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Transcriptional analyses and mapping of the ospC gene in Lyme disease spirochetes.

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Transcriptional Analyses and Mapping of the \textit{ospC} Gene in Lyme Disease Spirochetes

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In Lyme disease spirochetes, the \textit{ospC} gene encodes a 22.7-kDa protein referred to as either the pC or the OspC protein. Using a variety of electrophoretic approaches followed by Southern blotting and probing with oligonucleotide probes, we mapped the \textit{ospC} gene to a circular 26-kb plasmid. The \textit{ospC} gene represents the first gene to be mapped to a circular plasmid in Lyme disease spirochetes. The occurrence of this gene in isolates belonging to each of the three Lyme disease-associated species, \textit{Borrelia burgdorferi}, \textit{Borrelia garinii}, and the VS461 group, was evaluated. The \textit{ospC} gene was found to occur in all 21 isolates tested from each of the three species. Differential hybridization with a series of \textit{ospC} probes in both Northern (RNA) and Southern blot analyses demonstrated that there is sequence variability in the \textit{ospC} gene among isolates. While the gene was found to be present in all isolates, not all actively transcribed the gene. Transcriptional start site analyses suggest that the gene may be under the control of multiple promoters that are highly similar in nucleotide sequence.

The organization of the genetic material among members of the genus \textit{Borrelia} is unusual among eubacteria in that the genetic material is distributed between a linear chromosome and a series of both linear and circular plasmids (4, 5, 10). Although variable, \textit{Borrelia} plasmids account for a significant fraction of the total genetic material. In Lyme disease borreliae, all genes mapped to date have been found to reside on either the 1,000-kb linear chromosome (12, 28) or, in the case of the \textit{ospAB} operon, on a linear plasmid (5, 6, 8) that ranges in size from approximately 49 to 59 kb (4, 5, 13, 23). In general, the coding capacity of the plasmids remains largely undefined. It seems likely that in addition to \textit{ospAB}, other genes will also be mapped to plasmids.

Recently, a third outer surface protein (OspC) in Lyme disease spirochetes was identified, and its gene was sequenced (11). The physiological role or pathological significance of OspC has not yet been defined. Although somewhat variable among isolates, the molecular mass of OspC is approximately 22.7 kDa. OspC appears distinct from the 22-kDa proteins described by Luft et al. (14) and Simpson et al. (26). The level of expression of OspC has also been found to vary among isolates. Wilske et al. reported that OspC is expressed as a major protein in approximately 45% of European isolates and only rarely in North American isolates (31). For the three spirochete species, \textit{Borrelia burgdorferi}, \textit{Borrelia garinii}, and group VS461, known to be associated with Lyme disease (2, 15–17, 19, 29), the presence, degree of conservation, and map location of the \textit{ospC} gene have not yet been addressed.

In this study, we have mapped the location of the \textit{ospC} gene, determined its transcriptional start site(s), and compared the expression of the gene among isolates of Lyme disease spirochetes. The results presented here demonstrate that the \textit{ospC} gene is carried on a 26-kb circular plasmid. The \textit{ospC} gene represents the first \textit{Borrelia} gene to be mapped to a circular plasmid. Transcriptional start site analyses have revealed that the control region of the gene is similar among isolates of Lyme disease spirochetes and that the gene may be under the control of multiple promoters. Northern (RNA) blot analysis has also demonstrated that the transcriptional expression of the \textit{ospC} gene varies among isolates.

**MATERIALS AND METHODS**

Cultivation of \textit{Borrelia} isolates. The isolates investigated in this study are described in Table 1. All isolates were previously identified to the species level by 16S rRNA sequence analysis (1, 16, 17), through the use of 16S rRNA-directed species-specific oligonucleotide probes (19) or polymerase chain reaction primer sets (15), by DNA-DNA hybridization (2, 20), or by multilocus enzyme electrophoresis (9). The species nomenclature used in this paper is that proposed by Baranton et al. (2). Bacterial isolates were cultivated in BSKII medium at 34°C (3), harvested by centrifugation, and washed twice with phosphate-buffered saline prior to use in subsequent experiments.

