Analysis of the genetic determinant for production of the peptide antibiotic nisin

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The structural gene for the precursor of the peptide antibiotic nisin was isolated and characterized. As with other lanthionine-containing antibiotics, nisin is synthesized as a pre-propeptide which undergoes post-translational modification to generate the mature antibiotic. The sequence data obtained agreed with those of precursor nisin genes isolated by other workers from different Lactococcus lactis strains. Analysis of regions flanking the precursor nisin gene revealed the presence of a downstream open reading frame that may be involved in maturation of the precursor molecule. Nucleotide sequences characteristic of an IS element were located upstream of the nisin determinant. This element, termed IS904, is present in multiple copies in the genome of L. lactis. The nisin determinant of L. lactis is a component of a large transmissible gene block that also encodes nisin resistance and sucrose-metabolizing genes. Gene probe experiments indicated that the nisin/sucrose gene block was located in the chromosome. Furthermore, the copy of IS904 identified adjacent to the precursor nisin gene lies at, or very close to, one end of this transmissible DNA segment and may play a role in mediating its transfer between strains.

Introduction

Recent interest in the antagonistic properties of lactic acid bacteria has led to the discovery of a wide range of antimicrobial agents with relatively broad-spectrum activities (Klaenhammer, 1988). The one established antimicrobial from Lactococcus that is widely used in the food industry is nisin. The ability of this antibiotic to inhibit sporulation of spoilage bacteria has led to its extensive use as a preservative, particularly for dairy and canned food products (Hurst, 1981). In addition, nisin inhibition of both Listeria monocytogenes (Benkerroum & Sandine, 1988) and Clostridium botulinum (Tayler & Somers, 1985) suggests a potentially wider role in future protection of the food supply.

Nisin is a member of a unique group of small ribosomally synthesized peptide antibiotics which includes subtilin (Gross & Kiltz, 1973), epidermin (Allgaier et al., 1986) and gallidermin (Kellner et al., 1988). These antibiotics have in common a high proportion of the sulphur-containing amino acids lanthionine and \( \beta \)-methylanthionine. The most prominent structural feature of the molecules is the lanthionine rings formed by these atypical residues (Gross & Morell; Fig. 1a) and the term lantibiotic has been proposed for these related antibiotics. The presence of non-coded amino acids indicates that enzymic modification of a primary translation product occurs to generate the mature lantibiotic. Reactions between appropriately located serine and cysteine or threonine and cysteine residues in the precursor molecule have been proposed (Ingram, 1970) which would generate the lanthionine rings of the mature antibiotic. This sequence of events was confirmed by the cloning and nucleotide sequence analysis of a gene for the precursor of epidermin (Schnell et al., 1988). Genes for the precursors of other lantibiotics have recently been cloned and sequenced (Banerjee & Hansen, 1988; Buchman et al., 1988; Kaletta & Entian, 1989; Schnell et al., 1989) and in each case confirmation of this mode of synthesis was found. However, the mechanism of lantibiotic biosynthesis from the primary translation product has not been elucidated nor have the genes involved been identified.

The genetic determinant for nisin production is one component of a curable and transmissible gene block that also encodes nisin resistance and sucrose-metabolizing genes (Gasson, 1984; Steele & McKay, 1986).
The cloning and nucleotide sequence analysis of the nisin structural gene and the adjacent regions of DNA from a nisin-producing transconjugant of Lactococcus lactis MG1614 is reported. Buchman et al. (1988) and Kaletta & Entian (1989) have independently cloned and analysed a similar region from different nisin-producing culture collection strains of L. lactis and these results are compared.

Methods

Bacterial strains and plasmids. Lactococcus lactis FI5876 was constructed by conjugal transfer of the nisin determinants from NCFB 894 (Gasson, 1984) to the plasmid-free strain MG1614 (Gasson, 1983). Selection was made for transconjugants able to metabolize sucrose, as described by Gasson (1984) FI6016 is MG1614 carrying plasmid pFI172 (see below).

