An O-Phosphotransferase Catalyzes Phosphorylation of Hygromycin A in the Antibiotic-Producing Organism *Streptomyces hygroscopicus*\(^\dagger\)

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The antibiotic hygromycin A (HA) binds to the 50S ribosomal subunit and inhibits protein synthesis in gram-positive and gram-negative bacteria. The HA biosynthetic gene cluster in *Streptomyces hygroscopicus* NRRL 2388 contains 29 open reading frames, which have been assigned putative roles in biosynthesis, pathway regulation, and self-resistance. The *hyg21* gene encodes an O-phosphotransferase with a proposed role in self-resistance. We observed that insertional inactivation of *hyg21* in *S. hygroscopicus* leads to a greater than 90% decrease in HA production. The wild-type and the *hyg21* mutant were comparably resistant to HA. Using *Escherichia coli* as a heterologous host, we expressed and purified Hyg21. Kinetic analyses revealed that the recombinant protein catalyzes phosphorylation of HA (30 ± 4 μM) at the C-2′ position of the fucofuranose ring in the presence of ATP (Kₚ = 200 ± 20 μM) or GTP (Kₚ = 350 ± 60 μM) with a kₗₚ of 2.2 ± 0.1 min⁻¹. The phosphorylated HA is inactive against HA-sensitive ΔtolC *E. coli* and *Streptomyces lividans*. Hyg21 also phosphorylates methoxyhygromycin A and desmethylenehygromycin A with kₗₚ values similar to those observed with HA. Phosphorylation of the naturally occurring isomers of 5″-dihydrohygromycin A and 5″-dihydromethoxyhygromycin A was about 12 times slower than for the corresponding non-natural isomers. These studies demonstrate that Hyg21 is an O-phosphotransferase with broad substrate specificity, tolerating changes in the aminocyclitol moiety more than in the fucofuranose moiety, and that phosphorylation by Hyg21 is one of several possible mechanisms of self-resistance in *S. hygroscopicus* NRRL 2388.

Microorganisms that produce antibiotics are also equipped with resistance mechanisms to protect themselves from the toxic effects of their own molecules (4, 11). Three major mechanisms of resistance have been described in antibiotic-producing *Streptomyces* spp.: (i) antibiotic inactivation by chemical modification, (ii) target site modification, and (iii) antibiotic efflux by transporter proteins. The genes responsible for self-resistance are commonly present as components of antibiotic biosynthetic gene clusters and are often coregulated with them (14). A bacterium may make use of more than one of these intrinsic resistance strategies to prevent autotoxity. Self-resistance by enzymatic modification of an antibiotic is primarily mediated by O phosphorylation of hydroxyl groups and N acetylation of amino groups and is quite widely seen in aminoglycoside producers (4). These bacteria often doubly modify the antibiotic by phosphorylation and acetylation for optimal resistance, as in the neomycin producer *Streptomyces fradiae* (20) and the paromomycin producer *Streptomyces rimosus* (17). Inactivation by O glycosylation has been reported in producers of macrolides such as oleandomycin (21), spiramycin (6), and pikromycin (25). Modification can occur on the final product but is also observed on pathway intermediates/precursors during antibiotic biosynthesis. For example, *straA* in the streptomycin biosynthesis gene cluster encodes streptomycin-6-phosphotransferase, which acts on streptidine, an intermediate in the biosynthetic pathway, and produces an inactive, phosphorylated derivative (15, 22). The remaining biosynthetic pathway utilizes this phosphorylated precursor and leads to the production of 6′-phospho-streptomycin. An extracellular phosphatase is responsible for the formation of the active streptomycin (13).

Antibiotic inactivation by O phosphorylation is the second most prevalent mechanism, after N acetylation, by which many pathogenic bacteria have gained resistance to a large number of drugs, especially the aminoglycosides (19). Several O-phosphotransferases from clinical isolates have been overexpressed, purified, and characterized (24), and it has been proposed that the pathogens must have acquired these antibiotic inactivating kinases, as well as the other resistance genes, from antibiotic producers themselves by horizontal gene transfer (10, 11). Consequently, an understanding of the mechanisms of self-resistance in antibiotic-producing microorganisms is an important step in gaining insights into, and thereby combating, acquired resistance in pathogenic strains.

