Energy Coupling During Sulphur Compound Oxidation by Thiobacillus sp. Strain C

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SUMMARY

The addition of either sulphide or thiosulphate to an aerobic suspension of a Thiobacillus was followed within 10 sec. by the synthesis of intracellular ATP. The ATP formation accompanying sulphide oxidation was strongly inhibited by 0.1 mM 2,4-dinitrophenol, but that accompanying thiosulphate oxidation was unaffected. The presence of carbon dioxide had no effect on ATP formation or on the steady-state concentration attained. 2,4-Dinitrophenol did not significantly affect oxygen uptake nor the kinetics of thiosulphate oxidation but it inhibited carbon dioxide fixation with either sulphide or thiosulphate as substrate. It is concluded that sulphur-compound oxidation can be coupled to two different types of phosphorylation, only one of which is sensitive to 2,4-dinitrophenol. In the presence of 2,4-dinitrophenol and with thiosulphate as substrate, carbon dioxide fixation was not limited by the availability of ATP but might be limited by the availability of reduced nicotinamide adenine dinucleotides.

INTRODUCTION

Organisms of the genus Thiobacillus obtain energy from the oxidation of reduced inorganic sulphur compounds. The coupling of this energy to synthetic reactions has long been thought to be mediated by high energy phosphate compounds (Vogler & Umbreit, 1942); recent work supports the view that ADP, ATP and reduced coenzymes are involved (Trudinger, 1956; Vishniac & Santer, 1957; Milhaud, Aubert & Millet, 1957; Johnson & Peck, 1965). Peck (1960, 1962) used cell-free extracts of Thiobacillus thioparus, and suggested that thiosulphate was first split reductively to sulphide and sulphite, the sulphite being oxidized subsequently to sulphate by reactions which produce ADP by a substrate-level phosphorylation, with adenosine 5'-phosphosulphate as an intermediate. This phosphorylation is insensitive to 2,4-dinitrophenol (DNP; Peck & Fisher, 1962).

By using 14CO2-fixation as a measure of energy transfer in the presence or absence of DNP, we obtained results which suggested that both DNP-sensitive and DNP-insensitive phosphorylations accompany thiosulphate oxidation, but that DNP-sensitive phosphorylations are more important during sulphide oxidation (Kelly & Syrett, 1964a). We now present further evidence, from measurements of the ATP content of intact bacteria, for the occurrence of both DNP-sensitive and DNP-insensitive phosphorylation during sulphur-compound oxidation, and for the greater
importance of the former in sulphide oxidation. Our results also support the view that the reduced coenzymes necessary for CO$_2$-fixation are generated by an energy-dependent reduction by electrons from reduced cytochromes, similar to that shown in other autotrophs (Aleem, Lees & Nicholas, 1968) and recently reported for *T. novellus* by Aleem (1965).

A brief report of some of the present work has appeared elsewhere (Kelly & Syrett, 1964b) and the background to the problem was discussed more fully by Kelly (1965).

**METHODS**

*The organism.* *Thiobacillus* sp. strain c was grown as described previously in media containing 1-2 % (w/v) Na$_2$S$_2$O$_3$.5H$_2$O (Kelly & Syrett, 1964a). Organisms were harvested by centrifugation, washed and suspended in 0.1 M-sodium phosphate buffer (pH 7.0) to give dense suspensions usually containing about equiv. 3 mg. dry wt. organisms/ml.

