Curve (1) represents the baseline (oxidized minus oxidized). Curve (2) was obtained after addition of 10 mM-succinate to one of the cuvettes and exhaustion of dissolved O2 (state 5). Curve (3) was obtained following addition of 360 nmoles of antimycin A and reaeration of both cuvettes.](image)

Table 3. NADH-cytochrome c oxidoreductase activity in sonicated mitochondria

<table>
<thead>
<tr>
<th></th>
<th>Specific activities (nmoles/min./mg. protein)</th>
<th>Rate after 1 min. of reaction</th>
<th>Rate with rotenone added (150 nmoles/mg. protein)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Initial rate</td>
<td>Rate after 1 min. of reaction</td>
<td>Rate with rotenone added (150 nmoles/mg. protein)</td>
</tr>
<tr>
<td>Unsonicated mitochondria</td>
<td>45.8</td>
<td>37.7</td>
<td>25.8</td>
</tr>
<tr>
<td>Sonicated mitochondria</td>
<td>93.3</td>
<td>44.1</td>
<td>30.0</td>
</tr>
</tbody>
</table>
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membrane. In sonicated mitochondria, a higher NADH-cytochrome c reductase activity was observed, but this activity appeared to be unstable and declined after several minutes. Addition of excess rotenone to the assay system revealed similar rotenone-insensitive activity in both sonicated and unsonicated preparations (Table 3). The results suggest that rotenone-sensitive activity is associated with the respiratory chain, to which externally added NADH is inaccessible in intact mitochondria, while in sonicated mitochondria this inner membrane activity is rapidly lost. Both succinate-cytochrome c and ferricyanide reductase activities were inhibited by addition of antimycin A.

![Fig. 4. Low temperature difference spectrum of Tetrahymena mitochondria. The mitochondrial suspension contained 6.2 mg protein/ml. Reduction was achieved by addition of excess sodium dithionite and oxidation by aeration. The samples were introduced into pre-cooled cuvettes (path length 2 mm.) and immediately cooled by immersion in liquid N\textsubscript{2}.](image)

Extinction spectra of mitochondria. Extinction maxima in difference spectra at 560, 527 and 432 nm. (Fig. 3) represented the α, β and γ bands of cytochrome b, while those at 553 and 523 nm. indicated the presence of a c-type cytochrome. The classical extinction maximum for cytochromes (α + α\textsubscript{b}) at 605 nm. was not observed although a γ(α\textsubscript{b} + α) extinction was evident at 447 nm. The flavoprotein-non-haem iron trough was often observed at 465 to 470 nm. but not identified. Addition of antimycin A (sufficient for 50 % inhibition of state 3 respiration) followed by reaeration of the reduced suspension resulted in reoxidation of the 620 nm. band and of cytochrome c.
while cytochrome b remained reduced, indicating the presence of an antimycin A-sensitive site between cytochromes b and c. Part of the flavoprotein was reoxidized despite the reduction of cytochrome b, suggesting the presence of a flavoprotein oxidase(s).

Fig. 4 shows a low temperature difference spectrum (dithionite reduced minus oxidized) of the a-region. There was a peak shift of 2 to 3 nm. towards the blue end of the spectrum for all components other than flavoprotein, which appears at 475 nm. A minor component was present at 585 nm., probably corresponding to the 595 nm component often observed in room temperature spectra. The individual components

![Graph showing temperature difference spectrum](image)

Fig. 5. Carbon monoxide spectrum of mitochondria isolated from Tetrahymena. Each cuvette (path length 10 mm.) contained mitochondrial suspension (2.5 mg./ml.). The suspensions were reduced by addition of 10 mM-succinate to each cuvette, allowing time for exhaustion of dissolved oxygen. After recording a baseline (succinate reduced minus succinate reduced) (1) CO was passed through the test suspension, and the CO/reduced minus reduced spectrum was recorded (2).

were more sharply resolved than at room temperature, and a shoulder at 548 nm. indicated the presence of cytochrome cₙₙ.

**CO-spectrum.** In order to characterize further the cytochrome oxidase, the CO-reduced minus reduced difference spectrum of whole mitochondria was determined (Fig. 5). The preparation showed the spectrum, characteristic of cytochrome oxidase (Chance, 1957), although cytochrome b purified from Tetrahymena also reacts with CO (Yamanaka, Nagata & Okunuki, 1967).

