Analysis of an 8.1-kb DNA Fragment Contiguous with the 
Erythromycin Gene Cluster of Saccharopolyspora 
erthrea in the eryCI-Flanking Region 

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An 8.1-kb region of the Saccharopolyspora erythreae genome, significant for its contiguity to the known genes of the erythromycin biosynthetic gene cluster, was mutationally analyzed and its DNA sequence was determined. The region lies immediately adjacent to eryCI. The newly characterized region is notable for a large, 3.0-kb segment, predicted not to be translated, followed by four probable genes: a acetyltransferase gene, a protease inhibitor gene, a methyltransferase gene, and a transposase gene. Because the probable functions of the genes in this region are not required for erythromycin biosynthesis or resistance and because a deletion of a 6.0-kb portion of this region had no effect on erythromycin biosynthesis, this region marks the outside boundary of the erythromycin gene cluster. Therefore, eryCI represents the end of the cluster. These results complete the analysis of the erythromycin gene cluster and eliminate the possibility that additional sought-after pathway-specific structural or regulatory genes might be found within or adjacent to the cluster.

Microorganisms isolated from soil have been used for the production of bioactive molecules such as antibiotics for over a half century. The mycelial gram-positive actinomycetes are an exceptionally rich source of these compounds. Much effort has been devoted to genetic analysis of the antibiotic biosynthesis process (14), which has been greatly aided by the clustering of antibiotic biosynthesis and resistance genes in the chromosomes of these organisms. A related area of interest in this field concerns the need of the producing organism to control the level of production of these toxic molecules in its growth environment. In some antibiotic gene clusters this function is performed by a pathway-specific regulatory gene located within the cluster (2, 4, 10, 11, 20, 26, 35).

Erythromycin is a clinically important macroclide antibiotic; it is also the parent molecule for several commercially successful semisynthetic macrolide and ketolide derivatives (6). As such, it would be of both great biological and commercial interest to understand the regulatory mechanisms operating in Saccharopolyspora erythreae that control the level of production of this important antibiotic.

The erythromycin gene cluster was cloned prior to the advent of automated DNA sequencing; consequently, it was analyzed in fragments by several different laboratories. This could explain how the cluster had come to be referred to as “entirely known” (8, 9) when, in fact, significant information regarding a 6-kb region immediately downstream of eryCI was lacking. This oversight persisted, despite reports in the earlier literature that suggested that this region contained a sought-after pathway-specific regulatory gene(s) (12, 30). Furthermore, this region might have been predicted to contain additional early-stage deoxy sugars biosynthesis genes, for example, a dTDP-glucose synthase or a dTDP-glucose dehydratase, that had been found in the tylosin biosynthetic pathway gene cluster but not yet in the erythromycin gene cluster (19, 28).

In this study, then, we sought to determine the probable identities of the genes contained in this region contiguous with the erythromycin gene cluster and whether they are required for erythromycin biosynthesis. In so doing we completed the analysis of the erythromycin gene cluster, a project that had its beginnings 20 years ago with the cloning of the erythromycin resistance gene (29).

MATERIALS AND METHODS

Strains and plasmids. FL1347, a red variant derivative of S. erythreae ATCC 11635, and BC1, a commercial derivative of S. erythreae ATCC 11635, were used for this study. A list of the strains and plasmids constructed or used in this study is given in Table 1.

Chemicals, biochemicals, and growth media. Erythromycin A, thiostrepton, kanamycin sulfate, ampicillin, and tetrazolium chloride were purchased from Sigma (St. Louis, Mo.). The sporulation agar used for S. erythreae BC1 was E20A agar medium. E20A agar medium per liter of aqueous solution is 5 g of Bacto Soytone (Difco), 5 g of soluble starch, 5 g of CaCO3, 2.1 g of morpholinepropanesulfonic acid buffer, and 20 g of Bacto Agar. R2T2 medium (R2T2 medium [34] without the peptone) was used for protoplast regeneration for all strains of S. erythreae and as a sporulation medium for the FL1347 (red variant) strain of S. erythreae.

Plasmids constructed for DNA sequencing. The eryCI-flanking region was cloned in plasmid pH3 during an earlier study (32). A 6.7-kb HindIII-BglII fragment from pH3 was subcloned into BamHI-HindIII-digested pUC19 to make pFL2000.

