

A *Mycobacterium tuberculosis* IS6110 Preferential Locus (*ipl*) for Insertion into the Genome

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Received 19 June 1996/Returned for modification 30 August 1996/Accepted 5 November 1996

A 267-nucleotide *Mycobacterium tuberculosis* genomic sequence (*ipl*, the IS6110 preferential locus) which can harbor the insertion sequence IS6110 at six alternative locations has been identified in some three-quarters of the isolates tested. Only one IS6110 copy was observed at this locus in the *ipl*::IS6110⁺-containing isolates tested, and all insertions had the same orientation. The implications of this finding for IS6110 fingerprint typing methods is discussed in this work.

The insertion sequence IS6110 is found in the *Mycobacterium tuberculosis* complex (3, 9) and is usually present in several copies which are well dispersed in the genome (10, 12). It is generally assumed that there is very little discrimination in the selection of a target sequence by IS6110 upon transposition (2, 4, 11), and this fact has been utilized in the epidemiological characterization of strains by using IS6110 fingerprint (restriction fragment length polymorphism [RFLP]) differences between strains (2, 10). Philipp et al. (7), however, have reported an apparently nonrandom distribution of this element on the physical map of *M. tuberculosis* H37Rv. One locus, the direct repeat region, has a high frequency of carriage of IS6110 and has been proposed as a hot spot for integration of this element, although most of the copies are located at a single site (1). We report here another locus in which IS6110 was found in six different locations in a 267-nucleotide (nt) genomic region in some three-quarters of the isolates examined.

Our study collection comprised 84 isolates of *M. tuberculosis* collected between August and December 1992, from every culture-positive patient in Scotland (6), and isolates of *M. tuberculosis* H37Ra, *Mycobacterium bovis* (clinical isolate 10772 and strain BCG-Pasteur), and *Mycobacterium africanum* (clinical isolate 110). An anchored polymerase chain reaction (PCR) was used to amplify genomic sequences flanking IS6110 copies. Anchored PCR mixtures (final volume, 50 μ l) comprised 200 ng of genomic DNA, 0.25 μ M each primer, 50 mM KCl, 10 mM Tris-HCl (pH 8.0), 1.5 mM MgCl₂, 10% glycerol, and 0.5 U of *Taq* polymerase. The PCR mixtures were subjected to 40 cycles of amplification (1 min at 94°C, 2 min at 35°C, and 2 min at 72°C) following an initial denaturation at 94°C for 4 min and were completed by an extension of 8 min at 72°C. With M13 sequencing primer used as an arbitrary primer and the IS6110-specific primer IS6 as an anchored primer (Table 1), a 510-nt product was obtained from a clinical *M. tuberculosis* isolate (isolate 229). This 510-nt PCR product was sequenced by cycle sequencing and dye-labelled dideoxynucleotide chemistry on an ABI 373A automatic DNA sequencer and was found to comprise the expected right-hand end of IS6110 adjacent to a genomic sequence (Fig. 1). The IS6 primer in combination with a new primer (FL3) designed from the genomic sequence allowed the amplification (as described above but at an annealing temperature of 55°C) of a 194-nt product from isolate 229 as anticipated. Subsequent testing of

the whole collection with this primer pair identified other isolates with products of this length and also PCR-negative isolates which presumably lacked an IS6110 copy at this locus. However, many isolates yielded PCR products of other lengths suggestive of the possibility of different IS6110 insertion sites in this region. Sequencing of the longest of these products (isolate 91) allowed the design of a new primer, FL0, which was located in the genomic DNA just downstream from this particular IS6110 copy. PCR amplification using primers FL0 and FL3 with DNA from strains without IS6110 at this locus yielded a continuous sequence for this region (Fig. 1). This locus has been designated the IS6110 preferential locus (*ipl*).

Systematic screening of the culture collection by PCR using primers FL3 and IS6 to detect the presence of IS6110 in *ipl* and using product length to localize the positions of IS6110 copies in *ipl* identified six different IS6110 insertion sites. The exact location of the IS6110 copies in isolates selected from each of these six groups was determined by PCR amplification (using IS6 with FL3 and IS3 with FL0) and sequencing of these products to locate the ends of the IS6110 copies. These six different alleles were designated *ipl*1::IS6110 to *ipl*6::IS6110. The insertion sites were all located in a 267-nt section of *ipl* (Fig. 1). Of the 84 *M. tuberculosis* clinical isolates, 62 (74%) carried an IS6110 copy in *ipl*. The frequencies of the *ipl*::IS6110 alleles in these isolates ranged from 1 (1%) to 19 (23%) (Table 2). Sequence analysis indicated that three of the putative *ipl*1::IS6110 alleles had identical IS6110 insertion sites; two putative *ipl*3::IS6110 alleles also had identical insertion points. Not only were the points of insertion identical in each of these two groups of isolates, but so too were the duplicated nucleotides of the direct repeats at each end of the IS6110 copies which were generated by the transpositional insertion of the IS6110 element. Additional IS6110 copies were not observed in the nine *ipl*::IS6110 isolates in which the *ipl* locus was sequenced (Table 2). No occurrences of IS6110 in the opposite orientation were detected, since all PCR products exhibited the expected lengths and were not increased in size by the addition of IS6110 DNA.

