Growth of Sulphate-reducing Bacteria by Fumarate Dismutation

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SUMMARY

Desulfovibrio gigas and several strains of D. desulfuricans grew by fumarate dismutation in a sulphate-free medium. Two strains of D. desulfuricans grown in a chemically defined medium formed succinate, malate and acetate during fumarate dismutation. Sulphate reduction by these strains, though not by D. gigas, was almost completely inhibited in presence of fumarate as alternative electron acceptor. The anomalous behaviour of D. gigas was reflected to some extent by the hydrogen absorption coefficients for fumarate and sulphate reduction. Effects of fumarate media on the morphology of one strain are recorded.

INTRODUCTION

Under natural conditions the sulphate-reducing bacteria appear to utilize sulphate as terminal electron acceptor in the assimilation of organic substances, and have been shown (Postgate, 1951) to reduce certain other sulphur-containing anions in culture. Growth on pyruvate in the absence of sulphur compounds has been reported for strains of Desulfovibrio desulfuricans (Postgate, 1952) and Clostridium nigrificans (Postgate, 1963); Senez & Pascal (1961) and Baker, Papiska & Campbell (1962) obtained growth of various strains of D. desulfuricans on choline in absence of sulphate. Grossman & Postgate (1955) studied sulphate-free metabolism of malate and fumarate, not accompanied by growth, in D. desulfuricans strain El Agheila z. The present paper reports the discovery that Desulfovibrio gigas and a number of strains of D. desulfuricans are able to grow by dismutation of fumarate, i.e. by a Cannizzaro-type reaction which involves the simultaneous oxidation and reduction of the substrate.

METHODS

Organisms. Twenty-three strains of sulphate-reducing bacteria were obtained as freeze-dried ampoules from the National Collection of Industrial Bacteria (NCIB). The strain names are followed by the NCIB number.

1. Mesophiles (growth temperature 30°C).
   (a) Desulfovibrio gigas, a fresh-water strain (NCIB 9332).
   (b) D. desulfuricans, salt-water strains: El Agheila A, 8809; Norway 4, 8810; New Jersey sw-8, 8815; New Jersey sw-8, 8816; California 29:137:5, 8826; Texas 29:12:8, 8828; Australia, 8829; California 43:63, 8864; El Agheila 4, 8896; Venezuela, 8899; Aberdovey, 9492.

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(c) *D. desulfuricans*, fresh-water strains: Teddington r, 8312; Denmark, 8456; Woolwich, 8457; Byron, 8458.

(2) Thermophiles (growth temperature 55°). Strains of *Clostridium nigricans*, all fresh-water strains: Holland cr, 8856; Delft 48r, 8857; Delft 8r, 8859; Delft 13r, 8860; Delft 15r, 8861; unnamed strains 8706 and 8788.

**Maintenance of stock cultures.** Cultures were raised from the freeze-dried condition at 30° or 55°, as appropriate, in the medium of Baars (1980) containing 1.0 g. Difco yeast extract, and 5 mm-cysteine hydrochloride. Sodium chloride, 25 g./l., was added for salt-water organisms. Details of the preparation, sterilization and pH adjustment of this medium are given elsewhere (Saleh, Macpherson & Miller, 1964). Stock cultures of the mesophilic strains were maintained in Postgate’s modification of the medium C of Butlin, Adams & Thomas (1949; see Baker et al. 1962) containing cysteine, and NaCl where necessary. The thermophilic strains grew better in Baars’s medium + yeast extract, in which medium stock cultures were therefore maintained. Subcultures were made weekly. Stock and experimental cultures were grown in Pyrex test tubes or Erlenmeyer flasks plugged with cottonwool and incubated in McIntosh & Fildes anaerobic jars under an atmosphere of N₂. Frequent tests were made on stock and experimental cultures for aerobic and anaerobic contaminants (Postgate, 1958).

**Experimental media.** Two types of media were used:

1. A basal medium consisting of modified medium C with sodium lactate, Na₂SO₄ and MgSO₄ omitted, and containing 8 mm-MgCl₂.6H₂O. To this basal medium was added 50 mm-sodium lactate (L), 50 mm-sodium fumarate (F), or 50 mm-Na₂SO₄ (S), singly or in admixture. Since the basal medium contained yeast extract (Y), the various media derived from this basal medium were designated FY, LFY, FSY and LSY.