*Extraction and assay of ATP.* Samples (2-3 ml. of dense suspensions were extracted with 0.2 ml. 14.5 % (w/v) perchloric acid on ice for 10 min., then brought to pH 7.4 by the addition of 1.0 ml. of KOH of suitable concentration. The clear supernatant solution was pipetted from the perchlorate precipitate. ATP was measured by a firefly luminescence method (Strehler & Totter, 1952). The following were mixed in a 1.5 ml. cuvette at 20°: 0.5 ml. 0.1M-sodium arsenate+ sulphuric acid buffer (pH 7.4); 0.8 ml. distilled water; 0.2 ml. perchlorate extract; 0.04 ml. 20 % (w/v) MgSO$_4$.7H$_2$O. The reaction was initiated by rapidly adding 0.1 ml. of an extract of firefly tails in 0.1 M-arsenate+H$_2$SO$_4$ buffer (pH 7.4) containing 2 % (w/v) MgSO$_4$.7H$_2$O. The light emission peak was measured 20 sec. later in a ‘Locarte’ fluorimeter. The enzyme preparation and samples were kept on ice and allowed to warm only when assayed. Perchlorate depressed the luminescence, apparently by interfering with the enzyme reaction rather than by destroying ATP. In the presence of a constant amount of perchlorate, light emission was proportional to the amount of ATP in solution. ATP standards and samples were always assayed in the presence of equivalent amounts of perchlorate. Comparison of the ATP content estimated by this method with the content determined in similar samples extracted by boiling the organisms for 30 sec. in 90 % (v/v) ethanol in water, showed no significant difference. Bacterial extracts contained no substances other than perchlorate which interfered with the assay. DNP caused no interference. Data supporting these statements have been given elsewhere (Kelly, 1965).

*Simultaneous determination of ATP content and 14CO$_2$ fixation.* When the ATP content only was required, suspensions of bacteria were aerated in a rapid sampling device (Syrett, Bocks & Merrett, 1964), or were shaken in 50 ml. conical flasks. When the ATP content and 14CO$_2$-fixation values were required simultaneously, bacterial suspensions (2-2.6 ml.) were shaken in 50 ml. conical flasks sealed with vaccine stoppers. Substrates (KH$^{14}$CO$_3$; Na$_2$S$_2$O$_3$; Na$_2$S) were injected into the flasks and the bacteria killed by injecting perchloric acid. Samples of the neutralized extracts were mixed with equal volumes of 5 % (v/v) acetic acid in ethanol, and were used to estimate 14CO$_2$-fixation as described by Kelly & Syrett (1964a). The rest of the material was assayed for ATP. For time-course experiments it was better to set up a series of such flasks and to kill the bacteria after various time intervals.
rather than to take a series of samples from a single flask with syringes. The latter method gave erratic values for ATP.

Gas exchange was measured by Warburg manometry.

Oxidation of radioactive thiosulphate was followed by the sampling and chromatographic methods described by Kelly & Syrett (1966).

Chemicals. Na\textsubscript{2}\textsuperscript{14}CO\textsubscript{3} and Na\textsubscript{2}\textsuperscript{35S}SO\textsubscript{3} were obtained from the Radiochemical Centre, Amersham, Bucks; ATP (disodium salt; 3–4H\textsubscript{2}O; 99–100\% ) and desiccated firefly lanterns from the Sigma Chemical Company; hexokinase from L. Light and Co., Colnbrook, Bucks. Other reagents were ‘AnalaR’ or of the best quality marketed. Solutions of Na\textsubscript{2}S\textsubscript{2}O\textsubscript{3} and Na\textsubscript{2}S were always prepared immediately before use; washed crystals of Na\textsubscript{2}S were used.

RESULTS

ATP formation during sulphide and thiosulphate oxidation, and the effect of 2,4-dinitrophenol

The bacterial ATP content increased at least twofold within 10–20 sec. after the addition of sulphide or thiosulphate (Fig. 1; and Kelly & Syrett, 1964b). Low concentrations of DNP had little effect on the ATP increase during thiosulphate oxidation, while the same DNP concentrations severely depressed ATP formation after sulphide addition. Figure 1 demonstrates that this lower sensitivity of thiosulphate-dependent ATP formation to DNP inhibition was exhibited over a wide range of DNP concentrations. For example, 0.1 mM-DNP completely inhibited increase in ATP content when sulphide was oxidized, while the ATP formation with thiosulphate was unaffected by this concentration. Table 1 shows that treatment of the bacterial extracts with glucose and hexokinase virtually abolished the light emission, indicating the material determined by the firefly method was exclusively ATP.

