Mapping the Ligand-Binding Region of Borrelia burgdorferi Fibronectin-Binding Protein BBK32

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The cellular attachment and entry of pathogenic microorganisms can be facilitated by the expression of microbial adhesins that bind fibronectin. We have previously described a Borrelia burgdorferi gene, bbk32, that encodes a 47-kDa fibronectin-binding protein. In this study, the ligand-binding region of BBK32 from B. burgdorferi isolate B31 was localized to 32 amino acids. The bbk32 gene was cloned and sequenced from three additional B. burgdorferi isolates representing different genospecies of B. burgdorferi sensu lato. All four bbk32 genes encoded proteins having fibronectin-binding activity when expressed in Escherichia coli, and the deduced proteins shared 81 to 91% amino acid sequence identity within the ligand-binding domain. In addition, the ligand-binding region of BBK32 was found to share sequence homology with a fibronectin-binding peptide defined for protein F1 of Streptococcus pyogenes. The structural and functional similarity between the ligand-binding region of BBK32 and the UR region of protein F1 suggests a common mechanism of cellular adhesion and entry for B. burgdorferi and S. pyogenes.

Lyme disease remains the most prevalent vector-borne infectious disease in North America (17). The spirochete Borrelia burgdorferi was identified as the etiologic agent of Lyme disease in 1982 (1). Since then, at least 10 genospecies representing the complex B. burgdorferi sensu lato have been described (29). Only the genospecies B. burgdorferi sensu stricto, B. garinii, and B. afzelii are well established in causing disease in humans. Recently, however, B. burgdorferi strains resembling the newly described genospecies B. bissettii were isolated from Lyme disease patients in Slovenia (27). Human infections with B. burgdorferi are transmitted by ticks of the Ixodes subgenus. Once transmitted by tick bite, B. burgdorferi establishes a localized infection at the site of tick attachment. The spirochetes migrate in the skin, producing an oval rash termed erythema migrans in 80% of Lyme disease patients (6). Days to weeks later, the spirochetes enter the vasculature and disseminate to multiple tissue and organ sites. At this stage, the patient enters an early disseminated form of Lyme disease and may present with carditis, lymphadenopathy, meningitis, and migratory joint and muscle pain. Despite the presence of a strong host immune response, B. burgdorferi may persist in the host, and spirochetes may be isolated from the patient months to years after transmission. In this late stage of Lyme disease, the patient’s clinical manifestations may include cutaneous, musculoskeletal, and neurologic involvement. While early intervention with antibiotics is generally efficacious, this late form of Lyme disease remains the most prevalent vector-borne infectious disease in North America (17). The mechanisms by which B. burgdorferi invades and colonizes the host are poorly understood. For many bacterial pathogens, the initial step in host colonization involves the expression of adhesive molecules that mediate bacterial adherence to cells or to the extracellular matrix (20, 30). The capacity of B. burgdorferi to bind a wide variety of cells and extracellular matrix components indicates that these organisms may also express adhesive molecules (24). A common feature of several pathogenic bacteria, most notably Staphylococcus aureus and streptococci, is the expression of adhesins that bind fibronectin (10). Fibronectin is a large, dimeric glycoprotein that is produced by a broad range of cell types (22). It exists as a soluble molecule in body fluids and as an insoluble component of cell membranes and the extracellular matrix. Structurally, fibronectin is a mosaic protein composed of three types of protein modules that are organized into distinct functional domains. Through these functional domains, fibronectin can interact with a variety of macromolecules including fibrin, heparin, collagen, and integrins. Many of these functional domains are also targeted by adhesins expressed by pathogenic microorganisms.

Borrelia species also express adhesins that bind fibronectin (7, 13, 21, 28). We have previously reported on the identification of a gene, bbk32, which encodes a 47-kDa fibronectin-binding adhesin expressed by B. burgdorferi isolate B31 (21). BBK32 was localized to the outer surface of isolate B31, and the adhesin was found to interact specifically with the collagen-binding domain of fibronectin. The ability of recombinant BBK2 to inhibit binding of isolate B31 to immobilized fibronectin was also demonstrated. These results indicated that BBK32 is the primary fibronectin-binding adhesin expressed by B. burgdorferi.

