The Effects of Ribonucleic Acid and Deoxyribonucleic Acid on the Growth of Mycoplasma

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SUMMARY: The nucleic acid requirements of several Mycoplasma organisms were examined in a partially defined medium. The saprophytic *M. laidlawii* strain A did not grow in the basal medium alone (Razin & Knight, 1960a) but did when suitable concentrations of ribonucleic acid (RNA) and of deoxyribonucleic acid (DNA) were added. Too high a concentration of RNA inhibited growth; this inhibition was annulled by increasing the concentration of DNA. Similarly, too high a concentration of DNA inhibited growth, and this inhibition was annulled by RNA. Chemical and enzymic degradations of RNA showed that the growth-promoting effect could be brought about by a ribo-oligonucleotide but not by smaller fragments of the molecule. Similar degradations of DNA showed that the effective moiety was thymidine; thymine was less effective. The degradation of RNA abolished its growth-inhibitory activity. The growth-inhibitory activity of DNA was not affected by its degradation to oligonucleotides, and was only partially diminished by its degradation to nucleotides or nucleosides.

*Mycoplasma laidlawii* strain B grew in the basal medium when DNA alone was added. This nutritional requirement was also satisfied by thymidine, provided that some RNA was also present. The parasitic *Mycoplasma mycoides* var. *capri* resembled *M. laidlawii* strain B in responding to DNA alone, but differed from the saprophytic strains in its complete indifference to high DNA concentrations. Thymidine replaced DNA only to a certain extent when added together with RNA. A growth-promoting effect of DNA was also found with the L-phase of *Streptobacillus moniliformis*; thymidine then replaced DNA completely. The RNA/DNA antagonism was found with all the organisms examined. Possible explanations for this phenomenon are discussed.

Information about the effects of nucleic acids and their fragments on the growth of *Mycoplasma* organisms is very scanty. Edward & Fitzgerald (1952) found that certain bovine strains needed DNA to be added to a complex medium for primary culture; work by Crowther & Knight (1956) indicated that DNA acted here by annulling a growth inhibitory effect caused by the RNA present in the yeast extract used in the complex Edward medium (Edward, 1947). The growth-inhibitory activity of RNA and the activity of DNA in annulling this were destroyed by chemical or enzymic degradation of the polynucleotides to mononucleotides. No explanation of these observations could then be given, as practically nothing was known about the nutritional requirements of the *Mycoplasma* organisms in regard to nucleic acids and their constituent parts. The complex nature of the medium then used prevented analysis of such requirements. The partially defined medium for *Mycoplasma* organisms described by Razin & Knight (1960a) has now been found adequate for a
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closer examination of the effect of nucleic acids on the growth of these organisms. The RNA/DNA antagonism can be demonstrated with this medium; this has permitted further analysis of the phenomenon and a possible interpretation of it.

**METHODS**

*Organisms.* The *Mycoplasma* strains were those previously used (Razin & Knight, 1960a). The stable l-phase of *Streptobacillus moniliformis* was kindly given by Dr E. Klieneberger-Nobel (The Lister Institute for Preventive Medicine, London). *Mycoplasma laidlawii* strain A was used as the main test organism.

*Media.* A modified Edward medium (Butler & Knight, 1960b) was used for keeping stock cultures and for growing the organisms used as inoculum in the nutrition experiments. The same medium, solidified with 2% (w/v) agar, was used for viable counts (Butler & Knight, 1960a).

The nutrition experiments were carried out in the partially chemically defined medium described previously (Razin & Knight, 1960a; Table 1). This medium without added ribonucleic and deoxyribonucleic acids will be referred to as basal medium. For experiments with *Mycoplasma mycoides* var. capri and the l-phase of *Streptobacillus moniliformis* the basal medium was fortified by increasing the serum to 20% (v/v) and adding yeast extract (Oxoid) to 0·05% (w/v).