Escherichia coli P2392 and LE392 were the host strains used to isolate and propagate JEMBL3 recombinants (Maniatis et al., 1982). E. coli MC1022 [araD139 A(ara, leu)7697 lacZAM15 galU galK rpsL; Casadaban & Cohen, 1980] was the host strain for construction of plasmid pFI172. This plasmid, derived from the broad-host-range L. lactis vector pTG262 (Shearman et al., 1989), contains the cloned precursor nisin gene (Fig. 2b).

Microbiological techniques. L. lactis strains were propagated at 30 °C in M17 medium (Terzaghi & Sandine, 1975) supplemented with 0.5% (w/v) glucose instead of lactose. E. coli strains were propagated at 37 °C in L broth (Lennox, 1955).

Nisin production was detected by the plate diffusion assay (Tramer & Fowler, 1964), using as an indicator the nisin-sensitive strain MG1614 (Gasson, 1983).

Nisin resistance was determined by streaking L. lactis NCFB 894 (a nisin-producing strain) across the surface of an M17 agar plate containing 0.5% (w/v) glucose. After overnight growth (at 30 °C) the plate was inverted over chloroform for 10 min to kill the cells. Strains to be tested were streaked across the chloroform-treated NCFB 894 streak and the plates reincubated. Nisin sensitivity was indicated when growth of the test strain ceased just before the intersection of the streaks, in the area in which nisin produced by NCFB 894 had diffused into the surrounding agar. Nisin-resistant strains grew throughout the length of the whole streak.

For screening and selection of strains able to metabolize sucrose, bromocresol purple indicator (BCP) agar (McKay & Fowler, 1964), using as an indicator the nisin-sensitive strain MG1614 (Gasson, 1983) was used, with sucrose (0.5%, w/v) replacing lactose.

Transformation. Recombinant plasmids were recovered by transformation of either E. coli MC1022 by the method of Cohen et al. (1972) with the modification of Humphreys et al. (1979) or L. lactis MG1614 by electroporation using a Gene Pulser apparatus (Bio-Rad) as described by Shearman et al. (1989).

Molecular techniques. Plasmid DNA was isolated by the SDS-alkaline lysis method and purified by caesium chloride/ethidium bromide gradient centrifugation (Maniatis et al., 1982). Total genomic DNA from L. lactis FI5876 was prepared according to the method of Lewington et al. (1987). The plate lysate method for rapid small-scale isolation of bacteriophage λ DNA (Maniatis et al., 1982) was used for isolation of recombinant JEMBL3 DNA.

Restriction enzymes and other DNA-modifying enzymes used were obtained from Pharmacia, BRL or Amersham and used according to the supplier's recommendations. DNA ligations and agarose gel electrophoresis were done by standard procedures (Maniatis et al., 1982).

Genomic library construction and recombinant detection. Total DNA isolated from L. lactis FI5876 was partially digested with Sau3A and fragments were purified by sucrose density centrifugation. A genomic
library of $10^6$ recombinants was generated by their ligation with \( \text{aEMBL3} \) arms (according to the cloning manual supplied by Stratagene, USA).

Two different approaches involving DNA probes were used to screen the library for the precursor nisin gene. A 123 bp DNA fragment encoding the predicted 34 amino acids of precursor nisin (see Fig. 1b) was synthesized, the nucleotide sequence being based on \( E. coli \) codon usage. The fragment was assembled from two 70-nucleotide oligomers which were designed such that the 17 bases at the 3' end of each molecule were complementary. As a result of annealing the two strands, a 17 bp overlap was generated. Synthesis from the two free 3'–OH groups, using Klenow fragment of DNA polymerase I, filled in the single-stranded regions, resulting in a double-stranded 123 bp \( BsmH1/ EcoRI \) fragment (enzyme sites were included in the initial design of the oligomers). The fragment was cloned into the bacteriophage vector M13 and its sequence confirmed by the dyeoxy chain-termination method of Sanger et al. (1980).