The remaining portion of the eryCI-flanking region was cloned from pH3 by PCR to produce a fragment covering the region from the unique BglII site at the 3’ end of open reading frame (ORF) ORF2 (Fig. 1, site 11) to the 3’ end of ORF4. The reaction was performed according to the protocol of the manufacturer with the Expand Long Template PCR kit (Roche Molecular Biochemicals, Indianapolis, Ind.) by using kit buffer 3. The amplification reaction involved a heating step at 98°C for 4 min, followed by 30 cycles at 98°C for 30 s, 55°C for 1 min, and 68°C for 10 min. The PCR fragment was cloned into pCRtopo (Invitrogen, Carlsbad, Calif.) to form pH46. Plasmid pH2070 was constructed to obtain additional nucleotide (nt) sequence information on ORF3. The insert in pH2070 was produced by using primers K16-3 (5’-CGGAATTCCTTGTAGCCCGCTGGAATT-3’ [the EcoRI site is indicated in boldface]) and K17-3 (5’-CAAGGTTCATGCCGGCGCGTAGTCCC-3’ [the HindIII site is indicated in boldface]). The resulting
TABLE 1. Plasmids constructed or used in this study and strains used in this study

<table>
<thead>
<tr>
<th>Plasmid or strain</th>
<th>Parent plasmid</th>
<th>Plasmid insert or strain description</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>pMW3</td>
<td>pU702</td>
<td>Original clone of DNA from eryCI-flanking region of NRRL 32, 32</td>
<td>This study</td>
</tr>
<tr>
<td>pFB46</td>
<td>pCRTopo (Invitrogen)</td>
<td>2338 red variant strain</td>
<td>This study</td>
</tr>
<tr>
<td>pFL2000</td>
<td>pUC19</td>
<td>Contains two PCR fragments from ends of eryCI-flanking region</td>
<td>This study</td>
</tr>
<tr>
<td>pFL2063</td>
<td>pFL8</td>
<td>The Km&lt;sup&gt;+&lt;/sup&gt; gene from pUC4K, originally from Th903 (Amer- sham Pharmacia Biotech), cloned into pFL2063 between clone 1 and clone 2 for deletion of eryCI-flanking region</td>
<td>This study</td>
</tr>
<tr>
<td>pFL2067</td>
<td>pUC19</td>
<td>PCR product for resequencing of ORFL3</td>
<td>This study</td>
</tr>
</tbody>
</table>

Plasmids for deletion of eryCI-flanking region. Plasmid pFL8, which facilitates integrative transformation into S. erythraea and autonomous replication in Excherichia coli and Streptomyces lividans (which allows screening for clones in E. coli according to the blue or white color obtained with 5-bromo-4-chloro-3-indolyl- β-D-galactopyranoside), was constructed through the ligation of Msel-digested pJ487 (31) to Ndel-digested pBS+(Stratagene, Inc., La Jolla, Calif.). Msel and Ndel have compatible overlapping ends. Two smaller Msel fragments were eliminated from pJ487 during this step, with a portion of the unneeded Bsr terminator and a portion of the unneeded mec2 gene removed. The ligation mixture was transformed into E. coli DH5α-e, selecting for ampicillin resistance (Ap<sup>+</sup>), and plasmid DNA was prepared from a collection of primary transformants. This mixture of plasmid DNA was transformed into S. lividans TK21, selecting for thiotrepton resistance (Th<sup>+</sup>). DNA from the Th<sup>+</sup> Streptomyces lividans transformants was transformed back into E. coli, and the resulting plasmid was designated pFL6. To create pFL8, plasmid pFL6 was digested with BglII and then religated; this removed the polylinker that was originally carried on pJ487.

Plasmid pFL2067 was constructed to delete a 6.0-kb portion of the eryCI-flanking region from the chromosome. pFL2063 was first constructed from one 1.8-kb EcoRI-BglII DNA fragment (clone 1; Fig. 2A and C) carrying ORFL4 and part of ORFL5 generated by PCR with primer pair K16 (5′-CGAATTCGCA GGATGTCTGCG-3′ [the EcoRI site is indicated in boldface]) and K17 (5′-GAAAGCTTGCGAGAGTACGAT-3′ [the BglII site is indicated in boldface]). Another HinfIII-BglII DNA fragment (clone 2; Fig. 2A and C) was carried DNA from the entire eryCI gene and a part of ermE and was generated by PCR with primer pair K18 (5′-GAAAGCTTGCGAGAGTACGATCG-3′ [the BglII site is indicated in boldface]) and K19 (5′-GAAAGCTTGCGAGAGTACGATCG-3′ [the HindIII site is indicated in boldface]) (Fig. 2).