Pyrex glassware was used in all experiments; before use it was cleaned in chromic + sulphuric acid mixture and thoroughly rinsed in distilled water. The basal medium was dispensed in 0·5 ml. volumes into 6 in. x ½ in. test tubes; the solutions of nucleic acids or degradation products thereof to be tested were added to give a final volume of 10 ml. The nucleic acid solutions were sterilized by passage through sintered-glass filters or by steaming for 20 min. (Merrifield & Dunn, 1950). Sterilization of the nucleic acid degradation products was done by steaming only. For bioassay all solutions of nucleic acid degradation products were made up to known volumes so that their activities could be related to the initial amounts of nucleic acid from which they were made.

*Conditions of growth.* The inoculum used in the experiments was prepared as described previously (Razin & Knight, 1960a). In most experiments the inoculated test tubes were incubated vertically without agitation in air at 37°, but some experiments were done in T-shaped tubes (van Heyningen & Gladstone, 1958) which were rocked in a 37° water bath. Growth was usually estimated after incubation for 96 hr.

*Assessment of growth.* The methods described previously (Razin & Knight, 1960a) were used, generally the acid titration method.

*Chemicals.* Most of the inorganic and organic chemicals were analytical reagents. Ribonucleic acid, Na salt from yeast (Na-RNA) was the product of L. Light and Co. Ltd., Colnbrook, Bucks.; it was further purified by the method of Frisch-Niggemeyer & Reddi (1957). Deoxyribonucleic acid, Na salt from thymus gland (Na-DNA) was from the British Drug Houses Ltd., Poole, Dorset. Other samples of nucleic acids tested were: ribonucleic acid Na salt
from yeast (British Drug Houses Ltd.); deoxyribonucleic acid Na salt (thymus) and deoxyribonucleic acid Na salt (from herring roe; L. Light and Co. Ltd.). The purines, pyrimidines, ribonucleosides, deoxyribonucleosides, ribonucleotides and deoxyribonucleotides used were commercial products. Ribonuclease (RNase, crystallized, bovine pancreas) was from Armour Laboratories, Hampden Park, Eastbourne, Sussex, England. Deoxyribonuclease (DNAse, 25% activity of crystalline) was from L. Light and Co. Ltd. Freeze-dried Russell viper venom was the gift of the Wellcome Research Laboratories, Beckenham, Kent, through Mr H. Proom.

Analytical methods. The enzymic degradations of RNA and DNA were followed by estimating the total acid-soluble phosphorus liberated (McDonald, 1955; Butler, 1955), and by paper chromatography of the reaction mixtures (Markham, 1957). Whatman no. 1 paper and the isopropanol + ammonia system (solvent 8; Markham & Smith, 1952) were used. The spots were detected under ultraviolet irradiation. Commercial purines, pyrimidines, nucleosides and nucleotides served as markers.

All samples taken for bioassay from the enzymic degradation experiments were immediately frozen at $-20^\circ$.

Degradation of ribonucleic acid

Acid hydrolysis to oligonucleotides. This was done by the method of Merrifield & Woolley (1952). A Na-RNA solution (140 mg./ml.) was brought to pH 6-8 by adding $\text{n-NaOH};$ a sample was removed at zero time. An equal volume of concentrated HCl was then added to the remainder of the solution with stirring and the mixture placed in a water bath at 25°. After 1, 3, 7, 15 and 30 min. samples of the mixture were removed, diluted with 50 vol. ice-cold distilled water, neutralized with NaOH, and made to volume for bioassay. A sample of the RNA+6N-HCl mixture, incubated at 25° for 3 min., was dialysed in a cellophan bag (Visking, 0-85 in.) against distilled water for 24 hr. at room temperature. The dialysate and diffusate were tested for biological activity.

Degradation by RNase to oligonucleotides and pyrimidine mononucleotides. One mg. RNAse was added per ml. Na-RNA solution (50 mg./ml.). The mixture was brought to pH 8-0 and incubated at 37°. Samples for bioassay and acid-soluble phosphorus determination (McDonald, 1955) were removed at zero time and after 0-25, 1-5, 3-5, 6, 20 and 30 hr. The reaction mixture was re-adjusted to pH 8-0 when each sample was taken. The acid-soluble fraction of the RNA hydrolysate was obtained by mixing the sample of the reaction mixture, after incubation for 30 hr., with an equal volume of N-HCl. This produced a precipitate in the mixture which was centrifuged at 4°, the supernatant fluid neutralized by adding 6N-NaOH and kept for bioassay. Some of the reaction mixture, after incubation for 30 hr., was dialysed in a cellophan bag (Visking, 0-85 in.) against distilled water for 18 hr. at 4° and then for 6 hr. at room temperature. The dialysate and diffusate were separately collected and tested for biological activity. Paper chromatography of the reaction mixture at the end of the incubation period showed the disappearance of undegraded
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RNA and the appearance of spots whose $R_f$ values corresponded with those of oligo- and mono-nucleotides (Markham, 1957).