*Streptomyces hygroscopicus* NRRL 2388 produces the antibiotics hygromycin A (HA) and methoxyhygromycin A (compound 1) (Fig. 1). These antibiotics are structurally unrelated to the more widely known hygromycin B (9). The antimicrobial activity of HA arises from binding to the 50S ribosomal subunit and inhibition of the peptidyl transferase reaction of protein synthesis (7, 9). The structure of HA is composed of three distinct moieties, a 5-dehydro-α-L-fucofuranose (A subunit) attached by a glycosidic bond to (E)-3-(3,4-dihydroxyphenyl)-2-methylacrylic acid (B subunit), which in turn is attached by a peptide bond to 2L-2-amino-2-deoxy-4,5-O-methylene-neoisositol (C subunit). Feeding experiments with precursor compounds revealed that the biosynthetic route to HA has three convergent branches made of a combination of pentose sugar (A subunit), polyketide (B subunit), and aminocyclitol (C sub-

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The biosynthetic gene cluster of HA, which has recently been reported (9, 16), is 31.5 kb long and comprises 29 open reading frames (ORFs). Putative functions have been assigned to 26 of the hyg genes by sequence analysis. Hyg26 has been shown to be responsible for generation of the 5-dehydro-L-fucofuranose moiety by oxidation of L-fucofuranose. A hyg26 mutant has been shown to produce 5-dihydromethoxyhygromycin A (compound 2a), 5-dihydrohygromycin A (compound 2a), and (E)-3-(3-hydroxy-4-O-fucofuranosylphenyl)-2-methylacrylic acid (compound 4, 5-dihydrohygromycin A lacking the aminocyclitol) (see Fig. 1 for structures) (16).

With regard to self-resistance, analysis of the hyg gene cluster suggests that all three mechanisms outlined above may operate in S. hygroscopicus NRRL 2388. The hyg21 gene encodes a putative phosphotransferase that may inactivate HA by O phosphorylation. The hyg6 and hyg29 are methyltransferase homologs, and one of these may confer resistance by methylation of rRNA, the target for HA, while the other is presumed to be responsible for introduction of the methylene group in the aminocyclitol moiety. The hyg19 and hyg28 genes putatively encode a transmembrane protein and an ABC transporter, respectively, which may play a role in antibiotic efflux. There has been no experimental verification of these proposed roles for Hyg19, Hyg21, and Hyg28.

In the present study, we report a genetic and biochemical investigation of hyg21 and its gene product. We have shown that hyg21 is not an absolute requirement for hygromycin biosynthesis and that Hyg21 is a phosphotransferase that catalyzes the transfer of the γ-phosphoryl group of ATP to the 2-hydroxyl group of HA, abolishing its antibacterial property. We have also determined the steady-state kinetic parameters for turnover of HA and its analogs and have shown that Hyg21 can phosphorylate a range of substrates (Fig. 1) bearing a fucofuranose moiety, including compounds 1, 2, 3, 4, desmethylenhygromycin A (compound 5), 5-dihydromethoxyhygromycin A (compound 6), and antibiotic A201A (compound 7).

MATERIALS AND METHODS

Antibiotics and chemicals. All antibiotics and chemicals were purchased from Sigma-Aldrich (St. Louis, MO), unless otherwise indicated. HA was kindly supplied by Pfizer, Inc. Analogs of HA and compounds 1, 2a, 3a, and 4 (Fig. 1) were isolated from the wild type and a Δhyg26 blocked mutant strain of S. hygroscopicus (16). Compounds 5 and 6 were isolated from a hyg6 deletion mutant (unpublished data). A diastereomeric mixture of 5-dihydromethoxyhygromycin A (compounds 2a and 2b) was prepared by reduction of HA as described previously (16). A diastereomeric mixture of 5'-dihydromethoxyhygromycin A (compounds 3a and 3b) was prepared from methylenhygromycin A (compound 1) in the same way. Antibiotic A201A was a generous gift from A. Jimenez, Centro de Biología
Molecular Severo Ochoa, Madrid, Spain. [2-H]Hnpy-inositol (20.0 Ci/mmol) was procured from Moravek Biochemicals (Brea, CA). Primers for PCR were obtained from Integrated DNA Technologies (Coraville, IA). Enzymes for DNA manipulations was purchased from New England Biolabs (Beverly, MA). The prestained protein molecular weight markers in the sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) analysis were obtained from Bio-Rad (Hercules, CA).

Bacterial strains, plasmids, and culture conditions. Cosmid 17E3, based upon the SuperCos1 vector, was the source for hyg21 gene and has been described previously (16). The pCR4-TOPO vector and One-Shot MachI-T1 chemically competent Escherichia coli cells from Invitrogen (Carlsbad, CA) were used for cloning the hyg21 PCR product. The PET15b vector from EMD Biosciences (San Diego, CA) was used for the expression of N-terminal His-tagged Hyg21 protein. The PET-15b vector with 0.5-kb hyg21 ORF was designated PET15b-hyg21. E. coli BL21-CodonPlus (DE3), obtained from Stratagene (La Jolla, CA), was the expression host for His-tagged Hyg21 protein. E. coli BW25113/pLJ790 (a strain with the λ-red recombination plasmid), E. coli DH5α/pU773 (a strain with the aac(3)IV ampicillin resistance cassette plasmid), and E. coli ET12567/pUZ8002 (a nonmethylating plasmid donor strain for intergeneric conjugation) have been described previously (8) and were provided by the John Innes Institute (Norwich, UK). Bacterial strains, plasmids, and culture conditions. NRRL 2388 is the wild-type HA producer strain with the aac(3)IV gene and has been described previously (8).

