Sequential Expression of Macromolecule Biosynthesis and Candicidin Formation in *Streptomyces griseus*

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*Streptomyces griseus* did not produce the polyene macrolide antibiotic candicidin during the initial growth phase characterized by rapid RNA synthesis. The absence of candicidin production when RNA or protein synthesis was inhibited by rifampicin or chloramphenicol suggests a transcriptionally controlled late formation of the candicidin synthases. Phosphate levels in the medium control the rate of DNA, RNA and protein synthesis. Depletion of phosphate appears to trigger the onset of candicidin synthesis after a drastic reduction of the rate of RNA synthesis. Changes in the ATP pool during the fermentation suggest that ATP may be the intracellular effector controlling the onset of antibiotic synthesis.

**INTRODUCTION**

The ‘late’ expression of the synthesis of antibiotics and other secondary metabolites following intensive growth is a well-known phenomenon. It is equivalent to a differentiation process in which genetic information coding for growth and for antibiotic biosynthesis is sequentially expressed. In broad terms, two types of control mechanism are thought to be responsible, i.e. the antibiotic synthases are formed during growth but remain inhibited during the growth phase (post-translational control) or they are formed in a late phase subsequent to growth (transcriptional control). In most cases studied, the enzymes involved in antibiotic synthesis are repressed during the growth phase (Walker & Hnilica, 1964; Pruess & Johnson, 1967; Gallo & Katz, 1972). Although the intracellular factors that trigger onset of antibiotic synthesis are unknown, catabolite regulation (Gallo & Katz, 1972), inducer accumulation (Krupinski, Robbers & Floss, 1976) and phosphate (or ATP) control (Martin, 1976) have been proposed as control mechanisms.

The biosynthesis of many antibiotics is controlled by inorganic phosphate (Weinberg, 1974; Martín, 1976). In particular, the formation of the polyene macrolide antibiotic candicidin is a very suitable model system for the study of the control mechanisms involved in the onset of antibiotic synthesis because of its high sensitivity to phosphate (Liu, McDaniel & Schaffner, 1975; Martín & Demain, 1976). The possible involvement of ATP as the intracellular effector controlling antibiotic synthesis is suggested by the work of Janglova, Suchy & Vanek (1969) and J. F. Martín, P. Liras & A. L. Demain (unpublished observations). Since oxygen uptake and DNA synthesis follow diphasic patterns in cultures synthesizing polyene macrolide antibiotics (Martín & McDaniel, 1974; Martín & McDaniel, 1975a) it was of interest to study the relationship between intracellular ATP concentrations and the formation of the antibiotic as well as the synthesis of RNA and protein. The results obtained in this work indicate that formation of candicidin synthases occurs after a rapid growth phase characterized by a high rate of RNA synthesis and is probably mediated by the intracellular ATP level.
METHODS

Micro-organisms and growth conditions. Streptomyces griseus IMRU3570, the strain producing candicidin, was grown in a medium supporting high antibiotic production which contained soya peptone (25 g l−1), glucose (65 g l−1) and zinc sulphate (0.5 mm) (Martin & McDaniel, 1975a). The cultures were incubated at 32 °C in triple-baffled 250 ml flasks, containing 50 ml of medium, in a Gallenkamp orbital shaking incubator operating at 225 rev. min−1. The production flasks were inoculated with 5 ml of a 24 h inoculum grown in medium containing yeast extract (10 g l−1) and glucose (10 g l−1).

Synthesis of candicidin was determined as described previously (Martin & McDaniel 1975a). DNA content was measured in perchloric acid extracts after lipid and protein removal, according to Schneider (1947).

