Nutritional Studies on *Desulfovibrio desulfuricans* using Chemically Defined Media

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(Received 17 September 1962)

SUMMARY

A non-precipitating chemically defined medium containing lactate, sulphate and other inorganic salts supported repeated subculture of *Desulfovibrio desulfuricans*, strain Hildenborough; yields of bacteria were comparable with those obtained in media containing yeast extract or peptone. Addition of yeast extract, amino acid mixtures or ATP to the defined medium increased the crop. Growth on other organic substances was poorer than on lactate; amino acids were less efficient nitrogen sources than ammonia. Pyruvate served as electron acceptor for hydrogen uptake by resting organisms but did not support growth in sulphate-free medium.

INTRODUCTION

Numerous strains of sulphate-reducing bacteria representing three or more species have now been isolated, yet in relatively few cases have critical studies of their growth and nutrition been made. Butlin, Adams & Thomas (1949) developed a complex medium (medium C) giving rapid and abundant growth of unnamed strains of sulphate reducers, and demonstrated an iron requirement. Autotrophic growth in a chemically defined medium was claimed. Postgate (1951, 1953) obtained poor growth of *Desulfovibrio desulfuricans* strain Hildenborough in a simple glucose ammonia medium incorporating cysteine; the yield was improved somewhat by adding serine, ornithine and isoleucine, and markedly by adding yeast extract. Cysteine was shown by Grossman & Postgate (1953) to act by poising the redox potential and to be replaceable by Na₂S. A peptone yeast-extractmedium supporting massive growth of strain Hildenborough was devised (Postgate, 1951). This strain, one of those used by Butlin *et al.* (1949), was recently proved incapable of autotrophic growth (Mechalas & Rittenberg, 1960; Postgate, 1960), whereas an unnamed strain was stated by Sorokin (1960) to grow autotrophically in a defined medium. Autotrophic incorporation of CO₂ by strain Sylt 3 in presence of yeast extract was claimed by Stüven (1960). Grossman & Postgate (1955), who reviewed the earlier work on utilization of carbon sources, reported growth of strain El Agheila Z on malate, succinate, lactate or pyruvate in a medium containing yeast extract. Carbon sources were also reviewed by Postgate (1959), who pointed out that an unfavourable $E_h$ value, and marginal growth on impurities, may give rise to conflicting observations. Stimulatory effects in defined media by biotin (Wikén & Ghose, 1954) and ATP (Kadota & Miyoshi, 1960) have been reported. Studies on the inorganic nutrition of marine strains have been made by Kimata, Kadota, Hata & Tajima (1955b) and Hata (1960a, b) with a medium containing desalted
peptone; certain strains were unable to utilize inorganic nitrogen compounds (Kimata, Kadota & Hata, 1955a). Postgate (1956b) found that the halophile El Agheila Z required iron for utilization of sulphate, but not pyruvate, as terminal electron acceptor.

Certain of the contradictory reports on nutrition and metabolism of the sulphate reducers may reflect genuine differences between strains, while others may be attributable to ill-defined conditions of cultivation (e.g. presence of yeast extract or peptone; absence of E₄ control). The work reported here was concerned with the development of a non-precipitating chemically defined medium and its use in nutritional studies on Hildenborough, a strain on which several physiological and biochemical studies have been reported.

METHODS

Organism. Desulfovibrio desulfuricans strain Hildenborough (National Collection of Industrial Bacteria, NCIB, no. 8308), purified by Postgate (1953) from an impure culture obtained from Wealden clay at Hildenborough, Kent, was stored in a lyophilized condition. Frequent tests were made on subcultures for anaerobic (Postgate, 1953) and aerobic contaminants.

Preparation of media. All glassware was cleaned in chromic acid (potassium dichromate + sulphuric acid). Media were prepared from analytical (Analar) grade chemicals where available, and Pyrex-distilled water. ATP was obtained from Sigma Chemical Co. (St Louis, U.S.A.). Chemicals of doubtful purity were purified by recrystallization. Lactic acid and yeast extract were incorporated into media before autoclaving at 115° for 20 min.: the fall in pH during autoclaving indicated dissociation of the dimer of lactic acid. All other organic substances were sterilized separately in solution by passing through sterile Oxoid membrane filter disks, grade AP, 4 cm. diam. (Oxoid Ltd., London E.C. 4), supported in glass Seitz-type filters. Solutions of ferrous salts formed precipitates on autoclaving or filtering through grade EKS Seitz filter pads. Membrane-filtered solutions in 2.5 mM-H₂SO₄ showed no precipitation during storage for two months at 3°: stock solutions of FeSO₄ were therefore prepared in this manner. In all cases media were adjusted to pH 7.2-7.4 with sterile 15% (w/v) NaOH solution after autoclaving, using bromothymol blue indicator.