Degradation of RNA to nucleoside-5'-phosphates by phosphodiesterase from Russell viper venom. Purification of the phosphodiesterase from Russell viper venom was done by acetone fractionation of the crude venom according to the method of Sinsheimer & Koerner (1952). Purified enzyme preparation (1.5 ml.) was then added to 10 ml. of the RNA + RNase reaction mixture which had been pre-incubated at 37° for 30 hr., then MgCl$_2$·6H$_2$O was added to a final concentration of 0.025 M and the solution brought to pH 9.3 by N-NaOH. Further incubation was then carried out at 37° and samples for bioassay and acid-soluble phosphorus determinations (Butler, 1955) were removed at zero time and after 5.5 and 22 hr. The reaction mixture was readjusted to pH 9.3 at frequent intervals. Paper chromatography of the reaction mixture at the end of the incubation period showed spots with $R_f$ values corresponding to mononucleotides and nucleosides. The presence of the nucleosides may be explained by contamination of the enzyme preparation with phosphomonoesterase which was not completely eliminated by the purification method used (Privat de Garilhe & Laskowski, 1955).

Degradation to nucleosides by Russell viper venom. This was carried out by a modification of the method of Cohn (1957). Na-RNA (400 mg.) was dissolved in 10 ml. 0.01 M-MgCl$_2$·6H$_2$O solution. Freeze-dried Russell viper venom (20 mg.) was added and the mixture brought to pH 9.0 by N-NaOH. Incubation was at 37° and samples for bioassay and acid-soluble and inorganic phosphorus determinations (Butler, 1955) were removed at zero time and after 1, 6 and 24 hr. The mixture was readjusted to pH 9.0 at frequent intervals during incubation. Paper chromatography of a sample taken at the end of the reaction showed the presence of nucleosides only.

Alkaline hydrolysis to nucleoside-2' and -3'-phosphates. Na-RNA (450 mg.) was dissolved in 30 ml. 0.5 N-NaOH and the solution incubated at 25°. Samples were removed at zero time and after incubation for 0.25, 0.5, 1, 2, 4, 7.5 and 27 hr. The samples were neutralized immediately with N-HCl and made to volume for bioassay. Paper chromatography of a sample taken at the end of the reaction showed the complete disappearance of RNA and the appearance of mononucleotides.

Acid hydrolysis to purine bases and pyrimidine nucleotides. This was carried out according to Schmidt (1957), equal volumes of Na-RNA solution (28 mg./ml.) and 8 N-H$_2$SO$_4$ being mixed and placed in a boiling water bath for 1 hr.; the test tube containing the mixture was closed with a glass bulb. After cooling the solution was neutralized by 6 N-NaOH and tested for biological activity.

Degradation of deoxyribonucleic acid

Apurinic acid was prepared by a mild acid treatment of DNA (Tamm, Hodes & Chargaff, 1952).

Degradation by DNase to oligonucleotides. DNase (1 mg./ml.) was added to a 0.025 M-MgSO$_4$·7H$_2$O solution containing 50 mg. Na-DNA/ml. The solution
was adjusted to pH 6.8 and incubated at 37° for 25 hr. Samples for bioassay
and acid-soluble phosphorus determinations (McDonald, 1955) were taken at
zero time and after 2, 6 and 26 hr. The reaction mixture was readjusted to
pH 6.8 when samples were withdrawn. After incubation for 8 hr., more DNase
was added (0.5 mg./ml. solution). Paper chromatography of a sample taken at
the end of incubation showed spots corresponding mainly to oligonucleotides.
Some of the DNA + DNase mixture was taken at the end of the incubation
period and dialysed against distilled water at 4° for 48 hr. The dialysate and
diffusate were tested for biological activity. No precipitate was formed when
an equal volume of n-HCl was added to the DNA + DNase mixture at the end
of the incubation period, indicating that all the DNA was degraded to smaller
fragments (Schmidt, 1957).

