Selenium-dependent Growth and Glycine Fermentation by Clostridium purinolyticum

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Clostridium purinolyticum fermented glycine as a sole carbon and energy source according to the equation:

\[ 4 \text{Glycine} + 2\text{H}_2\text{O} \rightarrow 3 \text{Acetate} + 2\text{CO}_2 + 4\text{NH}_3 \]

The organism required adenine as a supplement and selenium compounds as micronutrients for growth. The molar growth yield on glycine was 6.5 g dry wt. Radiochemical and enzymic investigations revealed a new fermentation pathway for glycine in which 1 mol glycine was completely oxidized to CO₂ and the generated reducing equivalents were used to reduce a further 3 mol glycine to acetate via the glycine reductase system. This reaction was associated with the formation of ATP.

INTRODUCTION

The ability to use glycine as a sole carbon and energy source under strictly anaerobic conditions is restricted to just a few bacterial species. Cardon & Barker (1946) isolated Peptococcus (Diplococcus) glycophilus, an organism which ferments glycine to acetate, carbon dioxide and ammonia; in addition, variable amounts of hydrogen gas are released. Strains of Peptococcus (Micrococcus) anaerobius and Peptococcus (Micrococcus) variabilis grow on glycine without the formation of hydrogen (Douglas, 1951). Guillaume et al. (1956) reported that Clostridium histolyticum decomposes glycine according to the same fermentation balance as that found for P. glycophilus. To our knowledge there have been no further reports concerning the anaerobic degradation of glycine as a sole substrate by pure cultures, although there have been two reports concerning the fermentation of this amino acid by washed suspensions of rumen bacteria (Van den Hende et al., 1963; Wright & Hungate, 1967); in both cases glycine was decomposed mainly to acetate. In Clostridium sticklandii and Clostridium sporogenes glycine serves only as electron acceptor in the Stickland reaction (Stadtman, 1978; Costilow, 1977).

The pathway of glycine fermentation in P. glycophilus was elucidated by means of tracer experiments and enzymic investigations (Barker et al., 1948; Sagers & Gunsalus, 1961; Klein & Sagers, 1962). This bacterium grows only with glycine or several glycine-containing peptides (Cardon & Barker, 1947). Clostridium purinolyticum possesses a similar narrow substrate spectrum in decomposing only purines and some degradation products, e.g. glycine (Dürre et al., 1981). The results reported in this paper show that C. purinolyticum ferments glycine by a mechanism different from that reported for P. glycophilus. The proposed new pathway is energetically more favourable and includes glycine reductase and glycine decarboxylase as key enzymes.

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Organisms. Clostridium purinolyticum WA-1 (DSM 1384) was used throughout this study. Clostridium sticklandii (ATCC 12662; DSM 519) served as a control organism during the determination of glycine reductase activity and was kindly provided by Dr. A. C. Schwartz, Institute of Botany, University of Bonn, F.R.G.

Growth media. Clostridium purinolyticum was grown in a medium containing 100 mM glycine, 4 mM K$_2$HPO$_4$, 0.14 mM-MgSO$_4$, 6.3 μM FeSO$_4$, 29 μM-CaCl$_2$, 0.1 μM-MnSO$_4$, 0.1 μM-Na$_2$SeO$_3$, 0.1 μM-Na$_2$WO$_4$, 20 mM-KHCO$_3$, 2.3 mM-(NH$_4$)$_2$SO$_4$, 1.0 mM-adenine, 29.2 mM-thioglycollic acid and 0.1% (w/v) yeast extract (Difco). Clostridium sticklandii was grown in the medium described by Stadtman (1978). All media were prepared anaerobically (under an atmosphere of 100% N$_2$) by techniques described by Hungate (1969) and Bryant (1972). Traces of oxygen were removed from the gas stream by passage over a copper catalyst column heated to about 300°C. Culture tubes (16 ml) were from Belco Glass, Vineland, N.J., U.S.A., and butyl rubber stoppers from Arthur H. Thomas Co., Philadelphia, Pa., U.S.A. Cells for enzyme determinations and tracer experiments were grown in 3 liter carboys containing 20 ml medium. Anaerobic conditions were established by gassing with N$_2$ for 1 h before inoculation. In these cases sodium dithionite (35 mg l$^{-1}$) was used as a reducing agent instead of thioglycollic acid.