For PCR amplification the cycling conditions were a heating step at 95°C for 1 min, followed by 25 cycles at 95°C for 1 min, 65°C for 30 s, and 72°C for 2 min, with a final extension step at 72°C for 7 min. The reaction mixture contained 5 U of pfuTurbo polymerase in a final volume of 50 μL Mg<sup>2+</sup> was added to a final concentration of 2 mM.

After the amplification steps, the PCR products were run out on a 1% agarose gel and purified with the GeneClean II kit (Bio 101). The purified DNA was digested with EcoRI-BglII (from primer pair K16-K17) and HindIII-BglII (from primer pair K18-K19) in double digestions. After another purification step, the two PCR products were ligated in a three-component ligation reaction to create EcoRI-HindIII-digested pFL8ΔG (pFL8 with the unique BglII site removed) to form pFL2063. In a subsequent step, the 1.2-kb kanamycin resistance (Km<sup>+</sup>) gene from pUC4K (Amerham Pharmacia Biotech, Piscataway, N.J.) was ligated as a BamHI fragment into the BglII site between the two cloned PCR fragments in pFL2063. The ligation mixture was transformed into E. coli DH5α-e, and the cells were spread on Luria-Bertani plates containing ampicillin (100 μg/ml) and kanamycin (40 μg/ml). Km<sup>+</sup> and Ap<sup>+</sup> colonies were scored for the presence of the cloned PCR products by E.Nase digestion. The plasmids were digested with EcoRI, BglII, and HindIII in single and double digestions to verify the presence of the correct inserts. Several recombinant plasmids were obtained and were designated pFL2067 (Fig. 2A).

One-step integration and eviction of pFL2067. Plasmid pFL2067 was chromosomally integrated into S. erythraea by standard procedures (34), except that primary transformants were selected by using kanamycin (10 μg/ml) instead of thiotrepton. Primary transformants appearing on R2T2 plates with kanamycin were patched onto a series of three E20A plates containing thiotrepton at 15 μg/ml, kanamycin at 10 μg/ml, and no added antibiotic, respectively. Of 96 colonies tested, all 96 grew on the kanamycin-supplemented plates and the plates containing no drug. Ninety-one colonies grew on the thiotrepton-supplemented plates. The five putative Km<sup>+</sup> thiotrepton-susceptible (Th<sup>+</sup>) strains with gene replacements were designated FB50a to FB50c.

Southern blotting verification of deletion in FB50 strains. Chromosomal DNA from strains FB50a to FB50c was prepared (13) and digested with NolI. The Southern blotting procedure (Roche Molecular Biochemicals) was performed, with the prehybridization and hybridization steps carried out at 60°C (Fig. 2). Hybridizations were performed in the presence of an eryCI probe (for the size and the location of the probe, see Fig. 2C). The 4.4-kb hybridizing band, band A, which corresponds to the native eryCI-flanking region and which extends into the ermE gene, appears only in the parent strain, BC1. In FB50 deletion candidates, band A is changed to a 5.1-kb band, band B.

PCR verification of deletion in FB50 strains. The approach described above produces a unique PCR fragment that would be created only if the Km<sup>+</sup> gene was substituted for the deleted region. PCR primer pair K66-K69 (sequences below) was designed to amplify a 1.656-bp fragment that comprised the 1.266-bp Km<sup>+</sup> gene and 388 bp of DNA located on both sides of the resistance gene (Fig. 2). Primer K66 (5′-GGAATTCGCTCAACATGGCCGAGACAAA-3′ [where the boldface indicates EcoRI]) anneals in the eryCI-flanking region 175 bp upstream of the Km<sup>+</sup> gene. Primer K69 (5′-CAGACATCTGTGACTGATGCTGACGCC-3′ [where the boldface indicates SauI]) anneals in the eryCI gene 213 bp downstream of the Km<sup>+</sup> gene. Both primer sequences are present in the strain BC1 chromosome, but no amplified product is expected in the parent strain because the 2-min extension reaction used would not amplify a PCR fragment that large. The estimated size of each of the fragments in all three lanes is 1,692 bp.
FIG. 1. Schematic map of the eryCI-flanking region of the S. erythraea erythromycin biosynthetic gene cluster. The ORFs are shown as black boxes, with the direction of transcription indicated by arrows above the black boxes. The map positions (in kilobases) starting from the first nucleotide of the \textit{ermE} gene are given above the arrows. Above the map position scale are the locations of the restriction enzyme sites originally identified by Varà et al. (30) and refined in this study by DNA sequence analysis; compare this figure to Fig. 3 of Varà et al. (30). Restriction site abbreviations: G, BglII; B, BamHI; C, ClaI; E, EcoRI; H, HindIII; K, KpnI; P, PstI; S, SphI; T, SstI. The frame plot (percent G\+C content) for this region of the chromosome is shown at the top (3, 17). Below the ORFs are segments of the \textit{eryCI}-flanking region that are referred to in the text and are represented by eight grey boxes. The uppermost grey box, with a black border, represents the extent of the region subjected to DNA sequence analysis in this study. The scale (in base pairs) at the top of this grey box correlates to the numbering of the sequence with GenBank accession no. AF487998.
Plasmid construction, results of PCR and Southern blot analysis, and results of a diagrammatic analysis of the eryCI-flanking region deletion.