Degradation to deoxynucleotides by phosphodiesterase from Russell viper venom.
Phosphodiesterase preparation (1.5 ml.) was added to 5 ml. DNA + DNase
mixture which had been pre-incubated at 37° for 26 hr. The solution was
brought to pH 9.3 by n-NaOH and incubated at 37° for 24 hr. Samples for
bioassay and acid-soluble phosphorus determinations were taken at zero time
and after 6 and 24 hr. The solution was readjusted to pH 9.3 at frequent inter-
vals during incubation. Paper chromatography of samples taken during the
reaction period showed the disappearance of the oligonucleotides and the
appearance of mononucleotides. At the end of the reaction spots correspond-
ing to nucleosides also appeared.

Degradation of DNA to deoxynucleosides by Russell viper venom. To 6 ml.
DNA + DNase mixture, which had been pre-incubated at 37° for 26 hr., freeze-
dried Russell viper venom (10 mg.) was added, the solution brought to pH 9.3
by adding n-NaOH and further incubated at 37°. Samples for bioassay and
acid-soluble and inorganic phosphorus determinations were taken at zero time
and after 6 and 24 hr. During incubation the solution was readjusted to pH 9.3
at frequent intervals. A heavy white precipitate appeared at the end of the
reaction. Paper chromatography showed the complete conversion of the
oligonucleotides to nucleosides.

RESULTS

The effect of RNA and DNA on the growth of Mycoplasma laidlawii strain A
No growth of Mycoplasma laidlawii strain A took place in the basal medium
alone; the addition together of suitable concentrations of RNA and DNA
permitted growth. Too high a concentration of RNA inhibited growth; this
inhibition was overcome by raising the concentration of DNA; and vice versa.
Thus the growth inhibitory effects of RNA and DNA showed a mutual antago-
nism. The organism required RNA in higher concentrations than DNA
(Table 1). No growth took place with 1 µg. Na-RNA/ml.; 5 µg. of Na-RNA/ml.
allowed very poor growth. For optimal growth 50 µg. Na-RNA/ml. were
needed, whereas 1 µg. Na-DNA/ml. was sufficient for suboptimal growth and
maximal growth occurred with 25 µg./ml. (Table 1).

Experiments in which growth was estimated by the viable count technique
showed that the organisms survived in the medium without nucleic acids. Addition of RNA or DNA even in high concentrations did not kill the organisms (Fig. 1). The growth-promoting activities of RNA and DNA are not due to contaminating factors; purified or dialysed samples and nucleic acids from various sources showed the same effects.

Table 1. *The effects of RNA and DNA on the growth of Mycoplasma laidlawii strain A*

Test tubes contained 10 ml. basal medium supplemented with various amounts of Na-RNA and Na-DNA. Initial inoculum was $5 \times 10^4$ viable particles/ml. Acid production was determined after incubation at 37° for 96 hr. The numbers in heavy type indicate the optimal amounts of growth at different RNA and DNA concentrations.

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*The effect of the degradation of RNA on its growth-promoting and DNA-antagonistic activities*

Degradation of RNA to oligonucleotides by treatment with 6N-HCl for 3 min. at 25° (Merrifield & Woolley, 1952) did not significantly affect its growth-promoting activity. These active oligonucleotides passed through the cellophane during dialysis. Further hydrolysis of the oligonucleotides by longer treatment with acid rapidly abolished their growth-promoting activity.

Degradation of RNA by RNAse to oligonucleotides and pyrimidine mononucleotides (Davidson, 1957) decreased its growth-promoting activity. The products had no antagonistic activity against DNA 50 µg./ml. (Fig. 2). Dialysis of the degradation products showed the growth-promoting activity to be concentrated more in the diffusate than in the dialysate. The diffusate enabled even better growth than the undialysed RNA hydrolysate (Fig. 2). The acid-soluble fraction of the hydrolysate, which contains mononucleotides and small oligonucleotides (Schmidt, 1957), had the same activity as the diffusate.