Growth parameter measurements. The protein content of whole cells was determined according to Stickland (1951), and protein concentrations in cell-free extracts were measured by the method of Beisenherz et al. (1953). The dry weights of cells were determined on membrane filters and growth yields were calculated as described by Bauchop & Elsden (1960) and Stouthamer (1969). Turbidity was measured at 600 nm using a Bausch & Lomb Spectronic 88 or a Carl Zeiss PM 4 spectrophotometer.

Analytical methods. Acetate was determined by an enzymic procedure (Dorn et al., 1978), formate by the colorimetric method of Lang & Lang (1972), glycine according to Sardesai & Provido (1970), and ammonia as described by Da Fonseca-Wollheim et al. (1974). Carbon dioxide and hydrogen were measured with a Perkin-Elmer F20H gas chromatograph [oven temperatures, 185°C and 100°C, respectively; injection port temperatures, 185°C and 100°C, respectively; thermal conductivity detector temperature, 230°C; stainless steel columns 5.2 mm × 200 cm, filled with molecular sieve 5 Å, 18–50 mesh (E. Merck, Darmstadt, F.R.G.); carrier gas, N$_2$, at a flow rate of 2 ml min$^{-1}$; the method was adapted from that of Braun (1979)]. Adenine and hypoxanthine were separated by high-pressure liquid chromatography (Du Pont model 870) on a reversed-phase column (4-6 mm × 250 mm, packed with LiChrosorb RP-18, 10 μm). The purines were eluted with 100 mM-potassium phosphate buffer (pH 2.7) at a flow rate of 2 ml min$^{-1}$, and detected in a Kontron spectrophotometer (model 720 LC) at 205 nm.

Tracer experiments. Radioactive glycine, or glycine in the presence of radioactive sodium acetate or sodium bicarbonate, was fermented by resting cells of C. purinolyticum in 120 ml Warburg vessels. Washed cell suspension (5 ml; 4 μg, about 50) was pipetted into the main compartment which contained, in addition, 24.5 ml reaction buffer (composition similar to that of growth medium without glycine, adenine, KHCO$_3$, and yeast extract, but supplemented with 0.15 μM-thiamin). The centre well contained 3 ml KOH (10%, w/v) and the CO$_2$ (absorbed in KOH; 10%, w/v) was measured in a liquid scintillation counter (Philips; model PW4540). The scintillation mixture contained 5 ml Unisolve I (Koch-Light), 0.5 ml water and 0.1 ml sample.