(A) Plasmid map of pFL2067. Integrative plasmid derived from pFL8 (the construction of both plasmids is described in Materials and Methods).

(B) PCR and Southern blot verification of the deletion in FB50 strains. (Left panel) PCR was used as a confirmatory method to detect the deletion in the eryCI-flanking region in strains FB50a to FB50c. The approach produces a unique PCR fragment that would be created only if the gene for kanamycin resistance was substituted for the deleted region (see Materials and Methods). The size and presence of the amplification products in lanes 1 to 3 confirm that the deletion has occurred in strains FB50a to FB50c. Lane 4, BC1; lane 5, molecular size standard. (Right panel) Southern blot analysis of deletion candidates FB50a and FB50b. Lane 5, BC1; lane 6, Molecular size standard; lane 7, molecular size standard. (B) PCR and Southern blot verification of the deletion in FB50 strains. (Left panel) PCR was used as a confirmatory method to detect the deletion in the eryCI-flanking region in strains FB50a to FB50c. The approach produces a unique PCR fragment that would be created only if the gene for kanamycin resistance was substituted for the deleted region (see Materials and Methods). The size and presence of the amplification products in lanes 1 to 3 confirm that the deletion has occurred in strains FB50a to FB50c. Lane 4, BC1; lane 5, molecular size standard. (Right panel) Southern blot analysis of deletion candidates FB50a and FB50b. Lane 5, BC1; lane 6 and 7, deletion candidates FB50a and FB50b, respectively; lane 8, molecular size standard; lane 9, molecular size standard. (A) Plasmid map of pFL2067.


Table 2. Characteristics of \(\text{eryCI}\)-flanking region

<table>
<thead>
<tr>
<th>Feature</th>
<th>Size (bp)</th>
<th>Position (nt) in GenBank sequence</th>
<th>No. of amino acids</th>
<th>pI</th>
<th>Identity</th>
<th>Mol wt</th>
</tr>
</thead>
<tbody>
<tr>
<td>Noncoding region</td>
<td>3,037</td>
<td>5061–8097</td>
<td>NA*</td>
<td>NA</td>
<td>Noncoding region</td>
<td>13,553</td>
</tr>
<tr>
<td>ORFL1</td>
<td>387</td>
<td>4674–5060</td>
<td>129</td>
<td>9.68</td>
<td>Acetyltransferase</td>
<td></td>
</tr>
<tr>
<td>AT site</td>
<td>34</td>
<td>4470–4503</td>
<td>NA</td>
<td>NA</td>
<td>Attachment site</td>
<td></td>
</tr>
<tr>
<td>ORFL2 precursor protein</td>
<td>444</td>
<td>3144–3557</td>
<td>148</td>
<td>9.44</td>
<td>SSLI precursor protein</td>
<td>15,869</td>
</tr>
<tr>
<td>ORFL2 mature protein</td>
<td>336</td>
<td>3144–3479</td>
<td>112</td>
<td>6.77</td>
<td>SSLI mature protein</td>
<td>11,841</td>
</tr>
<tr>
<td>ORFL3</td>
<td>813</td>
<td>1654–2466</td>
<td>271</td>
<td>7.19</td>
<td>Methyltransferase</td>
<td>29,247</td>
</tr>
<tr>
<td>ORFL4</td>
<td>879</td>
<td>92–970</td>
<td>293</td>
<td>10.35</td>
<td>Transposase</td>
<td>33,026</td>
</tr>
</tbody>
</table>

* NA, not applicable.

