Streptomyces albus G Produces an Antibiotic Complex Identical to Paulomycins A and B

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(Received 8 December 1986)

An antibiotic complex active against multiply resistant strains of staphylococci and other Gram-positive bacteria was isolated from cultures of Streptomyces albus G. Silica gel and Sephadex LH-20 column chromatography gave two congeners with $M_r$ values of 786 and 772, which differed by one $-\text{CH}_2-$ group. The two homologues contained an isothiocyanate group, and proved to be identical with paulomycins A and B produced by Streptomyces paulus; the FAB mass spectra, in addition, proved the same two congeners to be present in proceomycin obtained from Streptomyces alboniger.

INTRODUCTION

Although very many independently isolated streptomycetes have been described, mainly in connection with their production of antibiotics, only a few have received repeated attention in the scientific literature for reasons other than antibiotic production. Nevertheless, information about other aspects of an organism (e.g. its ecology, physiology and genetics) may eventually provide a rich background within which to analyse antibiotic production. A notable example is provided by Streptomyces coelicolor A3(2), the subject of extensive genetic analysis for about 20 years before it was found to produce four different antibiotics, three of which have since become the focus of molecular genetic studies (see Hopwood et al., 1986).

In the case of S. albus G, discovered by Welsch (1936), some genetic analysis has begun, following the isolation of restrictionless mutants such as J1074 (Chater & Wilde, 1976, 1980), which facilitate its use as an acceptor of DNA from other sources. Protoplasts of mutants can be efficiently regenerated (Baltz & Matsushima, 1981), and can be transformed by several plasmid (Kieser et al., 1982; Rhodes et al., 1984; Bailey et al., 1986) and phage (Chater et al., 1982) vectors. The early studies of S. albus G, however, were primarily biochemical (reviewed by Welsch, 1962): they concerned its bacteriolytic activities, reflected in its renaming as S. griseus subsp. solvificiens (Pridham & Tresner, 1974), although adherence to the old name has persisted in the literature. Part of the lytic principle, a $\delta\delta$-carboxypeptidase, has recently been intensively studied (e.g. Joris et al., 1983). The early studies (Welsch, 1941) had shown that the enzymic lytic activities were accompanied by 'material of a lipoid nature specifically bactericidal for Gram-positive bacteria'. In the course of genetic studies, we noticed that some mutants of S. albus G were sensitive to a diffusible product of the wild-type cultures. In this study the product was purified and identified, and its activity against other Gram-positive bacteria was determined.

METHODS

Strains. The type strain of S. albus G was originally obtained from Dr R. J. Roberts. After treatment with N-methyl-N'-nitro-N-nitrosoguanidine (NTG) (Delić et al., 1970), a mutant (J1670) with increased production of the antibiotic was selected and used in this study.

Abbreviation: NTG, N-methyl-N'-nitro-N-nitrosoguanidine.
**Results and Discussion**

In the course of the genetic studies of the restriction and modification of certain bacteriophages by the SalGI (= SalI: Arrand et al., 1978) system of *S. albus* G (Chater & Wilde, 1980), we observed that growth of a restriction- and modification-deficient mutant (J1074) was inhibited by the parent strain. We found that most other streptomycetes were also inhibited (*S. rimosus* NRRL 2234 being a rare exception), as were *Staph. aureus* and *Bacillus cereus*, but not *Escherichia coli*. An unclassified wild-type *Streptomyces* strain (1193) was chosen as a particularly sensitive indicator strain to aid preliminary purification. Partially purified material (initially termed 'salgiomycin') was obtained which very strongly inhibited growth of strain 1193 and *Staph. aureus*.

In view of its unusually potent anti-staphylococcal activity, 'salgiomycin' was extensively purified as described in Methods from culture fluids of an over-producing mutant obtained after NTG mutagenesis. Its antimicrobial activity is indicated in Table 1.

The IR spectrum (in chloroform) of 'salgiomycin' displayed bands at 3400, 2920, 2030, 1735, 1730, 1690, 1640 and 1580 cm\(^{-1}\), of which the strong absorption band at 2030 cm\(^{-1}\) was indicative of a X=Y=Z cumulene such as N=C=S. UV maxima (\(A^{1}\%_\text{cm}\)) in ethanol were at 237 nm (154-8), 278 nm (109-6) and 324 nm (96-7). The H-NMR spectrum (90 MHz) indicated the presence of several C-CH\(_3\) functions (\(\delta_{1}20-140\)), one methyl group on an olefinic carbon (\(\delta_{1}90\)), protons of an acetate methyl group (\(\delta_{2}07\)), and one methoxy function (\(\delta_{3}44\)) along with additional resonances upfield and downfield from 5-0 p.p.m. The mass spectrum (electron impact) included fragments of \(m/z\) 514, 496, 371, 315, 143, 112, 111 and 93 with no evidence of a molecular ion.