An alternative probe involved the synthesis of a 17-nucleotide mixed oligomer that corresponded to a region of low codon degeneracy within the precursor nisin sequence. For amino acids 17 to 22 (Fig. 1b) a mixture of 16 different 17-mer contained all the possible codons that might be located in this stretch of the gene [ATGGG(A/C/G/T) TG(C/G)AA(T/C)ATG].

The mixed oligomer was end-labelled using T4 polynucleotide kinase (Maniatis et al., 1982) and the 123 bp fragment was labelled using the multimpr DNA labelling system (Amersham, UK). Both probes were used to screen the unamplified library. Because of the possibly limited homology between the probes and \( \text{aEMBL3} \) recombinant DNA in plaque hybridizations, conditions of low stringency were chosen. Hybridizations were carried out in 3 X SSC (1 X SSC is 0.15 M-NaCl/0.015 M-trisodium citrate, pH 7.0), 0.5% (w/v) SDS, 0.5% (w/v) skimmed milk powder, 0.01% (w/v) Ficoll, 0.01% (w/v) polyvinlypyrolidone, and 0.5 mg salmon sperm DNA ml\(^{-1}\). Hybridization and wash temperatures were 48°C for the 123 bp probe and 30°C rising to 33°C for the end-labelled mixed oligomer.

Nucleotide sequence analysis. This was performed by the dyeoxy chain-termination method (Sanger et al., 1980) using the modified T7 polymerase ‘Sequenase’ system (United States Biochemical Corp.) according to the supplier’s recommendations. The 17-nucleotide mixed oligomer (described above) was initially used in 10-fold excess as a primer in order to target the precursor nisin sequence. Further primers were synthesized from identified sequences as they became available. The sequences of both strands were determined (shown in Fig. 3) and the data compiled using the DNA analysis programs of the University of Wisconsin Genetics Computer Group (UWGCG: version 5.3, 1988).

The synthetic primers and probes used in this work were made on an Applied Biosystems DNA synthesiser (model 381A). The phosphoramidite method of oligonucleotide synthesis was followed according to the manufacturer’s instructions and using chemicals and columns supplied by them.

**Polymerase chain reaction (PCR).** The reaction mix consisted of: 25 \( \mu \)l 4 X reaction buffer (40 mm-Tris/HCl pH 8.3; 200 mm-KCl; 6 mm-MgCl\(_2\); 0.04% gelatin; 800 \( \mu \)M-dATP, dCTP and dTTP; 200 mm-dGTP; 600 mm-dCCTP); 20 pmol of each primer; 10 \( \mu \)l 10% Triton X-100; 3 units Taq DNA polymerase; 1 \( \mu \)g pF172 DNA. The final volume of 100 \( \mu \)l was overlaid with 100 \( \mu \)l sterile mineral oil. Reactions were carried out in a Techne Programmable Dri-Block. Primers corresponding to two 17-nucleotide regions within the precursor nisin gene were synthesized. For the specific amplification of DNA sequences lying between these primers (Fig. 3, coordinates 2242 to 2426) the programmable Dri-Block was run for 25 cycles at 90°C for 2 min, 55°C for 2.5 min and 70°C for 2.5 min. The amplified fragment was isolated from a gel and labelled as described above.

**Results and Discussion**

**Identification and mapping of precursor nisin clones**

A genomic library containing partial Sau3A fragments of \( L. lactis \) F15876 was constructed in \( \text{aEMBL3} \). The unamplified library was screened for precursor nisin sequences using two alternative probes (see Methods). Under the conditions used, a mixed 17-nucleotide oligomer failed to give a sufficiently strong signal to distinguish between homologous and non-specific hybridization. A 123 bp probe, based on a synthetic gene for precursor nisin, proved more successful and displayed weak homology with approximately 1% of the \( \text{aEMBL3} \) clones. DNA from six plaques which gave positive signals was isolated and analysed with restriction enzymes. All the DNAs had inserts which varied in size, although digestion with Sall (which cuts at the junctions between the \( \text{aEMBL3} \) arms and the inserted \( L. lactis \) DNA) generated some fragments of common size, indicating that a similar region of the \( L. lactis \) genome was present in the six clones (see Fig. 2a). In Southern transfer hybridization of these digestion patterns, a single SalI fragment displayed homology with the 123 bp probe (data not shown). Homology lay within a 4.4 kb fragment in clones 1, 2, 3 and 5, a 5.1 kb fragment in clone 4 and a 5.5 kb fragment in clone 6. From restriction enzyme analysis it was established that all these fragments were defined by the same SalI site within the \( L. lactis \) DNA, near the right-hand end of the different cloned inserts, and by the SalI site at the junction with the \( \text{aEMBL3} \) arms (Fig. 2a).