**Pyridine haemochrome derivatives.** Pyridine haemochrome derivatives were prepared from whole mitochondria in order to characterize the haems present. Fig. 6 shows the spectra of the derivatives (dithionite reduced minus ferricyanide oxidized). Peaks at 587 nm., 557 nm., 527 nm., and 551, 523 nm. correspond to the derivatives of
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Fig. 6. Difference spectra for pyridine haemochrome derivatives. The extract containing haem *a* and protohaem (Falk, 1964) was resuspended in distilled water, as was the pellet obtained after extraction (containing haem *c*). The difference spectrum of each fraction was obtained after addition of pyridine (1 ml.) and N-NaOH (0.5 ml.) to the preparations (4.5 ml.). To one cuvette 1 mg. of ferricyanide was added, and to the other 1 mg. of dithionite. (1) Haem *c*, (2) haem *a* + protohaem.

Table 4. *Steady-state levels of reduction of electron transport components in mitochondria isolated from Tetrahymena pyriformis*

<table>
<thead>
<tr>
<th>Substrate</th>
<th>10 mM-Succinate</th>
<th>10 mM-2-Oxoglutarate</th>
<th>10 mM-Pyruvate</th>
</tr>
</thead>
<tbody>
<tr>
<td>Electron transport</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>component</td>
<td>Cyt <em>b</em></td>
<td>Cyt <em>c</em></td>
<td></td>
</tr>
<tr>
<td>Wavelength pair</td>
<td>560 - 553</td>
<td>340 -</td>
<td>340 -</td>
</tr>
<tr>
<td>(nm.)</td>
<td>575 - 540</td>
<td>374 - 575 - 575 - 540</td>
<td>374 - 575 -</td>
</tr>
<tr>
<td>Reduction state 4 (%)</td>
<td>35.6 -</td>
<td>84.5 - 29.7 -</td>
<td>85 - 25 -</td>
</tr>
<tr>
<td>Reduction state 3 (%)</td>
<td>22.4 -</td>
<td>72.7 - 21.8 -</td>
<td>76 - 10 -</td>
</tr>
<tr>
<td>Not reduced in state 5 but reduced by excess dithionite %</td>
<td>35.0 -</td>
<td>55.0 - 10 -</td>
<td>50 - 10 -</td>
</tr>
</tbody>
</table>

One hundred per cent reduction of a component taken as anaerobic reduction level minus level in the absence of exogenous substrate (state 5 - state 2). ADP concentration was 0.1 mm. Absence of a figure indicates that if any reduction occurred it was below the level of detection. No reduction of the 620 nm. absorbing species (620 - 650 nm.) was detectable in states 3 or 4 with any of the substrates tested, and no extra reduction of this pigment was seen on adding dithionite to anaerobic suspensions. Temperature of incubation 25°.
haem a, protohaem and haem c, respectively. These figures coincide with those observed for mammalian mitochondria (Falk, 1964). Thus although no cytochrome a was seen in the 605 nm. region of difference spectra (Fig. 3) it seems that haem a was present in Tetrahymena mitochondria.

Steady-state reduction of electron transport components. Reduction levels of electron transport components in suspensions of isolated mitochondria (Table 4) were measured after addition of 3 mM substrate with ADP exhausted (state 4), after addition of substrate + ADP (state 3), and after exhaustion of the dissolved oxygen (state 5) (Chance & Williams, 1956). To detect the proportion of component not reducible by substrate, sodium dithionite was added to suspensions in state 5. Throughout these experiments the wavelength pairs used were as follows: 620 nm. absorbing species, 620 to 650 nm.; cytochrome b, 430 – 410 nm.; cytochrome c, 553 – 540 nm.; NADH, 340 – 374 nm. For cytochrome b the γ-band was chosen for greater sensitivity,
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measurements at 560 nm. were often interfered with by the $\alpha$-band of cytochrome $c$ (553 nm.). State 5 reduction for cytochrome $b$ by 2-oxoglutarate or by pyruvate was only 60% of that observed with succinate (Fig. 7). Cytochromes $c$ and $a$ were equally reduced with all three substrates. Endogenous respiration (washed mitochondria with no added substrate) gave a state 5 level of reduction similar to that observed for substrates other than succinate. Cytochrome $b$ was not completely reduced by substrate oxidation, as can be seen from the addition of dithionite. Cytochrome $c$ was 90% reduced and cytochrome $a$ fully reduced under these conditions. No reduction of cytochrome $c$ or the 620 nm. pigment was observed in states 3 and 4. Comparable results were obtained by observing difference spectra (succinate reduced minus oxidized) at successive time intervals after addition of substrate to one cuvette (Fig. 8).

Fig. 8. Steady-state spectra of mitochondria isolated from Tetrahymena. The mitochondrial suspension contained 4.1 mg. protein/ml. (path length 10 mm.). Curve (1), oxidized minus oxidized (baseline); curve (2), succinate reduced (state 4) minus oxidized (2 min. after addition of 10 mM-succinate); curve (3), 5 min. after addition of succinate, anaerobic (state 5) minus oxidized (state 2). Temperature of incubation 18°.