The *ipl* sequence without IS6110 was observed in all of the 22 *ipl*::IS6110-negative (*ipl*⁺) *M. tuberculosis* isolates, since PCR with primers FL0 and FL3 gave a product of 372 nt as predicted. Similarly, *M. bovis* (isolate 10772), *M. bovis* BCG-Pasteur, and *M. africanum* (isolate 110) each also gave a 372-nt product, which comprised the *ipl*⁺ sequence, with these primers. The *ipl* locus is therefore widespread in the *M. tuberculosis* complex. In all of these isolates, no PCR length polymorphisms in the 372 nt between primers FL0 and FL3 could be detected

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TABLE 1. PCR primers used in this study

Primer	Sequence	Description
IS3	5' GCTGCCTACTACGCTCAAC	nt 1272–1290 in IS6110 ^a
IS6	5' CAAGTAGACGGGCGACCTC	nt 162–179 in IS6110 ^a
FL3	5' GTGGCCTTGACGGTCTTCT	nt 355–373 in <i>ipl</i> ^b
FL0	5' CCCTTTGCGTCTCAGTGTC	nt 2–20 in <i>ipl</i> ^b
M13	5' GTAAAACGACGGCCAGT	M13 forward sequencing primer

^a EMBL sequence accession number X17348 sequence starting from IS6110 itself at nt 3 (9).

^b EMBL sequence accession number X95799.

by high-resolution agarose (3% Metaphor Agarose; FMC) electrophoresis, suggesting that these isolates had not undergone any insertion or deletion events (such as the gain or loss of IS6110 from this site, as the precise excision of IS6110 is expected to be a rare event). By using appropriate IS6110 and *ipl* primer combinations (IS6 and FL3; IS3 and FL0), *M. tuberculosis* isolates without a detected IS6110 at this locus were tested by PCR for IS6110 copies adjacent to this region, but none were found within the resolution of this technique (1 to 2 kb), suggesting that the *ipl* locus is restricted to the 267-nt region first noted, at least in this collection. While most of the isolates here were from British patients, a number were from patients from around the world, including *M. tuberculosis* H37Ra which carries *ipl1::IS6110* and was derived from an

TABLE 2. Frequencies of *ipl::IS6110* alleles in 84 *M. tuberculosis* isolates

Allele	No. of copies (%) as determined by ^a :		Insertion site sequence ^b
	PCR	Sequencing	
<i>ipl1::IS6110</i>	19 (23)	3 ^c	GTGAAGGAGGCAACCACCA
<i>ipl2::IS6110</i>	12 (14)	1	GTGTGTGCCGCGAGGTGGG
<i>ipl3::IS6110</i>	10 (12)	2	TGATAGGAGCGTGGCTTTC
<i>ipl4::IS6110</i>	10 (12)	1	GCCCTTTGCGTCT
<i>ipl5::IS6110</i>	10 (12)	1	GACCGGCCCAACCTCAACA
<i>ipl6::IS6110</i>	1 (1)	1	TCACCTGGTGTCTGGCTCA
<i>ipl::IS6110</i> negative	22 (26)	2	

^a Number (frequency) of alleles in the collection as determined by PCR product length and by DNA sequencing across the ends of the IS6110 copy.

^b IS6110 insertions resulted in direct repeats of the genomic sequence indicated (solid underlining), though in the case of *ipl4::IS6110* the length of this repeat was not determined.

^c *ipl1::IS6110* was also noted in *M. tuberculosis* H37Ra in addition to being noted in the three isolates sequenced in this collection.

isolate collected in 1905 from an American patient (8), implying that these alleles are widespread.

The reason for the strong preference of IS6110 to insert into this short region, *ipl*, of the *M. tuberculosis* chromosome is not clear. A number of features are of note however. (i) The preferred region is restricted to only 267 nt and does not seem