Enzyme assays. Cells were always harvested at the end of the exponential growth phase and stored anaerobically at −20°C. They were ruptured by French press treatment, and cell-free extracts were prepared as described by Leonhardt & Andreesen (1977). All tests were performed under strictly anaerobic conditions. Formate dehydrogenase (EC 1.2.1.−) was assayed according to Dürr et al. (1981), methylenetetrahydrofolate cyclohydrolase (EC 3.5.4.9) by the methods of Uyeda & Rabinowitz (1967a) and O'Brien & Ljungdahl (1972), methylenetetrahydrofolate dehydrogenase (EC 1.5.1.5) according to Uyeda & Rabinowitz (1967b) and O'Brien & Ljungdahl (1972), formyltetrahydrofolate synthetase (EC 6.3.4.3) as described by Rabinowitz & Pricer (1962) and O'Brien & Ljungdahl (1972), glycine formiminotransferase (EC 2.1.2.4) by the method of Uyeda & Rabinowitz (1965), serine hydroxymethyltransferase (EC 2.1.2.1) according to Barnard & Akhtar (1979), glycine decarboxylase (EC 2.1.2.10) as described by Sagers & Klein (1970) and glycine reductase using the radioactive test of Stadtman (1970). A second assay for glycine reductase was based on ATP formation during the reaction. It was carried out in 1 ml cuvettes (path length 1 cm) containing, in a total volume of 1 ml: 10 mM-N-tris(hydroxymethyl)methylglycine/KOH buffer (pH 8.25), 10 mM-MgCl$_2$, 20 mM-K$_2$HPO$_4$, 4 mM-ADP, 50 mM-N-acetyl-L-serine, 0.14 mM ATP, 20 mM-glycine, and 40 μg of cell-free extract. The reaction was started by tipping the Warburg flasks which were then closed with serum stoppers and attached to a reciprocal shaker in a water bath at 37°C. They were gassed with N$_2$ for 1 h and the reaction was started by tipping the glycerol solution from the side-arm into the main compartment. After 5–7 h the reaction was stopped by injecting 4 ml of 4 M-HCl. Subsequently, the flasks were shaken for 1 h to allow quantitative absorption of the CO$_2$ formed by the KOH. The reaction solution was centrifuged and acetate was separated from the other components by steam distillation. The purified acetate was degraded to CO$_2$ and methylamine according to Sakami (1955). The radioactivity of the methylamine (in H$_2$SO$_4$; 10%, w/v) and the CO$_2$ (absorbed in KOH; 10%, w/v) was measured in a liquid scintillation counter (Philips; model PW4540). The scintillation mixture contained 5 ml Unisolve I (Koch-Light), 0.5 ml water and 0.1 ml sample.
Selenium-dependent glycine fermentation

1 mM-NADP, 4 mM-AMP, 40 mM-glycine, 20 mM-dithiothreitol, 10 mM-glucose, 42 U hexokinase, 7 U glucose-6-phosphate dehydrogenase, and extract. The reaction was started by adding either extract or glycine and the formation of NADPH was monitored at 340 nm. l Serine dehydratase (EC 4.2.1.13) activity was determined in 1 ml cuvettes (path length 1 cm) containing, in a total volume of 1 ml: 100 mM-potassium phosphate buffer (pH 8.4), 100 mM-L-serine, 1 mM-FeSO₄, 10 mM-dithioerythritol, 55 U lactic dehydrogenase, 0.3 mM-NADH, and extract. The reaction was started by adding activated extract, obtained by incubating cell-free extracts with Fe(II) and dithiothreitol as described by Carter & Sagers (1972). The decrease in NADH was monitored at 366 nm. One unit of enzyme activity was defined as 1 μmol substrate transformed or product formed min⁻¹ at 37 °C.

RESULTS

Nutritional requirements

The growth curve of Clostridium purinolyticum on glycine always showed a distinct deflexion in the early-exponential growth phase (Fig. 1) unless the medium was supplemented with adenine (Fig. 2). Synthetic amino acid solution, vitamins, pyrimidines (uracil, cytosine and thymine), or biogenic amines such as spermine, agmatine, putrescine and cadaverine, in various concentrations could not substitute for adenine. Even guanine was not effective. Moreover, the growth rate decreased as soon as the adenine component of the yeast extract in the medium was exhausted. The requirement of C. purinolyticum for adenine could be satisfied using a 1 mM concentration (Fig. 2). The glycine concentration which gave the best growth yield was 100 mM. The organism showed no strict requirement for bicarbonate but in the presence of 0.2% (w/v) KHCO₃ the final turbidity of the culture was about 30% higher than in the unsupplemented culture. Higher concentrations proved to be somewhat inhibitory. Growth of C. purinolyticum on glycine was strictly dependent on the addition of the trace element selenium to the medium (Fig. 3), as was previously observed for growth on adenine (Dürre et al., 1981). Neither molybdenum nor tungsten could substitute for selenium. However, the addition of molybdenum and/or tungsten to the selenium-containing medium resulted in a further increase in growth. Maximum turbidity was reached at a concentration of 1 μM-selenite (Fig. 4) when the cultures were previously starved of this substance by at least five transfers (1%, v/v, inoculum each) in trace element-free medium. Obviously C. purinolyticum had the capability to concentrate selenium compounds very efficiently. If the organism was pre-cultured for several transfers in a medium supplemented with 0.1 μM-selenite, the final turbidity reached was similar to that obtained with 1 μM-selenite. For this reason a concentration of 0.1 μM-selenite was routinely chosen for all media. At 0.1 μM, selenomethionine was as effective as selenite, whereas selenocystine gave a lower stimulation. Chemically related elements such as tellurium and sulphur (in the form of tellurite, sulphate and cysteine) did not exhibit any stimulatory effect on growth (Fig. 4). The curves for sulphate and cysteine were identical to that for tellurite and are therefore not shown.