(extrapolated from a standard curve of the 1-kb ladder), which is very close to the known size of 1,656 bp. No amplified product of any size is observed in the lane containing strain BCI DNA (lane 4). The sizes and the presence of the ampliﬁcation products in lanes 1 to 3 conﬁrm that the deletion has occurred in strains FB50a to FB50c. PCR conditions were as follows: total chromosomal DNA was isolated from three deletion candidate strains (strains FB50a to FB50c and BCI) as described previously. The ﬁnal Mg\(^{2+}\) concentration was 1.5 mM. All other reaction components were added as described above. Five units of Taq polymerase (MBI Fermentas, Vilnius, Lithuania) were used in all reactions. The cycling program consisted of a heating step without Taq polymerase for 5 min at 95°C, followed by 25 cycles of 1 min at 95°C, 1 min at 60°C, and 2 min at 72°C. A ﬁnal extension cycle at 72°C for 7 min was performed to ﬁll in protruding ends.

Shake-ﬂask fermentation for production of erythromycin. Fermentations were performed on a Lab-line model 3525 incubator-shaker. Working stock spore suspensions were maintained at −80°C in 20% glycerol. Samples (25 μl) of spores from working stocks were transferred aseptically to capped test tubes (16 by 125 mm) containing 4 ml of sterile SCM broth (18). Test tube (seed) cultures were grown for 2 days at 32°C, with shaking at a 10° angle at 375 rpm (calibrated with a stroboscope) on a 1-in. rotary displacement. Soluble complete medium (SCM) cultures (25 ml) containing 1.3 ml of soybean oil were started from 2.5 ml of the seed cultures and were allowed to grow for 5 days at 32°C and 375 rpm with a 1-in. rotary displacement. After a 5-day fermentation, the ﬂasks were reweighed, and their weights were adjusted to the original weights by the addition of water to compensate for evaporation. The cultures were streaked onto agar plates to check for contamination.

Bioassay for precursor proteins. A double-agar layer system on a large plate (245-mm\(^2\) bioassay dish; catalog no. 431111; Corning Costar, Cambridge, Mass.) was used for the bioassay for erythromycin. The bottom agar layer consisted of 100 ml of tryptic soy broth (TSB) agar. Once the agar solidiﬁed (after it was allowed to sit for 30 min to 1 h at room temperature), a top agar layer was poured. Top agar consisted of 100 ml of TSB agar containing 250 μl of 1% triphenyltetrazolium red and a sufﬁcient quantity of T4 Bacillus subtilis spores to produce a confluent lawn of growth. The upper layer was allowed to solidify at room temperature for 1 h. Broth samples (15 μl) from random strain FBI1 and FL2067 were spotted directly onto 1/4-in. bioassay discs (Schleicher & Schuell, Keene, N.H.) and allowed to dry for 30 min. Broth samples from strains producing larger amounts (strains BCI and FB50) were ﬁrst diluted 1:10 before being bioassayed to keep the erythromycin concentrations within the values on the standard curves. Stock erythromycin A solution (100 mg/ml) was prepared in 95% ethanol. Standard erythromycin A solutions to be used for the bioassay were prepared at 25, 50, 100, and 250 μg/ml in Tris-EDTA (pH 8.0) buffer. The bioassay plate was incubated overnight at 37°C. After incubation, the diameters of the inhibition zones were measured and converted to concentrations by using the standard curve produced for each plate.

Thin-layer chromatography. Thin-layer chromatography was performed by a previously described method (34), except that only a 1-ml sample of broth, with the appropriate scaling down of the amounts of the reagents used, was assayed.

HPLC. For determination of erythromycin A and by-products, a method based on high-performance liquid chromatography (HPLC)—mass spectrometry (MS) was used. Measurements were carried out on an Agilent Technologies 1100 series LC/MSD system that comprised a capillary pump, a micro-vacuum degasser, an autosampler, a thermostatted column compartment, and an MSD detector. The LC-MSD system was used with an electrospray ionization source. Complete system control and data evaluation were done on an Agilent Technologies ChemStation for LC-MS. The HPLC separation was carried out on an Asahipak ODP-505, 5-μm column (125 by 2.0 mm; Agilent Technologies) with an ammonium formate buffer-acetonitrile mobile phase mixture. Electrospray ionization—MS in single-ion monitoring mode was used for detection.

For calibration, an erythromycin A solution of known concentration was used. Database searches and sequence alignments. The DNA sequence of the \(\text{eryCI}\)-flanking region was subjected to searches by use of the BLASTN, BLASTP, and BLASTX programs (1). Searches based on conserved protein family motifs were also performed.