2. A basal medium consisting of the chemically-defined medium of Macpherson & Miller (1963), also with lactate and sulphate omitted, and containing 0.25 mm-MgCl₂.6H₂O. Similar additions of lactate, fumarate or sulphate were made, the resulting media being designated F, LF, FS and LS.

FeSO₄ was added to all experimental media to 25 μM. This amount was sufficient for assimilatory sulphur and iron metabolism but insufficient for discernible growth by sulphate reduction to occur.

**Chromatography.** Dicarboxylic acids in culture filtrates were identified by descending chromatography on Whatman no. 1 paper using a tert-amyl alcohol + chloroform + water + formic acid (50+50+50+18.75, by vol.) solvent system or a di-isopropyl ketone + water + formic acid (50+50+18.8 by vol.) system. Oxaloacetate was searched for with a n-butanol + water + formic acid (95+100+5 by vol.) system (Magasanik & Umbarger, 1950) and a methylethylketone + acetone + water + formic acid (80+4+12+2 by vol.) system (Reio, 1959). Acetate, removed by steam distillation from the acidified culture filtrate, was identified by using n-butanol+1.5N-NH₄OH (1+1 by vol.) solvent, and by the standard chemical tests (Feigl, 1960).

**Sulphide estimation.** For estimation of sulphide formed in cultures, H₂S in the gas phase of the anaerobic jars was first displaced by a stream of N₂, absorbed in 1% (w/v) cadmium acetate solution and the precipitated CdS determined iodometrically (Wilson & Wilson, 1959). Dissolved and precipitated sulphide was then removed
Sulphate-reducing bacteria

from the acidified and heated cultures by a stream of N₂ and estimated as above.

Manometry. Measurement of hydrogenase activity was made at 87° in the Warburg respirometer following the procedure of Littlewood & Postgate (1956) except that 15% (w/v) KOH was used as absorbent for H₂S and CO₂.

Dry weight determinations. Dry weights for the sulphide estimations were determined by centrifuging portions of cultures at about 15,000 g for 10 min., washing and drying the bacteria at 105° to constant weight. For manometry, suspensions of organisms were standardized turbidimetrically and related to a calibration curve of strain Hildenborough (8808) for the purpose of calculating —Q₉₅ values.

RESULTS

Growth of various sulphate-reducing bacteria in fumarate media

Tubes of LFY and FY media were inoculated from stock cultures. The former medium was intended to test for the function of fumarate as terminal electron acceptor in place of sulphate, while growth in FY medium in absence of H₂ would indicate the simultaneous utilization of fumarate as electron donor, carbon source and electron acceptor. Where growth occurred, that in the fifth subculture into the experimental medium is shown rough-quantitatively in Table 1. Eleven mesophilic

Table 1. Growth of sulphate-reducing bacteria in fumarate media

Growth in the fifth subculture in experimental medium was recorded; that which was judged by eye to be about equal to growth in medium C was designated ++ +. For explanation of symbols denoting media see Methods.

<table>
<thead>
<tr>
<th>Organism</th>
<th>LFY medium</th>
<th>FY medium</th>
<th>Organism</th>
<th>LFY medium</th>
<th>FY medium</th>
</tr>
</thead>
<tbody>
<tr>
<td>D. gigas</td>
<td>-</td>
<td>++ +</td>
<td>D. desulfuricans</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>D. desulfuricans</td>
<td>-</td>
<td>+ + +</td>
<td>salt-water strains</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>El Agheila A</td>
<td>-</td>
<td>+ +</td>
<td>Teddington 8</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Norway 4</td>
<td>+ + +</td>
<td>+ + +</td>
<td>Denmark</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>New Jersey sw-8</td>
<td>+ - +</td>
<td>+ + +</td>
<td>Woolwich</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>New Jersey sw-8</td>
<td>-</td>
<td>+ + +</td>
<td>Byron</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>California 29:187:5</td>
<td>+ + +</td>
<td>+ + +</td>
<td>C. nigrificans</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Texas 29:12:6</td>
<td>-</td>
<td>+ + +</td>
<td>Holland cr</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Australia</td>
<td>+</td>
<td>+</td>
<td>Delft 48T</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>California 43:68</td>
<td>+ + +</td>
<td>+ + +</td>
<td>Delft 3T</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>El Agheila 4</td>
<td>-</td>
<td>+ + +</td>
<td>Delft 18T</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Venezuela</td>
<td>-</td>
<td>+ + +</td>
<td>Delft 15T</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Aberdovey</td>
<td>-</td>
<td>-</td>
<td>8706</td>
<td>-</td>
<td>-</td>
</tr>
</tbody>
</table>