Substrate utilization and fermentation products

The growth response of C. purinolyticum on glycine is shown in Fig. 2. Exponential growth occurred between 3 and 11 h after inoculation, as judged by the turbidity of the culture, its protein content and the dry weight of the cells. After reaching the stationary phase, cell lysis was observed. The protein : dry weight ratio of the cells was about 60–70% throughout the growth cycle, indicating that no storage material was formed. The doubling time was 3.7 h as calculated from the increase in turbidity. A maximum turbidity (A₆₀₀) of 1.14 was measured, corresponding to a dry weight of 0.4 mg ml⁻¹. The molar growth yield was 6.5 g dry wt (mol glycine)⁻¹. The fermentation products were acetate, NH₃ and CO₂. Formate and H₂ could not be detected. Glycine became depleted when the cells reached the stationary phase. Adenine was rapidly broken down, being degraded to hypoxanthine which could be detected in the medium. Subsequently, the cells took up this purine, probably utilizing it in biosynthetic reactions. The total purine content of the medium was exhausted after 13 h. The acetate level
Fig. 1. Growth curve of *C. purinolyticum* on glycine without addition of adenine to the medium. Turbidity was measured in anaerobic culture tubes with a Spectronic 88 spectrophotometer.

Fig. 2. Growth, substrate utilization and product formation by *C. purinolyticum* during growth on glycine. Cells were cultured in a 4 l fermenter under strictly anaerobic conditions. The volume of the gas phase in this apparatus was kept constant by a special construction and the pH was kept constant at 7.5. ○, glycine; □, adenine; △, hypoxanthine; ●, total purine content of the medium; ■, acetate; ▲, dry weight.

Fig. 3. Influence of trace elements on growth of *C. purinolyticum* on glycine. The medium used was that described in Methods with the omission of SeO₄²⁻, WO₄²⁻, and MoO₄²⁻. Data shown give the maximum turbidities reached after supplementation of the medium with various combinations of these elements (10⁻⁷ M) and inoculation with starved cells.

Fig. 4. Influence of increasing concentrations of selenium and tellurium compounds on growth of *C. purinolyticum* on glycine. The conditions were the same as described in Fig. 3. ○, selenite; ●, selenomethionine; □, selenocystine; ■, tellurite.
Selenium-dependent glycine fermentation

Table 1. Fermentation balance for glycine decomposition by C. purinolyticum

Resting cells were incubated with glycine in a Warburg vessel at 37 °C. For culture conditions and metabolite determinations, see Methods. Calculation of the oxidation–reduction (O/R) balance was performed according to the method of Whiteley & Douglas (1951) adapted to the definition of Wood (1961).