Degradation of the oligonucleotides, obtained by RNAse treatment, to nucleoside-5'-phosphates by Russell viper venom phosphodiesterase or to nucleosides by crude venom (Davidson, 1957) destroyed completely and very quickly their growth-promoting activity.

Alkaline hydrolysis of RNA to nucleoside-2' and 3'-phosphates (Davidson, 1957) abolished its growth-promoting activity very quickly. The mixture of the mononucleotides obtained had no growth-inhibitory activity even when
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added at a final concentration equivalent to 2500 µg. Na-RNA/ml. to the basal medium containing optimal concentrations of Na-RNA and Na-DNA (50 µg./ml. of each). Hydrolysis of RNA to purine bases and pyrimidine nucleotides by 1.5 N-H₂SO₄ at 100° completely abolished its growth-promoting activity.

Mixtures of purines and pyrimidines (adenine, guanine, uracil, cytosine), nucleosides (adenosine, guanosine, uridine, cytidine), nucleosides-3'-phosphates (yeast adenylic acid, guanylic acid, uridylic acid, cytidylic acid) and their-5'-phosphate isomers, were inactive in growth promotion, either alone or with 50 µg. Na-DNA/ml. The purines and pyrimidines were tested in concentrations of 5 and 25 µg./ml. and the nucleosides, and nucleotides in concentrations of 10 and 100 µg./ml.

![Fig. 1](image1.png)

**Fig. 1.** The effect of RNA and DNA on the growth of Mycoplasma laidlawii strain A. 1-Shaped tubes contained 10 ml. basal medium without nucleic acids (O) or supplemented with: 500 µg./ml. Na-RNA (△); 500 µg./ml. Na-DNA (▲); 500 µg./ml. Na-RNA + 500 µg./ml. Na-DNA (●); 50 µg./ml. Na-RNA + 50 µg./ml. Na-DNA (×). The tubes were rocked in a 37° water bath.

![Fig. 2](image2.png)

**Fig. 2.** The effect of degradation of RNA by RNAse on its growth-promoting activity for Mycoplasma laidlawii strain A. RNA was incubated with RNAse at 37° for 80 hr. Part of the resulting hydrolysate was dialysed against distilled water. Test tubes contained 10 ml. basal medium supplemented with 50 µg. Na-DNA/ml. and various concentrations of: undegraded Na-RNA (O); hydrolysed RNA (●); dialysate of the hydrolysate (△); diffusate of the hydrolysate (▲). Initial inoculum 1 x 10⁵ viable particles/ml. Acid production determined after incubation at 37° for 96 hr.

**The effect of the degradation of DNA on its growth-promoting and growth-inhibitory activities**

The mild acid treatment which removes the purine bases from the DNA molecule, giving the apurinic acid (Tamm et al. 1952) did not decrease the growth-promoting activity. High concentrations of apurinic acid showed, like DNA, growth-inhibitory activity.
Degradation of DNA by DNAses to a mixture of oligonucleotides (Davidson, 1957) did not affect its growth-promoting activity and did not remove the growth inhibition shown by high concentrations (Fig. 3). The growth-promoting material in the hydrolysate was found in the diffusate on dialysis, and was not precipitated by 0.5 N HCl. Thus, the growth-promoting activity of DNA is not dependent on the whole polynucleotide molecule.

Further degradation of the deoxyoligonucleotides to deoxynucleotides by Russell viper venom phosphodiesterase and to deoxynucleosides by the crude venom did not destroy the growth-promoting activity (Fig. 3). However, the activity in inhibiting growth at high concentrations was smaller. These results indicated that the growth-promoting effect of DNA resided in a moiety not greater than a deoxynucleoside. The following deoxynucleosides and deoxynucleotides were tested for growth-promoting activity in the presence of 50 µg Na-RNA/ml: deoxyadenosine, deoxyguanosine, deoxyuridine, deoxyctydine, thymidine, deoxyadenylic acid, deoxyctydyllic acid; the pyrimidine thymine was also tested. Only thymidine, and less effectively thymine supported
growth (Fig. 4). The growth response of Mycoplasma laidlawii strain A to thymidine was even better than to DNA. Another phenomenon shown in Fig. 4 is the interference by high concentrations of other deoxynucleosides and deoxynucleotides with thymidine activity.

