Nucleotide sequence of a repetitive element isolated from *Leptospira interrogans* serovar *hardjo* type *hardjo-bovis*

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A repetitive element from the genome of *Leptospira interrogans* serovar *hardjo* type *hardjo-bovis* ('*L. hardjo-bovis*') was identified, cloned and sequenced. Similar sequences were shown by hybridization to be encoded by a further eight of 32 other leptospiral serovars tested. An undefined number of repetitive elements were located in the *L. hardjo-bovis* genome; sequence degeneracy of the elements was observed and no significant open reading frames were identified within the AT-rich (60%) 1467 bp repetitive element. The termini encoded a GC-rich 8 bp repeat motif and two variants showed rearrangements centred on these motifs. The nucleotide sequences of the chromosomal regions flanking the repetitive elements were determined but showed no similarities, with one exception which had a GAAC repeat directly adjacent to both termini. Similar hybridization patterns were shown by Southern transfers of *L. hardjo-bovis* total genomic digests probed with the repetitive element. Oligonucleotide primer pairs designed from sequences internal to the repetitive element and adjacent chromosomal regions were used in polymerase chain reaction experiments. With one primer pair all *L. hardjo-bovis* isolates, but no other serovar, gave identical amplified products. Evidence that the repetitive element may have derived from an acquired insertion sequence that is now inactive and chromosomally fixed is discussed.

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**Introduction**

Leptospirosis is a zoonotic disease that affects cattle, swine, rodents and domestic pets (Ellis, 1986). Human cases are often a result of occupational or accidental contact with infected animals or due to ingestion of water contaminated with their urine (Lecour et al., 1989; Thiermann, 1984). The genus *Leptospira* comprises three species; *L. interrogans* isolates are recognized as pathogens whereas *L. biflexa* and *L. parae* are apparently free-living saprophytes (Johnson & Faine, 1984). The species *L. interrogans* contains over 170 serovars organized into 19 serogroups, differentiation being dependent upon microscopic agglutination tests and cross-absorption with group- and serovar-specific hyperimmune serum (Cole, 1984). Currently the detection and characterization of leptospires are done by culture isolation and serology, which are both time-consuming and difficult to perform. *L. interrogans* serovar *hardjo* type *hardjo-bovis* ('*L. hardjo-bovis*') is believed to be the major cause of milk drop syndrome and abortion in cattle in the UK and Australasia (Chappel et al., 1989). Ellis et al. (1982), however, reported the high incidence of *L. interrogans* serovar *hardjo* type *hardjo-prajitno* ('*L. hardjo-prajitno*') in aborted foetuses from random sampling and from herds with abortion problems in Ireland. Other serogroups, notably Canicola, Grippotyphosa and Sejroe, are found frequently in domestic animals and may have a pathogenic effect in cattle (T. W. A. Little & N. M. A. Palmer, personal communication). Restriction endonuclease analysis of bacterial DNA and DNA hybridization studies have been used extensively to establish epidemiological relationships between isolates and their hosts, as well as attempting to determine genetic relatedness amongst leptospires (Marshall et al., 1981; Robinson et al., 1982; Terpstra et al., 1986; Hata et al., 1988; Van Eys et al., 1988; Silbreck & Davies, 1989; Nielsen et al., 1989). Recently, detection of leptospires has been attempted using gene probes (Zuerner & Bolin, 1988) and the polymerase chain reaction (PCR) (Van Eys et al., 1989). It was our intention to develop gene probes/PCR reagents which could be applied directly to clinical samples such as urine or material from aborted foetuses and which would not only detect but also differentiate between important serovars.

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**Abbreviation:** PCR, polymerase chain reaction.

The nucleotide sequence data reported in this paper (consensus sequence) have been submitted to GenBank and have been assigned the accession number M57713.
Methods

Bacterial strains. All leptospires were from the Central Veterinary Laboratory culture collection; they were grown on EMJH medium (Ellis et al., 1982) and stocks were stored as freeze-dried samples. DNA of *L. hardjo-bovis* strain B215 was used for cloning experiments. Competent cells of *Escherichia coli* K12 strain DH5α (Life Technologies) were used as recipients in transformation experiments.