<table>
<thead>
<tr>
<th>Substrate and products</th>
<th>mmol (100 mmol substrate)⁻¹</th>
<th>mmol C (100 mmol)⁻¹</th>
<th>O/R balance</th>
<th>Balance of available H</th>
<th>mmol (100 mmol)⁻¹</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glycine</td>
<td>100-0</td>
<td>200-0</td>
<td>+1</td>
<td>6</td>
<td>600-0</td>
</tr>
<tr>
<td>Acetate</td>
<td>72-7</td>
<td>145-4</td>
<td>0</td>
<td>8</td>
<td>581-6</td>
</tr>
<tr>
<td>Ammonia</td>
<td>97-0</td>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>CO₂</td>
<td>47-5</td>
<td>47-5</td>
<td>+2</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Balance</td>
<td></td>
<td></td>
<td>96-5 %</td>
<td>1-05</td>
<td>96-9 %</td>
</tr>
</tbody>
</table>

Table 2. Distribution of radioactivity in the products of glycine decomposition by resting cells of C. purinolyticum in the presence of ¹⁴C-labelled substances

Conditions were as described in Methods. Glycine was used at a concentration of 1-2 mmol (per 30 ml) in each experiment; [1-¹⁴C]acetate, [2-¹⁴C]acetate and [¹⁴C]bicarbonate were added at concentrations of 0-11 μmol, 0-11 μmol and 0-68 μmol, respectively. The ¹⁴C balance was calculated according to the fermentation equation: 4 Glycine + 2H₂O → 3 Acetate + 2CO₂ + 4NH₃.

Specific radioactivity (d.p.m. mmol⁻¹)

<table>
<thead>
<tr>
<th>Labelled substrate</th>
<th>Substrate</th>
<th>CH₃ group of acetate*</th>
<th>COOH group of acetate*</th>
<th>CO₂</th>
<th>¹⁴C balance (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>[1-¹⁴C]Glycine</td>
<td>3.7 x 10⁵</td>
<td>41220 (12-3)</td>
<td>292730 (87-7)</td>
<td>201360</td>
<td>94-9</td>
</tr>
<tr>
<td>[2-¹⁴C]Glycine</td>
<td>3.7 x 10⁵</td>
<td>381760 (94-4)</td>
<td>22430 (5-6)</td>
<td>911690</td>
<td>94-3</td>
</tr>
<tr>
<td>[1-¹⁴C]Acetate</td>
<td>4.0 x 10⁹</td>
<td>14140 (8-5)</td>
<td>153200 (91-5)</td>
<td>91140</td>
<td>46-2</td>
</tr>
<tr>
<td>[2-¹⁴C]Acetate</td>
<td>4.0 x 10⁹</td>
<td>73660 (91-6)</td>
<td>6740 (8-4)</td>
<td>39790</td>
<td>21-7</td>
</tr>
<tr>
<td>NaH¹⁴CO₃</td>
<td>6.5 x 10⁹</td>
<td>5390 (52-2)</td>
<td>4920 (47-7)</td>
<td>600310</td>
<td>83-2</td>
</tr>
</tbody>
</table>

* Numbers in parentheses indicate the percentage of acetate labelled in the group specified.

increased during the course of the glycine fermentation, lowering the pH to a final value of about 6-9. Production of NH₃ and CO₂ was determined in another experiment. A typical fermentation balance is given in Table 1. These data are in accordance with the following fermentation equation:

\[
4 \text{Glycine} + 2\text{H}_2\text{O} \rightarrow 3 \text{Acetate} + 2\text{CO}_2 + 4\text{NH}_3
\]

As previously reported C. purinolyticum is also able to grow with formiminoglycine (Dürre et al., 1981). Fermentation experiments showed that this compound is decomposed according to the equation:

\[
\text{1 Formiminoglycine} \rightarrow 0.88 \text{Acetate} + 0.11 \text{Formate} + 0.95 \text{CO}_2
\]

(NH₃ was not determined).

Tracer experiments

Resting cells of C. purinolyticum were used to ferment [1-¹⁴C]glycine and [2-¹⁴C]glycine (Table 2). As a control, decomposition of glycine in the presence of radioactive [1-¹⁴C]acetate, [2-¹⁴C]acetate or [¹⁴C]bicarbonate was tested. Nearly all the radioactive glycine was directly fermented to the correspondingly labelled radioactive acetate. 'Missing'
Table 3. Specific activities of enzymes of C. purinolyticum participating in glycine metabolism

Culture conditions and assay systems were as described in Methods.