Table 1. Effect of treating thiobacillus extracts with hexokinase and glucose before assaying for ATP

Samples taken at intervals after addition of thiosulphate to dense bacterial suspensions were extracted with perchloric acid (see text). Two series of cuvettes were prepared, each cuvette containing in 0.9 ml. 50 \( \mu \)moles arsenate + H\textsubscript{2}SO\textsubscript{4} buffer (pH 7.4); 10 \( \mu \)moles MgCl\textsubscript{2}; 0.2 ml. bacterial extract; 20 \( \mu \)moles glucose. Series 1 received no further addition; series 2 received 0.5 mg. hexokinase. Both series were incubated at 28\textdegree C for 16 min., then the ATP estimation was made as usual by adding 0.1 ml. firefly extract to each cuvette. Mean values from duplicates or triplicates are given.

<table>
<thead>
<tr>
<th>Sample time after thiosulphate addition (min.)</th>
<th>ATP content (( \mu )moles/g. dry wt. bacteria)</th>
<th>Hexokinase</th>
<th>+ Hexokinase</th>
</tr>
</thead>
<tbody>
<tr>
<td>Initial</td>
<td>1.57</td>
<td>0.18</td>
<td></td>
</tr>
<tr>
<td>0.38</td>
<td>4.17</td>
<td>0.23</td>
<td></td>
</tr>
<tr>
<td>1.5</td>
<td>4.00</td>
<td>0.23</td>
<td></td>
</tr>
<tr>
<td>3.0</td>
<td>8.42</td>
<td>0.17</td>
<td></td>
</tr>
<tr>
<td>6.0</td>
<td>4.68</td>
<td>0.18</td>
<td></td>
</tr>
</tbody>
</table>
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ATP formation in the presence or absence of carbon dioxide

Initial experiments to measure ATP were made in the absence of CO₂. However, Fig. 2 shows that even the presence of CO₂ in high concentrations had little effect on the kinetics of the thiosulphate-dependent ATP increase. Since the presence of CO₂ did not abolish the increase in ATP content it was possible to measure simultaneously CO₂-fixation and ATP content of single samples taken over a period of time.

Simultaneous determination of ¹⁴CO₂ fixation and ATP content of bacteria oxidizing thiosulphate

The rate of ¹⁴CO₂-fixation during thiosulphate oxidation was not constant from zero time, but always increased to a steady rate 2–3 min. after substrate addition (Fig. 8). Since ATP formation was virtually instantaneous (Fig. 3) the phenomenon could not be attributed to a delay in thiosulphate metabolism. The addition of a second quantity of thiosulphate after the oxidation of the first resulted in renewed

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![Fig. 1](image1.png)

**Fig. 1.** Effect of various concentrations of 2,4-dinitrophenol on ATP content of thiosalbacillus strain c oxidizing thiosulphate or sulphide. 2·7 mg. dry wt. bacteria/ml. were shaken at 21° for 15 min. with the DNP concentration indicated, 8·8 μmoles Na₂S₂O₃/ml. (–x–x–) or 0·83 μmole Na₂S/ml. (–O–O–) were added. HClO₄ was added to the thiosulphate series after 6 min. and to the sulphide series after 1 min. Endogenous ATP content (+DNP) was about 0·70 μmole/g. dry wt.

![Fig. 2](image2.png)

**Fig. 2.** Effect of carbon dioxide on ATP concentrations during thiosulphate oxidation. Upper graph: 4·6 mg. bacteria in 2·4 ml. phosphate (pH 7·0) shaken in sealed 50 ml. flasks at 21°. One series (–O–O–) had 0·4 ml. 40% (w/v) KOH and a roll of filter paper in a centre well in each flask; the other series (–x–x–) had 4 μmoles NaHCO₃. Na₂S₂O₃ was injected at time 0. Lower graph: bacterial suspensions in a rapid-sampling device were aerated at 22° with CO₂-free air (–x–x–) or 1% (v/v) CO₂ in air (–O–O–) and received Na₂S₂O₃ at time 0.
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CO₂-fixation at an instantaneously high rate (Kelly, 1965). Possibly, therefore, the initially low rate represented the time required to build up a steady-state intracellular concentration of CO₂-acceptor molecules, such as ribulose diphosphate.