Genetic methods. Total genomic DNA was extracted from Leptospira species using the gentle lysis method of Zainuddin et al., 1982. After about 12 passages the DNA was sheared to predominantly between 5 and 10 kb in size. A method based on that described by Ohtsubo & Ohtsubo (1977) was used to identify repeat DNA sequences. The sheared DNA, approximately 50 μg in 500 μl TE in an Eppendorf tube (1.5 μl), was placed in a water bath and heated to 95°C for 5 min. The temperature of the water bath was reduced gradually to 37°C over a 30 min period by addition of ice. Buffer (30 μl x 10 concentration) and 2 units S1 nucleases (Life Technologies) were added to the sheared DNA and digestion of single stranded DNA continued at 37°C for up to 20 min. Samples of reaction mixture (100 μl) were taken at 5 min intervals, extracted twice with an equal volume of phenol/chloroform (1:1, v/v) and ethanol-precipitated after addition of sodium acetate to a final concentration of 300 mM (Maniatis et al., 1982). After centrifugation (13000 g, 4°C, 30 min), the DNA pellets were washed with 70% (v/v) ice-cold ethanol, air-dried, dissolved in TE (10 μl) and samples (5 μl) examined by agarose gel electrophoresis. DNA was extracted from agarose gels by freezing to –70°C and melting with TE (1.5 μl) containing up to 50 ng DNA were labelled with [32P]dCTP to a specific activity approaching 1 x 106 c.p.m. μg⁻¹ by random hexanucleotide priming (Amersham) as described by Feinberg & Vogelstein (1983). The labelled probe reaction mixture was denatured at 94°C for 2 min and added to filters along with fresh hybridization solution (5 ml per 100 cm² filter). Hybridization continued for 16 h at 42°C. Post-hybridization washes were sequentially with 2 x SSC, 2 x SSC containing 0.1% SDS, and 0.2 x SSC containing 0.1% SDS, all at 65°C for 30 min each. Filters were air-dried and exposed to X-ray film (Fuji- RX) for up to 16h.

**DNA nucleotide sequencing.** Double stranded DNA sequencing was done throughout using Sequenase version 2.0 and in accordance with the manufacturer’s (Cambridge Life Sciences) recommendations. Primers (24-mers) were synthesized on an ABI PCR-mate following the schedule shown in Fig. 2. The sequences obtained were compiled and analysed using staden plus version 5.0 software (Amersham).

PCR. The methods of Saiki et al. (1985) were followed. Primers were synthesized as described above and conditions for each pair of primers were determined empirically by titrating Mg²⁺ concentration and temperature for each cycle. Taq polymerase was supplied by Perkin-Elmer Cetus (Ampliqta) and dNTPs were supplied by Boehringer. Initial PCR experiments were carried out on purified DNA. Routine sample preparation was as follows. One millilitre of Leptospira culture was centrifuged (13000 g, 2 min), and the pellet was resuspended in 50 μl distilled H₂O and thoroughly vortexed in buffered phenol (Maniatis et al., 1982). The lysate was centrifuged (12000 g, 2 min) and 1-5 μl of aqueous phase and serial dilutions were transferred to the PCR reaction tubes. A HYBAID thermal cycler was used in all experiments. When using primers 3A (5’- CCGAAA-GAAGGGGCGCCAT 3’) and 3B (5’- CGATTTAGAAGGACTTG-CACAC 3’), oligonucleotides were used at a concentration of 50 pm in Saiki buffer containing 6 mM MgCl₂. Twenty-five cycles of 95°C for 2 min, 61°C for 3 min and 72°C for 2 min were used.