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Specific activity [units (mg protein)⁻¹]</th>
</tr>
</thead>
<tbody>
<tr>
<td>Glycine reductase (radiochemical test)</td>
<td>0.090</td>
</tr>
<tr>
<td>Glycine reductase (measured by ATP formation)</td>
<td>0.041</td>
</tr>
<tr>
<td>Glycine decarboxylase</td>
<td>0.012</td>
</tr>
<tr>
<td>Methylene tetrahydrofolate dehydrogenase (NADP as coenzyme)</td>
<td>1.1</td>
</tr>
<tr>
<td>Methenyl tetrahydrofolate cyclohydrolase</td>
<td>0.007</td>
</tr>
<tr>
<td>Formyl tetrahydrofolate synthetase</td>
<td>35.5</td>
</tr>
<tr>
<td>Formate dehydrogenase</td>
<td>1.0</td>
</tr>
<tr>
<td>Formiminotetrahydrofolate cyclodeaminase</td>
<td>present</td>
</tr>
<tr>
<td>Glycine formiminotransferase</td>
<td>3.2</td>
</tr>
<tr>
<td>Serine hydroxymethyltransferase</td>
<td>0.2</td>
</tr>
<tr>
<td>L-Serine dehydratase</td>
<td>1.8</td>
</tr>
<tr>
<td>Glycine reductase (C. sticklandii; radiochemical test)</td>
<td>0.06</td>
</tr>
</tbody>
</table>

Radioactivity as calculated from the ¹⁴C balance always appeared in the cells. Part of the substrate was completely oxidized to CO₂. Most of the added acetate was either taken up by the cells or oxidized to CO₂, since only 22–46% could be detected in the medium. The cells were able to convert CO₂ to acetate as shown by the results of the experiment performed with radioactive bicarbonate. It can be seen that acetate is derived directly from glycine. The small deviations from the theoretical value corresponded to the exchange determined in the control experiments with radioactive acetate. Only a very small amount of added ¹⁴CO₂ was converted to acetate with both carbon atoms equally labelled.

Enzyme studies

Activities of enzymes known to be involved in glycine metabolism were shown to be present in cell-free extracts of C. purinolyticum (Table 3). A high glycine reductase activity was detected as expected from the results of the tracer experiments. This enzyme converts glycine directly to acetate, releasing ammonia. Dithiothreitol was used as the electron donor. Furthermore, glycine-dependent ATP production could be observed during this reaction. Cell-free extracts of C. sticklandii showed a somewhat lower activity of the enzyme although the value was about 10-fold higher than those reported by Stadtman & Elliott (1956). This might be due to the more gentle storage conditions established by Barnard & Akhtar (1979) and also used by us. The activities determined for glycine decarboxylase, formyltetrahydrofolate synthetase, methenyltetrahydrofolate cyclohydrolase and methylene tetrahydrofolate dehydrogenase were of about the same order as known for purine-fermenting clostridia (Champion & Rabinowitz, 1977). Methylene tetrahydrofolate dehydrogenase reacted with both NAD and NADP as coenzymes but the activity measured with NADP was 3- to 10-fold higher than that with NAD. Uyeda & Rabinowitz (1967b) obtained similar results with crude extracts of C. cylindrosporum whereas the pure enzyme was specific for NADP. The test system for the glycine formiminotransferase also allows the detection of formiminotetrahydrofolate cyclodeaminase activity (Uyeda & Rabinowitz, 1965) which was also present in C. purinolyticum. Formate dehydrogenase, l-serine dehydratase, and serine hydroxymethyltransferase showed comparatively low activities. Preliminary results indicated the presence of a pyruvate:methylviologen oxidoreductase.
DISCUSSION