The fact that four out of six of these fragments were identical in size (clones 1, 2, 3 and 5) indicated that one end of all these clones was generated by digestion at the same Sau3A site in the original partial digestion of \( L. lactis \) F15876 chromosomal DNA. The Sau3A sites defining the end-points of clones 4 and 6 were located very near to those of the other clones (0.4 kb and 1.1 kb respectively; Fig. 2a).

Digestions with BglII and HincII and Southern transfer hybridizations of the restriction patterns generated confirmed the mapping of the six clones as shown in Fig. 2. Furthermore, from these data, it was possible to narrow down the region of homology with the precursor nisin probe to sequences lying within a HincII/SalI fragment at the extreme right-hand end of all six clones. It is possible that insertion of sequences lying to the right of those cloned in \( \text{aEMBL3} \) recombinant 6 (Fig. 2a) are detrimental to phase growth. The signal denoting homology with the synthetic probe was weak in plaque hybridizations due to the incomplete homology of this probe with the precursor nisin gene sequence (72%; see below). Screening the library with sequences from the cloned fragment, which would display 100% homology,
has identified additional recombinants from outside the region already cloned. These are being analysed.

**Precursor nisin gene – nucleotide sequence analysis**

The 5-5 kb SaI fragment from λEMBL3 recombinant 6 was cloned into pTG262 to generate pFI172 (Fig. 2) and subjected to nucleotide sequence determination. A 2.5 kb sequence identified from within this fragment is shown in Fig. 3. Three open reading frames (ORFs) were located in this region (Fig. 2b). The 171 bp ORF-2 (Fig. 3, coordinates 2215 to 2385) was identified as the gene for precursor nisin. It encodes a 57-amino-acid polypeptide of which the 34 C-terminal residues correspond precisely to those predicted for precursor nisin. Serine, threonine and cysteine residues were located at the appropriate positions within the structural gene so as to generate the lanthionine, β-methylanthionine and dehydro-residues present in mature nisin (Fig. 1). The N-terminal 23 amino acids represent a leader sequence involved in export of nisin through the cell wall, cleavage occurring between arginine and isoleucine residues of the precursor molecule during this process. Buchman et al. (1988) established that precursor nisin is synthesized as a pre-propeptide which undergoes post-translational modification. The conclusions drawn from our independent results and those of Kaletta & Entian (1989) support this mechanism for nisin biosynthesis.

A comparison of sequences obtained here with those published previously (Buchman et al., 1988; Kaletta & Entian, 1989), indicated that the genes for precursor nisin were identical. Furthermore, the similarities between strains extended into the regions flanking this structural gene. The start of an ORF is located 108 bp downstream of the precursor nisin gene (coordinate 2496, Fig. 3). This partially sequenced potential coding region extends for at least 1 kb downstream (data not shown). Buchman et al. (1988) have presented the first 190 bp of the equivalent ORF. In their strain, both this coding region and the intergenic sequence showed 100% homology with those obtained here. This identity between independently isolated L. lactis strains probably reflects the transmissible nature of this antibiotic determinant and may indicate the relatively recent dissemination of nisin biosynthesis genes.

