An Investigation of RNA Synthesis in *Anacystis nidulans* During Exponential Growth Using Techniques of RNA–DNA Hybridization

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**SUMMARY**

RNA isolated from exponential-phase cultures of *A. nidulans* was used to titrate denatured DNA over a wide range of RNA:DNA ratios. The following results were obtained: (i) 0.6% of *A. nidulans* DNA was complementary to purified rRNA and 0.062% was complementary to tRNA. (ii) Under the growth conditions employed (35 °C; mean generation time 4 h), unstable RNA accounted for 40% of the rapidly labelled RNA fraction but only 2% of the randomly labelled RNA fraction. The half-life of the unstable RNA was estimated to be 3% of the mean generation time. (iii) A wide variation in the abundance of unstable RNA species was observed; more than 80% of the labelled RNA in both rapidly and randomly labelled unstable RNA fractions was homologous to only 10% of the DNA that was actively transcribed (i.e. 1% of the total DNA). In turn, the actively transcribed DNA comprised only 10% of the total DNA since virtually all the unstable and readily hybridizable RNA fraction (> 99%) from rapidly and randomly labelled RNA would form stable hybrid with it. This indicated that the remaining fraction of the DNA (90%) was infrequently transcribed.

**INTRODUCTION**

Knowledge of macromolecule composition and synthesis in blue-green algae has arisen mainly from comparative studies with a limited number of species which emphasized the structural and functional similarities with the other prokaryotes (see Leach & Herdman, 1973). RNA content per cell of *Anacystis nidulans* increased with increased growth rate, such that at 35 °C each cell contained 5 x 10^-4 fg RNA when grown at a mean generation time of 10 h and 23 x 10^-1 fg RNA at a mean generation time of 3.3 h (Mann & Carr, 1974). RNA synthesis appeared to be modulated by growth rate in a manner comparable to that in heterotrophic bacteria (see Maaløe & Kjeldgaard, 1966) although the ratio of rRNA to tRNA remained constant over a range of division times (Mann & Carr, 1973). *Anacystis nidulans* also differed from heterotrophic bacteria in that the rate of increase of RNA content per cell, with increasing growth rate, paralleled the rate of increase of DNA content rather than exceeding it.

The control of RNA synthesis in blue-green algae is of particular interest since it has been demonstrated in several species, notably *A. nidulans* and *Anabaena variabilis*, that many of the enzymes are constitutive (see Carr, 1973). The synthesis of several groups of enzymes has been shown to be neither repressed nor derepressed by the presence or absence of their end-products. For example, in contrast to several bacterial species, the formation of the first enzyme in the synthesis of methionine, homo-O-transsuccinylase, was not repressed by

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methionine (Delaney, Dickson & Carr, 1973) in either wild-type or a methionine auxotroph of *A. nidulans*. There are some examples of apparent transcriptional control in this organism (Abeliovich, Kellenberg & Shilo, 1974; Ihlenfeldt & Gibson, 1975; Singer & Doolittle, 1975) although the caution expressed by Wan, Floyd & Hatch (1975) in attributing alteration in enzyme activity to control of enzyme synthesis rather than to persistent feedback inhibition in the unicellular blue-green alga *Synechococcus cedrorum* should be noted. Reduction in the rate of RNA synthesis following a shift-down in growth rate is accompanied by an increase in the concentration of guanosine 5'-diphosphate-3'-diphosphate (ppGpp), indicating a stringent response in the linking of RNA synthesis to cell growth (Mann, Carr & Midgley, 1975). It has been suggested that in *Escherichia coli* ppGpp is concerned in the control of stable RNA synthesis (Cashel, 1969; Travers, Kamen & Cashel, 1970).

This paper examines RNA synthesis in steadily growing cultures of *A. nidulans* and includes estimations of the content and relative rates of synthesis of the unstable and stable RNA fractions. The data is interpreted in terms of the transcriptional activity on the various complementary fractions of the genome.