Determination of RNA and protein synthesis. Incorporation of [2-14C]uracil (59 Ci mol−1) into 5 % (w/v) cold trichloroacetic acid (TCA)-precipitable macromolecules was done as previously described (Martin & McDaniel, 1975b; Martin & Demain, 1976) except that 0.1 μCi of [14C]uracil was added to 0.25 ml of culture supplemented with 2 mm unlabelled uracil. Incorporation of the 14C-labelled amino acid mixture (226 Ci mol−1) into 5 % cold TCA-precipitable material was done using 0.025 μCi in 0.25 ml of culture, supplemented with 0.05 mg unlabelled Casamino acids ml−1 (Difco). When measuring incorporation of labelled amino acids into protein, charged aminoacyl-tRNA was removed by solubilization in 5 % TCA at 90 °C for 20 min. Pulses of 10 min were used for RNA and protein synthesis determinations.

Preparation of phosphate-limited resting cells was carried out as previously described (Martin & McDaniel, 1975b; Martin & McDaniel, 1976).

Determination of phosphate utilization. Labelled phosphate [95 Ci (mg phosphate)−1] was added at inoculation time to control flasks and to flasks supplemented with 10 mm-unlabelled inorganic phosphate. Samples (250 μl) of the culture were filtered through Whatman GF/A glass-fibre filters and the filtrates were collected. Intracellular radioactive phosphate was measured on the dried filters, and extracellular radioactivity was determined on samples of the filtrates. Counting was done in a Packard Tricarb 3320 liquid scintillation counter. Colour quenching was corrected, if required, by the channels ratio method.

ATP determination. The method for determination of ATP was similar to that of Ball & Atkinson (1975). Samples (1 ml) were rapidly pipetted (within 15 s) into 0.2 ml of ice-cold 35 % (v/v) perchloric acid in order to ‘freeze’ the nucleotide pools prior to analysis. After 2 min at 0 °C, the extracts were rapidly frozen in liquid nitrogen and kept frozen until assayed. After thawing, the extracts were thoroughly mixed and centrifuged at 10000 g for 5 min. The supernatant fluids were neutralized with 0.4 ml of 2·6 M-KOH. The final pH was adjusted to 7.4 with 100 μl of 1 M-phosphate buffer pH 7.4. After at least 30 min at 0 °C, the KClO4 precipitate was removed by centrifugation. Recovery of externally added ATP was 85 %. ATP was determined by the luciferase reaction using the Packard Tricarb 3320 scintillation counter as a photometer (Stanley & Williams, 1969). One vial of crystalline luciferin—luciferase reagent (DuPont de Nemours and Co.) was dissolved in 3 ml of morpholinopropane sulfonic acid buffer (MOPS) pH 7.4. Before the enzyme reaction, the enzyme solution was diluted 1/10 into distilled water. A cuvette containing 200 μl of diluted enzyme solution, 500 μl of assay buffer (100 mm-glycine/NaOH buffer pH 7·4) and 500 μl of 15 mm-MgCl2 was placed in the belt of the scintillation counter and 10 to 100 μl of ATP extract were rapidly injected into the vial at the time it was entering the counting chamber. Standard ATP solutions, in the range 0·1 to 10 μM-ATP (10 pmol to 1 nmol per 100 μl sample), gave a reproducible linear response. Extracellular ATP, measured after Millipore filtration of culture broths, was always two orders of magnitude lower than the intracellular levels and was usually undetectable.

Chemicals. Chloramphenicol, rifampicin and ATP were purchased from Sigma. Luciferase was a product of DuPont de Nemours and Co. [2-14C]Uracil (59 Ci mol−1) and [32P]orthophosphate [95 Ci (mg phosphate)−1] were from The Radiochemical Centre, Amersham. Uniformly 14C-labelled L-amino acid mixture (226 Ci mol−1) was purchased from New England Nuclear Corporation. All reagents were of analytical grade.

RESULTS

Time sequence of macromolecule biosynthesis

The kinetics of the rate of RNA synthesis indicated a clear-cut separation of two phases, the growth phase (trophophase) and the production phase (idiophase) (Fig. 1). A high rate of RNA synthesis occurred during the first 15 h of fermentation and then decreased rapidly. No antibiotic was synthesized during the phase of rapid RNA synthesis. An increase in the content of DNA was also observed during this phase (Martin & McDaniel, 1975a).