Results

**Cloning the repetitive DNA element of *L. hardjo-bovis***

The sensitivity of gene probes and PCR may be enhanced if the test organism harbours multiple copies of the target sequence, such as rRNA genes or repeat motifs. A search for repeated sequences within the *L. hardjo-bovis* genome using the methods of Ohtsubo & Ohtsubo (1977) was done. Essentially, total genomic DNA from *L. hardjo-bovis* strain B215 was mechanically sheared to give fragments of average sizes between 5 kb and 10 kb, heated to 95°C for 5 min to denature the DNA and allowed to cool slowly to room temperature for 30 min. The annealed DNA was treated by S1 digestion to remove single-stranded DNA, run through an agarose (1%) gel and the predominant DNA bands visualized by UV transillumination after ethidium bromide staining. A diffuse band of about 1.5 kb was visible (Fig. 1). Similar experiments with *L. hardjo-prajitno* gave no predominant bands. A gel slice containing the DNA was

Nusieve agarose (FMC Biochemicals) gels and denaturing the DNA in 4 x volumes of distilled H₂O by heating to 94°C for 5 min. Samples (10 μl) containing up to 50 ng DNA were labelled with [3H]dCTP to a specific activity approaching 1 x 106 c.p.m. μg⁻¹ by random hexanucleotide priming (Amersham) as described by Feinberg & Vogelstein (1983). The labelled probe reaction mixture was denatured at 94°C for 2 min and added to filters along with fresh hybridization solution (5 ml per 100 cm² filter). Hybridization continued for 16 h at 42°C. Post-hybridization washes were sequentially with 2 x SSC, 2 x SSC containing 0.1% SDS, and 0.2 x SSC containing 0.1% SDS, all at 65°C for 30 min each. Filters were air-dried and exposed to X-ray film (Fuji-RX) for up to 16h.
Rapetitive element from *Leptospira interrogans* I.103

Fig. 1. Total genomic DNA from *L. hardjo-bovis* was sheared, denatured, annealed and treated with S1 nuclease to remove single-stranded DNA. Samples of reactions using DNA from strains B215 (lane A), 87/12/1 (lane B), 87/26/469 (lane C) and 88/23A/42 (lane D) are shown after electrophoresis through 1% (w/v) agarose. The arrow indicates the position of the repetitive element and relative mobilities of λ HindIII size markers are indicated.

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excised, and the DNA was recovered by freeze–thawing and purified by phenol/chloroform extraction and ethanol precipitation. Recessed strands of the isolated DNA were infilled using Klenow DNA polymerase and ligated to SmaI-cut pUC8 vector. The ligation mixture was used to transform *E. coli* DH5α and selection made for ampicillin resistance. Twelve white transformants were purified, plasmids extracted and each was shown by KpnI/BamHI double digestion to have an insert of about 1.5 kb; KpnI and BamHI were chosen because these restriction enzymes have recognition sites immediately flanking the SmaI site used for cloning. The 1.5 kb KpnI/BamHI fragments from four of these digests were purified from low-gelling-temperature agarose gels and used separately as hybridization probes against Southern transfers of preparations similar to those shown in Fig. 1. In each case a strong hybridization signal was observed corresponding to the diffuse DNA bands at about 1.5 kb (data not shown).

At the time of doing these experiments, Zuerner & Bolin (1988) published the cloning and restriction endonuclease mapping of a 1.4 kb repetitive element isolated from the *L. hardjo-bovis* genome. In their experiments, *L. hardjo-bovis* genomic DNA was digested with numerous restriction endonucleases; digestion with *NarI* gave a predominant 1-4 kb fragment which was cloned into the *AccI* site of pUC8. One recombinant made in this work, designated pVW300, was examined further by restriction endonuclease digestion and a map (Fig. 2) broadly similar to that described by Zuerner & Bolin (1988) established. The insert in pVW300 differed in that the locations of the *ClaI* and *XmnI* sites were identical (common recognition sequence) and the location of the *NruI* site was 100 bp not 300 bp from the outside end of the repetitive element. Assuming the elements described in this work and by Zuerner & Bolin (1988) were homologous, the variation of restriction maps indicated sequence polymorphism.