DNA manipulations and analyses. Plasmid DNA was isolated by using a QIaprep Spin Miniprep kit from Qiagen (Valencia, CA). DNA fragments from agarose gels were isolated by using Qiagen’s QIAquick gel extraction kit. The QIAquick PCR purification kit was used to clean up the PCR product. Genomic DNA from Streptomyces strains was obtained by using the Wizard Genomic DNA purification kit from Promega (Madison, WI). Automated DNA sequencing was performed at the DNA core facilities at Virginia Commonwealth University and Virginia Commonwealth University at the Expy proteomics database at the Expy proteomics server (12).

Cloning of hyg21 and construction of the expression plasmid. The 552-bp hyg21 gene was amplified from cosmid 17E3 by using the GC-Rich PCR system from Roche Diagnostics according to the manufacturer’s instructions. Primers (5'-CATATGCCGGAATCTGCCTGAGCCAGGC-3') and 5'-GGATCTCTGAGCGTCGCTGCAGAT-3') were designed to create an NdeI restriction site (in boldface) 3' to the hyg21 gene and have been described (9). Two-dimensional 1H correlation spectra were recorded on a Bruker AMX-400 NMR spectrometer. A standard HA sample was run under identical conditions. Two-dimensional 1H correlation spectra were recorded on a Bruker AMX-400 NMR spectrometer. A standard HA sample was run under identical conditions for comparison. Coupling constants (J) were expressed in hertz. Abbreviations for multiplicities are as follows: s, singlet; d, doublet; t, triplet; q, quartet; m, multiplet.

HA-P in D2O. Values for HA in D2O were as follows: 'H NMR (500 MHz, CDCl3), 6.73-4.5 Hz, 7.16 (s, 1H), 7.08 (s, 1H), 7.06 (s, 1H), 5.95 (d, J = 2.5 Hz, 1H), 5.31 (s, 1H), 4.97 (s, 1H), 4.69 (m, 1H), 4.63 (t, J = 4.5 Hz, 1H), 4.58 (d, J = 5.6 Hz, 1H), 4.36-4.31 (m, 2H), 4.20 (dd, J = 3.5, 9.5 Hz, 1H), 4.08 (dd, J = 5.0, 9.5 Hz, 1H), 3.95 (t, J = 4.5 Hz, 1H), 2.25 (s, 3H), and 2.12 (s, 3H).

HPLC and mass-spectrometry (MS) conditions. Reverse-phase HPLC analyses of the phosphorylation assays and the fermentation broths were carried out by using an Agilent 1100 Series HPLC system with either an analytical 5-μm Phenix C8 column (250 by 10 mm). The solvents used were A (10% methanol, 90% water, 0.05% formic acid) and B (90% methanol, 10% water, 0.05% formic acid). Gradient conditions for the analytical column were 100% A for 2 min, 100% to 36% A for 16 min, 36% to 0% A for 2 min, 0% for 8 min, to 100% A for 2 min, and 100% A for 10 min at a flow rate of 1 ml/min. Gradient conditions for the semipreparative column were 100% A for 2 min, 100% to 44% A for 16 min, 44% to 0% A for 2 min of NTA spin protocol. All of the protein purification steps were carried out at 4°C. Cells were disrupted by sonication, and the soluble fraction was applied to 1 ml of Ni-NTA agarose preequilibrated with lysis buffer. The column was washed with 50 ml of wash buffer (50 mM NaH2PO4, 300 mM NaCl, 20 mM imidazole [pH 8.0]), and the protein was eluted with elution buffer containing 250 mM imidazole in fractions of 500 μl. Each fraction was analyzed by SDS-PAGE using a 13% acrylamide gel for the presence of Hyg21. The eluted protein was dialyzed for 14 h against a buffer containing 50 mM Tris-HCl (pH 7.5), 20% glycerol and 5 mM 2-mercaptoethanol, with a replacement with fresh buffer after 4 h. The protein concentration was determined by Bio-Rad protein assay using bovine serum albumin as standard, and 50-μl aliquots of the purified protein were stored at −80°C until further use.

