Guanosine 5'-diphosphate 3'-diphosphate (ppGpp), guanosine 5'-diphosphate 3'-monophosphate (ppGp) and antibiotic production in *Streptomyces clavuligerus*

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*Streptomyces clavuligerus* produced cephamycin C under all nutrient limitations investigated in batch culture, whilst clavulanic acid was produced under phosphate and carbon limitations. Guanosine 5'-diphosphate 3'-diphosphate (ppGpp) was produced at very low levels in all fermentations, prior to the detection of cephamycin C in the fermentation broth. Guanosine 5'-diphosphate 3'-monophosphate (ppGp) was observed at high levels in all fermentations; it appeared following a downturn in nutrient levels and immediately prior to detection of isopenicillin N synthase (IPNS). ppGp was detected following nutritional shiftdown by amino acid depletion and it was not produced via degradation of ppGpp. The results point to a possible role for ppGp in the regulation of cephamycin production in *S. clavuligerus*.

**Keywords**: *Streptomyces clavuligerus*, stringent response, ppGpp, ppGp, antibiotics

**INTRODUCTION**

Streptomycetes produce a wide range of secondary metabolites, including antibiotics. A characteristic in the production of antibiotics is that they are produced when micro-organisms are growing at low specific growth rates. This implies that during batch culture, genes coding for the enzymes (synthases) necessary for antibiotic biosynthesis are repressed at high growth rates. The factor(s) that control the derepression of antibiotic synthases is probably the deficiency of one, or more, growth-limiting nutrients. At the moment very little is known about the nature and mode of action of the intracellular effector(s) that regulate antibiotic biosynthesis in streptomycetes. One potential effector molecule is guanosine 5'-diphosphate 3'-diphosphate (ppGpp).

When cells of bacteria, such as *Escherichia coli*, experience nutrient limitation, their growth rate decreases and they make a series of rapid adjustments to their metabolism. One response elicited is the stringent response, characterized by a reduction in the rate of stable RNA (i.e. rRNA, tRNA) accumulation and mRNA for ribosomal proteins and triggered by the binding of uncharged tRNA to ribosomes (Cashel, 1975). The stringent response is mediated by an accumulation of ppGpp, which is the product of the *relA* gene (Cashel & Rudd, 1987). In *E. coli*, growth rate is an inverse linear function of the intracellular ppGpp concentration, which suggests that ppGpp has a significant role in growth rate control (Sarubbi et al., 1988).

Several authors have shown that the stringent response and ppGpp production occur in a wide range of *Streptomyces* species (Riesenberg et al., 1984; Ochi, 1986; Strauch et al., 1991; Bascaran et al., 1991). Simuth et al. (1979) described a correlation between the onset of antibiotic production and ppGpp levels in *S. aureofaciens*. However, An & Vining (1978) had previously concluded that streptomycin production by *S. griseus* was not mediated by ppGpp. More recently, Ochi (1990a, b) has described the isolation of relC mutants that produced very low levels of ppGpp and were deficient in antibiotic production. However, Bascaran et al. (1991) isolated relC mutants of *S. clavuligerus* and could not find a correlation between antibiotic production and the stringent response. Moreover, Strauch et al. (1991) have concluded that increased levels of ppGpp alone are not sufficient to initiate antibiotic biosynthesis in *S. coelicolor*. Thus, there are conflicting interpretations concerning the role of ppGpp in the regulation of antibiotic biosynthesis in streptomycetes.

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**Abbreviations**: DCW, dry cell weight; IPNS, isopenicillin N synthase.
In order to study the role of ppGpp more fully we have initiated a physiological and molecular biology programme using the industrially relevant organism *S. clavuligerus*. In this paper we describe levels of ppGpp, ppGp (guanosine 5'-diphosphate 3'-monophosphate) and β-lactam antibiotics, during batch fermentation of *S. clavuligerus*.