strains dismuted fumarate; five of these grew less well or not at all in presence of lactate, for a reason not at present understood, while the converse is true for only one strain. No thermophilic strain grew in either medium. The trace amount of sulphate present was insufficient to support discernible growth when fumarate was omitted and lactate added.

Fumarate utilization in a chemically defined medium. To eliminate the possibility that fumarate dismutation was dependent on the presence of yeast extract, the two fresh-water strains which grew in LFY and FY media (Teddington 8 and Byron)
were later subcultured into both LF and F media. Heavy growth of both strains persisted through indefinite subculture in either medium.

Products of fumarate metabolism

Identification of metabolic end-products was done on filtrates of cultures of the fresh-water strains, since in these cases no desalting was necessary before chromatography. Succinate and acetate were present in F and LF cultures of Teddington R and Byron; traces of malate, confirmed by the colour reaction with ammoniacal \( \text{AgNO}_3 \) (Buch, Montgomery & Porter, 1952), were found in some cultures in F medium. Oxaloacetate was not detected. No growth of the Teddington R strain occurred in a chemically-defined medium containing succinate as sole carbon source, and sulphate; malate was dismutated in sulphate-free medium by this and certain other strains (Elford, Miller & Wakerley, in preparation).

Sulphide production during growth in presence of fumarate

It seemed of interest to determine whether sulphate reduction occurred in the presence of fumarate as an alternative electron acceptor. Strain Teddington R and \textit{Desulfovibrio gigas} were inoculated into 200 ml. portions of FSY medium and into controls of LSY medium. The cultures were analysed for sulphide before reaching the maximum extinction attainable in FY medium by the strain concerned. The results of the analyses are given in Table 2.

<table>
<thead>
<tr>
<th>Organism Grown in</th>
<th>Sulphide formed (( \mu \text{moles/mg. dry wt. bacteria} ))</th>
</tr>
</thead>
<tbody>
<tr>
<td>FSY medium</td>
<td>LSY medium</td>
</tr>
<tr>
<td>\textit{Desulfovibrio gigas}</td>
<td>27.3</td>
</tr>
<tr>
<td>\textit{D. desulfuricans} strain Teddington R</td>
<td>0.9</td>
</tr>
<tr>
<td>strain Byron</td>
<td>1.2</td>
</tr>
</tbody>
</table>

For explanation of symbols denoting media see Methods.

Uptake of hydrogen by organisms grown in fumarate media

The results of \( -Q_{h} \) determinations for fumarate and sulphate reduction are shown in Table 3. A curious feature was that \( Q_{h}^{\text{fumarate}} \) was more than ten times greater in strain Teddington R grown in LSY than in FSY medium. \textit{Desulfovibrio gigas} has already been reported (Le Gall, 1963) to be hydrogenase-positive; its \( -Q_{h} \) for sulphate reduction was markedly higher than those usually obtained with strains of \textit{D. desulfuricans}.

Morphology and pigmentation of strain Teddington R in fumarate media

The description of strain Teddington R lodged with the NCIB states that it is non-motile, with a tendency to pleomorphism. In the present work, organisms from cultures in lactate + sulphate medium (LS) were found to be normal vibrios or short spirilla, of mean dimensions about \( 1.8 \mu \times 0.7 \mu \), and some showed the typical progressive motility of, for example, the Hildenborough strain (NCIB 8303). To rule
out contamination of our stock, cultures were raised from single organisms, both motile and non-motile, obtained from a culture in LS medium. A roughly similar proportion of motile to non-motile organisms was found in all these clones.