In this study, we extended these earlier observations by mapping the ligand-binding region of BBK32 from B. burgdorferi isolate B31. In addition, we cloned the bbk32 gene from four isolates representing different B. burgdorferi genospecies into gene from four...
Mapping the fibronectin-binding Region of BBK32. The minimal region of BBK32 required to bind fibronectin was localized by creating a \textit{bbk32} gene fragment library using the Novatope system (Novagen, Madison, Wis.) and screening the library by ligand blotting. The \textit{bbk32} gene was amplified from \textit{B. burgdorferi} isolate B31 as previously described (21). The amplicon was purified using QIAquick spin columns (Qiagen, Valencia, Calif.), and 5 \mu g of \textit{bbk32} was digested with bovine pancreatic DNase I in 50 mM Tris-HCl (pH 7.5)—0.05 mg of bovine serum albumin/ml—10 mM MnCl\textsubscript{2}. The DNA fragments were separated on a 2% NuSieve agarose gel (FMC Bioproducts, Rockland, Maine); the 50- to 150-bp fragments were excised from the gel and purified using QIAquick spin columns. Blunt ends were created by treatment of the DNA fragments with T4 DNA polymerase. A 3' adenosine overhang was then added by \textit{Tth} polymerase, and the DNA fragments were ligated into the Novagen pScreen T vector, a pET vector derived peptide. As shown in Fig. 1A, expression of the various DNA is expressed as a peptide fused to a 37-kDa vector-derived terminator, and STAG (Novagen) were used to sequence both strands of the insert DNA. Among the 17 clones expressing the desired fragment of \textit{bbk32}, 5' and 3' ends were verified by DNA sequencing as described above. The clones were subcultured to LB broth for 8 h, and recombinant protein expression was induced for 2 h by the addition of isopropylthiogalactoside to final concentration of 0.3 mM. The bacteria were harvested by centrifugation and lysed in SDS-PAGE sample buffer, and proteins were resolved on a 12% polyacrylamide gel. Following electrophoretic transfer of the proteins to a nitrocellulose membrane, ligand blotting was performed as described earlier.

Using the pScreen T-vector expression system, the cloned DNA is expressed as a peptide fused to a 37-kDa vector-derived peptide. As shown in Fig. 1A, expression of the various \textit{pScreen/bbk32} constructs in \textit{E. coli} resulted in the production of recombinant proteins ranging in size from 48 to 54 kDa. With the exception of the AA 105-162 construct, the apparent molecular weight of the recombinant protein exceeded the molecular weight predicted for each construct. A similar observation has been made for recombinant proteins derived from different streptococcal and staphylococcal fibronectin-binding proteins (11). The AA 131-162 construct produced a 50-kDa protein that bound fibronectin, as demonstrated by SDS-PAGE and ligand blotting (Fig. 1). However, the level of fibronectin-binding activity displayed by this clone was much lower than those of the AA 131-167 and AA 105-162 constructs. Furthermore, this fibronectin-binding band appears to have migrated slightly faster than would be anticipated from the corresponding Coomassie blue-stained gel. Deletion of five amino acids from either the N-terminal (AA 136-162 construct) or C-terminal (AA 131-157 construct) end of amino acids 131 to 162 completely abolished fibronectin binding. All 17 clones possessed nucleic acid sequence encoding this segment of BBK32, indicating that this region is likely the principal fibronectin-binding domain of BBK32.

To further delimit the ligand-binding region of BBK32, we amplified, cloned, and expressed the segments of \textit{bbk32} encoding amino acids 131 to 162, 136 to 162, and 131 to 157 (AA 131-162, AA 136-162, and AA 131-157 constructs) and evaluated the fibronectin-binding activities of these clones by sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) and ligand blotting. The primers used to amplify these fragments of \textit{bbk32} are listed in Table 1. The amplification products were ligated into the pScreen T vector, and Novablu (DE3) competent \textit{E. coli} were transformed with each construct. Transformants were selected on LB agar plates supplemented with carbenicillin (50 \mu g/ml), and \textit{E. coli} clones possessing the corresponding Coomassie blue-stained gel. Deletion of five amino acids 131 to 162 of BBK32 were sufficient to mediate fibronectin binding. All 17 clones possessed nucleic acid sequence encoding this segment of BBK32, indicating that this region is likely the principal fibronectin-binding domain of BBK32.