With substrates other than 2-oxoglutarate and pyruvate (including succinate, DL-lactate, citrate, malate) no steady-state reduction of NAD was observed, indicating that these substrates were either unable to gain access to the mitochondrial matrix, or were not NAD-linked.

With tightly coupled mitochondria respiring in the presence of succinate, 2-oxoglutarate or pyruvate, changes in steady-state levels of reduction of cytochrome $b$ and NAD were induced by addition of 0.1 mM-ADP, both components becoming more oxidized (Fig. 7; Table 4).

Addition of antimycin A to mitochondrial suspensions oxidizing succinate (sufficient for 50% inhibition of state 3 oxidation) resulted in an increase in the reduction level of cytochrome $b$ in state 3 from 22 to 52% of the state 5 level. Since reduction was
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Fig. 9. Oscilloscope traces of reoxidation reactions of cytochromes b, e and oxidase. (a, b) $t_{0.5} = 14.2$ msec., mitochondrial protein, 11.2 mg./ml. (c, d) $t_{0.5} = 2.3$ msec., mitochondrial protein, 5.3 mg./ml.

Fig. 10. Oscilloscope traces for the reoxidation reactions of flavoprotein, measured spectrophotometrically and fluorometrically. In traces (a) and (b) the reactions were so slow that no oxidation was detectable during the flow, and half-times are given for these 'stopped-flow' reactions. In traces (c) and (d) the reactions are more complex, and it appears that fast-reacting component(s) were present whose oxidation was completed during the flow. Temperature of incubation 25°. $t_{0.5} = 15.9$ msec., mitochondrial protein 5.1 mg./ml.
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not 100%, a leak of electrons past the partially inhibited site was indicated. Both cytochrome c and 620 nm. absorbing species remained fully oxidized.

Kinetics of cytochrome and flavoprotein reoxidation. All the following experiments were carried out on suspensions of mitochondria containing 10 mM-succinate as substrate, and 2 μM-CCCP as uncoupling agent, with the regenerative flow apparatus of Chance (1954). When the respiring mitochondria had used up all the oxygen in the suspension, the state 5 → state 3 (uncoupled) transition was measured following rapid mixing of the anaerobic suspension with oxygen-saturated buffer. After mixing, the final concentration of oxygen in the mitochondrial suspension was 16 μM. Lowering of the added oxygen concentration to 3 μM did not noticeably alter the reaction velocities of the components. The oscilloscope traces obtained are shown in Fig. 9 and 10, together with the calculated k values for pseudo first-order reactions. The data shown in Fig. 9 indicate that the velocity of oxidation of the 620 nm. component was greater than the oxidation velocity of cytochrome c, which was greater than the oxidation velocity of cytochrome b.

Flavoprotein oxidation (Fig. 10) was measured spectrophotometrically (465 to 510 nm.) and fluorometrically (excitation at 436 nm., 460 nm. and 475 nm., emission at 520 nm.). Spectrophotometric measurements suggested a slowly reoxidizing component (t1 = 400 msec.). Similarly, fluorometric measurements by excitation at 475 nm. detected a slow reacting component only (t1 = 1200 msec.). However, excitation at 460 nm. and 436 nm. indicated biphasic reactions, consisting of a very fast reacting component (t1 = 100 msec.) and a more slowly reacting component(s) (t1 = 900 msec., 436 nm.; t1 = 600 msec., 460 nm.). Since the oxidation of the fast component was completed during the flow (compare with oxidation of cytochrome b in Fig. 9) it is apparent that this component(s) was able to react with oxygen faster than cytochrome b, and thus its reoxidation could not proceed via cytochrome b. Fluorometric measurements of flavoprotein oxidation are more reliable than spectrophotometric measurements, being less susceptible to interference from non-haem iron.

DISCUSSION

Oxidation of 2-oxoglutarate, together with demonstration of control with this substrate, and the presence of sites I and II as suggested by electron transport inhibitors, suggests that the NAD-linked electron transport pathway to oxygen was intact in isolated mitochondria. The slow oxidation of citrate, isocitrate and malate suggests a permeability barrier for these substrates. Although endogenous NAD underwent reduction on addition of 2-oxoglutarate and pyruvate to mitochondrial suspensions, externally added NADH was not oxidized. Rat liver mitochondria do not oxidize exogenous NADH (Lehninger, 1955), but mitochondria isolated from several eukaryotic micro-organisms, including Prototheca zopfii (Lloyd, 1965), Polyomella caeca (Evans & Lloyd, 1966), Hartmanella castellanii (Lloyd & Griffiths, 1968), Saccharomyces carlsbergensis (Ohnishi, Kawaguchi & Hagihara, 1966) and Aspergillus niger (Watson & Smith, 1967), are able to, making Tetrahymena unusual in this respect.

