A Defined Medium for Mycoplasma Strain Y

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(Accepted for publication 13 May 1969)

SUMMARY

In a defined medium for Mycoplasma strain Y, a mixture of the diacetoxysuccinoyl esters of monoolein and monopalmitin replaced unesterified fatty acids and serum protein fractions, enabling the minimal growth requirements to be determined. Other Mycoplasma strains did not grow in media in which these esters replaced fatty acids and protein fractions.

INTRODUCTION

Good growth of Mycoplasma strain Y was obtained in a medium containing defatted bovine serum albumin to bind fatty acids and a heat-stable serum protein fraction to disperse cholesterol (medium C2, Rodwell, 1969). The amino acid, peptide and growth factor requirements of strain Y were determined in a protein free medium (medium D, Rodwell, 1967) in which fatty acids were added in growth limiting concentrations. Growth in medium D was poor and might have been limited by nutrients other than fatty acids, while the protein supplements might have contributed unrecognised nutrients in medium C2. A completely defined medium giving good growth of strain Y is described in this paper. Some observations on the nutrition of other strains are also reported.

METHODS

Organisms. Strain Y, isolated from a goat (Laws, 1956), resembles Mycoplasma mycoides in its nutrition and metabolism (Rodwell, 1960, 1967), but differs in that its growth is not improved by aeration. The following strains were also used: v5, GLADYSDALE and KH3J of M. mycoides; the CHU strain of M. mycoides var. capri; strains N29 and L2917, isolated from bovine arthritis, and s6 (M. gallisepticum).

Growth assays. These were performed as described previously (Rodwell, 1969). Cultures of strain Y were incubated vertically without agitation at 37°; cultures of the other strains were incubated in the same way and also in an inclined rack which was rotated at 10 rev./min. (rotated tube cultures).

Growth was measured by turbidity at 660 m\(\mu\); by the incorporation of \(^{3}\text{H}\) thymidine (Rodwell, 1969), or by protein determination, by the method of Lowry, Rosebrough, Farr & Randall (1951) with crystalline bovine serum albumin dried to constant weight as standard, in cells washed in 0.25 M-sodium chloride + 0.02 M-sodium phosphate (pH 7.4) + 0.01 M-magnesium sulphate.

In some cases, an accumulation of pyruvate during growth suggested a defect in the pyruvate oxidase system, and cells were then examined manometrically for pyru-
vate oxidase activity. The preparation of cell suspensions, and amino acid supplement for manometric experiments, were described previously (Rodwell, 1967).

Media. Media C, C1 and C2 were as described (Rodwell, 1967, 1969). Medium C3 had the same composition as medium C except that the concentrations of glycerol and of 'fatty acid poor' BSA were increased to 0·02 M and 2·7 g./l. respectively, that of sodium phosphate was decreased to 0·1 M and sodium DL-lactate (0·03 M) and fraction C (0·8 g./l.) were added.

Alanyl peptides were obtained from Yeda Research and Development Co., Rehovoth, Israel. TEM-4T was a product of Hachmeister Co., Pittsburg, Pennsylvania. It had a fatty acid composition of 28 mole palmitic acid, 19 mole stearic acid, 42 mole oleic acid and smaller amounts of palmitoleic and linoleic acids per 100 mole fatty acid. Other medium components were as described previously (Rodwell, 1969).

Synthesis of diacetoxy succinoyl esters of monoglycerides. Monopalmitin was synthesized by reaction of DL-isopropylidene glycerol with palmitoyl chloride (Baer & Fischer, 1945). The product (m.p. 76-76·5°) was chromatographically pure by thin layer chromatography (t.l.c.). Monoolein was synthesized by direct esterification of glycerol (Hartman, 1957). The product (m.p. 33·5-34·5°) contained no diglyceride or other impurities detectable by t.l.c.