Phosphorylation assay conditions. Initial enzyme assays were performed with −10 μg of purified enzyme in a 100-μl reaction mixture of 50 mM Tris-HCl (pH 7.5), 5 mM 2-mercaptoethanol, 10 mM MgCl2, 200 μM substrate, and 1 mM ATP. The reaction mixture was incubated at 30°C for 2 h and frozen at −20°C. Hyg21 activity was studied by reverse-phase high-pressure liquid chromatography (HPLC) analysis by monitoring the appearance of phosphorylated product, which had a shorter retention time than the substrate. For kinetics studies, the reaction volume was scaled up to 600 μl with −14 μl of enzyme. The assays were carried out for 15 min, during which phosphorylation activity was linear. The ATP concentration was fixed at 400 μM, and the antibiotic concentration was varied from 5 to 100 μM. To determine kinetic parameters for ATP and GTP, the concentrations of these were varied, and the concentration of HA was fixed at 90 μM. In all cases, the samples were analyzed by HPLC, and the amount of product formed was determined from its peak area using an HA standard curve as reference. The rate of phosphorylation was reported as the nanomoles of product formed per minute. Kinetic data were plotted in a Lineweaver-Burk format and determined using NONLIN software as the kinetic parameters. The pH profile was obtained using 100 μM HA and 1,000 μM ATP at 30°C for 15 min in the following buffers: 50 mM potassium acetate, pH 5; 50 mM Tris-maleate, pH 6 to 6.5; and 50 mM Tris-HCl, pH 7, 7.5, 8, or 9. The effect of temperature on Hyg21 activity was checked using 100 μM HA and 1,000 μM ATP in 50 mM Tris-HCl (pH 7.5) at room temperature (23°C) and at 30, 37, and 50°C in a 15-min assay.

Expression and purification of His-tagged Hyg21 protein. Ten reaction mixtures (5 ml each) with a final composition of 200 μM HA, 1,000 μM ATP, 10 mM MgCl2, 5 mM 2-mercaptoethanol, and 0.5 mg of enzyme in 50 mM Tris-HCl (pH 7.5) were incubated at 30°C. The assay mixtures were pooled and filtered by using Milllex-LCR 0.45-μm-pore-size filter unit from Millipore, and the phosphorylated HA (HA-P) was purified by semi preparative reversed-phase HPLC. Fractions containing the purified product were pooled and dried by rotary evaporation. The dried sample was dissolved in deuterium oxide for structural elucidation by nuclear magnetic resonance (NMR) analysis. 1H NMR spectra were recorded on Nicolet NM-500 MHz (modified with a Tecmag Libra interface) instruments calibrated using residual deuterated solvent as an internal reference. Two-dimensional 1H correlation spectra were recorded on a Bruker AMX-400 NMR spectrometer. A standard HA sample was run under identical conditions for comparison. Coupling constants (J) were expressed in hertz. Abbreviations for multiplicities are as follows: s, singlet; d, doublet; t, triplet; q, quartet; m, multiplet.
Insertional inactivation of hyg21. The hyg21 gene of S. hygroscopicus NRRL 2388 was replaced by the apramycin resistance cassette using the PCR-targeted Streptomyces gene replacement method (8). The Expand High Fidelity kit from Roche was used for PCR amplification of apramycin resistance cassette from pJ773 with the primers hyg21_KO_Forw (5'-ACCCCTGAGCCGAGGTTAAAGGACA CAGAAAACATGATTTCCGGGATGTCGACC-3') and hyg21_KO_Rev (5'-TCCAGCGGACCGGGCCGCACTGATAGGCTTG ATGCTGTC-3') (boldfacing indicates homology to the nucleotide sequence flanking hyg21, while the italicized portions of the sequences are homologous to pJ773). Gene replacement was carried out first in cosm I 17E3 and then in the genome of S. hygroscopicus. Gene replacement in the resulting mutant strain, Δhyg21, was confirmed by PCR amplification of chromosomal DNA using the outer primers, hyg21_Out_Forw (5'-CCGGTTTCGTCAGTACACG ATG-3') and hyg21_Out_Rev (5'-CATGCGGACCTGCTGAC-3') and by sequencing the PCR product.

Fermentation conditions for S. hygroscopicus strains. S. hygroscopicus NRRL 2388 and Δhyg21 mutant strains were cultivated as reported earlier (9). To obtain cell lysates, mycelium was collected from 100 ml of culture, suspended in 5 ml of methanol, and shaken on an orbital shaker at room temperature for 6 h. The suspension was then subjected to a brief sonication, and cell debris was removed by centrifugation.

Feeding of 2-3H-labeled myo-inositol. Feeding studies using radiolabeled precursors were carried out in 5 ml of production medium inoculated with 200 μl of seed culture. [2-3H]myo-inositol was added after 24 h to a final concentration of 0.4 μCi/ml.