**METHODS**

*Culture conditions.* *Anacystis nidulans* (Culture Collection of Algae, Indiana University, strain 625) was grown in medium C (Kratz & Myers, 1955) modified by the addition of Tris (0.4 g l⁻¹) and reduction of the KH₂PO₄ concentration to 7.0 × 10⁻⁵ M. The medium was adjusted to pH 7.8 with 1 M-HCl before autoclaving. Experimental cultures (1 l) were maintained at 35 ± 0.1 °C and gassed with air/CO₂ (95:5, v/v) at 50 ml min⁻¹. Illumination was provided by two 75 W reflector lamps (Cryselco, Kempton Works, Bedford) placed 12 cm from either side of the culture vessel; the incident light intensity was 1.94 klux. These conditions resulted in a mean generation time of 230 to 240 min. Growth was measured at 650 nm using a Gilford spectrophotometer.

*Preparation of nucleic acids.* Nucleic acids were isolated as described by Pigott & Midgley (1968). ³²P-labelled DNA was prepared by allowing a continuous uptake of [³²P]orthophosphate (specific radioactivity 100 Ci g⁻¹, final concentration 1 µCi ml⁻¹) for six generations of exponential growth. Cells were harvested by centrifuging (100000 g for 15 min at 4 °C), mixed with a larger quantity of unlabelled wet packed cells and suspended (100 to 200 mg wet wt ml⁻¹) in 0.1 M-EDTA/0.4 M-NaCl, pH 8.0. The suspension was treated with lysozyme (500 µg ml⁻¹; Sigma, grade 1) for 30 min at 37 °C (Craig, Leach & Carr, 1969) before proceeding with cell lysis as described by Pigott & Midgley (1968). DNA yield was poor [0.48 mg (g wet wt of packed cells)⁻¹], due possibly to incomplete cell lysis.

Total RNA was prepared from exponential-phase cultures with an extinction *E*₆₅₀ of 0.5 ± 0.05. Total RNA, containing radioactivity in the stable RNA fractions only, was prepared from late exponential-phase cultures. These cultures were supplemented with a 2000-fold excess of unlabelled orthophosphate and incubated for a further cell generation time before harvesting in order to chase radioactivity from the unstable RNA fraction. Rapidly labelled RNA was prepared by allowing rapidly growing cultures (*E*₆₅₀ 0.4) to incorporate [³²P]orthophosphate (specific radioactivity 30 Ci g⁻¹, final concentration 3 µCi ml⁻¹) for 4 min. The cells were then poured into crushed ice and harvested at 4 °C.

The supernatant obtained from the ribosome sedimentation step (Pigott & Midgley, 1968) was used to prepare tRNA. After DNAase treatment and deproteinization, the RNA was purified by repeated fractionation on polylysine keiselguhr columns using a linear gradient between 0.4 and 4.5 M-NaCl in phosphate buffer pH 7.2. The pancreatic ribonuclease-
sensitive material eluting as a sharply defined peak at 1 M-NaCl was taken as the purified tRNA.

**RNA–DNA hybridization.** The nitrocellulose filter technique of Gillespie & Spiegelman (1965) was used as described by Pigott & Midgley (1968). All filters were presoaked in 10x SSC buffer (0.15 M-NaCl/0.015 M-trisodium citrate, pH 7.3) for 5 days before use. This procedure reduced non-specific binding of RNA during hybridization. Filters were loaded with 0.1 to 100 µg denatured DNA. Measurements on filters taken through the hybridization procedure showed that in no case was more than 1% of the initial DNA load lost; the average loss was 0.3%. After the hybridization incubation (16 h at 66°C) the filters were washed on either side with 50 ml 6x SSC buffer and then incubated for 1 h at 37°C in 2 ml 1x SSC buffer containing 10 µg pancreatic ribonuclease (EC 3.1.4.22; BDH). The results were corrected for the efficiency of the hybridization reaction and for non-specific adsorption of any RNA which had resisted pancreatic ribonuclease digestion, using filters containing no DNA (Pigott & Midgley, 1968).