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**Distribution of sequence homologous with the repetitive element within leptospires and apparent constant chromosomal location in *L. hardjo-bovis***

DNA was extracted from single isolates of 32 serovars representative of 15 serogroups and used as target in dot-blot experiments with the repetitive element isolated as a KpnI/BamHI 1.5 kb fragment from pVW300 as probe. Hybridizing serovars were *javanica* and *pot* (serogroup Javanica), *zanoni* (serogroup Pyrogenes), *balcanica*, *hardjo-bovis*, *istrica*, *saxkoebing* and *sejroe* (serogroup Sejroe) and *tarassovi* (serogroup Tarassovi). Serovars which did not hybridize were *australis*, *bratislava* and *muenchern* (serogroup Australis), *ballum* (serogroup Ballum), *bataviae* (serogroup Bataviae), *celledoni* (serogroup Celledoni), *cynopteri* (serogroup Cynopteri), *grippotypohsa* (serogroup Grippotypohsa), *hebdomadis* (serogroup Hebdomadis), *icterohaemorrhagiae Vi*+ and Vi- (serogroup Icterohaemorrhagiae), *mini* (serogroup Mini), *panama* (serogroup Panama), *mozdoc*, *pomona* and *proechimys* (serogroup Pomona), *pyrogenes* (serogroup Pyrogenes), *hardjo-prajitno*, *medanensis*, *recreo*, *ricardi*, *roumanica*, *trinidad* and *wolffii* (serogroup Sejroe).

Zuerner & Bolin (1988) suggested that there were about 40 copies of the repetitive element in the *L. hardjo-bovis* genome. To determine whether this was true of the isolates in the Central Veterinary Laboratory collection, the repetitive element was used to probe Southern transfers of total genomic BamHI digests. The restriction endonuclease *BamHI* was used because the repetitive element lacks a recognition site for this enzyme. An example of the results obtained is shown in Fig. 3. With *L. hardjo-bovis*, the number of discrete hybridizing bands was difficult to estimate. In all, 30 isolates of *L. hardjo-bovis* were confirmed by dot-blot hybridization to encode the repetitive element; 12 of these, from the UK (8), Ireland (2) and Zimbabwe (2), were analysed by
Southern hybridization of genomic digests and shown to have similar hybridization patterns although, as can be seen from Fig. 3, interpretation was subjective. L. saxkoebing is serologically similar to L. hardjo-bovis and a potential cause for concern for misdiagnosis by serological testing. It has been isolated from fox, badger, woodmouse and cat but never from cattle in the UK (N. M. A. Palmer & J. Redstone, personal communication).

Sequence analysis of the repetitive element

It was considered unlikely that the method for cloning the repetitive element used in this work would give a full-length sequence, because of probable infidelity of S1 nuclease digestion. Furthermore, it was considered important not only to establish the true ends of the repetitive element but also to determine if commonality of sequences existed between chromosomal regions adjacent to the repetitive elements. Therefore, repetitive elements with flanking chromosomal regions were cloned for use in sequencing experiments. To do this, total genomic DNA of L. hardjo-bovis was digested to completion with BamHI, ligated with dephosphorylating BamHI-cut pUC18 and used to transform E. coli DH5α. Transformants were hybridized with the KpnI/BamHI 1·5 kb insert from pVW300 in colony hybridization experiments to detect recombinants harbouring the repetitive element. Recombinant plasmids from 20 transformants (pVW311–330) shown to encode the repetitive element were extracted, digested with BamHI and demonstrated by agarose gel electrophoresis to have inserts ranging from 5 kb to 13 kb in size.