The pathways of electrons from succinate or NADH to added mammalian cytochrome c may be similar to those in mammalian mitochondria. Thus although mammalian cytochrome c is not reoxidized by the oxidase of Tetrahymena (Kobayashi,
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1969, it is able to act as an artificial electron acceptor in a similar manner to ferri-
cyanide, possibly reacting with the electron transport chain in the region of Tetra-
hymena cytochrome c. Much of the NADH-cytochrome c reductase activity is rote-
none-insensitive and may be associated with an outer membrane pathway as is the case
with mammalian mitochondria (Parsons, 1966). Up to 40 % of the NADH-cytochrome
c reductase activity is rotenone-sensitive, suggesting an inner membrane pathway via
site I. The partial rotenone sensitivity of NADH-cytochrome c reductase is difficult
to reconcile with the inability of intact mitochondria to oxidize NADH.

The most unusual feature of the cytochrome components as observed in difference
spectra was the absence of a cytochrome oxidase band at 605 nm. Tetrahymena mito-
chondria are the only ones of those yet studied which exhibit this feature. Eichel
(1954), Kobayshi (1965), van de Vijver (1966) observed that homogenates or cell-free
preparations of *Tetrahymena pyriformis* were unable to reoxidize reduced mammalian
cytochrome c, and concluded that cytochrome oxidase was absent from Tetrahymena
mitochondria. It appears that all organisms which possess cytochrome \((a+a_3)\)
\(\alpha\)-band 605 nm.) are able to reoxidize reduced mammalian cytochrome c \(\lambda\) (Lloyd,
1965; Ohnishi *et al.* 1966; Evans & Lloyd, 1966; Watson & Smith, 1967; Lloyd &
Griffiths, 1968). Pyridine haemochrome characterization of the component haems
indicated the presence of haem \(a\). It seems possible that a cytochrome oxidase was
present which, though atypical, contained haem \(a\). In addition, evidence provided by
the kinetic studies suggested that the 620 nm. component reacted with oxygen at a
velocity greater than that of cytochromes \(b\) and \(c\).

Both inhibitor studies and kinetic data suggested the presence of a flavoprotein
oxidase, since flavoprotein was reoxidized even in the presence of antimycin A, and at
a greater velocity than cytochrome \(b\).

The observation that the state 5 reduction level of cytochrome \(b\) observed with other
substrates was only 60 % that with succinate suggested the presence of two pools of
cytochrome \(b\), one reducible by succinate alone:

\[
\text{Succinate} \rightarrow \text{cyt } b \\
2\text{-Oxoglutarate} \rightarrow \text{NAD} \rightarrow \text{cyt } b \rightarrow \text{cyt } c \rightarrow \text{cyt oxidase} \rightarrow \text{O}_2 \\
\text{Pyruvate}
\]

Reduction of cytochrome \(c\) and oxidase could not be detected in states 3 and 4.
Chance & Williams (1956) observed a stepwise sequence in the reduction levels of
electron transport components during steady-state respiration, from NAD, the most
reduced component, to cytochrome oxidase, the most oxidized. In state 4 coupled
mitochondria exhibit some degree of inhibition of respiration at coupling sites, and
on addition of ADP, cytochrome \(a\) becomes more reduced and all other components
more oxidized as the inhibitory effect is relieved. This 'crossover point' is observed
between cytochromes \(c\) and \(a\) in rat liver mitochondria. The crossover point in
Tetrahymena mitochondria was on the oxygen side of cytochrome \(b\), indicating at
least one coupling site between cytochrome \(b\) and oxygen. Since cytochromes \(c\) and \(a\)
were maximally oxidized in either steady state, no change was observed on addition
or exhaustion of ADP.
Mitochondria from Tetrahymena

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EXPLANATION OF PLATE

Fig. 1. Thin section of mitochondria isolated from Tetrahymena pyriformis, showing outer membrane (om), inner membrane (im), and cristae (cr). Fixed with KMnO₄. Unstained. × 50,000.

Fig. 2. Thin section of T. pyriformis; organism from a late exponential phase culture. Nucleus (n), mitochondria (m), endoplasmic reticulum (er) and pellicle (p) are visible. Fixed with KMnO₄. Unstained. × 26,500.

Fig. 3. Mitochondria isolated from T. pyriformis. Unfixed and stained with K phosphotungstate. The stain has entered the intermembrane space and filled the tubes of the cristae. It has not entered the matrix (m). × 26,500.