Disk diffusion assays to test the bioactivity of HA-P. The bioactivity of HA-P was determined by antibiotic diffusion method using an efflux pump deficient ΔtolC E. coli strain as test organism. Fifteen milliliters of LB agar was overlaid with 1.5 ml of culture. The plate was examined for the appearance of zones of inhibition. The assay was also repeated with Hyg21-catalyzed phosphorylation of HA.

Hygromycin sensitivity of S. hygroscopicus and the Δhyg21 strain. The MIC50 of HA for the wild type and the Δhyg21 mutant were determined for the agar plate dilution method. Portions (30 μl) of spore suspensions of both the strains (2,000 CFU/ml) were plated on 3-ml ISP2 agar plates containing 200, 300, or 400 μg of purified HA-P. After incubation for 15 h at 37°C and 30°C, the plates were examined for the zones of inhibition after 15 h of incubation at 30°C.

Sequence analysis of Hyg21. The hyg21 ORF is 552 nucleotides long, with a putative translation product of 183 amino acids and a theoretical molecular mass of 19.87 kDa. A search of the NCBI protein sequence database using the web-based BLASTP program revealed that the deduced amino acid sequence is similar to the deduced amino acid sequence of hyg21.

Optimal pH for Hyg21 activity was measured across a range of pH values from 6 to 9. The fastest rate of reaction, determined by the amount of product formed per minute, was in the range of pH 7.5 and 8. Activity was markedly decreased at lower pH values and almost absent at pH 5. The effect of temperature on activity was also studied across a range of 23 to 50°C. In a 15-min assay, product formation was maximal at 30°C. The enzymatic activity was maintained for several hours at 50°C.

Bioassay for antibiotic activity of HA-P. The activity of HA-P was tested in a disk diffusion assay against an HA-sensitive ΔtolC E. coli strain that has an impaired efflux mechanism due to the absence of the outer membrane component of the efflux pump (5). As seen in Fig. 3, distinct zones of inhibition were seen with HA. In contrast, no zones of inhibition were seen with HA-P, even at 60 μg. Similar observations of antibacterial activity for HA but not HA-P were observed in disk diffusion assays against HA-sensitive S. lividans. These observations are consistent with the hypothesis that phosphorylation abolishes the antibiotic potency of HA and also indicate the likely function of hyg21 as a self-resistance determinant in S. hygroscopicus NRRL 2388.

Site of phosphorylation in HA. 1H NMR analyses of purified HA (5 mg) and HA-P (2 mg) were carried out to determine the site of phosphorylation by Hyg21. The chemical shifts of GSGKS. This glycine-rich motif is often found in nucleotide-binding proteins and is known to form a flexible loop that interacts with ATP or GTP (18, 23). On the basis of these observations we hypothesized that the hyg21 gene encodes a kinase involved in the O phosphorylation of HA.

Hyg21-catalyzed phosphorylation of HA. The PCR-amplified hyg21 ORF was cloned into E. coli expression vector pET15b to give pET15b-hyg21 plasmid and overexpressed in BL21-CodonPlus cells as an N-terminal His6-tagged protein. A large fraction of the expressed protein formed inclusion bodies under the given expression conditions, but it was possible to recover substantial amounts of soluble protein using a 1-liter culture and affinity chromatography purification on a Ni-NTA agarose column. SDS-PAGE analysis of the purified preparation demonstrated a prominent protein band with a relative molecular mass of 22 kDa, a finding consistent with the mass of Hyg21 (~20 kDa) containing a His6 tag (~2 kDa). In order to determine the biochemical function of Hyg21, the recombinant protein was incubated with HA and ATP at 30°C for 2 h. Reversed-phase HPLC analysis of the reaction mixture after incubation revealed a significant decrease in the HA peak and the appearance of a new peak with a shorter retention time (Fig. 2B). In a control incubation without Hyg21, this new peak was absent, and the levels of HA did not alter (Fig. 2A). The shorter retention time for the new peak was consistent with increased polarity of HA through the addition of a phosphoryl group. MS analysis of the reaction mixture showed that the compound under the new peak had a molecular mass of 592 ([M+H]+) (Fig. 2C). This increase of 80 atomic mass units from that of HA (512 [M+H]+) corresponds to the addition of a single phosphoryl group to the molecule. These data support the hypothesis that hyg21 encodes a phosphotransferase that carries out monophosphorylation of HA.