A series of subclones of pVW300 were made in pUC18 (see Fig. 2). Use was made of pUC forward and reverse primers for initial sequencing whilst oligomer primers
Repetitive element from *Leptospira interrogans* 1105

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Fig. 4. Nucleotide sequence of the *L. hardjo-bovis* repetitive element from clone pVW300. The presumptive termini were determined by sequencing across the repetitive element/chromosome junction of 20 genomic fragments encoding the element (pVW311–300); the termini were determined from the consensus sequence.
The repetitive element was AT-rich (60%), as were the flanking chromosomal regions. No large open reading frame within the repetitive element was identified; indeed the largest open reading frame was 291 bp. As for potential for secondary structure, no large repeat motifs were identified. However, an 8 bp motif, GGC GGCCC, was located at both termini (12–19 and 1461–1454). The motif, which contains the recognition sequence for NarI used by Zuerner & Bolin (1988) for identification and subsequent cloning of the repetitive element, is unusually GC-rich; there are very few sites for other restriction endonucleases with GC-rich recognition sequences. Examination of the terminal sequences derived from pVW321 and pVW324 with the consensus sequence revealed a right end for left end substitution centred on the GC-rich motif on the one hand and a reiteration of the GC-rich motif with a loss of the terminal 8 bp on the other hand. Flanking chromosomal regions, of about 250 bp either side of the repetitive element (data not shown), were analysed for any significant commonality but none was observed. One repetitive element (pVW312), however, shared a 4 bp (GAAC) inverted repeat immediately flanking the element. A search of the GenBank and EMBL databases revealed no significant homology with any other sequences except for poly(A) and poly(T) tracts found in many eukaryotic sequences.

Evidence for conservation of chromosomal location of repetitive elements

The evidence so far indicated that the repetitive elements in L. hardjo-bovis (and presumably L. saxkoebing) were degenerate sequences and possibly fixed in
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Fig. 6. PCR reactions using primer pair 3A/3B of cell lysate samples (1 μl) from *L. hardjo-bovis* strains B215 (lane 13), Ellis 2 - Northern Ireland (lane 12), 87/12/1 Zimbabwe (lane 11), 87/32.Hfr9 – UK (lane 10), 88/23/136– UK (lane 9), *L. saxkoebing* strains 766V1 (lane 8) and 776V2 (lane 7), and *L. hardjo-prajitno* strains 1– UK (lane 6), 2– UK (lane 5), Ellis 6 - Northern Ireland (lane 4) and Ellis 7 - Northern Ireland (lane 3). Lane 2 is a negative control and lane 1 contains size markers (1 kilobase ladder; Life Technologies). The amplified product (187 bp) is labelled AP whilst excess oligonucleotide primers and primer dimers are labelled OP.

specific chromosomal locations. To test this latter hypothesis, pairs of synthetic oligomer primers were made homologous to internal regions of the repetitive element on the one hand and homologous to flanking chromosomal regions on the other. These pairs of primers were used in PCR experiments with 30 different *L. hardjo-bovis* isolates. In principle, if the sequences complementary to the primers were juxtaposed in the test isolates, then a discrete amplified product of the desired size would be observed. One primer pair was designed using sequences derived from pVW324. Primer 3A (5' CCGGAAGAAAGGCGCCAT 3') was homologous with the variant left end of the repetitive element whilst primer 3B (5' CGATTTAGAAGGACTTGGCACA 3') was homologous to flanking chromosomal sequence and the predicted size of amplified product was 187 bp. All 30 *L. hardjo-bovis* isolates collected from the UK, Ireland and Zimbabwe were tested and gave the predicted amplified product. Furthermore, in Southern hybridization experiments, the amplified products were shown to be homologous with the PCR product amplified using pVW324 as template. The primers 3A and 3B did not give amplified product with *L. hardjo-prajitno* (8 isolates) or *L. saxkoebing* (4 isolates) or with single isolates representative of 29 other serovars. An example of the results is shown in Fig 6.