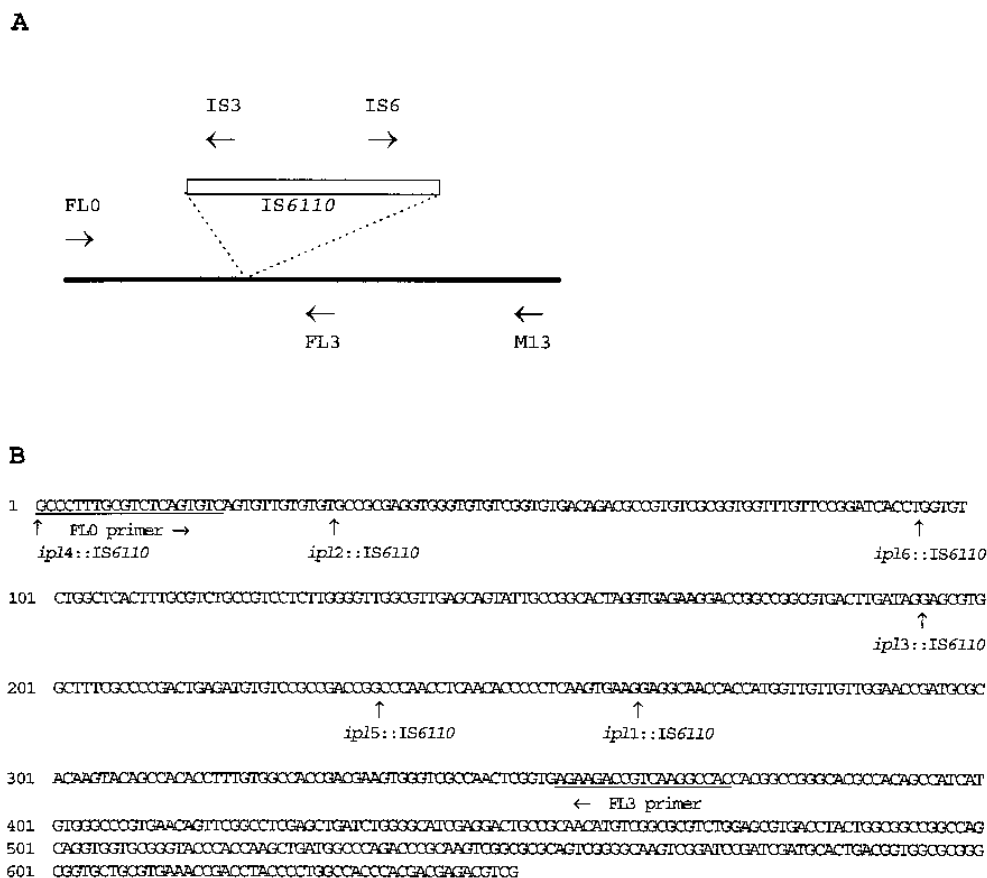


FIG. 1. DNA sequence of the *ipl* locus (EMBL accession number X59799) with the locations of IS6110 insertions shown. (A) Schematic of locations of IS6110-specific primers IS3 and IS6 and genomic primers FL0 and FL3 and presumptive location of primer M13 in isolate 229. (B) IS6110 insertions are indicated by vertical arrows (EMBL accession numbers X98148 to X98158), and primers FL0 and FL3 are indicated by underlining.

to extend beyond this, at least to a distance of 1 to 2 kb in either direction. Miller et al. (5) have noted a 25- to 50-fold preferential insertion of Tn9, a composite transposon which is flanked by *ISI* copies, into a 1-kb region of the *Escherichia coli lac* operon, so such localization is not without precedence. (ii) None of the isolates tested carried more than one copy of *IS6110* in *ipl*. (iii) The six insertions all had the same orientation with respect to the chromosome. It was not possible to determine whether *ipl* was in a coding region of the genome because of the short length of sequence that was available. However, it did contain a number of open reading frames, so it is possible that the influence of externally initiated transcription on the transpositional activity of *IS6110* is a governing feature of *IS6110* orientation. (iv) Although only 10 *ipl::IS6110* insertions have been sequenced, the insertions in the other *ipl::IS6110* isolates have been located by PCR (using high-resolution agarose electrophoresis), and these mapped to the six sites identified by sequencing (Table 2). The frequency of the six alleles in our strain collection varies greatly.

The most popular method at present for the differentiation of clinical *M. tuberculosis* isolates is *PvuII* RFLP of *IS6110*-containing fragments (10). This method characterizes isolates with dissimilar RFLP patterns as unrelated and conversely characterizes isolates with similar patterns as related. If the location of *IS6110* copies in the genome is random, then the occurrence of similar RFLP patterns in unrelated isolates will be a rare event, since *IS6110* insertions could occur anywhere in the (comparatively large) genome (7). However, if *IS6110* insertion occurs in preferred regions of the genome, then the variety of RFLP patterns that can be generated will be reduced, especially if these regions are short, as in this study. In this case, the likelihood of unrelated isolates having coincidentally similar patterns would be increased. As the ratio of preferred to randomly selected *IS6110* insertion sites increases, so does the discriminatory power of *IS6110* RFLP typing decrease. It remains to be seen where the balance of these probabilities lies.

Nucleotide sequence accession numbers. The sequences described in this work have been assigned EMBL accession numbers X59799 and X98148 to X98158.

This work was supported by Aberdeen University's Research Committee and ACTR, Aberdeen Royal Hospitals-NHS Trust Endowment Fund, and benefited from the use of the SEQNET facility, CCL, Daresbury Laboratory, Warrington, United Kingdom.

We acknowledge the extensive assistance of B. Watt and the staff of SMRL, City Hospital, Edinburgh, United Kingdom.

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