The mapping of the fibronectin-binding regions of BBK32 and the development of a gene fragment library allowed the construction of expression plasmids for the various constructs. The expression plasmids were then transformed into \textit{E. coli} for the production of recombinant proteins. SDS-PAGE analysis indicated that the recombinant proteins migrated slower than expected, with molecular weights lower than those predicted for each construct. A similar observation has been made for recombinant proteins derived from different streptococcal and staphylococcal fibronectin-binding proteins (11). The AA 131-162 construct produced a 50-kDa protein that bound fibronectin, as demonstrated by SDS-PAGE and ligand blotting (Fig. 1). However, the level of fibronectin-binding activity displayed by this clone was much lower than those of the AA 131-167 and AA 105-162 constructs.

Seventeen colonies expressing fibronectin-binding activity by ligand blotting were selected for DNA sequence analysis. Plasmid DNA was purified from each of the 17 clones using QIAquick spin columns, and the insert DNA was sequenced by dye terminator cycle sequencing on an ABI 373 sequencer (PE Biosystems, Foster City, Calif.). Vector-specific primers, T7 primers, and STAG (Novagen) were used to sequence both strands of the insert DNA. Among the 17 clones expressing fibronectin-binding activity, the smallest \textit{bbk32} insert encoded amino acids 131 to 167 (AA 131-167 construct). Nucleic acid sequence from a second clone encompassed amino acids 105 to 162 of BBK32 (AA 105-162 construct). The sequence overlap between these two clones suggested that amino acids 131 to 162 of BBK32 were sufficient to mediate fibronectin binding.

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<thead>
<tr>
<th>BBK32 construct</th>
<th>Forward primer</th>
<th>Reverse primer</th>
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<tbody>
<tr>
<td>AA 131-162</td>
<td>5'-CAAGGAAGTTAATTCCT</td>
<td>5'-CCCTACAGGATTCTATAG</td>
</tr>
<tr>
<td>AA 131-157</td>
<td>5'-CAAGGAAGTTAATTCCT</td>
<td>5'-TATAAGAACATCTATAG</td>
</tr>
<tr>
<td>AA 136-162</td>
<td>5'-TCCCTTAAGCGGTTAAGT</td>
<td>5'-CCCTAAATCAGAATCTATAG</td>
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\textit{E. coli}, and determined the degree of sequence conservation and functional activity of the expressed proteins.

\textbf{TABLE 1. Oligonucleotide primers used for amplification of \textit{bbk32} gene fragments}

\begin{tabular}{|l|l|l|}
\hline
BBK32 construct & Forward primer & Reverse primer \\
\hline
AA 131-162      & 5'-CAAGGAAGTTAATTCCT & 5'-CCCTACAGGATTCTATAG \\
AA 131-157      & 5'-CAAGGAAGTTAATTCCT & 5'-TATAAGAACATCTATAG \\
AA 136-162      & 5'-TCCCTTAAGCGGTTAAGT & 5'-CCCTAAATCAGAATCTATAG \\
\hline
\end{tabular}
region of BBK32. A striking feature of this sequence is the relatively high number of acidic amino acid residues. This sequence characteristic has been noted for other bacterial fibronectin-binding proteins (11).

Functional activity and sequence analysis of BBK32 from different genospecies of *B. burgdorferi* sensu lato. Having defined the ligand-binding region of BBK32, we sought to establish whether BBK32 is functionally and genetically conserved among isolates of *B. burgdorferi* sensu lato. To this end, we cloned and expressed *bbk32* from four isolates representing different genospecies of *B. burgdorferi*: isolate B31, *B. burgdorferi* sensu stricto; isolate IP90, *B. garinii*; isolate ACA1, *B. afzelii*; and isolate DN127, *B. bissettii*. As previously described for isolate B31 (21), the *bbk32* gene was amplified from each isolate and cloned into the pMalc2 expression vector (New England Biolabs, Inc., Beverly, Mass.), in which the gene of interest is expressed as a peptide fused to a 42-kDa vector-derived peptide. In each case, the transformation of *E. coli* with a *bbk32* construct derived from either B31, IP90, ACA1, or DN127 resulted in the expression of an 80-kDa recombinant fusion protein that bound fibronectin, as determined by SDS-PAGE and ligand blot analysis (Fig. 2). The appearance of lower-molecular-weight bands having fibronectin-binding activity in Fig. 2 likely represents proteolysis of the overexpressed recombinant proteins. These results demonstrate that each *B. burgdorferi* isolate possesses a *bbk32* gene that was capable of encoding a functional protein upon expression in *E. coli*. In contrast, when spirochete lysates of B31, IP90, ACA1, and DN127 were tested by SDS-PAGE and ligand blotting for fibronectin-binding activity, only two of these four *B. burgdorferi* isolates expressed a fibronectin-binding protein (Fig. 3). As observed previously (21), isolate B31 expressed relatively high levels of a 47-kDa fibronectin-binding protein (BBK32), whereas ACA1 fibronectin-binding activity was visible as a faint band of 45 kDa. No fibronectin-binding activity was detected for the IP90 and DN127 lysates by ligand blotting. The lack of fibronectin-binding activity observed for isolates IP90 and DN127, despite the presence in both cases of a *bbk32* gene capable of encoding a functional protein, suggest that *bbk32* expression may be tightly regulated during in vitro cultivation of these isolates.