Glycine seems to be a difficult substrate to utilize for growth. This might be due to its interference with cell wall biosynthesis (Schleifer et al., 1976). As already mentioned, besides C. purinolyticum, only two anaerobic bacteria are known to ferment this amino acid as sole carbon and energy source. The striking nutritional features of C. purinolyticum during growth on glycine are the requirements for adenine and selenium compounds. Omission of adenine from the medium did not prevent growth completely, indicating that the organism is able to synthesize purines de novo. The deflecting growth curve suggests that the rate of purine synthesis is limiting under these conditions. Since C. purinolyticum can use purines as a carbon and energy source (Dürre et al., 1981), degradation and synthesis of these compounds must be regulated very carefully. The thiamin requirement of most purine-utilizing anaerobes (J. R. Andreesen, P. Dürre, H. Schiefer & R. Wagner, unpublished results) might be due to the restricted biosynthetic capacity in the common reactions of purine and thiamin formation (Newell & Tucker, 1968). Clostridium purinol-yticum may satisfy its requirement for purines by converting adenine first into hypoxanthine which is subsequently used for the production of purines needed for the synthesis of nucleic acids and coenzymes. The detection of hypoxanthine in the medium during growth may be an indication that the first reaction is performed by an exo-enzyme or a periplasmic enzyme resulting in an imbalance between the rates of production and utilization of hypoxanthine.

Only a few instances of a similar strict requirement for selenium compounds have been described. Both C. sticklandii and C. sporogenes need selenite for the utilization of glycine as an oxidant in the Stickland reaction (Turner & Stadtman, 1973; Costilow, 1977). Selenite is an essential nutrient for Peridinium cinctum fa. westii (Lindström & Rodhe, 1978). The last two organisms have therefore been suggested for use in a bioassay for selenium (Costilow, 1977; Lindström, 1980). During decomposition of purines, C. purinolyticum also shows a requirement for selenium which, however, can be partly satisfied by molybdate and tungstate (Dürre et al., 1981). The extremely low concentrations of selenium which proved to be effective for growth on glycine point to a high-affinity selenium transport system in this organism.

The dependence on the trace element selenium first gave rise to the idea that a glycine reductase might be involved in glycine fermentation, because selenium was shown to be an essential component of this enzyme complex (Turner & Stadtman, 1973). The reaction catalysed by glycine reductase was discovered by Stadtman & Elliott (1956) while investigating the degradation of glycine and other amino acids in a Stickland reaction by C. sticklandii. They reported that the direct reduction of glycine to acetate was concomitant with the formation of ATP from ADP plus inorganic phosphate. The results of the tracer experiments (Table 2) and the enzyme assays (Table 3) prove that in vivo the acetate formation from glycine in C. purinolyticum does involve such an enzyme. Furthermore, energy production during the reaction could be observed. Hence, the pathway of glycine fermentation operating in C. purinolyticum is different from that known for Peptococcus glyciphilus. The latter decarboxylates glycine to methylenetetrahydrofolate which is condensed with a second molecule of glycine, yielding serine. Serine is subsequently converted to pyruvate from which acetate is formed (Sagers & Gunsalus, 1961; Klein & Sagers, 1962). Only 1 mol ATP is synthesized during the degradation of 4 mol glycine. The fermentation follows the equation (Cardon & Barker, 1946):

\[4 \text{Glycine} + 2\text{H}_2\text{O} \rightarrow 3 \text{Acetate} + 2\text{CO}_2 + 4\text{NH}_3\]