![Fig. 3. Simultaneous measurement of ¹⁴CO₂-fixation and ATP content during thiosulphate oxidation by thiobacillus strain c. Upper: ATP content; and lower: ¹⁴CO₂ fixed by dense bacterial suspensions shaken in sealed flasks into which KH¹⁴CO₃ and Na₂S₂O₃ were injected at time 0. Temp. 21°C; pH 7.0.](image)

**Fig. 4. Effect of 2,4-dinitrophenol on ATP content (upper graph) and ¹⁴CO₂ fixation (lower graph) by thiobacillus strain c oxidizing thiosulphate. Dense bacterial suspensions received KH¹⁴CO₃ and Na₂S₂O₃ at time 0, after pre-incubation with no DNP (− × − × −), 5 × 10⁻⁵ M-DNP (− O − O −) or 5 × 10⁻⁴ M-DNP (− ○ − ). Temperature 21°C; pH 7.0.**

Effect of 2,4-dinitrophenol on ¹⁴CO₂-fixation and ATP content with thiosulphate or sulphide as substrate

DNP depressed both the initial low rate of ¹⁴CO₂-fixation and the subsequent steady high rate during thiosulphate oxidation (Fig. 4). The effect of DNP on ¹⁴CO₂-fixation bore no obvious relationship to the ATP content of bacteria oxidizing thiosulphate, since the initial rapid increase in ATP and the subsequent values were
not significantly altered by DNP (Fig. 4). Figure 5a confirms that no simple relationship existed between ATP content and $^{14}$CO$_2$ fixed after oxidation of thiosulphate for a set time, with a range of DNP concentrations. $^{14}$CO$_2$-fixation was 90% inhibited by a DNP concentration (0.2–0.8 mM) which only slightly lowered the ATP content. Even a DNP concentration which completely inhibited CO$_2$-fixation (0.75 mM) only decreased the increase of ATP content to 50% of the control level. However, the depression of the ATP content by DNP in bacteria oxidizing sulphide was similar to the relative inhibition of CO$_2$-fixation during

![Graph](image)

Fig. 5. Effect of various concentrations of 2,4-dinitrophenol on the ATP content (–○–○–) and $^{14}$CO$_2$ fixed (–×–×–) by Thiobacillus strain c after oxidizing thiosulphate or sulphide: (a) Thiosulphate oxidized for 6 min. ATP content in excess of the endogenous value of 0.8 μmole/g. dry wt. is shown. (b) Sulphide oxidized: ATP data are those obtained from two experiments after exposure of the thiobacilli to Na$_2$S for 60 or 90 sec. $^{14}$CO$_2$ data are derived from 1 to 7 separate experiments, and are values for total fixation during complete sulphide oxidation. Both values are expressed as % of control values. Vertical lines indicate the extreme values around the mean for $^{14}$CO$_2$ fixation. ATP values were calculated after subtraction of the endogenous ATP content in the absence of sulphide.

sulphide oxidation: Fig. 5b shows the inhibition of $^{14}$CO$_2$-fixation in bacteria allowed to oxidize equal amounts of sulphide completely. These values were derived from several experiments, and are to be compared with the ATP content of bacteria which were allowed to oxidize sulphide for a set time in two experiments. Clearly the formation of ATP during sulphide oxidation was at least as sensitive to DNP as was the coupled CO$_2$-fixation.