**Expression of cloned L. lactis DNA**

Plasmid pFI172, carrying the cloned precursor nisin gene, was introduced into L. lactis MG1614 by electroporation to generate strain FI6016. When tested for expression of genes associated with the nisin/sucrose gene block, this strain did not produce active nisin. Plasmid-mediated traits in L. lactis strains are often spontaneously lost as a result of deletion (Gasson et al., 1987). Such deletion events do not appear to have affected the recombinant plasmids in this study. Plasmid pFI172 DNA isolated from both E. coli and L. lactis backgrounds had identical restriction enzyme profiles (data not shown), indicating that the precursor nisin gene was intact.

One obvious reason for the absence of nisin production by the cloned nisin structural gene is the lack of...
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maturation enzymology in strain FI6016. Nucleotide sequence analysis of the cloned L. lactis DNA in pFI172 suggested that this plasmid is unlikely to encode complete genes for enzymes involved in modification of precursor nisin. There is no previous evidence to suggest that these genes are linked to the nisin structural gene and the possibility exists that this maturation function is genetically separate, perhaps fulfilling other functions in the cell. Assuming that the cloned nisin precursor gene is expressed by L. lactis carrying pFI172, its failure to produce nisin supports the likely genetic linkage of maturation genes and the nisin structural gene. Whilst expression of the pFI172-encoded nisin precursor gene in L. lactis has yet to be demonstrated in vivo, the inclusion of one terminus of the nisin/sucrose gene block (see later) proves the presence of all upstream regions likely to be involved in its transcription.

A potential hairpin loop is located in the small intergenic gap between ORFs 2 and 3 (Fig. 3). It is not followed by the run of Ts which characterizes rho-independent terminators (Rosenberg & Court, 1979) and the suggestion has been made that these inverted repeat sequences (IRs) represent a processing site for a large polycistronic message in which ORF-3 is transcribed by readthrough from the upstream precursor nisin gene (Buchman et al., 1988). From the 1 kb sequence determined to date no similarity has been found with sequences in the EMBL or GenBank databases to suggest a function for ORF-3. However, the potential linkage with the precursor nisin gene, at the level of transcription, may indicate that this downstream coding region is involved in the production of nisin.

The presence of determinants for nisin resistance and sucrose metabolism on pFI172 was tested. L. lactis FI6016 did not exhibit either of these genetically linked phenotypes. Other DNA fragments flanking the precursor nisin gene have been isolated and are being investigated for the presence of these genes together with those involved in the maturation of precursor nisin.

IS904 – nucleotide sequence analysis

On examination of nucleotide sequences upstream of the precursor nisin gene, a region was found which exhibited many of the hallmarks characteristic of IS elements. IRs of 39 bp were found which would generate an element of 1241 bp (Fig. 3) designated IS904 (registered with the Plasmid Reference Center Registry; Lederberg, 1987). The only ORF of significant length (>300 bp) within these IRs was ORF-1 (coordinates 625 to 1386; Fig. 3), which would encode a protein of 253 amino acids. When this sequence was compared with those in the databases, significant homology was found with the transposases of insertion sequences IS3 (1258 bp; Timmerman & Tu, 1985) and IS600 (1264 bp; Matsutani et al., 1987). These latter two elements are present in multiple copies in the genomes of the Gram-negative organisms E. coli and Shigella dysenteriae, respectively. The L. lactis protein is smaller than those encoded by IS3 and IS600, but the homology extends throughout the sequences (47% homology with IS3 and 49% homology with IS600 when conservative changes are considered as matched; see Fig. 4). The 109 C-terminal amino acid sequence encoded by a similarly located ORF was presented in the results of Buchman et al. (1988). Only one difference at the amino acid level was noted between this latter sequence and the one presented here. This involved the substitution of a serine with a threonine residue (amino acid 202, Fig. 4). A conservative change of this type may not functionally alter the activity of the putative transposase and could be accomplished by a single point mutation at the nucleotide level.