The increase in the rate of protein synthesis lagged behind the increase in RNA synthesis,
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Fig. 1. Kinetics of the synthesis of RNA (10 min pulses) (▲), protein (10 min pulses) (□) and candididin (○).

Fig. 2. Effect of chloramphenicol (100 µg ml⁻¹) on candididin synthesis. ○, Control; chloramphenicol added at (h): ▲, 10; △, 18; ▲, 20; ■, 25; □, 32; ●, 43. Similar curves were obtained with rifampicin at 10 µg ml⁻¹.

reaching a maximum rate at 30 h and decreasing slowly thereafter (Fig. 1). Thus, considerable protein synthesis occurred during the phase of antibiotic production, whereas there was little RNA turnover in this phase. Total dry weight also increased during the idiophase, this increase representing mainly storage materials plus antibiotic (Martin & McDaniel, 1975a).

Antibiotic production began at about 20 h, following the rapid decrease in RNA synthesis. Thereafter, antibiotic production was linear for at least 40 h.

‘Late’ formation of antibiotic synthases

The late onset of candididin synthesis may be due to a delayed formation of candididin synthases or to a release from inhibition of growth-associated antibiotic synthases. We checked these two possibilities by studying the effect of rifampicin and chloramphenicol on candididin synthesis. At the concentration used (100 µg ml⁻¹), chloramphenicol completely inhibited protein synthesis within 40 min of its addition but did not produce lysis of the culture. Rifampicin was used at 10 µg ml⁻¹ which completely inhibited RNA synthesis within 40 min; higher concentrations produced cell lysis. We found that little or no candididin was formed if chloramphenicol or rifampicin was added before 18 h, but increasing amounts of candididin accumulated when either RNA or protein synthesis was inhibited at 18, 20, 25, 32 and 43 h. The results presented in Fig. 2 show the effect of chloramphenicol but the results with rifampicin were not significantly different. Both chloramphenicol and rifampicin permitted a residual synthesis of candididin for a few hours, probably due to the existing pool of mRNA for candididin synthases and the pool of pre-formed synthases.

These results suggest that antibiotic synthases are formed late. However, it could be that the enzymes formed early were inhibited by the phosphate in the medium until this had been depleted. Moreover, addition of chloramphenicol or rifampicin to the batch culture may...
have retarded the rate of phosphate utilization. To check this possibility, phosphate-limited resting cells were prepared at different times during the fermentation. The synthesis of candididin by phosphate-limited resting cells was compared with candididin production by similar cells in which protein synthesis was inhibited by chloramphenicol. No candididin synthesis by phosphate-limited cells was observed in chloramphenicol-treated cells prepared from a 9 h batch culture (Fig. 3a). Control cells without chloramphenicol were derepressed for candididin synthesis after suspension in the phosphate-free medium, even though there was no candididin at 9 h in the batch fermentation. In phosphate-limited cells prepared from a 16 h batch culture (Fig. 3b) some candididin was produced in chloramphenicol-treated cells, which indicates that at this time some candididin synthases were already formed, even though they were not expressed in the batch culture. Higher rates of synthesis of candididin by chloramphenicol-treated cells obtained after 24, 36 or 60 h of fermentation suggest that at this time a large part of the candididin synthases are already formed (Fig. 3c, d, e). It seems, therefore, that no candididin synthases are formed in the first hours of fermentation, suggesting transcriptional control of its ‘late’ formation.