In a large number of experiments, DNP never inhibited oxygen uptake during sulphide oxidation. Thiosulphate oxidation by dilute suspensions was also unaffected or stimulated by DNP (Kelly & Syrett, 1963, 1964a). By using thiosulphate labelled in the outer ('reduced') atom with $^{35}$S, 0.1 mM-DNP was found to have little effect on the kinetics of oxidation of the labelled atom by dense bacterial suspensions (in which oxidation was probably rate-limited by the concentration of dissolved oxygen). Figure 6 shows that labelled thiosulphate rapidly disappeared, with the parallel
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formation of sulphate and the transitory accumulation of labelled trithionate, which was unaffected by DNP. The subsequent oxidation of the trithionate to sulphate was, however, slightly retarded by DNP (see Kelly & Syrett, 1964a).

![Graph](image)

Fig. 6. Oxidation of Na₂(³⁴S₂SO₄) by thiobacillus strain c with and without 0.1 mM-DNP. 250 ml flasks shaken at 26°C contained in a final 12 ml equiv. 33 mg dry wt. bacteria; 1 mmole sodium phosphate (pH 7.0); with or without DNP, 47 µmoles Na₂(³⁴S₂SO₄) (28 µc) were added at zero time. Samples (1 ml.) were pipetted into 1 ml ethanol, centrifuged, and the supernatant fluids analysed chromatographically. With and without DNP, symbols are: S₂O₄²⁻ -O-O- and -●-●-; SO₄²⁻ -×-×- and ---+--+; S₄O₆²⁻ -△-△ and -▲-▲.

Table 2. Effect of 2,4-dinitrophenol on ¹⁴CO₂-fixation coupled to the oxidation of sulphur, sulphide and thiosulphate by thiobacillus strain c

<table>
<thead>
<tr>
<th>DNP concentration (m)</th>
<th>Initial rate of sulphur oxidation (µl/hr flask)</th>
<th>Oxygen consumed in sulphur oxidation (µmoles flask)</th>
<th>¹⁴CO₂ fixed/µmole or µg atom substrate oxidized*</th>
<th>¹⁴CO₂ fixed/µmole or µg atom substrate oxidized (Counts/100 sec.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>58</td>
<td>13-84</td>
<td>46,400 27,200 172,500 5,040 10,880 17,250 100 100 100</td>
<td>58</td>
</tr>
<tr>
<td>1 x 10⁻⁵</td>
<td>58</td>
<td>13-80</td>
<td>26,800 -- -- 2,920 -- -- 58 -- --</td>
<td></td>
</tr>
<tr>
<td>5 x 10⁻⁵</td>
<td>62</td>
<td>14-42</td>
<td>22,000 18,900 126,000 2,280 7,560 12,600 45 69 73</td>
<td></td>
</tr>
<tr>
<td>1 x 10⁻⁴</td>
<td>54</td>
<td>13-17</td>
<td>13,600 -- -- 1,540 -- -- 31 -- --</td>
<td></td>
</tr>
<tr>
<td>2 x 10⁻⁴</td>
<td>57</td>
<td>12-37</td>
<td>7,000 7,700 59,000 850 3,080 5,900 17 28 34</td>
<td></td>
</tr>
</tbody>
</table>

* Calculated for sulphur oxidation as 1-5 µmoles O₂ = 1 µg atom S°, from: S° + H₂O + ¹⁴O₂ = H₂SO₄.

Comparison of the effect of 2,4-dinitrophenol on ¹⁴CO₂-fixation with sulphide, thiosulphate or elementary sulphur as substrate

DNP had virtually no effect on the oxidation of elementary sulphur, but inhibited CO₂-fixation more than when sulphide or thiosulphate was oxidized (Table 2). The significance of this result in understanding the energy-coupling process is uncertain.
The relative amounts of \( \text{CO}_2 \) fixed during the oxidation of 1 \( \mu \)mole thiosulphate or sulphide and 1 \( \mu \)atom sulphur (calculated from oxygen consumed) were 100, 63 and 29, respectively, in this experiment.