Next, we evaluated the degree of sequence conservation in the ligand-binding region of BBK32 by sequencing the cloned gene from isolates IP90, ACA1, and DN127. For each pMalc2/*bbk32* construct, three representative clones were sequenced by dye terminator cycle sequencing using an ABI 373 DNA...
FIG. 4. Alignment of BBK32 amino acid sequences from isolates representing different genospecies of *B. burgdorferi* with the ligand-binding region (amino acids 131 to 162) of BBK32 from isolate B31. The genospecies designations for the isolates are as follows: B31, *B. burgdorferi* sensu stricto; IP90, *B. garinii*; ACA1, *B. afzelii*; and DN127, *B. bissetti*. Sequence dissimilarity is indicated with a single-letter amino acid code. Sequence identity is shown as a period. Shaded amino acids residues represent sequence identity between the ligand-binding regions of BBK32 from isolate B31 and the UR region of protein F1 from *S. pyogenes*.

sequence identity among all four isolates.

A BLAST search of GenBank for sequences homologous to the BBK32 ligand-binding region from isolate B31 produced a single match with an in vivo-expressed protein described for *B. burgdorferi sensu stricto* isolate N40 (3). The N40 amino acid sequence was identical to the BBK32 ligand-binding region of *B. burgdorferi sensu stricto* isolate N40 (3). The N40 amino acid sequence motifs LSGESGEL and IESNEID were conserved among all four isolates.

We also aligned the BBK32 ligand-binding region from isolate B31 with *B. burgdorferi* sensu lato isolates sequenced in our study. Like BBK32, the protein F1 and the UR region of protein F1 from *S. pyogenes* mediate attachment of Lyme disease spirochetes to human cells.

Burgdorfer et al. (14) have demonstrated that the indirect interaction of protein F1 with β1 integrins via a fibronectin bridge can induce internalization of adherent *S. pyogenes* by the cell. The cellular uptake of *S. pyogenes* through the indirect interaction of protein F1 with integrins may provide this pathogen with a mechanism of penetrating tissue barriers and establishing deep tissue infections. Likewise, the ability of BBK32 to bind fibronectin, and indirectly integrins, may provide *B. burgdorferi* with a similar mechanism of cellular adherence and entry. The observation that β1 and β3 integrins may be involved in the adherence of *B. burgdorferi* to human cells lends support to this hypothesis (2). Furthermore, the ability of *B. burgdorferi* to invade cultured cells and survive intracellularly has been documented by electron microscopy, by confocal microscopy, and by antibiotic protection assays performed with ceftriaxone (5, 8, 12). Despite support from these in vitro studies, the in vivo capacity of *B. burgdorferi* to invade and persist within cells has proven difficult to establish. The potential intracellular existence of *B. burgdorferi*, however, may help explain the persistent nature of *B. burgdorferi* infections and the occasional failure of antibiotic therapy. We are currently investigating the role of BBK32 in cell adherence and invasion by expressing bbk32 in a heterologous host.

Nucleotide sequence accession numbers. The GenBank accession numbers for the bbk32 sequence from *B. burgdorferi* isolates B31, IP90, ACA1, DN127, and N40 are AE000788, AF213178, AF213179, AF213180, and U82107, respectively.

REFERENCES


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