Thus the growth-promoting effects of RNA and DNA are shown, respectively, by the following moieties, namely, oligonucleotides probably less than tetranucleotides from RNA, and thymidine (thymine) from DNA; these appear to supply the minimal nutrient requirements. Smaller fragments are inactive. However, there still remains to be explained the antagonistic growth inhibitory effects shown by undegraded RNA and DNA.

Possible mechanism of the RNA/DNA antagonism

We chose as a working hypothesis the idea that high concentrations of RNA inhibit growth promoted by DNA by interfering with the enzymic degradation of DNA by the organism, which is a pre-requisite for the utilization of the DNA-thymidine, the real biosynthetic precursor. If this idea is correct then RNA should not have growth-inhibitory effects when the part of DNA which is the essential nutrient requirement of the organisms, i.e. thymidine, is included in the medium. This was found to be the case; Fig. 5 shows that high concentrations of RNA had no inhibitory activity when added to medium containing 10 μg. thymidine/ml. An experiment run simultaneously showed strong growth inhibition by high RNA concentrations when 50 μg. Na-DNA/ml. replaced thymidine in the medium.

However, the growth inhibitory effect of high concentrations of DNA or its degradation products cannot be explained by interference with RNA degradation by the organism, since DNA had the same growth inhibitory effects when added in high concentrations to the mixture of ribo-oligonucleotides (obtained by RNase action) which apparently supply the minimal nutrient requirement of Mycoplasma laidlawii strain A as regards RNA.

The effects of RNA and DNA on the growth of Mycoplasma laidlawii strain B

Table 2 shows that Mycoplasma laidlawii strain B, which is closely related to M. laidlawii strain A, differs from the latter in not requiring RNA. Addition of DNA itself enabled optimal growth when added in sufficient quantities. When the concentration of DNA was suboptimal (5 μg./ml.) the addition of RNA improved growth. Apparently RNA supplies in this case components which are common to it and to DNA. As may be seen from Table 2 high concentrations of DNA inhibited the growth of M. laidlawii strain B; RNA did not overcome this inhibition. RNA showed the DNA-antagonizing effect when added in high quantities to low DNA concentrations, e.g. 5 μg./ml.; higher concentrations of DNA (50 μg./ml.) overcame this RNA inhibitory effect. Thymidine satisfied completely the nutritional requirement of M. laidlawii strain B for DNA, provided RNA was also present. Optimal growth occurred with 10 μg. thymidine/ml. and 50 μg. Na-RNA/ml.; even 1 μg. thymidine/ml. + 50 μg. Na-RNA/ml. supported good growth.
The effects of RNA and DNA on the growth of Mycoplasma mycoides var. capri

In contrast to the saprophytic Mycoplasma strains tested, *M. mycoides* var. *capri* grew very poorly in the basal medium + Na-RNA + Na-DNA (50 μg./ml. each). Increasing the concentration of serum from 10 to 20 % (v/v) and addition of 0.05 % (w/v) yeast extract (fortified medium) enabled good growth to occur. These changes in the composition of the basal medium did not mask the requirements for nucleic acids. Practically no growth of *M. mycoides* var. *capri* took place in this fortified medium without added DNA; optimal growth occurred only with high DNA concentrations. In contrast to the findings with the saprophytic *M. laidlawii* strains A and B, DNA did not show any growth-inhibitory effect in the highest concentration tested with *M. mycoides* var. *capri* (Fig. 6). With this organism thymidine replaced DNA only to a certain extent when added together with RNA. Increasing the concentration of thymidine depressed growth rather than improved it (Fig. 7). No requirement for RNA by *M. mycoides* var. *capri* could be shown in the fortified medium; however, RNA improved growth when added to suboptimal concentrations of DNA. High concentrations of RNA had the DNA-antagonistic effect (Fig. 6).
Table 2. The effects of RNA and DNA on the growth of Mycoplasma laidlawii strain B

Test tubes contained 10 ml. basal medium supplemented with various concentrations of Na-RNA and Na-DNA. Initial inoculum $7 \times 10^6$ viable particles/ml. Acid production determined after incubation at $37^\circ$ for 96 hr.