**Control by phosphate of macromolecular synthesis and candididin production**

To study whether phosphate is involved in the time-sequence of macromolecular synthesis and antibiotic production, fermentations were carried out with and without added phosphate (10 mM). In the soya peptone medium, 5 mg cell dry wt ml⁻¹ was reached in 60 h, accompanied by candididin production amounting to about 600 μg ml⁻¹ (Fig. 4). In the presence of added phosphate, growth (as cell dry weight) was doubled but no candididin was formed (Fig. 4). Addition of 10 mM-phosphate also resulted in an increased rate of RNA synthesis, protein synthesis and DNA content (Fig. 5).
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Fig. 4. Effect of phosphate on (a) cell dry weight and (b) candicidin synthesis. ○, Control; ●, medium supplemented with 10 mm-potassium phosphate at inoculation time.

Fig. 5. Effect of phosphate on the synthesis of (a) RNA (10 min pulses), (b) protein (10 min pulses) and (c) DNA. ○, Control; ●, medium supplemented with 10 mm-potassium phosphate at inoculation time.

Phosphate depletion and onset of candicidin synthesis

Martin (1976) suggested that phosphate might be exhausted before the onset of antibiotic synthesis. We followed the uptake of $^{32}$P]phosphate in control cultures (unsupplemented, Fig. 6a) and in cultures supplemented with 10 mm-phosphate (Fig. 6b). $^{32}$P]Phosphate was rapidly utilized in control cultures. Only small amounts of extracellular phosphate remained in the broth at 13 h, 2 to 3 h before the onset of candicidin synthesis in this experiment. This was also the time when candicidin synthases start to be formed (Figs 2 and 3). $^{32}$P]Phosphate is accumulated as intracellular phosphate in control cultures. Extracellular phosphate levels
remained quite low during most of the production phase. In cultures supplemented with 10 mM-phosphate, there was fast phosphate utilization during rapid growth, followed by slower utilization thereafter. Extracellular levels of phosphate in supplemented cultures were much higher throughout the fermentation.

If the depletion of phosphate triggers the onset of candicidin synthesis, it would be expected that cultures supplemented with increasing concentrations of phosphate would be depleted later and later, thus delaying the onset of candicidin synthesis. This was found to be the case (Fig. 7).
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![Graph](image)

Fig. 9. Changes in the intracellular ATP pool during candicidin batch fermentation inoculated with spores. (a) ○, Cell dry weight; □, candicidin. (b) ATP concentration calculated as μmol (g dry wt)^{-1} (△) or nmol ml^{-1} (▲).

Intracellular ATP levels and onset of candicidin synthesis

As a control in intermediary metabolism, ATP is a potential effector in phosphate-mediated control of candicidin synthesis. To study this possibility, we followed intracellular ATP levels during candicidin fermentation (Fig. 8). The ATP level per g of cells decreased sharply during the first 10 h of fermentation, before the onset of antibiotic synthesis. The ATP generated via rapid respiration at this time (Martin & McDaniel, 1975a) was actively used in the synthesis of macromolecules (RNA and protein) and thus decreased rapidly. As long as phosphate does not become limiting, the ATP pool is partially replenished, but after phosphate becomes limiting, ATP pools are kept necessarily low and the synthesis of macromolecules is severely limited. The level of the ATP pool remained at a very low level throughout the production phase.

The high ATP level immediately after inoculation of the production medium reflects the ATP pool of cells actively growing in the inoculum medium. In order to speed up antibiotic production, fermentations are usually inoculated with a heavy, fast-growing inoculum to shorten the trophophase. Therefore, to get a better understanding of the role of ATP, we inoculated the production medium with spores (Fig. 9). The rapid growth phase lasted for about 30 h under these conditions and antibiotic synthesis started after 25 h. When expressed per unit of cell dry weight, the ATP pool increased during the lag phase and decreased sharply during growth prior to the onset of antibiotic synthesis (Fig. 9); it then reached a rather low steady-state level during the production phase. Thus, independently of the type of inoculum used, ATP levels appear to be high during growth and low during the phase of antibiotic production.