The presence of 5 mM 2-mercaptopethanol in assays of HA-P was found to be necessary for maintaining optimal activity. The optimal pH for Hyg21 activity was measured across a range of pH values from 6 to 9. The fastest rate of reaction, determined by the amount of product formed per minute, was in the range of pH 7.5 and 8. Activity was markedly decreased at lower pH values and almost absent at pH 5. The effect of temperature on activity was also studied across a range of 23 to 50°C. In a 15-min assay, product formation was maximal at 30°C. The enzyme retained two-thirds of its activity at room temperature and 37°C but very little at 50°C.
the two samples were virtually identical, with the exception of the proton signal for the C2\textsuperscript{\textgreek{m}} methine (H2\textsuperscript{\textgreek{m}}), which is shifted downfield from 4.36 ppm for HA to 4.69 ppm in HA-P. In each case the H2\textsuperscript{\textgreek{m}} assignment was confirmed by a \textsuperscript{1}H correlation spectroscopy experiment, which revealed a strong cross peak with the distinctive anomeric H1\textsuperscript{\textgreek{m}} resonance. The chemical shift change for the H2\textsuperscript{\textgreek{m}} alone is consistent with phosphorylation of the C2\textsuperscript{\textgreek{m}} hydroxyl substituent (Fig. 1).

**Substrate specificity of Hyg21.** The substrate specificity of Hyg21 was explored by carrying out enzyme assays with various analogs of HA (compounds 1 to 6), A201A (compound 7), and puromycin (compound 8) (Fig. 1). HPLC-MS analyses revealed that, with the exception of compound 8, all of these compounds were partially or completely converted to new products with shorter retention times and with masses increased by 80 AMU (Table 1). Notable among these observations was that antibiotic A201A (compound 7), which is much larger than HA but has structurally similar fucofuranose and (E)-3-hydroxyphenyl)-2-methylacrylic acid moieties, was also a substrate for Hyg21. In addition, compound 4, which lacks the aminocyclitol and thus is significantly smaller than HA, is also a substrate. In both cases the substrate contains a fucofuranose with a C2-hydroxyl substituent. This moiety is missing in compound 8, despite its structural similarities with both HA and antibiotic A201A, and in this case phosphorylation with Hyg21 is not observed. These observations demonstrate that Hyg21 exhibits broad substrate specificity and that the fucofuranose moiety with C2-hydroxyl group is a key structural element required for phosphorylation. The latter inference is consistent with the observation that in HA, Hyg21 catalyzes phosphorylation at C2 of this moiety.

HA phosphorylation was also carried out by substituting ATP with GTP, dTTP (TTP), and CTP. Besides ATP, product formation was seen only with GTP. There was no reaction with either TTP or CTP.

The rate of phosphorylation was determined by measuring the amount of product formed per minute (Table 1). The reaction was linear during the first 15 min of incubation under the given assay conditions. Under these conditions $k_{\text{cat}}$ values of 2.2 min\textsuperscript{-1} and $K_m$ values of approximately 30 μM were obtained for HA. The reaction rate $k_{\text{cat}}$ was the same with either ATP or GTP, although ATP had a lower $K_m$. Similar but slightly slower $k_{\text{cat}}$ rates were observed for ATP-dependent phosphorylation of compounds 1 and 5 by Hyg21. These data...
and similar $K_m$ values indicated that the methylene group of HA, which is present as a methyl group in compound 1 and absent in compound 5, is not important for binding and catalysis.

Compounds 2a (a 5''-dihydro analog of HA) and 3a (a 5''-dihydro analog of methoxyhygromycin A [compound 1]) have previously been identified in fermentation of the Δhyg26 S. hygroscopicus mutant (16). We produced a diastereomeric mix of compound 2a and the other 5''-dihydroisomer (compound 2b) by chemical reduction of HA. HPLC analysis of Hyg21-catalyzed phosphorylation of this mixture (Fig. 4) revealed much faster reaction rates with compound 2b than with compound 2a. Apparent kinetic values for the reaction of each isomer in the mixture were obtained and revealed that compound 2b was processed with reaction rates ($k_{cat} = 1.68 \text{ min}^{-1}$) similar to those of HA, compound 1, and compound 5. In contrast, the reaction rate is more than 10-fold slower ($k_{cat} = 0.11 \text{ min}^{-1}$) with the isomer compound 2a, which appears to be a shunt metabolite in the HA biosynthetic pathway. A similar set of observations was made with a diastereomeric mixture of compounds 3a and 3b (obtained by chemical reduction of compound 1). The phosphorylation rate for the compound 3a isomer is slow ($k_{cat} = 0.11 \text{ min}^{-1}$), but much faster ($k_{cat} = 1.34 \text{ min}^{-1}$) for compound 3b. In contrast to the $k_{cat}$ values, much