**METHODS**

Organisms and growth conditions. *S. clavuligerus* NRRL 3585 was grown at 28 °C as a batch culture in a 16 litre Bioengineering L1523 fermenter with a working volume of 12 litres, or in 1 litre triple-baffled flasks containing 200 ml medium. The pH was maintained at 6·9 with 50 mM MOPS. In all cultures the chemically defined medium (FM) was as described by Aharonowitz & Demain (1979), except that under carbon-limiting conditions 4 g glycerol l −1 was used; under nitrogen-limiting conditions asparagine (2 g l −1) was replaced by 0·54 g NH₄Cl l −1 and under phosphate-limiting conditions 0·35 g K₂HPO₄ l −1 (1·5 mM) was used. A linear response of biomass concentration to an increase in the individual limiting substrate concentration was taken as proof that all nutrients, other than the required limiting nutrient, were in excess. Spores of *S. clavuligerus* were used to inoculate seed medium (Aharonowitz & Demain, 1979) and incubated in triple-baffled Erlenmeyer flasks for 48 h in a rotary shaking incubator at 200 r.p.m. and 28 °C. The resulting seed cultures were washed with medium of the same composition as used for each of the fermentations before being used as inoculum at 25·5% (v/v). Growth was estimated by measuring the dry cell weight (DCW).

Assay of nucleotide pools. Samples of culture (100 ml) were removed from the fermenters and filtered through 90 mm diameter filter papers (Millipore, 0·45 pm pore size), extracted with cold 1 M formic acid for 1 h, centrifuged for 10 min at 6000 g, and the supernatants filtered through nitrocellulose (Gelman, 0·45 pm pore size). The filtrates were freeze-dried and resuspended in 500 μl deionized distilled water. The intracellular concentrations of nucleotides were assayed by HPLC on a Partisil 10 SAX column (Whatman) with a gradient of 7 mM K₂HPO₄, pH 4·0, to 0·5 M K₂HPO₄/0·5 M Na₂SO₄, pH 5·4, at a flow rate of 1·5 ml min −1. Nucleotides were detected at 254 nm. The concentrations of nucleotides were expressed relative to mycelial dry weight.

Antibiotic determinations. Samples of filtered fermentation broths were assayed for β-lactam antibiotics by HPLC on a Nucleosil C18 ODS 25 cm column (Jones Chromatography). The column was pre-conditioned with 60 ml 4·5 mM tetraethylammonium bromide (TBAB) in 0·05 M K₂HPO₄ (pH 3·0)/methanol (85:15, v/v) at 1·5 ml min −1 and then equilibrated with 0·15 mM TBAB in 0·05 M K₂HPO₄ (pH 3·0)/methanol (95:5, v/v). Samples (20 μl) of culture filtrates were analysed at a flow rate of 1·5 ml min −1 and antibiotics were detected at 220 nm.

Nutrient consumption. Glycerol in the fermentation broth was determined by the method of Bok & Demain (1977). Asparagine was determined according to Plummer (1971). Ammonium was measured by the method of Weatherburn (1967) and phosphate was determined by the method of Golterman et al. (1978).

Western blotting. Isopenicillin N synthase (IPNS) expression was determined by Western blot analysis. Cells (10 ml) from the fermenter were washed and resuspended in 10 ml 0·05 M Tris/HCl buffer, pH 7·2, containing 0·1 mM dithiothreitol, 0·01 mM disodium EDTA and 1 mM phenylmethylsulfonyl fluoride. Cells were broken by passage through a pre-cooled French press at 70–80 MPa. Cell debris and unbroken cells were removed by centrifugation at 10000 g for 20 min at 4 °C. Protein content of the cell-free extract was determined according to the method of Bradford (1976). Proteins and prestained low-range molecular mass markers (Bio-Rad) were separated by SDS-PAGE (12%, w/v, acrylamide), using the method of Laemmli (1970). After SDS-PAGE, proteins were transferred to nitrocellulose filters at 30 V overnight in a Trans-Blot apparatus (Bio-Rad) according to the manufacturer’s instructions. Transferred proteins were probed with IPNS polyclonal antibody as a whole serum (1:500 dilution); the membranes were washed with 0·01 M Tris/HCl, pH 8·0, and 0·15 M NaCl and incubated for 1 h with a 1:3000 dilution of goat anti-rabbit IgG alkaline phosphatase conjugate (Sigma), and washed again. The membranes were then developed with Western blue reagent (Promega). Polyclonal antibodies were prepared in rabbits to a peptide sequence derived from the *S. clavuligerus* IPNS protein. The peptide was conjugated to bovine serum albumin and this conjugate was used for immunization of New Zealand White rabbits. Cell-free extracts of *E. coli* DS941, containing the IPNS gene on pMZR100, were run on each polyclonal gel to verify IPNS expression. Antisera and *E. coli* DS941 were kind gifts of Dr B. Barton, SmithKline Beecham Pharmaceuticals, Worthing, UK.