Nucleotide sequence accession number. The nucleotide sequence of the \(\text{eryCI}\)-flanking region has been submitted to the GenBank database and has been given accession number AF487998.

RESULTS

Cloning of \(\text{eryCI}\)-flanking region. The \(\text{eryCI}\)-flanking region described in this report is 8.1 kb and extends from the 3′ end of \(\text{eryCI}\) to the 3′ end of ORFL4. By using the traditional \(\text{ery}\) gene cluster mapping index, the region extends from −1.25 kb to −9.4 kb (Fig. 1). For GenBank deposit AF487998, the nucleotide numbering proceeds from left to right, from ORFL4 to \(\text{eryCI}\) (Fig. 1). The DNA used for sequencing of the \(\text{eryCI}\)-flanking region was subcloned from plasmid pMW3 (32).

DNA sequence analysis of \(\text{eryCI}\)-flanking region. The \(\text{eryCI}\)-flanking region (Table 2) begins with a 3,037-bp region next to \(\text{eryCI}\) for which the BLASTX and BLASTN programs found no significant sequence similarities. FramePlot analysis (3, 17) of this region does not depict clearly deﬁned ORFs, in contrast to the region to the right, within the \(\text{ery}\) cluster, where ORFs are densely packed and can be clearly visualized (Fig. 1). One potentially signiﬁcant hairpin-loop structure (12-bp stem, two mismatches) was found between nt 7670 and 7703, and a 12-bp perfect direct repeat was between nt 7789 and 7825; both of these were close to the 3′ end of the \(\text{eryCI}\) gene.

ORFL1, the reverse complement of nt 4674 to 5060 (GenBank accession no. AF487998), shows an AGAAGC ribosome binding site 7 bp upstream from a GTG start codon, followed by 129 amino acid (aa) residues (molecular weight, 13,553; pI 9.58) and a TGA stop codon. Analysis of the deduced amino acid sequence of ORFL1 with the BLASTP program revealed that it has a large central domain (aa residues 27 to 107) characteristic of the GNAT family of acetyltransferases. This family of acetyltransferases is known to play an important role in antibiotic resistance, particularly to aminoglycoside antibiotics (7). Two aminoglycoside antibiotic resistance genes from \(\text{Acinetobacter}\) showed the highest degree of homology [aminoglycoside 6′-N-acetyltransferase from \(\text{Acinetobacter baumannii}\) (Protein Information Resource [PIR] accession no. I39502) and the aminoglycoside 6′-N-acetyltransferase-\(\text{I}\) protein from an \(\text{Acinetobacter}\) sp. (PIR accession no. I39505)]. The analysis with the BLASTP program also revealed 34% identity over a 45-amino-acid region to the sequence of a predicted protein from the kasugamycin biosynthesis gene cluster (an-
The GNAT family also includes the histone acetyltrans-
ferases of eukaryotes. These enzymes catalyze the acetylation of
lysine residues in histones, leading to transcriptional activa-
tion; however, the analysis with the BLASTP program did not
identify this specific type of protein as homologous to ORFL1.
Considering the close proximity of ORFL1 to the erythromycin
gene cluster, it might have been thought to be a tailoring
enzyme, but to the best of our knowledge, no acetylated tai-
lored products of erythromycin have been reported, and our
results show that ORFL1 plays no role in the biosynthesis of
erthyromycin A. ORFL1 is followed by a gap of 169 bp of
noncoding DNA until an att site is reached at nt 4470 to 4503,
and then another 881 bp separates the att site from the start
codon for ORFL2.

ORFL2, a precursor protein of 148 aa residues with a mole-
cular weight of 15,869 and a pl of 9.48, showed significant
sequence similarity to substitution inhibitor-like (SSIL) precursor
proteins (proteaceous inhibitors of proteolytic enzymes)
from a wide variety of actinomycetes. The predicted mature
protein of ORFL2 would have 112 aa, a molecular weight of
11,841, and a pl of 6.77. The strongest match for ORFL2 (40%-
identity over 242 aa) was to another SSIL protein previously
identified in S. erythraea by Pereda et al. (24) as ORF4 (re-
tested to in this report as ORFR4) in the eryK-flanking region
of the ery gene cluster. Analysis with the BLAST program
revealed that ORF2 has significant homologies to many other
SSIL precursor proteins from other actinomycetes. SSIL pre-
cursor proteins are widely distributed in nature; however, little
is known about their biological role or significance. One SSIL
protein-coding sequence from Streptomyces venezuelae has
been used commercially to provide the signal sequence for the
large-scale production of secreted tumor necrosis factor alpha
(25).