**Southern blot analysis of the \textit{ospC} gene.** DNA was isolated as previously described (7). Whole-cell DNA was fractionated by electrophoresis in 0.35% agarose gels with TAE buffer (40 mM Tris-acetate, 1 mM EDTA [pH 8.5]) at 0.4 V/cm for 48 h. Fractionated DNA was transferred to GeneScreen membranes by use of the VacuGene vacuum blot system as instructed by the manufacturer (Pharmacia-LKB). The transferred DNA was fixed to the membranes by UV cross-linking with a Stratalinker (Stratagene) as instructed by the manufacturer. \textit{ospC} hybridization probes were synthesized on the basis of the \textit{ospC} sequence previously presented for the Lyme disease isolate PKo1 (11). Both Adam et al. (1) and Wallich et al. (28) have identified this isolate as belonging to the VS461 group of Lyme disease spirochetes. Oligonucleotide probe sequences are presented in Table 2. All sequences given in this paper are listed 5' → 3'. Oligonucleotide probes were 5' end labeled with [γ-32P]ATP by use of polynucleotide kinase by standard methods. Radioabeled probes were purified from unincorporated label by passage through G25 spin columns as instructed by the manufacturer (Boehringer Mannheim). The blots were pre-
hybridized in hybridization buffer (16) for 1 to 4 h at 37°C. After prehybridization, the buffer was removed and radioactive probes at 10 ng/ml in fresh hybridization buffer were added to the hybridization bottles. Hybridization was conducted for 16 to 24 h at 37°C. The blots were washed twice in 2× SSC (1× SSC is 0.15 M NaCl plus 0.015 M sodium citrate)–0.1% sodium dodecyl sulfate (SDS) at 37°C (10 min each) and once for 1 h in 0.1× SSC–0.1% SDS at 37°C, wrapped in cellophane, and exposed to Kodak XAR5 film at −70°C with intensifying screens.

**Two-dimensional agarose gel electrophoresis.** For determination of the conformation of the DNA molecule carrying the *ospC* gene, two-dimensional gel electrophoresis was performed (21, 22). DNA from isolates R-IP21 and VS461 was electrophoresed in 0.35% agarose gels in the first dimension for 18 h at 0.4 V/cm with TAE buffer. The gels were then equilibrated in second-dimension running buffer (15 μM chloroquine in TAE buffer) by soaking for 5 h. After the gels were rotated 90° relative to the direction of electrophoresis in the first dimension, electrophoresis was performed for 18 h at 0.4 V/cm. Prior to ethidium bromide staining, the gels were soaked in H2O for several hours to remove the chloroquine. Fractionated DNA was transferred to GeneScreen membranes for hybridization analysis with *ospC* oligonucleotide probes as described above.

**Transcriptional start site and Northern blot analyses.** Total cellular RNA was isolated as previously described (25). All reagents were either prepared with diethylpyrocarbonate-treated H2O or treated directly with diethylpyrocarbonate by standard methods. The transcriptional start site(s) for the *ospC* gene was determined by reverse transcriptase primer extension. Fifteen micrograms of total cellular RNA was incubated with 2 pmol of 5'-end-labeled primer pC13, pC52, pC67, or nc20 in 10 μl of annealing buffer (100 mM KCl, 5 mM Tris-HCl [pH 7.5]) at 85°C for 4 min and then at 30°C for 4 to 6 h. Fifteen microliters of reverse transcription mixture (0.9 mM each dATP, dTTP, dCTP, and dGTP; 83 mM Tris-HCl [pH 8.3]; 8 mM dithiothreitol; 8 mM MgCl2; 83 μg of bovine serum albumin per ml) and 50 U of avian myeloblastosis virus reverse transcriptase were added, and the reaction mixtures were incubated for 1.5 h at 47°C. For digestion of the RNA, 1 μl of 0.5 M EDTA and 1 μl of RNase (0.5 μg/μl) were added and the mixture was incubated for 0.5 h at 37°C. One volume of 2.5 M ammonium acetate was added to each tube, and each was extracted with an equal volume of phenol–chloroform–isoamyl alcohol. The extension products were precipitated with ethanol, washed with 70% ethanol with 2% sodium dodecyl sulfate, redissolved in 10 μl of formamide loading buffer (50% formamide, 10 mM EDTA, 0.025% bromophenol blue, 0.025% xylene cyanol), heated at 85°C for 5 min, and loaded onto 7% polyacrylamide sequencing gels (8 M urea, 83 mM Tris-borate, 1 mM EDTA [pH 8.0]). The approximate sizes of the extension products were determined by use of a DNA sequencing ladder. After electrophoresis, the gels were transferred directly to 3MM paper (Whatman) and exposed to Kodak XAR5 film for 3 h.

Northern blot analysis was used to assess the expression of the *ospC* gene and to determine the nucleotide length of the transcript. Fifteen micrograms of RNA sample was electrophoresed through 1.2% agarose–formaldehyde gels as previously described (27). The RNA was transferred to GeneScreen membranes and hybridized as described above for the Southern blot analysis. The length of the *ospC* transcript was determined by use of an NA2 nucleic acid analyzer as instructed by the manufacturer (Bethesda Research Laboratories).