Reproducibility. Figs 1–6 show representative results from single experiments, all of which were repeated twice, and all parameter determinations were simultaneously carried out at least twice. Fermenter-grown cultures were sampled every hour, but for clarity not all sample points are shown in the figures. Fig. 7 shows representative results from single experiments carried out at least twice.

**RESULTS**

Production of cepharnycin C and clavulanic acid in batch culture

Fermentation medium (FM). Cepharnycin C was initially detected after 30 h of fermentation, and production continued until early stationary phase. Onset of the stationary phase corresponded to the depletion of asparagine (Fig. 1a). Clavulanic acid was not detected, due to the repressive levels of phosphate (20 mM) maintained during the course of the fermentation. The biosynthesis of clavulanic acid is repressed at phosphate concentrations above 10 mM (Romero et al., 1984). Fig. 1(b) shows that the pool of ppGpp remained constant below the level of detection [<3 pmol (mg DCW) −1] until 28 h, when a rapid burst was observed, reaching 23 pmol (mg DCW) −1. The pool returned to basal levels within 60 min and remained at this level for the rest of the fermentation. At 20 h, corresponding to the onset of the exponential growth phase, another nucleotide was detected which had an HPLC retention time identical to that of pure ppGp. Pure ppGp and the new nucleotide were shown by HPLC analysis to be completely hydrolysed after incubation with bacterial alkaline phosphatase yet resistant to hydrolysis after incubation in 0·3 M KOH. Further characterization
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When *S. clavuligerus* was grown as a shake-flask culture, appearance of ppGp production at 16 h [45 pmol (mg DCW)] corresponded to the start of exponential growth and a rapid downturn in asparagine levels (Fig. 2). Peaks of ppGp were also observed at 24 h [316 pmol (mg DCW)] and at 43 h [280 pmol (mg DCW)], the latter corresponding to late exponential growth and depletion of asparagine. Production of ppGpp was similar to that reported by An & Vining (1978) for *S. griseus*. Peaks were observed at 24 h [37 pmol (mg DCW)] and, also after 44 h, [97 pmol (mg DCW)], corresponding to the late exponential growth phase.

**Nitrogen limitation.** Cephamycin C was initially detected after 34 h and production followed a growth-associated pattern (Fig. 3a). Exhaustion of ammonium at 48 h corresponded to the onset of the stationary phase, and was followed by the cessation of cephamycin C production after 50 h. Clavulanic acid was not detected during the fermentation. Fig. 3(b) demonstrates that the pool of ppGpp remained at a basal level until 32 h when a rapid burst was detected, reaching 17 pmol (mg DCW). The ppGpp level decayed rapidly over the following 60 min, returning to basal levels, and remained at that level for the rest of the fermentation. Production of ppGp was noted between 22 and 28 h, at levels up to 1-75 nmol (mg DCW)

Fig. 1. Fermentation profile during batch-culture growth in FM. *S. clavuligerus* was grown as 12 litre cultures in a Bioengineering fermenter at 28 °C and 500 r.p.m. Dissolved O₂ tension above 50% air saturation was maintained during the course of the fermentation. (a) Growth as dry cell weight (●), cephamycin C (△), asparagine (■) and glycerol (◇). (b) Intracellular concentration of ppGpp (○) and ppGp (▲).
ontset of the exponential growth phase. The production of ppGpp and ppGp occurred prior to the detection of cephamycin C in the fermentation broth. ppGpp was not detected during the fermentation.