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### TABLE 1. Summary of the kinetic data for Hyg21 phosphorylation

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Compound</th>
<th>Mass [M+H]+</th>
<th>Mean ± SD</th>
</tr>
</thead>
<tbody>
<tr>
<td>HA</td>
<td></td>
<td>512 592</td>
<td>2.23 ± 0.13</td>
</tr>
<tr>
<td>Methoxyhygromycin A</td>
<td>1</td>
<td>514 594</td>
<td>1.96 ± 0.13</td>
</tr>
<tr>
<td>Desmethylenehygromycin A</td>
<td>5</td>
<td>500 580</td>
<td>1.89 ± 0.08</td>
</tr>
<tr>
<td>5''-Dihydrohygromycin (natural isomer)</td>
<td>2a</td>
<td>514 594</td>
<td>0.15 ± 0.03</td>
</tr>
<tr>
<td>5''-Dihydrohygromycin (non-natural isomer)</td>
<td>2b</td>
<td>514 594</td>
<td>1.62 ± 0.1</td>
</tr>
<tr>
<td>5''-Dihydromethoxyhygromycin A (natural isomer)</td>
<td>3a</td>
<td>516 596</td>
<td>0.13 ± 0.03</td>
</tr>
<tr>
<td>5''-Dihydromethoxyhygromycin A (non-natural isomer)</td>
<td>3b</td>
<td>516 596</td>
<td>1.31 ± 0.08</td>
</tr>
<tr>
<td>5''-Dihydrodesmethylenehygromycin A (natural isomer)</td>
<td>6</td>
<td>502 582</td>
<td>ND*</td>
</tr>
<tr>
<td>5''-Dihydrodesmethylenehygromycin A (non-natural isomer)</td>
<td>4</td>
<td>363 443</td>
<td>ND</td>
</tr>
<tr>
<td>A201A</td>
<td>7</td>
<td>803 883</td>
<td>ND</td>
</tr>
<tr>
<td>ATP</td>
<td></td>
<td></td>
<td>198 ± 19</td>
</tr>
<tr>
<td>GTP</td>
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<td>356 ± 62</td>
</tr>
</tbody>
</table>

* ND, not determined.
smaller changes in $K_m$ were observed for the various substrates. These kinetic data indicate that Hyg21 can process HA analogs in which the 5-keto group is reduced, but the efficiency of the process is determined by the stereochemistry at the C5. Kinetic studies for substrates 4, 6, and 7 were precluded because of low availability of the substrates and/or the presence of impurities. A comparison of reaction completion with HA and compound 7 carried out under standard conditions indicated that the rate of conversion of compound 7 was at least fivefold slower.

**Generation and analysis of a Δhyg21 S. hygroscopicus mutant.** The hyg21 gene in S. hygroscopicus was replaced by the apramycin resistance gene to assess its role in HA biosynthesis and resistance. A gene replacement strategy was used because the hyg21 is well separated from its flanking genes, and its orientation relative to these indicates that it is not part of a polycistronic transcript (Fig. 5).

The Δhyg21 mutant strain was cultured, and the fermentation broth was analyzed for the production of HA. HA was detected, indicating that hyg21 was not essential for the biosynthetic process. Nonetheless, the yields were dramatically reduced (>90%) from the 0.8 g of HA/liter obtained in the wild-type strain. Detailed LC-MS analyses of fermentation broth of Δhyg21 also revealed the presence of compounds 1, 3a, 5, and 6. The same compounds could be detected by radioactive scintillation counting when the mutant was grown in the presence of tritiated myo-inositol (which is converted into the aminocyclitol portion of HA) (Fig. 6). Although the levels of compounds 1, 3a, 5, and 6 could not be quantitated, the analyses indicated that they were similar in both the wild type and the Δhyg21 mutant, in contrast to the observations made for the levels of HA.

The low amounts of HA in the fermentation broth of Δhyg21 raised the possibility that it is the HA-P that is exported from the S. hygroscopicus by an efflux pump and subsequently converted to HA by an extracellular phosphatase. In such a case HA might be expected to accumulate inside the cells of Δhyg21. To investigate this possibility, the mycelium of Δhyg21 was lysed by methanol treatment, and the lysate was analyzed by HPLC. However, HA could not be detected in the cell lysate, and it was estimated that nearly all of the HA made by this strain is found in the fermentation broth. Also, no HA-P could be detected in either the cell lysate or the fermentation broth of the wild-type strain.

An agar plate dilution assay was carried out to assess whether the Δhyg21 strain had increased sensitivity to HA...
compared to the wild-type strain. The MIC\textsubscript{95}s for both strains were found to be 400 \mu g/ml.