Reaction of diacetyl D-tartaric acid anhydride (m.p. 133·4°) with equimolar amounts of the monoglycerides at 110° in the absence of a cosolvent gave a complex mixture of products. Reaction in boiling chloroform gave a less complex mixture, while in the presence of pyridine it proceeded rapidly and gave only two major products. A solution of monoolein (2 mmoles) in 10 ml. dry chloroform was concentrated by distillation to 5 ml. Diacetyl tartaric acid anhydride (2·1 mmoles) and dry pyridine (4 mmoles) were then added. After 30 min. at 65°, a thin layer chromatogram—solvent chloroform + methanol + acetic acid + water (84+15+5+2 by vol.)—of the reaction mixture showed, in decreasing order of mobility, a small amount of unreacted monoglyceride, traces of unknown products, a very heavy spot (fraction OT/1) and a smaller amount of another product (fraction OT/2). Fractions OT/1 and OT/2 were purified by chromatography on a column of silicic acid. The column (40 g. silicic acid) was prepared and washed in chloroform + methanol (100+2 v/v), and the sample (1 g. reaction products) applied in the same solvent. The components were eluted with 600 ml. chloroform + methanol (100+2) which eluted unreacted monoolein, traces of unknown products and fraction OT/1; and 300 ml. chloroform + methanol (100+5) which eluted fraction OT/2. Fractions (25 ml.) were collected and examined by t.l.c., and those containing the pure components OT/1 and OT/2 were combined. The residues after evaporation of solvents were dissolved in methanol, and pyridine was removed by stirring with an excess of Dowex 50 (H+) ion exchange resin washed and suspended in methanol. The yields of fractions OT/1 and OT/2 were about 50 and 25 % respectively of the starting materials. The corresponding monopalmitin derivatives, which migrated like OT/1 and OT/2 on t.l.c. plates, were prepared in the same way (fractions PT/1 and PT/2). The m.p. of fractions OT/1 (-8° to -6°) and PT/1 (34° to 36°) were similar to published values for 1-O-(2′(R),3′(R)-diacetoxy succinoyl)-3-O-oleoylglycerol (IIA), and the palmitoyl homologue (IB) respectively (Birnbaum, 1955). It was suspected that fractions OT/2 and PT/2 might be 1,2-di-O-(2′(R), 3′(R)-diacetoxy succinoyl)-3-O-oleoylglycerol(IIA), as and the palmitoyl homologue (IIIB). IIA and 1B would have one titratable carboxyl group and would consume five equivalents of alkali on saponification, while IIA and
IIB would have two titratable carboxyl groups, and consume nine equivalents of alkali on saponification (Fig. 1).

Neutralization equivalents were determined by potentiometric titration of solutions in 50% (v/v) aqueous ethanol with aqueous sodium hydroxide, and saponification equivalents after saponification in aqueous ethanolic sodium hydroxide at 80° for 1 hr. The values found (Table I) agree closely with those calculated for IA and IB, and IIA and IIB.

\[
\begin{align*}
\text{H}_2\text{C}-\text{O}-\text{R}' & \quad \text{H}_2\text{C}-\text{O}-\text{R}' \\
\text{HC}-\text{OH} & \quad \text{HAC}-\text{O}-\text{R}^2 \\
\text{H}_2\text{C}-\text{O}-\text{R}^2 & \quad \text{H}_2\text{C}-\text{O}-\text{R}^2 \\
\text{IA} & \text{R}'=\text{oleoyl} \quad \text{IIB} & \text{R}'=\text{palmitoyl} \\
\text{IIA} & \text{R}'=\text{oleoyl} \quad \text{IIB} & \text{R}'=\text{palmitoyl}
\end{align*}
\]

Fig. 1. Structures of diacetoxysuccinoyl esters of monoglycerides.

Table 1. Neutralization and saponification equivalents of diacetoxysuccinoyl esters of monoolein and monopalmitin

<table>
<thead>
<tr>
<th>Fraction</th>
<th>Compound</th>
<th>Neutn. equiv.</th>
<th>Sapon. equiv.</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td>Calc. Found</td>
<td>Calc. Found</td>
</tr>
<tr>
<td>OT/1</td>
<td>1-O-(2'(R),3'(R)-diacetoxysuccinoyl)-3-O-oleoylglycerol</td>
<td>573 567</td>
<td>114 114.5</td>
</tr>
<tr>
<td>OT/2</td>
<td>1,2-di-O-(2'(R),3'(R)-diacetoxysuccinoyl)-3-O-oleoylglycerol</td>
<td>395 400</td>
<td>87.7 88.7</td>
</tr>
<tr>
<td>PT/1</td>
<td>1-O-(2'(R),3'(R)-diacetoxysuccinoyl)-3-O-palmitoylglycerol</td>
<td>547 552</td>
<td>109 111</td>
</tr>
<tr>
<td>PT/2</td>
<td>1,2-di-O-(2'(R),3'(R)-diacetoxysuccinoyl)-3-O-palmitoylglycerol</td>
<td>381 375</td>
<td>84.8 86.0</td>
</tr>
</tbody>
</table>