ORFL2 (precursor form) is the reverse complement of nt
3144 to 3587. It begins with an ATG start codon and is fol-
lowed by a 36-aa signal peptide, as determined by SignalP
computer analysis (21). SignalP analysis of ORF4, the sub-
tilisin-like inhibitor from the study of Pereda et al. (24), shows
that it also has a classic signal sequence. No ATG or GTG start
codon was found close to the ORFR4 signal sequence, indi-
cating that the gene is either not translated or not translated
by the usual mechanisms. We also determined the nucleotide
sequence for the 5′ flanking region of ORFR4 and found it to
be in agreement with the results of Pereda et al. (24). Our
interpretation of ORFR4, if it is translated, indicates that an
additional 6 aa would exist on the 5′ end of the mature protein
described by Pereda et al. (24), which would then be preceded
by a 37-aa signal peptide. Another similarity between these
two genes is the stem-loop structures following both of these
ORFs. The stem-loop structure (nt 2543 to 2573) is farther
away from the end of ORFL2 than it is from the end of
ORFR4. A large gap of 673 bp follows the end of ORFL2
before the gap reaches the 3′ end of ORFL3.

The predicted amino acid translation for ORFL3 contains
271 aa and has a molecular weight of 29,247 and a pl of 7.19.
From the analysis with the BLASTP program, the sequences
with the two highest degrees of homology to ORFL3 (nt 1654
to 2466) were for uncharacterized proteins: the putative S-
adenosyl-l-methionine methyltransferase from Sinorhizobium
mellotit, (38% identity over 231 aa; the protein with GenBank
accession no. CAC49637) and the probable O-methyltrans-
ferase from Mesorhizobium loti (34% identity over 199 aa;
the protein with GenBank accession no. pBqg94). The analysis
with the BLASTP program showed a domain in the predicted
amino acid sequence of ORFL3 that is common to the UbiE
and COQ5 family C-methyltransferases and a domain common
to the PCMT family of O-methyltransferases.

O-Methyltransferases are known to play a role in antibiotic
biosynthesis, either through the methylation of deoxysugars or
through the polyketide portion of the structure (see reference
22 and references therein). One of the high-scoring ORFL3
homologs found by analysis with the BLASTP program was to
a C-5-O-methyltransferase, the product of the aveD gene of the
avermectin biosynthetic pathway in Streptomyces avermitilis
(38% identity over 283 aa; GenBank accession no. JG6531
and ZIP accession no. JG6531) (15). The protein encoded by
aveD, in turn, shows significant homology to that encoded by
eryG, the O-methyltransferase in the ery gene cluster (23, 33).
Only one methyltransferase, the product of the eryG gene, has
been found to be required for erythromycin A biosynthesis.
Although ORFL3 is close to the ery gene cluster, it is not required
for erythromycin A biosynthesis (see “Mutational analysis
of eryCI-flanking region” below).

The deduced product of the ORF4 gene, a protein of 293
aa residues with a molecular weight of 33,026 and the reverse
complement of nt 92 to 970, shows a very high degree of
sequence similarity to a probable transposase for IS1533 from
Mycobacterium tuberculosis (57% identity over 264 aa; ZIP
accession no. D70668). ORF4 has a homolog, ORFR6, in the
eryK-flanking region. Interestingly, both transposases are
located one ORF downstream of a substitisin-like inhibitor and
are transcribed in the same direction. The sequences of the
ORFL4 and ORFR6 transposases, however, do not share a
significant degree of homology to one another, and only the
sequence of ORFR6 shows similarity to the transposase in the
region between eryAI and eryAII (5).

Mutational analysis of eryCI-flanking region. Plasmid
pFL2067 (Fig. 2) was constructed so that a 6.0-kb portion of the
eryCI-flanking region encompassing the area between −1.5
and −7.5 kb (between nt 1925 and 7888) could be deleted. The
deletion removed the 3′ half of ORFL3, all of ORFL2 and
ORFL1, and the 3-kb noncoding region next to eryCI (Fig. 1;
see the grey bar labeled 6.0-kb deletion). Deletion strains FB50
and FL2067, for which the deletions were confirmed by PCR
and Southern blot analysis (Fig. 2), were grown in shake flasks
for erythromycin production (see Materials and Methods).
The results of the agar plate bioassay (see Materials and Meth-
ods) revealed that the ability of the strains to make antibiotic
was not noticeably affected, indicating that ORF1, 2, and 3
are not required for erythromycin production. Further analysis
of fermentation broth extracts by thin-layer chromatography
and HPLC showed that the bioactive material produced by the
deletion mutants migrated and eluted identically to erythrom-
ycin A (data not shown).