Upstream of the coding region for the putative transposase of IS904, homology with IS3 and IS600 is insignificant, but some similarity is again found at the ends of the elements. A second element (IS629), present in multiple copies, has been identified in S. dysenteriae. It is 1310 bp in length, and is closely related to IS600 and thus to IS904 and IS3. Nucleotide sequences for the ends of IS629 have been determined (Matsutani et al., 1987) and these, together with the IRs of IS600 and IS3, are compared with those of IS904 in Fig. 5. The two ends of all four elements terminate with the dinucleotide 5'-TG followed by a purine. Homology is most pronounced between IS904 and IS3, both of which are flanked by 39 bp, imperfect IRs, containing 8 and 11 mis-matches respectively. The IRs of the two elements resident in S. dysenteriae are about 10 bp shorter, but they contain an internal region made up of sequences common to all eight termini (Fig. 5). A second region within the longer IRs of IS904 is also present in IS3. These sequence homologies within the IRs might be expected, as these regions are specifically recognized by the transposases of the respective elements, which have also been shown to be related. Examination of IS904 sequences adjacent to the IRs revealed the presence of a 10 bp direct repeat of the extreme end of IR-L. This truncated IR lies 44 bp in from the left end of the element (Fig. 3; coordinates 229 to 238).

As a consequence of the process of transposition, the target site is usually duplicated and is present as directly repeated sequences flanking the newly inserted element. Both IS3 and IS600 generate a 3 bp duplication of the target site upon transposition (Yoshioka et al., 1987; Timmerman & Tu, 1985; Matsutani et al., 1987). The sequence TATT is found at either end of the IRs of IS904 (Fig. 3). Whilst this may represent a duplication of the target site of a maximum of 4 bp, further junctions
resulting from insertion of this putative element into a region of known sequence would be necessary to establish the nature of the target site in the L. lactis genome.

Sequences upstream of the start of ORF-1 were analysed for potential transcriptional and translational start signals. No significant homology was found with the 3'-OH end of the 16S rRNA of L. lactis (Ludwig et al., 1985), which would be indicative of a ribosome-binding site. Two possible regions were located which shared some homology with the promoter consensus sequences proposed for Gram-positive bacteria (Graves...
Fig. 4. Comparison of amino acid sequences of putative transposases encoded by IS904, IS3 and IS600. Amino acids which are the same or functionally equivalent in all three sequences are indicated by the shaded areas. (The conservative changes I, L, V and M; D and E; Q and N; K and R; T and S; A and G; F and Y are considered acceptable.) 'Padding', required to maximize homology, is indicated by dashes in the three sequences.

Consensus

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<tr>
<th></th>
<th>TGR</th>
<th>YRGA.AC</th>
<th>YTAAG.GAR</th>
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<tr>
<td>IS904 IR-L</td>
<td>TGG AAAGGTTATAAAAAC TAGCACC AAGG TTAAAGAA</td>
<td></td>
<td></td>
</tr>
<tr>
<td>IS904 IR-R</td>
<td>TGG AAAGTCAAGGAAAAC TAGCACC GGAG TTAAAGAA</td>
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</tr>
<tr>
<td>IS3 IR-L</td>
<td>TGA TCTTACACGCAATAG TGGAAC CGGC CTAAGTGAG</td>
<td></td>
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<tr>
<td>IS3 IR-R</td>
<td>TGA TCTTACACGCAATAG TGGAAC AGGC CTAAGTGAG</td>
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</tr>
<tr>
<td>IS600 IR-L</td>
<td>TGA GTTAGCGTGATTTAG CGGACAC TCC</td>
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<tr>
<td>IS600 IR-R</td>
<td>TGA GTTAGCGTGATTTAG CGGACAC TAC</td>
<td></td>
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</tr>
<tr>
<td>IS629 IR-L</td>
<td>TGA ACCGCCCGGAAATCC TGGAGAC T</td>
<td></td>
<td></td>
</tr>
<tr>
<td>IS629 IR-R</td>
<td>TGA ACCGCCCGGAAATCC TGGAGAC T</td>
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Fig. 5. Comparison of left (IR-L) and right (IR-R) inverted repeat sequences at the termini of IS904 with other related elements. 5' to 3' single-stranded sequences are shown. Matches between the nucleotides of IR-L and IR-R of each element are indicated by dashes. The consensus sequence is given above the eight termini; R = purine and Y = pyrimidine. Boxed regions indicate regions common to all the IRs.