**RESULTS**

**Titration of denatured DNA with stable RNA**

The fraction of the stable RNA capable of forming pancreatic ribonuclease-resistant hybrids with DNA was determined for a series of increasing RNA:DNA input ratios. The fraction of the denatured DNA input hybridized to rRNA (Fig. 1a) or tRNA (Fig. 1b) as a function of the RNA:DNA input was measured. The titration curves for both rRNA and tRNA are comparable with those normally found for bacterial stable RNA (Kennell, 1968; Pigott & Midgley, 1968). The DNA sites capable of forming true hybrids with rRNA became saturated at a RNA:DNA input ratio of 1:80; those sites capable of accepting tRNA became saturated at a ratio of 1:14. When complementary sites were fully saturated, 0.64% of the DNA was hybridized to rRNA and 0.062% to tRNA indicating that 0.7% of the *A. nidulans* genome codes for stable RNA.

**Titration of denatured DNA with total RNA**

The fraction of randomly labelled total RNA capable of forming pancreatic ribonuclease-resistant hybrids is shown as a function of the RNA:DNA input in Fig. 2. Although non-specific adsorption of RNA increased more rapidly than true hybrid formation as the RNA input was increased, reproducible results that agreed within 10% were obtained even at high RNA:DNA ratios. Non-specific adsorption was kept within acceptable limits by the use of presoaking techniques combined with thorough washing of the filters both before and after the pancreatic ribonuclease incubation.

To distinguish messenger and stable RNA fractions, *A. nidulans* DNA was titrated with total randomly labelled RNA and, in another experiment using the same hybridization procedure, with an RNA extract differing only in that the radioactivity in the unstable RNA fraction had been removed by chasing with a 2000-fold excess of unlabelled orthophosphate for at least 4 h. The difference between the two curves (Fig. 2) provides an estimate of the contribution made by the unstable RNA fraction to the titration of the DNA at that RNA:DNA input ratio.

The titration curves show that the stable RNA species saturate their complementary DNA sites at considerably lower RNA:DNA input ratios than do the unstable RNA species. Not only do the unstable RNA species form a smaller proportion of the input RNA, but they are also complementary to a larger fraction of the DNA than are the stable RNA species. Thus a significant contribution to the hybrid from the unstable RNA species is not apparent.
Fig. 1. Titration of denatured DNA with stable RNA fractions: (a) ribosomal RNA; (b) transfer RNA. The percentage of filter-bound DNA that formed hybrids is plotted against the RNA:DNA input ratio. Results were corrected as described in Methods.

Fig. 2. Titration of denatured DNA with randomly labelled total cell RNA. ●, Percentage of labelled RNA input forming hybrid plotted against RNA:DNA input. ○, Percentage of labelled RNA input forming hybrid, when the total cell RNA contains radioactivity in stable RNA only, plotted against RNA:DNA input (corrected to a maximum of 98%, see text). The difference between these curves gives an estimate of the fraction of input RNA that is unstable RNA. Results were corrected as described in Methods.
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Fig. 3. Titration of denatured DNA with rapidly labelled total cell RNA. ●, Percentage of labelled RNA input forming hybrid plotted against RNA:DNA input. Dashed line shows the titration of stable RNA, reproduced from Fig. 2, corrected to a maximum contribution of 60% (see text). Results were corrected as described in Methods.

until the RNA:DNA input ratio is 0.2. The unstable RNA fraction may therefore also be described as that RNA fraction which hybridizes efficiently with DNA at high RNA:DNA ratios. Such ‘readily hybridizable’ RNA fractions are commonly equated with messenger RNA (Kennell, 1968). In this instance an additional criterion, that of mRNA instability, has been observed. The slight plateau beginning at the input ratio of 0.2 signifies that most of the unstable RNA fraction is complementary to a relatively small fraction of the DNA. However, as total saturation is not achieved even at high RNA:DNA ratios, infrequent unstable RNA species must still be contributing to hybrid formation.