In this study we also determined that ORFL4, the presump-
tive transposon homolog, would have been deleted in an ear-
erlier study (deletion +457; see Fig. 1 in reference 34). Since the
+457 deletion was reported to have no effect on erythromycin
production, we can now conclude that ORFL4 is also not required for erythromycin production.

DISCUSSION

The erythromycin gene cluster had come to be referred to in the literature as “entirely known” (8, 9), although a significant segment of DNA contiguous with the cluster adjacent to eryCI had been left essentially uncharacterized (34).

The only information available regarding the eryCI-flanking region was presented in two reports that mapped erythromycin biosynthetic genes to this region by complementation analysis (30) and suggested the presence of pathway-specific regulatory genes through plasmid insertion experiments (12, 30). Since publication of those reports, no mutational analyses concerning this region have been reported and no DNA sequence information had been published or deposited.

The earlier results were not further developed, although any gene in this region that played a regulatory or structural role in erythromycin biosynthesis would have been considered part of the cluster. Also, there was significant potential commercial value in knowledge of this region. The potential was heightened because two genes predicted to play a role in the early steps of erythromycin deoxysugar biosynthesis were still unidentified (27, 28) and because no pathway-specific regulatory genes had yet been found for Em.

Ironically, we found that the erythromycin gene cluster does in fact end with eryCI and that there are no additional erythromycin biosynthetic or regulatory genes in the 8.1-kb eryCI-flanking region. Hence, references to eryCI as the boundary gene stand without need for correction. In this study it was shown that ORFs in this region code for enzymes that are not known to be required for erythromycin biosynthesis or resistance. Furthermore, deletion analysis showed that this region is not required for production of erythromycin.

Interestingly, pathway-specific regulatory genes have recently been reported for two other macrolide antibiotics; pikromycin has one regulator (35), and tylosin has five regulators (2). In each case it was reported that these regulatory genes are contiguous with or contained within their respective clusters. Therefore, the results reported here show that there are further differences between macrolide antibiotic gene clusters with respect to the number of regulatory genes that they contain.

In an earlier study dealing with this region, the eryD24 locus, which defined a putative regulatory gene controlling deoxysugar biosynthesis, was mapped by complementation analysis (30) immediately downstream of eryCI (Fig. 1). The evidence presented in this report shows clearly that a deoxysugar biosynthesis or regulatory gene is not located in or near this area.

Two other previous results are relevant to this report. They concern reports of significant strain improvement effects caused by plasmid insertions in this area of the chromosome (12, 30). When the first plasmid, plasmid pWHM14 (30), is inserted into the eryCI-flanking region, it was reported to cause significant overproduction of erythromycin (Fig. 1). The 1-kb cloned fragment (the Vara fragment) responsible for the insertion overlapped portions of ORFL1 and ORFL2 and the region between them. The second plasmid, plasmid pWHM55, contained a 5.3-kb fragment (the Hanel fragment) that was also reported to cause a significant increase in the level of erythromycin production. The Hanel fragment overlapped the complete eryCI gene, the 3-kb noncoding region next to eryCI, and all of ORFL1 (Fig. 1). The DNA sequence information reported here does not provide any further insights into the strain improvement effects reported earlier, for example, how the effects might be rationally explained or further exploited.

Some of the results from earlier studies that also dealt with the eryCI-flanking region were not further clarified by this study. For example, the eryB locus that Vara et al. (30) mapped downstream of eryCI was beyond the region analyzed in this study. If the mapping for the eryB locus done by Vara et al. (30) is correct, it will represent a rare case in which an essential antibiotic biosynthesis gene falls well outside the bounds of the antibiotic biosynthesis gene cluster. Additional DNA sequence analysis and mutational analysis (up to \(10\) kb) will be needed to answer this question.

Finally, the eryA34 mutation, originally genetically mapped downstream of eryCI (32), was not found within the 8.1-kb region analyzed in this study. It could conceivably fall within the uncharacterized region farther to the left, up to \(22\) kb, but since the only known structural eryA locus is now known to be on the right (other) side of \(ermE\), this earlier conclusion, based on less definitive genetic mapping data, is likely to be incorrect.

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REFERENCES