**DISCUSSION**

Analysis of the deduced amino acid sequence of the \(hyg21\) gene in the HA biosynthetic gene cluster led to a hypothesis that it encodes a protein that phosphorylates HA and thus provides a resistance mechanism to the producing organism by antibiotic inactivation. The first component of this hypothesis has been unequivocally verified by in vitro assays in which Hyg21 HA-P selectively at the C2'-OH position. Hyg21 also phosphorylates a number of HA analogs, and the site of modification in these may also be the C2'-OH. Some of these analogs are observed at various levels in fermentations of the wild-type \(S.\ hygroscopicus\) (9, 16). We have also observed that these analogs, in particular methoxyhygromycin A and desmethylhygromycin A, have some level of antibacterial activity, albeit less than that observed for HA. Presumably, resistance to these compounds, as well as to HA, can be provided in part by the Hyg21-catalyzed phosphorylation.

The enzyme-catalyzed process was most efficient with HA, a finding consistent with phosphorylation occurring with the final biosynthetic product, rather than with an intermediate. Nonetheless, the rate of reaction with HA was relatively slow (\(k_{\text{cat}}\) of \(\sim 2\) min\(^{-1}\)). Analyses of \(S.\ hygroscopicus\) fermentations reveal that the majority of the HA product is found in fermentation media, indicating an efficient export process. We see no evidence for any HA-P in the fermentation broth or in the cell lysate, and our analyses of the HA biosynthetic gene cluster to date have not revealed a candidate phosphatase that would convert HA-P to HA (the \(hyg25\) gene product appears to be involved in the conversion of myo-inositol-1-phosphate to myo-inositol, a step in the biosynthesis of the aminocyclitol moiety of HA). Thus, all evidence indicates that it is HA that is exported and that HA-P may be generated from residual HA in the cell. In such a case, a moderately slow rate of phosphorylation by Hgy21 would allow most of the HA biosynthetic product to be exported rather than become trapped intracellularly as HA-P.

The results from these studies do not provide unequivocal evidence for our hypothesis that phosphorylation of HA is a resistance mechanism in the producing organism. We do observe that HA and not HA-P has antibacterial activity against other bacteria. However, it remains to be determined whether the loss of antibacterial activity is associated with poorer penetration into the organisms or inhibition of binding to the ribosomal target. In addition, the \(\Delta hyg21\) mutant retains significant resistance to HA. This observation likely reflects the presence of other operational mechanisms of resistance. The most likely of these are methylation of the tRNA, catalyzed by the \(hyg29\) gene product, and an efficient HA export process, catalyzed by the \(hyg19\) and \(hyg28\) gene products. Characterization of the remaining putative resistance determinants in the HA biosynthetic gene cluster is currently under way. Finally, the observation that insertional inactivation of \(hyg21\) reduces the production of HA to almost negligible amounts is also curious. A polar effect from replacement of the gene cannot be ruled out but seems unlikely given the strategy used and the organization of genes surrounding \(hyg21\) (Fig. 5). This result was consistently observed in all of the four mutant colonies that were examined. A significant decrease in HA production would be predicted in this mutant if the Hgy21-catalyzed phosphorylation process occurred at an earlier step in the biosynthetic process, with the later stages occurring with phosphorylated intermediates. Early modification would ensure that the resistance mechanism is in operation prior to assembly of the final product and has precedent in the biosynthesis of streptomycin (15) and paromomycin (17). The slow turnover of HA might also indicate that a pathway intermediate, and not the
final product, may be the preferred substrate for Hyg21. However, we do not have any genetic evidence for a phosphatase that would be responsible for conversion of the HA-P to HA, we detect only HA in fermentations of the wild-type strain (see above), and HA was the preferred substrate for Hyg21 in our kinetic studies. These observations argue against Hyg21-catalyzed phosphorylation of a pathway intermediate as a step in the biosynthetic process. In the case of tylosin biosynthesis it has been shown that disruption of any of the four genes involved in the biosynthesis of the mycaminose sugar, which is the first sugar that gets attached to tylactone (the polyketide aglycone), abolishes not just tylosin production but also the accumulation of tylactone (3). In a similar way, HA-P may be required for stimulating the HA biosynthetic pathway by positive feedback regulation, accounting for the decreased production in the Δhyg21 mutant.

In conclusion, we have shown that Hyg21 catalyzes phosphorylation of HA and related compounds, leading to the loss of antibacterial activity. Thus, hyg21 is the first of the several possible resistance determinants in S. hygroscopicus NRRL 2388 that has been functionally characterized. The roles of Hyg6 and Hyg29 (methyl transferases) and of Hyg19 and Hyg28 (antibiotic efflux proteins) as additional resistance determinants are currently being investigated.

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REFERENCES