Cultivation. Lyophilized bacteria were inoculated initially into the medium C of Butlin et al. (1949); thereafter subcultures were made weekly into the experimental defined medium and growth experiments inoculated from these stock cultures. During later work stock cultures were maintained in the final version of this medium (the ‘standard medium’, the development of which is described in Results): lactic acid, 100 mM; KH₂PO₄, 2.5 mM; NH₄Cl, 10 mM; Na₂SO₄, 50 mM; CaCl₂, 0.5 mM; MgSO₄·7H₂O, 0.25 mM; trace elements B, Co, Cu, Mn, Mo, Zn each 0.05 mg/l. The medium was adjusted to about pH 6-5 with NaOH and autoclaved. When cool, sterile FeSO₄·7H₂O solution was added to 25 μM and the medium adjusted to pH 7.2-7.4.

Both stock and experimental cultures were grown in Pyrex test tubes plugged with cotton-wool and containing 10 ml. medium. Cultures were incubated at 30° in McIntosh & Fildes’ anaerobic jars under an atmosphere of 99% H₂ + 1% CO₂ (v/v). Except as indicated below, 1.0 mM-Na₂S was added immediately before
inoculation to poise the redox potential at about −300 mV. Triplicate cultures were used in all experimental treatments.

*Estimation of growth.* Attempts to observe early exponential growth in studies on nutrient concentrations by haemacytometer counts and optical density readings gave meaningless results. Dry wt. determinations on large samples withdrawn from 500 ml. cultures gave somewhat better results (typical standard deviations of a single determination on each of three cultures were 73.5% after 24 hr., 35.8% after 48 hr., 21.4% after 72 hr., and 6.0% after 96 hr. of incubation); however, this method was unsuitable for comparing growth in large numbers of cultures. Late exponential and post-exponential growth were therefore estimated by the optical density method described by Postgate (1951) and, when required, equivalent values for dry wt. organism/ml. suspension were read from a calibration curve. Na₃S was omitted from these cultures owing to its darkening effect, and neither cysteine nor ascorbic acid was added since they might serve as nutrients. Instead, a relatively large inoculum (0.2 ml.) from a 4-day-old stock culture was used: this was of sufficient size to prevent failures of growth while not giving excessive carry-over of nutrients, and the maximum standard deviation of the results obtained on days 3–5 (Figs. 1–3) was 7.9%.

Growth in tubes containing Na₃S was estimated by eye when it appeared to be complete; cultures showing doubtful growth were examined microscopically.

*Manometry.* Measurement of hydrogenase activity was made at 32° following the procedure of Littlewood & Postgate (1956) except that 15% (w/v) KOH was used as absorbent for H₂S and CO₂.

**RESULTS**

*Development of the 'standard medium'*

*A non-precipitating defined medium.* Medium C of Butlin et al. (1949) had the disadvantages for our purposes that it formed a precipitate when autoclaved, thus losing an uncertain proportion of nutrients (including Fe²⁺: Dr J. R. Postgate, personal communication), it contained yeast extract, and sodium lactate which is not available in Analar grade, and its content of lactate and sulphate were not optimal (Dr J. R. Postgate, personal communication). The medium was modified in the following ways. The major nutrients were supplied as Analar reagents and their concentrations adjusted to molarities for convenience. Thus 50 mM-lactic acid, 2.5 mM-KH₂PO₄, 0.5 mM-CaCl₂ and 25 mM-Na₂SO₄ were used. 10 mM-NH₄Cl was tentatively supplied as sole nitrogen source despite the findings of Kimata et al. (1955a) that certain strains of *Desulfovibrio desulfuricans* cannot utilize ammonia. To minimize precipitation the divalent cation content was decreased by using 0.25 mM-MgSO₄.·7H₂O: there appear to be no published results which justify the use of 8–13 mg.-ion Mg²⁺/l. in medium C. In addition, an arbitrary trace element mixture was incorporated giving a final concentration of 0.05 mg./l. each of B, Co, Cu, Mn, Mo and Zn. This medium, adjusted to about pH 6.5 with NaOH and autoclaved at 115° for 20 min., did not precipitate. After addition of sterile FeSO₄ solution to 25 μM and adjustment to pH 7.2, the medium supported growth of strain Hildenborough through serial subculture, giving crops equivalent to about 380 μg. dry wt./ml. suspension.
Effect of concentration of the defined medium. To determine whether the over-all concentration of the defined medium was optimal, growth was observed at $\frac{1}{2}$, 1, 2 and 4 times the original concentration. The results (Fig. 1) show that medium $\times 4$ was inhibitory, while growth in medium $\times 2$, though eventually outstripping that