**RESULTS AND DISCUSSION**

Mapping of the *ospC* gene to a circular plasmid. In this study, we sought to map the *ospC* locus in a variety of isolates of each of the three Lyme disease-associated spirochete species (2, 9, 15–17, 19, 20, 29). Total DNA from the isolates listed in Table 1 was isolated and fractionated in 0.35% agarose gels. After Southern transfer, the blots were probed with an *ospC* oligonucleotide probe (pC13). Two or three hybridization bands that differed in intensity and relative ratio among isolates were observed. The ethidium bromide-stained gel and the corresponding Southern blot of representative isolates are presented in Fig. 1. The detection of more than one band suggested that the probe was either hybridizing with an *ospC* gene carried on multiple DNA species or cross-hybridizing with non-*ospC* but closely related sequences. To differentiate between these possibilities, we used a second probe (pC609), which complements a region in the 3' end of the *ospC* gene, to rescreen some of the blots. This probe hybridized with the same bands as the pC13 probe. A third probe (pC67) also yielded identical hybridization results. Both the pC67 and the pC609 probes,
however, did not hybridize with the ospC gene in all isolates, suggesting sequence variability in this gene (see below). Nonetheless, the use of multiple probes, all of which hybridized with the same DNA bands, demonstrates that the multiple hybridization signals observed were due to the detection of multiple ospC-carrying DNA molecules.

The detection of multiple hybridizing bands and their relative migration rates in agarose gels suggested that the gene may be carried on a circular plasmid. Circular plasmids can be found in three distinct conformational states. Form I is the supercoiled form. Form II consists of relaxed molecules that comigrate with supercoiled plasmids that have been nicked. Form III represents plasmids that have been linearized by a double-strand break. The hybridization pattern suggested a circular plasmid of 26 kb, with the supercoiled form (I) migrating with an apparent size of 12 kb and the nicked or relaxed form (II) migrating with an apparent size of 44 kb (Fig. 1B). Furthermore, upon restriction of the DNA from several isolates with EcoRV, Southern blotting, and probing with the pC67 probe, only one hybridization band was observed (data not shown). These results further support the idea that one copy of the ospC gene was present and being carried on different conformational forms of the same plasmid. While the detection of multiple hybridizing bands in undigested DNA preparations is highly suggestive of a circular plasmid, we further addressed this question by using two-dimensional gel electrophoresis. Linear and circular plasmids, which cannot be distinguished from each other by conventional agarose gel electrophoresis, can be conclusively differentiated via this approach. For these analyses, new DNA preparations were gently isolated (i.e., vortexing and pipetting were minimized during isolation) from VS461 group isolates R-IP21 and VS461 to reduce shear damage to the circular plasmids in the DNA preparations. After standard electrophoresis of the DNA in the first dimension, the gels were equilibrated in chloroquine. Chloroquine intercalates into the DNA, causing an unwinding of the double helix. The electrophoretic mobility of linear molecules is not significantly affected by chloroquine treatment. These molecules migrate along an apparent diagonal axis in two-dimensional chloroquine gels. However, the electrophoretic mobility of negatively supercoiled plasmids is retarded by chloroquine treatment and, as a result, these plasmids migrate off the diagonal axis during electrophoresis in the second dimension (22). Three DNA species were found in ethidium bromide-stained gels to migrate off the diagonal axis, indicating that these molecules were covalently closed and negatively supercoiled (Fig. 2A). The gels were then blotted and subjected to Southern blot analysis with the pC67 hybridization probe. A hybridization signal was detected associated with one of the molecules running off the diagonal axis (Fig. 2B), demonstrating that the ospC gene is carried on a single negatively supercoiled plasmid. In contrast to the results of the Southern blot analysis described above, only one band was found to hybridize with the probe. Hence, the elimination of vortexing from the DNA isolation procedure significantly reduced shearing of the circular plasmids and, as a result, only the supercoiled form of the ospC-carrying plasmid was observed. The size of the circular plasmid was obtained by measuring the linearized form (Fig. 1) and found to be 26 kb.