Carbon limitation. Cephamycin C was detected after 28 h and production followed a growth-associated profile (Fig. 4a), ceasing after 58 h of fermentation. Depletion of glycerol caused the onset of the stationary phase at 48 h. Clavulanic acid was initially detected after 36 h and followed a growth-dissociated profile, with levels remaining low during the course of the fermentation. Fig. 4(b) shows that the pool of ppGpp remained at a basal level until a burst was detected at 25 h, reaching 10⁻⁵ pmol (mg DCW)⁻¹, and decaying back to basal level within 60 min. ppGp was initially detected at 22 h [32 pmol (mg DCW)⁻¹], corresponding to the onset of the exponential growth phase and 6 h prior to the appearance of cephamycin C in the medium. Production of ppGp was also detected between 30 and 32 h [10 pmol (mg DCW)⁻¹], 4 h before clavulanic acid was detected in the fermentation broth. ppGpp was not detected during the fermentation.

Phosphate limitation. Cephamycin C and clavulanic acid were produced in a growth-dissociated manner, and were initially detected after 46 h, corresponding to the late exponential growth phase (Fig. 5a). Phosphate depletion from the medium after 52 h corresponded to the onset of the stationary phase. Phosphate limitation severely decreased the biomass levels; the decrease was considerably greater than reported by Lebrihi et al. (1987). However, the kinetics of cephamycin C and clavulanic acid production were very similar to the kinetics described by these authors. Fig. 5(b) shows that a burst of ppGpp was initially seen at 24 h [325 pmol (mg DCW)⁻¹], at the onset
of exponential growth. At 42 h, a burst of ppGp was noted [350 pmol (mg DCW)⁻¹], 4 h before cephemycin C and clavulanic acid were detected in the medium. pppGpp was not detected during the fermentation.

**Stringent response in S. clavuligerus**

In order to determine whether ppGp was produced by *S. clavuligerus* during the stringent response, cells growing exponentially in FM+1% Casamino acids were transferred to the same medium without Casamino acids. Fig. 6 shows that maximal ppGp levels were detected 10 min after shiftdown [118 pmol (mg DCW)⁻¹] and preceded the typical transient accumulation of ppGpp observed 15 min after shiftdown [35 pmol (mg DCW)⁻¹]. These kinetics are not consistent with a precursor–product relationship as would be expected if ppGp were merely a degradation product of ppGpp. We have been unable to detect ppGpp or ppGp during nutrient shiftdown to carbon or phosphate limitation.

In order to assess further the possibility that ppGp was a degradation product of ppGpp, authentic pure standard was incubated at 25 °C and 37 °C for up to 2 h in Tris/acetate/salts buffer (pH 8.0), with or without cell-free extracts derived from early and late exponential cells of *S. clavuligerus*. Nucleotide extraction and resolution by HPLC were performed as described in Methods. We were unable to detect any significant hydrolysis of ppGpp.

**Western blot analysis of IPNS**

*S. clavuligerus* grown in FM produced IPNS at 21 h (Fig. 7a), in nitrogen-limiting (Fig. 7b) and carbon-limiting (Fig. 7c) media at 23 h and in phosphate-limiting (Fig. 7d) conditions at 43 h. Under all nutrient conditions the appearance of the enzyme was immediately preceded by production of ppGp in the cells. There was no correlation with the timing of ppGpp production. It is interesting to note that under nitrogen and phosphate limitation there was an apparent increase in amount of IPNS, as determined by band intensities, compared with growth in FM or under carbon limitation.

**DISCUSSION**

In contrast to the prolonged lag phase exhibited by *S. clavuligerus* after inoculation into specific nutrient-limited media (Figs 3–5), a reduced lag period was seen following
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from the nutritionally rich seed medium to the growth-limiting defined medium.