![Fig. 1](image)

Fig. 1. Growth of *D. desulfuricans* strain Hildenborough in the experimental defined medium at original concentration (○) and at $\frac{1}{2}$ (×), 2 (▲) and 4 (▼) times this concentration.

![Fig. 2](image)

Fig. 2. Effect on growth of *D. desulfuricans* strain Hildenborough of decreasing one component of the defined medium by one-half. ○ = full medium, × = $\frac{1}{2}$-FeSO$_4$, ▲ = $\frac{1}{2}$-lactate, ▼ = $\frac{1}{2}$Na$_2$SO$_4$. 

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in \( \times 1 \), was slower, suggesting an unfavourably high osmotic pressure or a toxic level of a nutrient. It thus appeared that growth might be more effectively improved by an increase in the concentration of one or two nutrients.

**Effect of concentration of individual components.** In the experiment illustrated in Fig. 2 one of the major nutrients (lactate, \( \text{KH}_2\text{PO}_4 \), \( \text{NH}_4\text{Cl} \), \( \text{MgSO}_4 \), \( \text{CaCl}_2 \), \( \text{Na}_2\text{SO}_4 \) or \( \text{FeSO}_4 \)) was decreased to one-half of its original concentration, all others being present at full concentration. For the sake of clarity the growth curves for \( \frac{1}{2}\text{KH}_2\text{PO}_4 \), \( \frac{1}{2}\text{NH}_4\text{Cl} \), \( \frac{1}{2}\text{MgSO}_4 \) and \( \frac{1}{2}\text{CaCl}_2 \) are omitted from the figure; they were all very close to that for full medium. The results suggested that growth in the full medium was limited by both lactate and sulphate. Thus the lactate and sulphate concentrations were separately increased \( 1 \frac{1}{2} \times, 2 \times \) and \( 3 \times \) in all combinations. It appeared that \( 2 \times \text{lactate} + 2 \times \text{sulphate} \) produced the most rapid growth; \( 3 \times \text{lactate} + 2 \times \text{sulphate} \) gave a slightly greater crop but a much lower growth rate. The medium with \( 2 \times \text{lactate} + 2 \times \text{sulphate} \) was therefore selected as the standard medium for all further experimentation with the Hildenborough strain and for maintenance of stock cultures. Details of preparation are given in Methods.

Yields equivalent to about 480 \( \mu \)g. dry wt. bacteria/ml. suspension are obtained in the standard medium. Soon after the onset of the stationary phase the number of bacteria remaining in suspension appears from microscopic examination greatly to decrease, while mucin containing large numbers of embedded organisms is precipitated, so that optical density determinations become misleading.
Perpetuation of strain Hildenborough in the standard medium. The organism has been serially subcultured more than 50 times in the standard medium without noticeable decrease in rate or abundance of growth. Inoculations of 2 μl. (equivalent to about 0·8 μg. dry wt. organism or a loopful of culture) from young cultures into 50 ml. medium have always in our experience produced normal growth provided that Na₂S, cysteine or ascorbic acid is incorporated. Lyophilized organisms originally grown in the medium can be successfully revived in it.