RESULTS

Replacement of proteins and fatty acids by diacetoxysuccinoyl esters of monoglycerides

BSA and unesterified fatty acids were replaced for growth of strain Y in medium C1 by TEM-4T, a complex mixture of diacetoxysuccinoyl esters of mono- and diglycerides (Fig. 2). The minimum concentration of TEM-4T required for maximum growth was about 50 mg./l. This amount would provide, after complete hydrolysis, a total of 85 μmoles fatty acids. With this concentration of TEM-4T the concentration of cholesterol had to be increased before any growth occurred. The medium was...
then slightly turbid, and some highly refractile microcrystals formed during subsequent incubation.

TEM-4T was replaced (Fig. 2) by an equimolar mixture of IA and IB and also by a mixture of IIA and IIB. The concentrations of the mixtures, and the proportions of these to cholesterol, were varied: the results shown in Fig. 2 are for the optimum concentrations. Growth with the synthetic homologues was not quite as good as with TEM-4T. They appeared to be more toxic in that the growth-inhibitory concentration of the mixtures was a little lower than with TEM-4T, but the growth was almost as good as with TEM-4T, and the results obtained by turbidity measurement were confirmed by thymidine incorporation. The yields of cell protein at maximum turbidity were 0.20, 0.18 and 0.16 mg./ml. culture for media containing TEM-4T, IA + IB, and IIA + IIB respectively.

Growth of strain Y was obtained also when aqueous dispersions of monoolein and monopalmitin were substituted for TEM-4T. Media with these were very turbid and growth had to be measured by thymidine incorporation. Growth was slower and the amount of thymidine incorporated was about 60% of that in a parallel culture in TEM-4T medium.

Nutritional requirements of strain Y

Since it is unlikely that TEM-4T would be contaminated with amino acids and water soluble growth factors, requirements for these were in most cases determined in a medium E in which it replaced fatty acids and protein fractions.

Vitamins and coenzymes. Requirements for thiamine, riboflavin and nicotinamide (or nicotinic acid) in a partially defined medium were described previously. Coenzyme A was essential in the TEM-4T medium and could not be replaced by pantothenate, pantethine or pantetheine. Pyruvate accumulated during growth in a partially defined medium (medium B2, Rodwell, 1963). This medium probably contained growth-limiting amounts of coenzyme A or other utilizable precursor contaminating one or other of the undefined components. Mycoplasmas grown in it lacked pyruvate oxidase activity, but oxidised pyruvate normally in the presence of coenzyme A (Fig. 3). The abrupt increase in the rate of oxygen uptake with pyruvate as substrate (Fig. 3) did not indicate a change in the rate of pyruvate oxidation, but was due to the accumulation, and subsequent oxidation, of lactate (Rodwell, 1967). With growth-limiting amounts of coenzyme A, growth was accompanied by cellular lysis, presumably because of a failure to incorporate fatty acids.

Growth stimulation by α-lipoic acid in a partially defined medium was found previously (Rodwell, 1960). The effect of α-lipoic acid on growth in medium C2 is shown in Fig. 4. While it stimulated growth in primary culture, it had little or no effect on growth in subculture in the same medium. The pyruvate oxidase activity of mycoplasmas harvested after 20 hr incubation from replicate subcultures in media with and without α-lipoic acid was examined (Fig. 5). Mycoplasmas from the deficient culture had no pyruvate oxidase activity, but oxidized it normally in the presence of α-lipoic acid. There was no growth response to α-lipoic acid in medium E either in primary culture, or in subculture. Thus it appears that the functioning of the pyruvate oxidase system was not essential for growth of this strain—at least in non-aerated cultures.

Pyridoxin or other vitamin B6 derivatives were not required in the TEM-4T medium.
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in which alanine was provided by L-alanyl-L-alanyl-L-alanine (L-ala₃). Strain Y grew slowly without alanine or alanyl peptides (see below) and, under conditions where growth was made dependent on alanine synthesis, i.e. in the absence of alanine or of alanyl peptides, growth was stimulated by pyridoxamine (Fig. 6). Growth under these conditions was no better when other derivatives of pyridoxin, including pyridoxal phosphate, were substituted for pyridoxamine. Requirements for biotin, folinic acid or vitamin B₁₂ were not found in any medium.