& Rabinowitz, 1986). One of these lies entirely within a region of dyad symmetry whilst the other coincides with one end of this repeated sequence (Fig. 3). The two complementary IRs could form a stem structure of 21 bp with two mis-matches (AG" = -19 kcal mol\(^{-1}\); -79.5 kJ mol\(^{-1}\)). The role of similar secondary structures in regulation of transposition has been implicated for a number of other IS elements (see Timmerman & Tu, 1985). A different arrangement of sequences exists for the elements IS3 and IS600, in which a second small ORF, with appropriate transcription initiation sites, is located upstream of the transposase coding region. It has been suggested that expression of the transposase occurs as a result of transcription of a bicistronic mRNA (Timmerman & Tu, 1985). The L. lactis IS904 sequences lying between IR-L and the start of ORF-1 bear no resemblance to those in the equivalent regions of IS3 and IS600. The ability of IS904 to transpose and the significance of the internal inverted and directly repeated sequences is at present unknown.

One of the properties of these discrete sequences is their ability to influence the level of expression of nearby
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Indeed, it has been demonstrated that IS3 can switch on a silent gene in E. coli (Zafarullah et al., 1981). The possibility exists that IS904 may be involved in expression of the downstream nisin determinants. An alternative explanation for this arrangement of sequences in L. lactis FI5876 may be simply the fortuitous insertion of IS904 at this site. However, partial sequence data of Buchanan et al. (1988) indicate that this element is resident in a similar location in L. lactis ATCC 11454. It will be interesting to see whether this element is also present on the plasmid carrying the nisin determinant in L. lactis 6F3 (Kaletta & Entian, 1989).

The homology between sequences of IS904 and those encoded by IS3 and IS600 supports the idea that these elements have evolved from a common ancestor. There are other examples of transposable elements traversing the barrier between Gram-negative and Gram-positive organisms. Tn917, isolated from Enterococcus faecalis (Perkins & Youngman, 1984), is a member of a large class of antibiotic resistance transposons, the prototype of which is Tn3 from Escherichia coli (Heffron, 1983). The only other IS element to be described in L. lactis to date is ISSIS (Polzin & Shimizu-Kadota, 1987), which also displays homology with IS elements located in both Gram-negative and other Gram-positive organisms (see Delecluse et al., 1989). The divergence of IS904 from the other related elements would seem to have occurred a long time ago. Possible transcription initiation signals in IS904 show no similarity to those of IS3 and IS600, suggesting a difference between the respective hosts in the expression of IS-encoded genes. Furthermore, despite the similarity in the amino acid sequences of the putative transposases, the ORF in IS904 tends to follow codon usage characteristic of other L. lactis genes.

Hybridization analysis

The introduction of nisin genes by conjugal transfer into L. lactis MG1614 was investigated using DNA probes for the nisin structural gene (probe 3), for IS904 (probe 2) and for a region located on the opposite side of this element to the precursor nisin gene (probe 1; Fig. 6b). DNA from the nisin-producing strain FI5876 and the non-nisin-producing recipient strain MG 1614 was digested with PvuII and PvuII/HindIII. MG1614 digests are in the odd-numbered tracks and FI5876 digests are in the even-numbered tracks. P1, P2 and P3 refer to the 32P-labelled probes used in the three hybridizations, the numbers corresponding to the probes shown below the map of FI5876.

MG1614 digests (Fig. 6a, tracks 1 and 3), indicating that the non-nisin-producing strain lacked the precursor nisin gene. FI5876 thus gained these sequences as a result of the conjugation event in which the nisin/sucrose gene block was transferred. Probe 2 was derived from sequences within IS904 and was isolated as a HindIII/HincII restriction fragment (Fig. 6b). This probe hybridized to many bands in digests.
of both FI5876 and MG1614 DNA, indicating that this element was present in at least seven copies within the *L. lactis* MG1614 genome (Fig. 6a, tracks 5 and 7). An extra band was present in hybridizations with FI5876 and, significantly, the additional band, in both digests, was the same size as the band displaying homology to the precursor nisin gene probe, i.e. an 8 kb fragment in *PvuII* digestions and a 4.3 kb fragment in *PvuII/HindIII* digest (Fig. 6a: compare tracks 6 and 2 and tracks 8 and 4).