Effect of nitrogenous supplements in the standard medium

Typical growth in the standard medium and the original defined medium is shown in Fig. 3 which also illustrates the stimulatory effect of adding 1·0 g./l. Difco yeast extract (the concentration in medium C) to the standard medium. Since the lactate and ammonia concentrations in the standard medium were considered to be optimal, it was inferred that certain substances present in yeast extract might constitute more acceptable carbon or nitrogen sources, or serve as growth factors. The effects of the following supplements were therefore examined: Postgate’s (1951) mixture of serine + ornithine + isoleucine; the mixture of 18 amino acids employed by Kadota & Miyoshi (1960) as a nitrogen source; 18 amino acids + 0·1 g. ATP/l. (Kadota & Miyoshi, 1960); ATP alone. Table 1 shows that ATP, as well as mixtures of metabolites, markedly improve growth in the standard medium.

Table 1. Maximum growth of Desulfovibrio desulfuricans strain Hildenborough in standard medium with various nitrogenous supplements

<table>
<thead>
<tr>
<th>Supplement</th>
<th>Growth (μg. dry wt. bacteria/ml.)</th>
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<tr>
<td>None</td>
<td>470</td>
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<tr>
<td>Serine + ornithine + isoleucine (each 1·0 g./l.)</td>
<td>500</td>
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<tr>
<td>18 amino acids</td>
<td>620</td>
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<tr>
<td>18 amino acids + ATP 0·1 g./l.)</td>
<td>690</td>
</tr>
<tr>
<td>ATP (0·1 g./l.)</td>
<td>680</td>
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<tr>
<td>Yeast extract (1·0 g./l.)</td>
<td>560</td>
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Tests of alternative nutrients and electron acceptors

Carbon sources. Substrates to be tested as carbon sources were added to lactate-free standard medium so as to give 150 mg.-atom C/l., no allowance being made for the partial unavailability of DL-forms. Difco yeast extract was tested at 1·0 g./l. Toxicity tests (tubes containing lactate + substrate under trial) were also inoculated; a few amino acids proved inhibitory and their concentration was subsequently halved. Inoculation was with bacteria washed in C-free medium. Despite addition of Na₂S, only pyruvate, the monohydric alcohols and a few sugars supported growth. A ‘training’ method was therefore tried, in which the media under test were initially supplemented with lactate. Subcultures were made into media without a lactate supplement, and substrates were considered to be utilized when growth continued thereafter at an apparently unimpaired rate through five subcultures. In this way growth was obtained in several more substrates as shown in Table 2, though lags
Nutritional studies on D. desulfuricans of up to 5 weeks occurred in some cases before growth appeared in the first subculture into lactate-free medium, and growth was only marginal (denoted ±) in certain others. The first attempt to ‘train’ the organism to utilize glycerol failed, though at a second attempt fairly heavy growth occurred. No substrate supported such abundant growth as lactate.

Table 2. Utilization of carbon sources by Desulfovibrio desulfuricans strain Hildenborough

Substrates were at a concentration giving 150 mg.-atom C/l. unless otherwise stated.

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Glucose</th>
<th>Fructose</th>
<th>Lactose</th>
<th>Lactose</th>
<th>Sucrose</th>
<th>Methanol</th>
<th>Ethanol</th>
<th>Propanol</th>
<th>Glycerol</th>
<th>Acetate</th>
<th>Citrate</th>
<th>Lactate</th>
<th>Pyruvate</th>
<th>Oxaloacetate</th>
<th>Malate</th>
<th>Fumarate</th>
<th>Succinate</th>
<th>Succinamide</th>
<th>DL-alanine</th>
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<td>L-aspartic acid</td>
<td>L-asparagine</td>
<td>L-citrulline</td>
<td>L-cysteine HCl</td>
<td>L-cystine</td>
<td>L-glutamic acid</td>
<td>L-glutamine</td>
<td>Glycine</td>
<td>L-histidine HCl</td>
<td>DL-isoleucine*</td>
<td>L-leucine</td>
<td>L-lysine HCl</td>
<td>DL-methionine*</td>
<td>L-phenylalanine*</td>
<td>L-proline</td>
<td>DL-serine</td>
<td>DL-threonine*</td>
<td>DL-tryptophane</td>
<td>DL-valine</td>
<td>Yeast extract (1·0 g./l.)</td>
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Nitrogen sources. The mixture of 18 amino acids as prescribed by Kadota & Miyoshi (1960), and the individual acids at 5 mg.-atom N/l., were tested as nitrogen sources in ammonia-free standard medium; asparagine, citrulline and cysteine were also tested. Growth in the mixture was about as abundant as in 5 mm-NH₄Cl, while all the individual amino acids except cysteine, glycine, histidine, methionine and tyrosine supported slight growth.