The proportion of unstable RNA in the total RNA input may be measured at an RNA:DNA input of 0.2. At this ratio the most abundant unstable RNA species begin to saturate their complementary DNA sites. Some 4.8% of the RNA input forms stable hybrids under the conditions employed, of which 2.9% is contributed by the stable RNA fraction. Thus nearly 2.0% of the total RNA is unstable, and the stable RNA species comprise 98% of the total RNA in a rapidly growing culture of A. nidulans. For this reason the hybridization curve for total cell RNA containing radioactivity only in stable RNA fractions was corrected to a maximum value of 98% in accounting for the efficiency of the hybridization technique (Kennell, 1968).

In a similar manner to that described for randomly labelled RNA, the fraction of the radioactivity in rapidly labelled RNA that formed hybrids was determined as a function of the RNA:DNA input ratio (Fig. 3). Again the titration curve reaches a plateau at an RNA:DNA input ratio of 0.2 though it is more pronounced than that seen in Fig. 2. Estimates made at this input ratio indicate that about 40% of the $^{32}$P present in hybrids is contributed by the unstable RNA fraction. The dashed line in Fig. 3 represents the hybridization of total cell RNA containing $^{32}$P in stable RNA only. The curve is reproduced from Fig. 2 but the maximum contribution was set at 60%, the estimated proportion of stable RNA in the rapidly labelled fraction.

Frequency of transcription of mRNA coding sites

The data contained in the DNA titration curves for rapidly and randomly labelled RNA Figs 2 and 3) may be expressed in terms of the relative activity of the sites within the genome which give rise to the unstable RNA fraction. Figures 4(a) and (b) were generated using
Fig. 4. The fraction of mRNA complementary to a fraction of the DNA input for (a) randomly labelled RNA, (b) rapidly labelled RNA. The values were derived from Fig. 2 and Fig. 3 respectively according to the method of Kennell (1968). The dashed line in Fig. 4(b) was constructed assuming that corrections to account for the effect of nucleic acid concentrations on the efficiency of the hybridization reaction were required to the full extent described by Kennell (1968) and Kennell & Kotoulas (1968).

a simple arithmetical procedure devised by McCarthy & Bolton (1964) as modified by Kennell (1968) and essentially depict saturation curves for the titration of DNA with the unstable RNA fractions of rapidly or randomly labelled RNA. The results indicate that nearly all the unstable RNA detected by the hybridization assay (greater than 99%) was complementary to only 10% of the genome (Fig. 4). Furthermore, the larger fraction of this RNA (80 to 90%) is complementary to less than 1% of the DNA or 10% of the actively transcribed genes.

No fraction of the unstable RNA appeared to be effectively more stable than the rest. When DNA was titrated against total RNA containing 32P in the stable RNA fraction only, a uniform saturation plateau was observed even at high RNA:DNA ratios (Fig. 2). If a significant fraction of the mRNA species were relatively stable, they would be expected to contribute to the hybrid formed. There is also little difference between the frequency distribution for rapidly labelled and long labelled unstable RNA (Fig. 4). This does not exclude the possibility that infrequent transcripts may be relatively stable but are present at concentrations too low to be detected by the hybridization assay (Pigott & Midgley, 1968; Kennell, 1968).

When one strand of DNA is transcribed and 10% of the DNA is complementary to virtually all the RNA detected, only 20% of the potential gene material is expressed. It is unlikely that the quiescent gene material is incompetent in forming hybrid with complementary RNA (Nygård & Hall, 1964; Kennell, 1968). Therefore it appears that 80% of the potential gene material is very rarely expressed in steadily growing A. nidulans cultures.

When DNA is filter-bound the efficiency of hybridization is a function of the nucleic acid concentration. This dependence is most pronounced when infrequent RNA species form a large fraction of the total RNA input, as in rapidly labelled RNA, and when the RNA:DNA input ratio is high, 1:10 (Kennell & Kotoulas, 1968). Using filters bearing various amounts of DNA, the effect of RNA concentration was assessed by comparing samples of the same RNA:DNA input ratio. For each RNA:DNA input the maximum attainable percentage
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of $^{32}$P incorporation into hybrid was measured. Although this procedure may lead to an underestimate of the maximum efficiency of hybrid formation at high RNA:DNA input ratios, the interpretation of the titration in terms of the relative activity of the mRNA sites within the genome is not unduly affected (see Fig. 4b).