In LF medium the morphology was much the same as in LS medium. In F medium the organisms were rods of up to 8.8 μm × 0.7 μm, often with no detectable curvature, a few being motile. In FS medium the rods were longer (up to 6.2 μm), occasionally motile, with a tendency to form chains (non-motile) of up to four organisms. Only normal black colonies developed when these morphologically aberrant organisms were inoculated into Postgate's (1953) solid test medium.

Organisms from all these media showed a strong absorption band at about 558 mμ typical of reduced cytochrome c₅₉ in the Hartridge reversion spectroscope, and gave a positive reaction to the desulfoviridin test (Postgate, 1959).

Table 3. Hydrogen absorption coefficients of Desulfovibrio organisms suspended in phosphate buffer

<table>
<thead>
<tr>
<th>Medium in which grown</th>
<th>Desulfovibrio gigas</th>
<th>D. desulfuricans</th>
<th>Hydrogen absorption coefficients</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Q₅⁴₁H₄</td>
<td>Q₅⁴₃H₄</td>
<td>QH₄</td>
</tr>
<tr>
<td>LSY</td>
<td>677</td>
<td>240</td>
<td>20</td>
</tr>
<tr>
<td>FSY</td>
<td>605</td>
<td>338</td>
<td>72</td>
</tr>
<tr>
<td>LFY</td>
<td>*</td>
<td>*</td>
<td>4</td>
</tr>
<tr>
<td>FY</td>
<td>*</td>
<td>*</td>
<td>9</td>
</tr>
</tbody>
</table>

* Not determined.

For explanation of symbols denoting media see Methods.

DISCUSSION

Grossman & Postgate (1955) reported growth of Desulfovibrio desulfuricans strain El Agheila z in an FSY-type medium; organisms suspended in phosphate buffer metabolized fumarate in absence of sulphate. Two strains have now been shown to grow by fumarate dismutation through indefinite subculture in a chemically-defined medium containing only trace amounts of sulphate; this nutritional pathway appears to be of common occurrence amongst the mesophilic sulphate-reducing bacteria, including the aberrant Norway 4 strain of D. desulfuricans (Miller & Saleh, 1964) and the recently-described species D. gigas (Le Gall, 1968). The detection of succinate, malate and acetate as end-products of growth suggests the existence of a metabolic pathway: fumarate → malate → lactate → pyruvate → acetate at the expense of reduction of some of the substrate to succinate (compare Grossman & Postgate, 1955). Hydrogen is not necessary for this sulphate-free growth. The mode of fumarate oxidation in sulphate-reducing bacteria is clearly different from that in Acetobacter xylinum, investigated by Benziman & Abeliowitz (1964), a characteristic of which is oxaloacetate accumulation.

Fumarate can act as an alternative electron acceptor to sulphate for growth reactions in Teddington R and Byron (the only strains of D. desulfuricans examined in this connexion) even when sulphate is present: sulphide production was almost completely suppressed in presence of fumarate, and succinate appeared among the
end-products of metabolism. Such preferential use of fumarate would be expected from theoretical considerations, though Grossman & Postgate (1955) found that small additions of fumarate to suspensions of strain El Agheila z organisms in phosphate buffer under H₂ increased the rate of sulphate reduction, in contrast to its effect on strains Teddington R and Byron in a nutrient medium under N₂. In the case of D. gigas, however, sulphide production per unit dry wt. of bacteria formed was roughly the same in FSY as in LSY medium (see Table 2), and a relatively small amount of succinate appeared on chromatograms of culture filtrates. The much lower values for \( \text{Q}^{3+}_{\text{H}2} \) than for \( \text{Q}^{3+}_{\text{H}4} \) in strain Teddington R, and the converse for D. gigas (Table 8), may have a bearing on this observation, though Teddington R organisms grown in FSY medium actually had an unusually high \( \text{Q}^{3+}_{\text{H}2} \) value while only minute traces of sulphide were produced during growth in this medium. It appears that in the case of the sulphate reducers the behaviour of resting organisms towards oxidizable or reducible substrates may sometimes be quite different from that during growth.

The original description of strain Teddington R as non-motile is evidently incorrect; we have confirmed that it is pleomorphic, though not in lactate+sulphate medium.

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REFERENCES


Sulphate-reducing bacteria