Organic terminal electron acceptors. Pyruvate and fumarate at non-toxic concentration (50 mm), in standard medium without Na₂SO₄ and with one-half of the normal MgSO₄ concentration, were tested for ability to serve as terminal electron acceptors. No growth occurred, though the abundant growth of Escherichia coli (NCIB, no. 7271) obtained in these media under aerobic conditions indicated that the sulphate concentration was sufficient for assimilatory metabolism. Manometric experiments showed that washed organisms in non-nutrient buffer absorbed hydrogen for the reduction of 15 mm-sodium pyruvate, \(-Q_{H_2} = 28.5 \mu l./mg.\) dry wt./hr. or about one-tenth of the value obtained with equimolar sulphate as electron acceptor. No detectable hydrogen uptake occurred with sodium fumarate.
DISCUSSION

Kadota & Miyoshi (1960) reported that they were unable to grow certain sulphate-reducing bacteria in defined media unless abnormally large inocula were used or organic substances such as peptone, yeast extract or mixtures of amino acids added. For 'normal' growth Desulfovibrio desulfuricans strain Hildenborough required, in a basal ammonia-free medium containing lactate, ATP and mineral salts, the addition of peptones or 18 amino acids: growth was greatly diminished when five of the amino acids were omitted. The effect of adding an ammonium salt was not studied. Our experiments have shown that ammonia is a more acceptable nitrogen source for strain Hildenborough than organic nitrogen compounds. In this respect the organism appears to differ from the marine strains examined by Kimata et al. (1955a) which were unable to utilize ammonium salts. A lactate medium of suitable composition, containing no other organic substance and with ammonia as sole nitrogen source, supports vigorous growth through indefinite subculture and thus appears to be a complete medium for the Hildenborough strain. Small transfers (2 µl. into 50 ml., the smallest tested by us) grow normally in such a medium poised at a suitable redox potential. The addition of mixtures of metabolites such as yeast extract or an amino acid mixture to the standard defined medium stimulates growth, despite their being poor carbon sources. Nevertheless, the crop in the standard medium without any organic addition (about 480 µg. dry wt. organism/ml. suspension) compares favourably with the value of 800 µg./ml. in medium C and 500–520 µg./ml. in medium C with additional lactate and sulphate (Dr J. R. Postgate, personal communication). The growth of Hildenborough, like that of the halophile Maizuru 1 (Kadota & Miyoshi, 1960) is stimulated by ATP in defined media.

The survey of substrate utilization by strain Hildenborough presented difficulty. Postgate (1959) drew attention to the necessity of poising the redox potential especially of media to which the organism is unaccustomed; even so, in our experience, cases of apparent utilization by the 'training' technique are not always repeatable, or growth dies out after a few subcultures. Growth on most carbon sources is extremely scanty, and may take several weeks to appear, so that marginal growth on impurities cannot always be ruled out. Thus utilization of substrates—other than acetate, the utilization of which by a sulphate reducer would be highly interesting (Selwyn & Postgate, 1959)—does not promise to be a useful method of characterizing these organisms. The growth obtained with 1·0 g. yeast extract/l. as carbon source (up to 120 µg. dry wt./ml.) indicates that reports of substrate utilization or 'autotrophic' growth by Desulfovibrio desulfuricans in media incorporating yeast extract should be viewed with caution. Several strains, chiefly halophilic, are able to dismutate pyruvate in sulphate-free media (Postgate, 1952, 1956a). The Hildenborough strain has been shown to be incapable of growth by this means, in keeping with Postgate's (1952) findings. Neither has it the ability possessed by certain marine strains (Sisler & ZoBell, 1951; Grossman & Postgate, 1955) to reduce fumarate with hydrogen. There are thus some grounds for supposing that the Hildenborough strain may differ from marine strains nutritionally as well as in the ways described by Ochynski & Postgate (1968).
Nutritional studies on D. desulfuricans

We are indebted to Dr J. R. Postgate, Dr G. H. Booth and our colleagues in the Microbiology Section for valuable discussions. The paper is published by permission of the Director, National Chemical Laboratory.

REFERENCES