During batch culture of S. clavuligerus, grown under various nutrient conditions in a 16 litre laboratory-scale fermenter, ppGpp was produced either at the beginning of, or during, exponential growth, and prior to the appearance of cephamycin C and clavulanic acid in the fermentation broth. We did not observe ppGpp at the end of exponential growth in any fermenter-grown culture. These results differ from those of Bascaran et al. (1991) for S. clavuligerus; however, these workers used a different growth medium from that used in this study, and they did not sample the cultures until 28 h post-inoculation and may have missed a burst of ppGpp. In all fermentations we consistently detected, by HPLC, the production of a new nucleotide before the appearance of antibiotics in the medium. The new nucleotide was hydrolysed by snake venom phosphodiesterase to pG3'p, which was then further hydrolysed by ryegrass 3'-nucleotidase. This enzyme hydrolysis pattern was identical to that of pure ppGp and hence we attributed the nucleotide to ppGp. It is interesting to note that in contrast to ppGpp, where single peaks are observed, multiple peaks of ppGp are produced; the reason(s) for this is presently unclear.

The production of ppGp was found not to be due to the degradation of ppGpp. It is known that the 3'-β-phosphate residue of ppGpp is sensitive to acid hydrolysis and could provide a non-enzymic route for ppGp production. Figs 3(a) and 5(b) specifically show that ppGpp is not hydrolysed during the acid extraction conditions employed in this study. This was confirmed when we were unable to detect any significant hydrolysis of pure ppGpp following incubation with cell-free extracts of S. clavuligerus.

It has been reported that ppGp is produced following induction of the stringent response in E. coli (Pao & Gallant, 1979) and Bacillus subtilis (Nishino et al., 1979). A similar observation was reported by Strauch et al. (1991) for S. coelicolor following a nutrient shiftdown. We have also observed ppGp production in S. clavuligerus following nutrient shiftdown from an amino-acid-containing medium to a minimal medium. Following shiftdown of S. clavuligerus to a carbon- or phosphate-limiting minimal medium we were unable to detect production of ppGp or ppGpp. This suggests that production of ppGp and ppGpp in S. clavuligerus is in response to an alteration in levels of nitrogen available to the cells.

Immunoblots with polyclonal antibodies to IPNS (Fig. 7a-d) showed that the enzyme is produced following a reduction of the nitrogen or carbon concentration in the growth medium and immediately after production of ppGp. It remains to be elucidated whether there is a positive relationship between production of ppGp and the appearance of IPNS. Earlier work has shown that the biosynthesis of cephamycin C (Brana et al., 1986) and clavulanic acid (Romero et al., 1984) are strongly regulated by nitrogen sources. On the basis of data from the fermentation experiments and nutrient shiftdowns, it is intriguing to speculate that the production of ppGp may

**Fig. 7.** Western immunoblot analysis of proteins in cell-free extracts of S. clavuligerus: (a) FM, (b) nitrogen limitation, (c) carbon limitation and (d) phosphate limitation. Samples were transferred to nitrocellulose membranes, which were incubated with a 1:500 dilution of IPNS antiserum followed by a 1:3000 dilution of goat anti-rabbit lgG alkaline phosphatase conjugate; 50 μg protein was loaded in all experiments. Times indicated are hours of fermentation. The control (lane C) in each experiment was cell-free extract derived from E. coli DS941 expressing pMZRl00, which contains the IPNS gene. Molecular mass standards are prestained low range (Bio-Rad). The molecular mass of IPNS is 35 kDa.

Inoculation into the nutritionally more favourable FM (Fig. 1a, Fig. 2). This difference is attributed to a period of adjustment required by the organism following transfer
be involved in nitrogen regulation of antibiotic biosynthesis in streptomycetes.

Many questions are now raised concerning the role of ppGp: how it is produced and its relationship with ppGpp and the protein biosynthetic machinery in *S. clavuligerus*. Preliminary experiments have demonstrated that 70S ribosomes prepared from *S. clavuligerus* are able to produce ppGpp and ppGp; however following a high-salt wash, ppGpp production is significantly diminished. After further salt washing the ribosomes still retain the activity to produce levels of ppGp similar to those observed from control ribosomes (results not shown); it is therefore tempting to suggest that ppGp synthesis is not overtly associated with any ppGpp synthetic activity. These results suggest that the ppGp levels observed in our fermentation experiments are physiologically representative and that production is associated with the ribosomes, possibly triggered by nitrogen depletion.

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