Occurrence and conservation of the ospC gene in Lyme disease spirochetes. Wilske et al. have reported that OspC is expressed in 45% of European isolates and only rarely in North American isolates (31). To evaluate the occurrence of the ospC gene, we performed Southern blotting of undigested whole-cell DNA from a variety of isolates of the three spirochete species associated with Lyme disease. The results obtained with representative isolates are presented in
FIG. 2. Two-dimensional gel electrophoresis of DNA from Lyme disease isolate VS461 and Southern blot analysis with the pC67 probe. Electrophoresis of DNA from isolate VS461 was conducted in the first dimension in 0.35% agarose gels. The gels were then equilibrated in 15 µM chloroquine, rotated 90°, and electrophoresed in the second dimension as described in the text. The DNA was visualized by ethidium bromide staining (A) and then transferred to GeneScreen membranes for Southern blotting as described in the text with the radiolabeled pC67 oligonucleotide probe (B). In both panels, the position of the loading well is indicated by a circle, and that of the 26-kb plasmid is indicated by an arrow. The direction of migration in each dimension is also given. DNA molecular size markers are given in kilobases.

Fig. 1. Detection of the ospC gene in a given isolate was found to be dependent on the probe used. While the pC609 probe was found to exhibit differential hybridization with different isolates, this probe was not used to screen all isolates and hence will not be discussed further. Most B. burgdorferi isolates and B. garinii G1 and R-IP90 were hybridization negative with the pC67 probe but hybridization positive with the pC13 probe. B. burgdorferi 25015 and Illinois 1 were exceptions in that they hybridized with both the pC67 and the pC13 probes. We have demonstrated the peripheral relationship of these isolates to others of B. burgdorferi via 16S rRNA sequence analysis (15). In contrast to B. garinii R-IP90 and G1, VS461 group isolates and B. garinii G25, 20047, and VS102 were hybridization positive with both the pC13 and the pC67 probes. Hence, while the phenotypic expression of the ospC gene appears most pronounced in European isolates, the ospC gene itself is present in all three Lyme disease-associated spirochete species. Furthermore, it consistently maps to an approximately 26-kb circular plasmid in all isolates tested. The variability in hybridization among isolates indicates that the ospC gene sequence may be somewhat variable. A similar variability has been noted for outer surface protein genes ospA and ospB (7, 13, 18, 30).

Northern blot analysis of the ospC transcript. To determine in which isolates the ospC gene was being transcriptionally expressed, we performed Northern blot analysis with the pC13 and pC67 hybridization probes. Consistent with the Southern blot analysis discussed above, the detection of the ospC transcript in a given isolate was dependent on the probe used. ospC transcript size was constant among isolates, being approximately 700 to 730 nucleotides (Fig. 3). The ospC coding sequence alone (excluding 5' and 3' non-coding sequences) would generate a 636-base transcript. While the ospC gene was detected by Southern blot analysis in all isolates with either the pC13 or the pC67 probe, we did not detect the transcript in several isolates by Northern blot analysis with these same probes. B. burgdorferi CA12 and 1352, B. garinii N34, and VS461 group isolates J1, R-IP21, and UM01 were all found to carry the ospC gene but not to express it under the growth conditions used in these experiments. The rRNA bands in the hybridization-negative preparations were found to show good integrity via ethidium bromide staining, providing indirect evidence that the lack of hybridization was not the result of template degradation. Furthermore, using these same RNA preparations, we were able to detect the ospAB transcript. Factors associated with the transcriptional activation of the ospC gene have not yet been defined. In addition, several VS461 group isolates that produced the ospC transcript did not exhibit detectable amounts of OspC protein, as determined by inspection of Coomasie-stained sodium dodecyl sulfate-polyacrylamide gel electrophoresis whole-cell protein profiles (data not shown).

FIG. 3. Detection of the ospC transcript by Northern blot analysis. Fifteen micrograms of total cellular RNA was fractionated in 1.2% agarose–formaldehyde gels, vacuum blotted onto GeneScreen membranes, and hybridized with the pC13 probe as described in the text. The species designation for each isolate is indicated by the prefixes Bb for B. burgdorferi, Bg for B. garinii, and (VS) for VS461 group isolates. The positions of 23S and 16S rRNAs (approximately 2,900 and 1,540 bases, respectively) are indicated.
No correlation between while typic expression of the ospA and ospB total termination. VS102, 20 bases sequence of use promoter, a and some sequence of translational control.

It has been demonstrated that the phenotypic expression of the ospA and ospB genes in some isolates can be influenced by the number of in vitro passages (24). While in this study we did not systematically monitor OspC expression over a large number of passages in a given isolate, isolates of both high and low passage numbers were used. No correlation between passage number and OspC expression was observed.