DISCUSSION

The variation in abundance of individual RNA species in E. coli has been noted by McCarthy & Bolton (1964) and Kennell (1968). Our results shows that such variation is apparent in A. nidulans and, in common with E. coli, probably arises from differences in the frequency of transcriptional events on the various gene sites rather than from variation in the stability of individual transcripts. As the fraction of the genome complementary to stable RNA is relatively small the proportion of rapidly labelled RNA that is stable RNA would suggest that the stable RNA cistrons are among the most frequently transcribed fractions of the genome. Crowding of RNA polymerase molecules on E. coli rRNA cistrons (Mueller & Bremer, 1968) influences the expression of rRNA and mRNA cistrons during purine deprivation (Varney, Thomas & Burton, 1970; Smith & Midgley, 1973).

Variation in abundance of RNA species was, however, most apparent in the unstable RNA fraction of A. nidulans; virtually all of this fraction detected was complementary to only 10% of the DNA. Thus only 20% of the potential gene material of A. nidulans is actively transcribed. Furthermore 90% of the unstable RNA was complementary to only 10% of this actively transcribed DNA fraction (i.e. 2% of the potential gene material). Since the variation in abundance is continuous, each transcript having a characteristic rate of synthesis, one may attribute the variation to differential rates of initiation of new RNA chains. The initiation mechanism may be governed by the affinity of RNA polymerase for the promoter site (Scaife & Beckwith, 1966) of individual cistrons. Thus even when no other form of regulation is present the maximum rate of expression of a cistron would be governed by an aspect of the promoter site involved in the RNA polymerase initiation mechanism (cf. Travers, 1974). An additional or alternative explanation may be the reiteration, to varying degrees, of genes which lead to differential rates of mRNA synthesis. The wide range of mutation rates leading to auxotrophic strains of A. nidulans could also be interpreted as being a consequence of multiple gene sites coding for enzyme proteins. It is noteworthy that we report here a more extensive gene duplication for stable RNA cistrons in A. nidulans than is reported to exist in E. coli (Avery, Midgley & Pigott, 1969).

If the unstable and readily hybridized RNA fraction is equated with mRNA the results allow a description of the mRNA pool present in rapidly growing A. nidulans. The relative flux of labelled precursors into unstable RNA would suggest that 40% of the RNA synthesized is mRNA. Being unstable, however, this fraction is not accumulated and thus represents only 2% of the total cellular RNA. From these data, the average half-life of the unstable RNA species in the pool is estimated, assuming an exponential mode of decay, as approximately 7 min or 3% of the mean generation time of the culture. Comparable results were obtained for Anabaena variabilis by Leach & Carr (1974) who estimated the average half-life of the unstable RNA fraction to be 10 min or 3% of the generation time.

Finally two aspects of the present study require elaboration. Anacystis nidulans has been shown to contain a ‘small’ labile mRNA pool. This characteristic is a necessary prerequisite of the ‘stop/go’ mechanisms of induction and repression (Jacob & Monod, 1961) as distinct from the regulation of transcriptional frequency suggested above. In view of the apparent lack of such regulation in A. nidulans (Carr, 1973), this rapid turnover may
indicate the requirement to effectively extend the size of the ribonucleoside phosphate pool as an insurance against lack of precursor in unfavourable growth conditions.

A large fraction of the genome of *A. nidulans*, some 80% of the potential gene material, manufactures no detectable RNA product in the growth conditions employed. Although the non-transcribed DNA may be ascribed roles relating to gene punctuation and regulation, or as an insurance against lack of precursor in unfavourable growth conditions, these appear insufficient to account for the size of the quiescent fraction.

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**REFERENCES**


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