Discussion

The repetitive element of *L. hardjo-bovis* described in this work is most probably the same as that described by Zuerner & Bolin (1988). There are differences between the restriction endonuclease maps but this is most probably explained by the considerable degeneracy of sequences amongst the many copies of the element encoded by the host cell. The element has a limited distribution within leptospires, being found in two serovars of the Javanica serogroup, one of two serovars tested of the Pyrogenes serogroup and five of twelve serovars tested belonging to the Sejroe serogroup. Although this survey is limited, it may indicate relatedness if, for example, the element evolved from a progenitor gene within an ancient leptospire and, therefore, serovars harbouring the element are clonally derived. Alternatively, a more likely explanation is that the element is a classic IS element or transposon (Calos & Miller, 1980) which has infected the leptospiral gene pool. Infection by such an element may be historically recent and random and, thus, may not indicate clonal relatedness. The evidence to suggest that the element is (or was) an IS element comes from sequence analysis. An 8 bp GC-rich sequence was found repeated about 12 bp in from the apparent ends of the element and these could be target sequences for transposition; repeated termini are a feature of mobile genetic elements (Calos & Miller, 1980). The fact that novel forms of the termini were identified (pVW321 and pVW324) suggests that these may have arisen by incorrect crossing-over between adjacent termini, perhaps brought together during transposition. One repetitive element was flanked by a 4 bp repeat, a feature common to many mobile genetic elements such as Tn3 (Hefron, 1983) and Tn10 (Kleckner, 1979), which have 5 bp and 9 bp repeats, respectively. No genetic systems are yet available in leptospiral species to test whether these repetitive elements are mobile. Certainly, there was no evidence of spontaneous genetic rearrangements of any of the cloned repetitive elements in *E. coli*; but *E. coli* is probably a non-permissive host.

The sequences of the repetitive elements were highly degenerate. Accumulation of many mutations may indicate ancient acquisition of the elements and explain the lack of a significant open reading frame to encode a transposase or any other gene function. Also, prelimi-
inary data from PCR experiments suggest that the elements may be fixed in relation to flanking chromosomal sequences. Indeed, the same repetitive element/chromosomal alignment in isolates of *L. hardjo-bovis* from such geographically separated regions as the UK and Central Africa were observed. It is tempting to speculate, therefore, that *L. hardjo-bovis* is derived clonally from an ancestral leptospire in which the repetitive element, historically as a transposable element, entered the genome. However, the evidence to support this is as yet very tenuous. A significant feature of this repetitive element is the high copy number in *L. hardjo-bovis* and also *L. saxkoebing*. It is generally assumed that saturating numbers of *IS* elements and transposons are potentially lethal to a cell and self-regulation mechanisms to repress transposition have evolved (Berg, 1980). It is possible that a derepressed burst of transposition saturated the chromosome with many copies of the element and gave rise to the *L. hardjo-bovis* clone. Alternatively, these repetitive elements may not be *IS* elements or transposons, degenerate or otherwise, but rather function as target sequences for intracellularization during transformation, or perhaps encode some cryptic activity or act as recombinational hot-spots. This may explain how recognized restriction endonuclease polymorphs of *L. hardjo-bovis* have arisen (Thiermann et al., 1986). Alternatively, it is possible that one or more copies of the element may still possess an active transposase which may act in trans to mobilize elements and thus generate restriction endonuclease polymorphisms.

Nielsen et al. (1989) used cross-hybridization of total genomic DNA to analyse genetic relatedness of leptospiral serogroups. *L. hardjo-bovis* and *L. hardjo-prajitno* were separated into two distinct groups because they showed very little homology. Our findings support those of Nielsen et al. (1989) because *L. hardjo-bovis* harboured the repetitive element which, as this work shows, contributes significantly to cross-hybridization. The question arises as to the contribution the repetitive element makes to grouping by cross-hybridization. If all *Leptospira* serogroups harbouring the repetitive element were clonally derived, these serogroups may have arisen from one progenitor. It is possible that differences in terms of pathogenicity, host range, serology and biochemistry may be attributable to the mutational effects of the repetitive element.

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### References