**Transcriptional start site analyses of the ospC gene.** To identify the putative promoter(s) of the ospC gene, we performed transcriptional start site analyses by using reverse transcriptase primer extension. By using two independent primers (pC13 and pC52), the same primer extension termination sites were observed in all isolates which had been by Northern blot analysis to be expressing the ospC gene. The major termination sites mapped approximately 18 to 22 nucleotides upstream from the translational start codon. The results obtained with a representative isolate of each of the three Lyme disease-associated species and the pC52 primer are presented in Fig. 4. An appropriately spaced consensus promoter sequence (5'-TTG-AAA-A) that we refer to as P1 and a consensus TATA box sequence (5'-TATTAA) were identified upstream (on the basis of the ospC sequence for isolate PKo) (Fig. 5). Hence, the transcripts derived from the use of promoter P1 possess a 5'-untranslated leader sequence of approximately 20 nucleotides, with the T residue located 20 bases from the translational start codon serving as the major transcriptional start site (TS1). Using the RNA secondary structure analysis program of Zucker (contained in the PCGENE software package) (32), we analyzed sequences 5' of the translational start codon for regions of secondary structure that could conceivably be responsible for the termination sites observed in the primer extension analysis. However, significant secondary structure elements were not observed, supporting the identification of TS1 as the major transcriptional start site. Through a consensus promoter sequence homology search, Fuchs et al. (11) identified a putative promoter upstream of P1 that we refer to as P2. This sequence is not optimally spaced to serve as the promoter for TS1. However, when both the pC13 and the pC52 primers were used, a second transcriptional start site (TS2) was observed up-stream of the TS1 site, implicating active transcription from a promoter 5' of P1 (Fig. 4). In *B. burgdorferi* 3028, a minor extension product intermediate in size in comparison with those derived from P1 and P2 was observed. The sequence of this product is not appropriately spaced from a consensus promoter sequence, so this product most likely represents a premature termination product. P2 appears appropriately spaced from TS2 to serve as a promoter for this transcriptional start site. Visualization of the band corresponding to TS2 required a longer exposure of the separating gel to film (approximately 6 to 10 h) than that required for the detection of TS1. The exposure presented in Fig. 4 was chosen since it allowed visualization of both transcriptional start sites without overexposure of TS1. Both P1 and P2 are identical in sequence over a 7-base stretch (5'-TTG-AAA-A) and have appropriately spaced −10 sequences (Fig. 5). The core sequences of P1 and P2 and their respective −10 sequences are separated by 15 and 13 bases, respectively. While transcriptional start sites were observed appropriately spaced from both P1 and P2, it is possible that only the more distal promoter, P2, was used and that the shorter transcript was the result of an RNA processing event. While the data presented here are suggestive of active transcription from more than one promoter, RNA processing cannot be ruled out.

Upon analysis of the PKo ospC sequence (11), we noted a third potential promoter sequence (P3) upstream of and identical to the 7-base core sequences of P1 and P2. Over a 16-base stretch, P3 exhibits 88% sequence similarity to P2. To determine whether this putative promoter sequence was transcriptionally active, we used a third primer (nc20) in the primer extension analysis. This primer complements a sequence in the ospC gene 5' of TS1. While the 5' end of this 20-base primer complements the first 6 nucleotides of transcripts derived from the expression of promoter P2, these transcripts should not serve as a template for primer extension, since the 3' end of this primer does not hybridize.
Hence, any extension products should be derived from hybridization with a transcript that uses a more distal 5' promoter. In two isolates (B. burgdorferi 3028 and B. garinii VS102), a weak transcription start site, not previously observed when primers pC13 and pC52 were used, was detected (data not shown). Hence, promoter P3 may be used at a low level in the transcription of the ospC gene in some isolates. The low level of transcription from this promoter may be a reflection of the spacing between the −35 and −10 sequences (11 bases). None of the three transcriptional start sites described above was observed in isolate R-IP21, which was shown not to produce the ospC transcript by Northern blot analysis, demonstrating that the start sites detected were not the result of spurious or nonspecific primer extension from non-ospC transcripts. The role of each promoter, the extent to which each is used, and the factors influencing promoter selection in vivo are currently undefined.

The ospC gene represents the first Borrelia gene to be mapped on a circular plasmid. The plasmid is approximately 26 kb in size. On the basis of hybridization results with a variety of oligonucleotide probes, this gene appears to exhibit some variability in its coding sequence among the Lyme disease isolates tested. The organization of the upstream regulatory regions, however, appears similar among isolates with a potential multiple-promoter arrangement. The results presented here may provide further insight into the genetic coding capacity of the plasmids found in Lyme disease isolates. Since the plasmids represent approximately 150 kb of genetic material, it is likely that additional genes will be localized on plasmids and shown to encode products that may be important in the pathobiology of Lyme disease.

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