The numbers in heavy type indicate the optimal amounts of growth at different RNA and DNA concentrations.

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Fig. 7. The effect of thymidine on the growth of Mycoplasma mycoides var. capri. Experimental conditions as in Fig. 6. 1 = 50\(\mu g./ml\). Na-RNA; 2 = 20\(\mu g./ml\). thymidine; 3, 4, 5 = 50\(\mu g\). Na-RNA/ml.+1, 10 or 100\(\mu g\). thymidine/ml., respectively; 6, 7 = 50\(\mu g\). Na-RNA/ml.+50 or 300\(\mu g\). Na-DNA/ml., respectively.

The effects of RNA and DNA on the growth of the stable L-phase of Streptobacillus moniliformis

It seemed interesting to determine whether the RNA/DNA antagonism could be demonstrated in organisms different from Mycoplasma spp. and indeed this RNA/DNA antagonism was shown with the L-phase of Streptobacillus moniliformis. This organism grew on the fortified basal medium when DNA was added; RNA was not required (Fig. 8). Optimal growth occurred even with small concentrations of DNA (5\(\mu g./ml\.)); higher concentrations of
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DNA decreased the amount of growth. RNA showed a clear DNA-antagonistic effect. Increasing the concentrations of RNA in the medium increased the concentration of DNA required for optimal growth. Thymidine (1 µg./ml.) replaced DNA. In contrast to Mycoplasma mycoides var. capri 20 µg. thymidine/ml. were sufficient for optimal growth without any addition of RNA. It should be remembered that the fortified basal medium contained small amounts of RNA from the yeast extract.

DISCUSSION

The Mycoplasma organisms were found to be capable of using for growth undegraded RNA and DNA which are presumably degraded to smaller fragments before they are used for biosynthesis. Mycoplasma mycoides var. mycoides contains RNase (Plackett, 1957), but there is no information yet about the presence of RNase and DNase in other Mycoplasma organisms. However, DNases are known to be widespread in many groups of micro-organisms (Catlin & Cunningham, 1958). The possibility that RNA and DNA were degraded before utilization by enzymes present in the serum of the medium was ruled out by showing that growth was not affected by boiling the serum for 30 min. before its addition to the medium.

The nutritional requirements for nucleic acid fragments were best defined with Mycoplasma laidlawii strain A. This organism requires as minimal
essential nutrients thymidine (or less effectively, thymine) which can be derived from DNA, and one or more small oligonucleotides (di- or tri-nucleotides?) derived from RNA. The requirement for the ribo-oligonucleotide seems to be obligatory; it was not replaced by mononucleotides, nucleosides, purines or pyrimidines. This requirement for an oligonucleotide for growth would appear to indicate a deficiency in the biosynthetic ability of \( M. \text{laidlawii} \) strain A which is, at most, rarely found in bacteria.

All the Mycoplasma organisms tested were found to require something supplied by DNA. Thymidine replaced DNA with all the organisms tested except with \( M. \text{mycoides} \) var. \( \text{capri} \). Thymidine seems to play an important role in the biosynthesis of DNA by these organisms. The formation of purine deoxyribosides from purine bases and thymidine, by the action of a deoxy-nucleoside phosphorylase of Mycoplasma organisms (Lynn, 1957), supports this assumption. The much better growth response obtained with the deoxyriboside thymidine than with its pyrimidine base is analogous with other findings; better utilization of nucleosides than their purine or pyrimidine bases has been described for several other micro-organisms (Loring & Pierce, 1944; Hoffmann & Pavcek, 1952; MacLeod, Hogenkamp & Onofrey, 1958).