Amino acid requirements. Absolute requirements for all amino acids except glutamic acid, aspartic acid and cystine were reported previously in medium D (Rodwell, 1967). This was confirmed in the TEM-4T medium in which alanine was provided by the tripeptide (L-ala₃). In contrast to the results obtained earlier, there was good growth when L-alanine at a concentration of 1 mM was substituted for alanyl peptides, and slow growth in the absence of alanine or alanyl peptides (Fig. 6). Higher concentrations of L-alanine (8 mM) inhibited growth, and this inhibition was partly reversed by increasing the glycine concentration. Growth with the tripeptide (L-ala₃) and tetrapeptide (L-ala₄) was a little faster than with the free amino acid, but the dipeptide L-alanyl alanine inhibited growth.
Minimal medium. The inorganic requirements, and those for fatty acids, sterol, spermine, nucleic acid precursors and glycerol have been discussed previously and were not considered further. The composition of the minimal medium giving the best growth of strain Y is listed in Table 2.

Fig. 4. Effect of α-lipoic acid on growth of Mycoplasma strain Y. Cultures in medium C2. ■, With DL-α-lipoic acid (0.2 μg ml.); ○, no α-lipoic acid.

Fig. 5. Effect of α-lipoic acid deficiency on pyruvate oxidase activity of Mycoplasma strain Y. Mycoplasmas were from subcultures for 20 hr in medium C2 with α-lipoic acid (α-lipoic acid cells), and without α-lipoic acid (deficient cells)—see Fig. 4. Manometer flasks contained: Na₂HPO₄+ KH₂PO₄, pH 7.4, 200 μmoles; sodium pyruvate, 20 μmoles; DL-α-lipoic acid, 10 μg.; amino acids 3.5 mg.; α-lipoic acid cells, 2.2 mg.; deficient cells, 3.8 mg.; KOH in centre well; fluid volume 2.5 ml. ■, Deficient cells with α-lipoic acid; ×, deficient cells without α-lipoic acid; ○, deficient cells with α-lipoic acid but no pyruvate; □, α-lipoic acid cells with α-lipoic acid; △, α-lipoic acid cells without α-lipoic acid.

Nutrition of other strains of Mycoplasma

Mycoplasma mycoides strain v5 did not grow in media in which TEM-4T replaced fatty acids and the protein fractions, even with the addition of a pronase digest of BSA to provide peptides and other growth factors. Several strains, including v5, were tested for growth in medium C2; none grew well, or in rotated tube cultures in medium C2 modified for aerobic growth conditions by increasing the glycerol concentration and adding lactate. Two strains of M. mycoides (v5 and GLADYSDALE), the bovine arthritis strains (N29 and L2917), and a strain of M. mycoides var. capri (CHU) grew well in rotated tube cultures in medium C3, the growth rate and yield of organisms being about the same as that obtained with strain Y in medium C2. Mycoplasma
Table 2. Composition of defined medium E for growth of *Mycoplasma strain Y*

<table>
<thead>
<tr>
<th>Additions</th>
<th>Concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Na₂HPO₄</td>
<td>0.14 M</td>
</tr>
<tr>
<td>KCl</td>
<td>0.01 M</td>
</tr>
<tr>
<td>MgSO₄</td>
<td>0.004 M</td>
</tr>
<tr>
<td>Glucose*</td>
<td>0.1 mM</td>
</tr>
<tr>
<td>Spermine</td>
<td>0.2 mM</td>
</tr>
<tr>
<td>Cholesterol†</td>
<td>5.0 mM</td>
</tr>
<tr>
<td>Glycerol</td>
<td>0.04 mM</td>
</tr>
<tr>
<td>Diacetoxyuccinoyl esters of monoolein and monopalmitin (or TEM-4T)†</td>
<td>(or 50 mg./l. of TEM-4T)</td>
</tr>
<tr>
<td>Adenine, guanine, uracil</td>
<td>10.0 mg/l. each</td>
</tr>
<tr>
<td>Thymine</td>
<td>50 mg/l.</td>
</tr>
<tr>
<td>Coenzyme A³, riboflavin*, nicotinamide, thiamine</td>
<td>1.0 mg/l. each</td>
</tr>
<tr>
<td>DL-α-lipoic acid</td>
<td>1.0 mg/l. each</td>
</tr>
<tr>
<td>L-asparagine, L-glutamine*, L-arginine, L-lysine, L-histidine, L-leucine, L-isoleucine, L-proline</td>
<td>1.0 mm each</td>
</tr>
<tr>
<td>DL-phenylalanine, DL-valine, DL-methionine, DL-serine, DL-threonine, DL-tryptophan, glycine</td>
<td>2.0 mm each</td>
</tr>
<tr>
<td>L-cysteine*</td>
<td>1.3 mM</td>
</tr>
<tr>
<td>L-alanyl-L-α-1-1anine</td>
<td>0.3 mM</td>
</tr>
</tbody>
</table>