Although C. purinolyticum ferments glycine according to the same overall balance, the pathway proposed is completely different (Fig. 5). This scheme is in accordance with all the results from the tracer experiments and enzyme assays. 1 mol glycine is oxidized to CO₂, thus providing energy via the formyltetrahydrofolate synthetase reaction. The enzyme of C. purinolyticum was not assayed by formate production; however, all other known enzymes catalyse ATP formation during this reaction (Himes & Harmony, 1973) so that a similar
process can be assumed to function in this organism. The reducing equivalents produced during the oxidation of 1 mol glycine are transferred to a glycine reductase which reduces 3 mol glycine to 3 mol acetate and 3 mol ammonia, concomitant with the formation of ATP. The stoichiometry of glycine reduction and ATP formation is not quite clear. In *C. sticklandii* a ratio of 1:1 was reported (Stadtman et al., 1957). However, Barnard & Akhtar (1979) obtained a lower proportion which they attributed to the presence of active ATPase in the cell-free extracts. The determination of an exact stoichiometry can only be performed with lysates free of adenylate kinase (Tanaka & Stadtman, 1979). These factors may be responsible for the lower glycine reductase activities obtained with the ATP assay (Table 3).

According to the proposed pathway, the molar growth yield of 6.5 g dry wt (mol glycine)$^{-1}$ determined in *C. purinolyticum* will be equivalent to a $Y_{ATP}$ of 6.5 g dry wt (mol ATP)$^{-1}$ because a maximum of 4 mol ATP can be gained from 4 mol glycine (1 mol during the formyltetrahydrofolate synthetase reaction and 3 mol during the glycine reductase reaction). Although $Y_{ATP}$ was originally thought to be a biological constant, many studies have indicated that it may vary in the range 4.7–28.5 in anaerobic bacteria, depending on, for example, the size of the carbon skeleton of the substrate (Stouthamer, 1979). From this point of view it is probable that *C. purinolyticum* forms ATP via the reductase system as indicated by the enzymic studies. Thermodynamic data show that a free energy change ($\Delta G^{\circ}$) of $-207.5$ kJ (4 mol glycine)$^{-1}$ [i.e. $-49.6$ kcal (4 mol glycine)$^{-1}$] can be calculated for the reaction:

$$4 \text{Glycine} + 4\text{H}_2\text{O} \rightarrow 3 \text{Acetate}^- + 2\text{HCO}_3^- + 4\text{NH}_4^+ + \text{H}^+$$

(all data taken from Thauer et al., 1977). This corresponds to a value of $-51.9$ kJ (mol glycine transformed)$^{-1}$ (i.e. $-12.4$ kcal mol$^{-1}$). Because the formation of 1 mol ATP must be associated with a free energy change of $-41.8$ to $-50.2$ kJ mol$^{-1}$ (i.e. $-10$ to $-12$ kcal mol$^{-1}$), the formation of 4 mol ATP during the reaction described above would result in a thermodynamic efficiency of more than 80%. Up to now, only values up to 70% are known (McCarty, 1964). Further investigation is necessary to clarify the stoichiometry of glycine.
Selenium-dependent glycine fermentation

It would be helpful to investigate the fermentation of formiminoglycine in detail, because the formimino group of this compound is obviously oxidized to formate and CO₂ via tetrahydrofolate derivatives, whereas the remaining glycine is reduced to acetate. This scheme implies a 1:1 ratio between the two reaction sequences, which would allow a more precise determination of the amount of energy gained. The glycine-serine pathway is probably only used for biosynthetic purposes, as indicated by the tracer experiments and the low activities of both serine hydroxymethyltransferase and L-serine dehydratase. In Clostridium acidiurici, an organism known to generate energy via this pathway, the activities are 400 and 8 times higher, respectively (Carter & Sagers, 1972; Champion & Rabinowitz, 1977).

Fermentation of glycine as sole carbon and energy source via a glycine reductase was presumed to occur in rumen bacteria (Van den Hende et al., 1963). The authors based this conclusion only on the fact that no CO₂ was released during the decomposition of glycine. In contrast, Wright & Hungate (1967) assumed that suspensions of rumen micro-organisms fermented glycine mainly in the same manner as P. glyciphilus, although the data from their tracer experiments pointed to a high participation of a glycine reductase.

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REFERENCES