For the definitive structural assignment the availability of FAB mass spectrometry proved invaluable. First, evidence of the presence of homologues with \(M_{r}\) values of 786 and 772, corresponding to molecular formulae \(C_{34}H_{46}N_{2}O_{7}S\) and \(C_{33}H_{42}N_{2}O_{7}S\) respectively, was
readily obtained, based on (i) positive quasi-molecular peaks generated in glycerol–thioglycerol and alkali salts

\[
m/z 795 \ [M + Na]^+; \ m/z 903 \ [M + Na + thioglycerol]^+; \ (R = H)
\]
\[
m/z 809 \ [M + Na]^+; \ m/z 917 \ [M + Na + thioglycerol]^+; \ (R = CH_3)
\]
\[
m/z 811 \ [M + K]^+ \ (R = H)
\]
\[
m/z 825 \ [M + K]^+ \ (R = CH_3)
\]

and (ii) negative radical ions \(m/z\) 772 (R = H) and \(m/z\) 786 (R = CH_3). Subsequent formation of \(m/z\) 754 and \(m/z\) 768, and of \(m/z\) 753 and \(m/z\) 767 respectively, after the loss of an H_2O molecule and a proton, further verified the two \(M_1^q\) values. At this point, a close resemblance was apparent between ‘salgiomycin’ and the paulomycin A and B complex produced by \(S.\) paulus (Wiley et al., 1984), and the fragmentation patterns allowed the deduction of structural features which could be compared and matched with the independently obtained paulomycin data. These data are presented in Fig. 1. In the positive ion FAB mode mass spectra, the quasi-molecular ion (1) gave as prominent fragments \(m/z\) 515 and \(m/z\) 497 assigned to the monoglycoside (2) following the loss of the deoxysugar and a proton back transfer. This pair of peaks establishes the carbon skeleton common to both congeners. The removal of paulic acid \([\text{CH}_3\text{CH}=\text{C}(\text{NCS})\text{COOH}]\) and a molecule of H_2O resulted in the formation of fragments \(m/z\) 372 and \(m/z\) 354 respectively. The moiety of the esterified dideoxysugar cleaved off without the glycosidic oxygen atom (3) was observed as \(m/z\) 273 and \(m/z\) 259 relative to either of the acyl homologues present as part of the pertinent ester group as well as \(m/z\) 241 and \(m/z\) 227 (4) after a loss of CH_3OH. Evidence for the latter structural feature comes from the negative ions observed at \(m/z\) 101 (R = CH_3) and \(m/z\) 87 (R = H) consistent with the 2-methylbutyryl and isobutyryl carbon chains as the only structural difference between the two components of the complex.

Occurrence of a pyranose ring in the dideoxysugar was also supported by fragments \(m/z\) 143 (5) and \(m/z\) 111 (6) in the electron impact spectrum (Fig. 2).

Finally, the negative ions \(m/z\) 142, \(m/z\) 59 and \(m/z\) 58 seen in the FAB spectrum complete the structural assignment as being ascribed to \(\text{CH}_3\text{CH}=\text{C}(\text{NCS})\text{COO}^-\), \(\text{CH}_3\text{COO}^-\) and NCS^- respectively. A sample of proceomycin, an antibiotic complex from \(S.\) alboniger (kindly provided by Dr H. Kawaguchi), gave a similar FAB mass spectrum with different proportions of its two components.

These observations show that the two congeners from \(S.\) albus G with \(M_1^q\) values of 786 and 772 are identical to those in proceomycin (Tsukiu et al., 1964) and also to those in the antibiotic U-43,120 (Wiley, 1976), subsequently resolved into paulomycin A and B (Wiley et al., 1984). These antibiotics represent a class of antimicrobials that may include senfolomycins A and B (Mitscher et al., 1966). At present, there is a lack of information about the sequence of reactions involved in the assembly of these metabolites, although continued work on paulomycins (Wiley et al., 1986) indicates that they may be of more than academic interest. The
finding that a representative of this class is produced by *S. albus G* opens up the possibility of a genetic approach to the study of the biosynthetic pathway and its regulation. This could exploit the large collection of single and multiple auxotrophs in our collection of *S. albus G* derivatives (Chater & Wilde, 1980 and unpublished), among which two isoleucine-valine auxotrophs (including the parent of J1074) and a guanine auxotroph are non-producers (K. F. Chater, unpublished). It is pertinent in this context to note that *S. albus G* strain J1074 is one of the few streptomycetes which have been successfully transformed with large pieces of *Streptomyces* DNA cloned in *E. coli* using bifunctional cosmid vectors (Rhodes et al., 1984), and that the only documented case in which expression of the pBR322 *tet* gene was usable for vector selection in a streptomycete also used this strain (Chater et al., 1982).

We thank Professor D. A. Hopwood for providing J. M. with the opportunity to work at the John Innes Institute and Drs D. Noble, J. Herrmann and W. P. Blackstock, Glaxo, Greenford, Middlesex for carrying out FAB mass spectrometry. Thanks are also due to Dr H. Kawaguchi of the Bristol-Banyu Research Institute, Tokyo, for a sample of proceomycin, Dr P. F. Wiley for antibiotic U-43,120, Dr J. B. McAlpine for critical reading of the manuscript and the Upjohn Company Infectious Diseases Research Office (Mr R. L. Keene) for specimens of paulomycin A and B. This work was funded in part by the Office of Research and Sponsored Projects of the Northwestern University.
REFERENCES