* Added from sterile solutions to previously autoclaved medium. Glutamine and cysteine solutions were freshly prepared.

† Aqueous dispersions prepared by the addition of solutions in ethanol to stirred water at 65° for cholesterol, or at 20° for TEM-4T and homologues.

‡ Added before autoclaving from a 0.1 mg./ml. solution containing dithiothreitol, 1.0 mg./ml. in phosphate buffer, pH 7.0.

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**Fig. 6. Effect of L-alanine and L-alanyl peptides on growth of Mycoplasma strain Y.**

Medium E contained TEM-4T and L-alanine, peptides and pyridoxamine as indicated. ×, without L-ala, peptides or pyridoxamine; ▽, without L-ala or peptides; □, L-ala, 1 mm; ■, L-ala, 8 mm; △, L-ala₂, 1 mm in ala; ○, L-ala₃, 1 mm in ala; ▲, L-ala₄, 1 mm in ala.
mycoides strain KHj grew poorly, while the s6 strain of M. gallisepticum gave no growth. The strains of M. mycoides and the bovine arthritis strains grew in medium C3 as extremely long, branched filaments.

Glycerol was essential for the Mycoplasma mycoides strains and for the strain of M. mycoides var. capri. Exogenous glycerol was not essential for the bovine arthritis strains, but they grew poorly without it, and the cells underwent lysis. Requirements for other nutrients were not determined, apart from the observations already reported for strain v5 (Rodwell, 1960).

**DISCUSSION**

The protein supplements might have contributed unrecognized nutrients for growth of strain v in medium C2. To eliminate these it was necessary to substitute fatty acids in a non-toxic, chemically defined form, which would at the same time disperse cholesterol. Mixtures of ‘Tweens’ (polyoxyethylene sorbitan mono esters of fatty acids) were tried without success. Lund & Shorb (1966) described a partially defined medium in which TEM-4T provided fatty acids for growth of Mycoplasma strain J. TEM-4T contains a balanced mixture of saturated and unsaturated fatty acids, and gave good growth of strain v. The concentration required for optimum growth was much greater than in the medium of Lund & Shorb, and the cholesterol concentration had to be increased in proportion. It is possible that an interaction between TEM-4T and cholesterol modifies the surface active properties of the glycerides and disperses cholesterol. TEM-4T was in turn replaced by synthetic homologues, enabling the minimal nutritional requirements to be determined.

TEM-4T and the synthetic homologues do not seem to be completely non-toxic for mycoplasmas. Strain v5 required both BSA (at an increased concentration) and fraction C (or PC) for good growth, and these could not be replaced by TEM-4T. While strain v5, and the other strains which grew well in medium C3, may require unrecognized nutrients (supplied by the protein supplements), it is also possible that TEM-4T is not a suitable source of fatty acids for them.

Other strains might be grown in partially defined media by including more complex mixtures of nucleic acid precursors (such as are required by Mycoplasma laidlawii strain B, Tourtellotte, Morowitz & Kasimer, 1964), by altering the proportions of amino acids, or by adding arginine in high concentration as an energy source for growth of non-fermentative strains having the arginine dihydrolase pathway (Barile, Schimke & Riggs, 1966).

I wish to thank Professor J. M. Swan for advice on the synthesis and nomenclature of the diacetoxy succinoyl esters of monoglycerides, Dr H. Birnbaum for supplying me with a sample of TEM-4T and Mr B. Zwolak for his capable technical help.

**REFERENCES**

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