DISCUSSION

The results of Fig. 1 confirm earlier results obtained in fermenters where a rapid increase in respiration rate and of DNA content was observed during the rapid growth phase (Martin & McDaniel, 1975a). The increase in the rate of RNA synthesis during the period of non-limited growth followed by the rapid decrease before the onset of antibiotic synthesis, suggest a sequential expression, first of genetic information coding for biosynthesis of macromolecules and then for candicidin production. Similarly, a drastic decline in RNA synthesis has been observed in *Streptomyces orientalis* at the initiation of vancomycin production (Mertz & Doolin, 1972) and in *Streptomyces antibioticus* before the beginning
of actinomycin formation (Jones & Weissbach, 1970; Jones, 1975). This decrease in RNA synthesis is manifested as a general decrease of the major ribosomal RNA species. However, the changes in RNA could have been due either to a decreased rate of RNA synthesis or to increased degradation (Jones, 1975). Our data obtained by pulse-labelling experiments on in vivo RNA synthesis (Fig. 1) suggest that the changes in RNA are mainly due to decreased de novo RNA synthesis after 15 h of fermentation.

Candicidin synthases were formed only after 18 h of fermentation (see Figs 2 and 3). Thus, formation of antibiotic synthases appears to be repressed during the rapid growth phase. Similar control at the transcription level has been described in the late formation of several antibiotic-synthesizing enzymes (Walker & Hnilica, 1964; Pruess & Johnson, 1967; Matteo et al., 1975; Gallo & Katz, 1972; Matern, Brillinger & Pape, 1973). The results of Fig. 4 confirm previous results indicating that phosphate addition (10 mM) completely inhibits antibiotic synthesis and greatly increases cell dry weight.

The formation of antibiotic synthases following a decrease in the rate of RNA synthesis might indicate a transcriptional control, e.g. by modification of the template specificity of the RNA polymerase, as occurs in the control of late expression of sporulation genes (Losick & Sonenshein, 1969). Changes in the structure of the sigma subunit of the RNA polymerase have been postulated to alter the template specificity during sporulation.

Extracellular phosphate depletion occurred at the same time as the onset of candicidin synthesis (Fig. 6). In support of phosphate-mediated control of antibiotic synthesis is the fact that the onset occurred later at higher concentrations of phosphate (Fig. 7). A similar result was described by Mertz & Doolin (1972), i.e. the presence of excess phosphate delayed the synthesis of vancomycin until 96 h of fermentation. Moreover, phosphate addition to a candicidin-producing culture resulted in an immediate inhibition of candicidin synthesis (Liu et al., 1975; Martin & Demain, 1976).

Some clues as to how phosphate can control differential expression of antibiotic synthesis can be derived from the work of Zilling et al. (1975) on the phosphorylation of DNA-dependent RNA polymerase of E. coli by a phage T₇-induced protein kinase. The β' subunit of E. coli RNA polymerase (and to a lesser extent the β subunit) is phosphorylated and its activity is lost. Phosphorylation of host RNA polymerase seems to be required for the termination of early transcription of phage T₇.

Whether the intracellular pool of ATP is the effector mediating the phosphate effect has not been definitively established although the results of Figs 8 and 9 provide some evidence in support of this mechanism. Moreover, phosphate addition to candicidin-producing cultures resulted in an immediate two to threefold increase in the ATP pool level, and simultaneous inhibition of antibiotic synthesis (Martin, 1976; J. F. Martin, P. Liras and A. L. Demain, unpublished observations).

Several other reports point to the involvement of ATP in controlling the onset of antibiotic production. This was first suggested by Silaeva et al. (1965) and Janglova et al. (1969). They reported that ATP levels were lower in high-producing strains than in low-producing strains of Bacillus brevis and Streptomyces aureofaciens. A recent report of Fynn & Davison (1976) also points to the possible involvement of ATP in the regulation of antibiotic biosynthesis.
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REFERENCES