These results indicated that the nisin-producing strain FI5876 had gained a segment of DNA that included the precursor nisin gene and the upstream copy of IS904. Because of the involvement of IS elements in gene transfer it was of interest to determine whether this copy of IS904 marks one end of the gene block.

Probe 1 was isolated as a *PvuII/XbaI* fragment from sequences upstream of IS904. This probe hybridized to a 2.5 kb *PvuII/HindIII* fragment and to an 8 kb *PvuII* fragment in FI5876 digests (Fig. 6a, track 10), indicating that these sequences lie within the same *PvuII* fragment as probes 2 and 3. MG1614 DNA also displayed homology to probe 1, although the size of the hybridizing fragments differed from those in FI5876 digests (Fig. 6a, tracks 9 to 12). The 2.5 kb *PvuII/HindIII* fragment of FI5876, containing IR-R of IS904, is absent and is replaced by a larger band in the equivalent MG1614 digest (Fig. 6a, tracks 11 and 12). The absence of the IS904 junction band in the MG1614 digestions suggests that a copy of this element is not present in the equivalent region of the MG1614 chromosome. A second weak signal was evident after hybridization of these digestions with probe 1 (Fig. 6a, tracks 9 to 12). The additional bands were the same size for the two strains and there were no bands of equivalent size hybridizing to probe 2 (Fig. 6a, tracks 5 to 8). Thus, while a second region of the *L. lactis* chromosome displayed a lesser degree of homology with probe 1, these sequences are not considered to be involved in the DNA rearrangements under investigation in this study.

The data obtained from Southern blot hybridization experiments are consistent with maps for the equivalent regions of FI5876 and MG1614 chromosomes as shown in Fig. 6(b). The involvement of IS904 in transfer of a chromosomal nisin/sucrose gene block is implicated by the conclusion that this element lies at, or very close to, the terminus of sequences acquired by nisin-producing transconjugants. IS elements have the ability to mobilize DNA sequences lying between two copies of the transposable sequences. Many examples of the evolution of these composite transposons in *E. coli* have been described (Berg *et al.*, 1982; Kleckner, 1983; Iida *et al.*, 1983). A similar arrangement of the sequences described here would result in the nisin/sucrose gene block being bounded by a copy of IS904 at each end. Transfer might then be achieved by a conservative transposition event, mediated by the transposase of IS904, in which the entire gene block is excised (see Berg *et al.*, 1988). Alternatively, the two copies of IS904 may be substrates for homologous recombination, the outcome again being excision of the gene block from the chromosome in circular form. Conjugation functions mediating transfer between strains may be located within this hypothetical intermediate structure. This would be a similar arrangement to that found in the well-characterized streptococcal conjugative transposons, Tn916 and Tn919 (Franke & Clewell, 1981; Clewell & Gawron-Burke, 1986; Fitzgerald & Clewell, 1985), in which a major segment of the element is devoted specifically to a conjugative function (Senghas *et al.*, 1988). A chromosomally located transposable sex factor involved in directing high-frequency plasmid transfer in *L. lactis* has been described (Fitzgerald & Gasson, 1988). The strains used in this study encode this independent conjugation function and it is possible that it, or other as yet uncharacterized conjugal systems, may be involved in transfer of the nisin/sucrose gene block. The various possibilities are being investigated.

The production of the peptide antibiotic nisin, together with the necessarily associated nisin resistance determinant, clearly puts these cells at a selective advantage over other susceptible cells in the same environment. IS-mediated events tend to occur at low frequency, but their activity has the effect of increasing the plasticity of the bacterial genome in their vicinity. The presence of IS904 adjacent to the nisin determinants of *L. lactis* may provide a means by which these beneficial traits can evolve and spread throughout populations.

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### References