**DISCUSSION**

These results show clearly that the addition of sulphide or thiosulphate to an aerobic suspension of *Thiobacillus* sp. strain c was followed within 10 sec. by the synthesis of ATP. But whereas the ATP synthesis which accompanied the thiosulphate oxidation was quite unaffected by the presence of 0.1 mm-2,4-dinitrophenol (DNP), that which accompanied the sulphide oxidation was strongly inhibited. These observations support the view that in this Thiobacillus, ATP can be synthesized by at least two different mechanisms, only one of which is inhibited by DNP. The sensitive mechanism appears to be the major one coupled to sulphide oxidation, at least in its early stages. At present the most probable interpretation is that the mechanism sensitive to DNP inhibition is similar to oxidative phosphorylation in mitochondria which is also inhibited by similar concentrations of DNP (Loomis & Lipmann, 1948). When thiosulphate is oxidized, the mechanism of ATP formation which is insensitive to DNP may well be that which Peck (1960) showed could accompany the oxidation of sulphite to sulphate by cell-free extracts of *Thiobacillus thioparus*. With these extracts, Peck showed that sulphite could be formed by the reductive cleavage of thiosulphate. Our results certainly suggest that the DNP-insensitive phosphorylation studied by Peck takes place in intact organisms and that therefore sulphite is an intermediate in thiosulphate oxidation. Whole organisms of *Thiobacillus* sp. strain c do not oxidize exogenous sulphite, possibly because it does not enter the organisms, so that the coupling of ATP formation to sulphite oxidation cannot be observed directly. From our results we cannot conclude anything about the origin of sulphite. It might be formed by reductive cleavage of thiosulphate, as Peck suggests, or by oxidation through a polythionate (London & Rittenberg, 1964).

ATP is required to drive the reactions of the Calvin cycle by which carbon dioxide appears to be fixed by Thiobacillus (Trudinger, 1956; Aubert, Milhaud & Millet, 1957). One might have expected, then, that in the presence of carbon dioxide the steady-state value of ATP would be lower. That this was not so presumably indicates that ATP formation coupled to thiosulphate oxidation was so rapid that the ATP concentration was maintained at its maximum value in spite of utilization by the carbon-dioxide fixation reactions.

When sulphide is the substrate, the effect of DNP is to decrease both the steady-state concentration of ATP and the rate of fixation of carbon dioxide and it is possible that the rate of carbon-dioxide fixation is limited by the amount of ATP available (Fig. 5b). But this cannot be so when thiosulphate is the substrate. Here the effect of DNP on carbon-dioxide fixation, while less marked than when sulphide is the substrate, is nevertheless far greater than its effect on the steady-state level of ATP (Fig. 5a). In the presence of DNP the rate of carbon dioxide fixation must be limited by something other than the availability of ATP. This limiting factor could be the availability of reduced nicotinamide adenine dinucleotides, whose formation might require NAD(P) reduction with electrons from cytochromes reduced during thiosulphate oxidation (Aleem, 1965). Certainly, NAD reduction by reduced cyto-
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Chromes in mitochondria is dependent on ATP or a high energy compound (Chance, 1961; Danielson & Ernster, 1968) and is inhibited by DNP (Chance & Hollunger, 1968).

We have concluded earlier (Kelly & Syrett, 1964a) from the effect of DNP on carbon-dioxide-fixation that both DNP-insensitive and DNP-sensitive phosphorylations accompany thiosulphate oxidation. The decrease in the steady-state concentration of ATP at higher DNP concentrations with thiosulphate as substrate (Fig. 5), may indicate such a DNP-sensitive mechanism. However, the DNP concentrations producing the decrease are too high, in comparison with those affecting sulphide-dependent ATP synthesis, for this conclusion to be accepted with confidence. It is now clear, from the results presented in this paper, that the rate of carbon-dioxide-fixation is not a satisfactory index of ATP content when thiosulphate is the substrate. It may well be that thiosulphate oxidation is accompanied by both types of phosphorylation, but neither our results on carbon-dioxide-fixation nor those on ATP formation demonstrate this. Similarly, when sulphide is oxidized, it seems clear that the ATP formed initially is synthesized by a DNP-sensitive mechanism, but this does not rule out the participation of DNP-insensitive phosphorylation at a later stage in sulphide metabolism, perhaps at the level of sulphite oxidation.

REFERENCES