Strains A and B of \( M. \text{laidlawii} \) differ serologically, but cannot be distinguished by cultural and biochemical properties (Edward, 1950). Our finding that strain B differs from strain A by not requiring RNA (or oligonucleotide) as a nutrient is the first clue for a biochemical difference between these otherwise closely related strains.

We have found only one other example of the RNA/DNA antagonism in the literature, namely with \( \text{Lactobacillus bijdus} \) for which Skeggs, Spizizen & Wright (1950) found that DNA (or derived nucleosides or nucleotides) was a nutrient requirement and that RNA antagonized growth promoted by DNA. In our case, however, the picture is more detailed. \( M. \text{laidlawii} \) strain A needs its minimal nutrient requirements to be supplied by moieties of both DNA and RNA. When these moieties are supplied as undegraded RNA and DNA the antagonistic effect is shown by high concentrations of the polynucleotides. The growth inhibition caused by high concentrations of RNA or DNA may be due to interference with the preparation from these polynucleotides of the moieties required for biosynthesis. The growth inhibitory effect of RNA can be explained by its interference with preparatory degradation of DNA by the organisms (e.g. to release thymidine as a minimal biosynthetic precursor). RNA may inhibit competitively the degradation of DNA by the DNAse of the organisms; inhibition of microbial DNAses by RNA has been described (Bernheimer & Ruffier, 1951; Catlin & Cunningham, 1958). Indirect evidence in favour of this theory is the inability of RNA to inhibit growth when thymidine, which is the part of DNA needed by \( M. \text{laidlawii} \) strain A, is added to the medium instead of DNA. The observation that the degradation of RNA destroys its DNA-antagonistic activity also fits the theory; the whole RNA molecule is apparently necessary for an effective competition with DNA at the active sites of the organism's DNAse.

The growth inhibition caused by high DNA concentrations cannot, however,
be explained in the same way, namely by interference with RNA utilization, since inhibition by DNA could be shown with organisms which did not require added RNA (or oligonucleotides) for growth. High DNA concentrations inhibited also the growth of *Mycoplasma laidlawii* strain A when ribo-oligonucleotides were supplied instead of undegraded RNA. Degradation of DNA to deoxynucleotides and deoxynucleosides did not abolish the growth inhibitory activity. This might be explained by our observation that high concentrations of deoxynucleotides and deoxynucleosides interfered with the growth-promoting activity of thymidine, probably by competition at the active sites of biosynthetic enzymes. The interference of non-essential nucleotides and nucleosides with the utilization of the essential ones has been described, e.g. for *Escherichia coli* (Cohen & Barner, 1957), a marine Flavobacterium (MacLeod *et al.* 1958) and for *Lactobacillus acidophilus* (Siedler & Schweigert, 1959). The annulment of the growth inhibitory effect of high DNA concentrations by RNA might be explained by the interference by RNA with DNA degradation and therefore with the formation of interfering nucleotides and nucleosides. RNA could thus fulfil a regulatory function. The possibility that DNA might interfere with preparatory degradation of RNA by *Mycoplasma laidlawii* strain A, through inhibition of the RNase of this organism, should also be kept in mind; inhibition of RNase action by DNA was reported by McDonald (1955).

The RNA/DNA antagonism seems to be a general phenomenon and is not limited to Mycoplasma organisms. We observed it with the 1-phase of *Streptobacillus moniliformis* and Skeggs *et al.* (1950) described it with *Lactobacillus bifidus*. It looks as if other micro-organisms which require some part of DNA as a precursor in biosynthesis, and which can degrade DNA for this purpose, might be inhibited by RNA when DNA is given in the culture medium as the source of this nutritionally essential moiety.

This work was begun by the late Stuart Crowther who supplied an invaluable basis from which our work was continued. It is our deep regret that his premature death prevented his exploiting it. We wish to acknowledge with sincere thanks the supply of cultures of *Mycoplasma* species by Dr D. G. ff. Edward and our valuable discussions with him. Similarly we wish to thank Dr E. Klineberger-Nobel. One of us (S.R.) is indebted to the Friends of the Hebrew University of Jerusalem for the award of a Michael and Anna Wix Trust Fellowship. A grant from the Agricultural Research Council in support of this work is gratefully acknowledged.

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