Decorin Binding by DbpA and B of *Borrelia garinii*, *Borrelia afzelii*, and *Borrelia burgdorferi* Sensu Stricto

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**Background.** Decorin adherence is crucial in the pathogenesis of Lyme borreliosis. Decorin-binding proteins (Dbp) A and B are the adhesins that mediate this interaction. DbpA and B of *Borrelia garinii*, *Borrelia afzelii*, and *Borrelia burgdorferi* sensu stricto (ss) differ in their amino acid sequence, but little attention has been paid to the potential difference in their decorin binding.

**Methods.** We expressed recombinant DbpA and DbpB of *B. garinii*, *B. afzelii*, and *B. burgdorferi* ss and studied their binding to decorin. We also generated recombinant *Borrelia* strains to study the role of DbpA and DbpB in the adhesion of live spirochetes to decorin and decorin-expressing cells.

**Results.** Recombinant DbpA of *B. garinii* and DbpB of *B. garinii* and *B. burgdorferi* ss showed strong binding to decorin, whereas DbpA of *B. burgdorferi* ss and both DbpA and DbpB of *B. afzelii* exhibited no or only minor binding activity. DbpA and DbpB of *B. garinii* and *B. burgdorferi* ss also supported the adhesion of whole spirochetes to decorin and decorin-expressing cells, whereas DbpA and DbpB of *B. afzelii* did not exhibit this activity.

**Conclusions.** Dbp A and B of *B. garinii* and *B. burgdorferi* ss mediate the interaction between the spirochete and decorin, whereas the same adhesins of *B. afzelii* show only negligible activity.

Lyme borreliosis (LB) is a tickborne infectious disease caused by *Borrelia burgdorferi* sensu lato (Bbsl) which includes *B. garinii* (Bg), *B. afzelii* (Ba), and *B. burgdorferi* sensu stricto (Bsss) [1]. Late manifestations of LB are associated with different tissue tropism of borrelia genospecies. Bg has the tendency to cause neuroborreliosis, whereas Ba is mostly associated with chronic skin disorders, such as acrodermatitis chronica atrophicans and Bsss with Lyme arthritis. The molecular mechanisms behind this phenomenon are unknown.

Bbsl has several surface proteins that mediate attachment to different tissues and molecules of the host. It can bind to host plasminogen and plasminogen activators [2] and to various extracellular matrix and cell surface molecules [3]. Several adhesins have been identified in Bbsl, which mediate adherence to decorin [4], glycosaminoglycans (GAGs) [5], integrins [6, 7], fibronectin [8–10], laminin [9, 11, 12], and collagen [9, 13].

Decorin-binding proteins (Dbp) A and B are 2 well-characterized adhesins of Bsss [4, 14]. DbpA and B are expressed during mammalian infection [15], and they mediate bacterial attachment to the proteoglycan decorin, which is associated with collagen fibers in the extracellular matrix [16]. The decorin binding site of DbpA has been mapped to a conserved peptide motif (EAKVRA) [17]. Decorin consists of a collagen-binding core protein and a single GAG side chain that can be either dermatan sulphate or chondroitin-6-sulphate...
[18]. The binding target of Dbps is the GAG side chain, especially dermatan sulphate, in decorin [19].

Decorin is widely expressed throughout the body. High concentrations are detected in the skin, joints, and endothelium [20]. Decorin binding activity has an important role in borrelial dissemination, and it is also suggested that tissues rich in decorin might serve as a protective niche for Bbss [21]. In mice, for example, mutant strains lacking either one or both of the proteins are infective, but their dissemination to distant organs is less efficient than is the dissemination of the wild-type strain [22].

Thus far, all studies concerning the biological functions of Dbps have been performed with Bbss, and little attention has been paid to the potential differences in the decorin binding activity among the 3 genospecies. In amino acid sequence comparison, only 40–60% similarity was seen among the Dbps of the genospecies [23]. Variation in decorin binding activity may be a factor that in part determines the different tissue tropism of the 3 genospecies. It may also have a role in immune evasion and persistence of LB. The aim of this study was to characterize and compare the biological activities of the Dbps of Bg, Ba, and Bbss. To accomplish that, we took 2 approaches: (1) we recombinantly expressed individual Dbps and characterized their decorin binding properties and (2) generated recombinant borrelia strains in a surface protein–deficient background strain (B313) [24] to study the role of Dbps in the adhesion of live borrelia to decorin and decorin-expressing cells.

METHODS

Bacterial Strains and Culture Conditions

Low passage wild-type Bbss N40, Bbss B31, Bbss HB-19, Bbss SH–2-82, Bg SBK40, Bg SBK46, Bg Å218, Bg 387, Ba A91, Ba 570, Ba bo23, and Ba pKo and high-passage Bbss B313 and Ba 1082 were used in this study. All Bg and Ba strains were isolated from skin biopsy samples from Finnish patients except for Bg Å218, which is a tick isolate, and Ba pKo, which was kindly provided by Volker Fingerle (National Reference Centre for Borrelia, Bavarian Food and Health Safety Authority). Bbss wild-type strains were kindly provided by Sven Bergström (University of Umeå, Sweden), and B313 by Thomas Kamradt (Deutches Rheuma-Forschungszentrum). The spirochetes were cultivated in Barbour-Stoenner-Kelly II medium [25] at 33°C and were passaged weekly. Commercial *Escherichia coli* host strains (INV*F* [Invitrogen]; Novablu Giga Singles and BL21(DE3)-pLysS [Novagen]; M15 [Qiagen]) were used in cloning and expression of recombinant proteins.

Cloning and Expression of Recombinant Dbps

*E. coli* expressing DbpA and B of Bbss N40 were constructed with pET-30 Ek/LIC vector system (Novagen) according to the manufacturer’s instructions. The primers used in polymerase chain reaction (PCR) amplification of the *dbpA* and *B* are listed in Table 1. The recombinant strains expressing DbpA and B of Bg SBK40 and Ba A91 are described elsewhere [23, 26]. The proteins were purified under native conditions with use of Ni-NTA agarose beads (Qiagen) and were dialyzed against phosphate-buffered saline (PBS).

Generation of Recombinant B313 Strains

Recombinant B313 strains expressing DbpA and B of Bbss N40, Bg SBK40, and Ba A91 were constructed using previously described methods [27]. Primers used in PCR amplification of *dbpAB* operon are listed in Table 1. The resulting recombinant borrelia strains were named B313/dbpAB/Bbss, B313/dbpAB/Bg, and B313/dbpAB/Ba, and the control strain containing only pBSV2 was named B313/pBSV2.

Characterization of the Recombinant Borrelia Strains

The presence of DbpA and B was shown by Western blot assay using polyclonal anti-Dbp antiserum (Medprobe) and anti-rabbit IgG (Santa Cruz Biotechnology). Anti-flagellin B antibody (H9724; gift from S. Bergström) with anti-mouse IgG (Santa Cruz Biotechnology) was used as a control.

Surface localization of the adhesins was shown with proteinase-K treatment. The bacteria were washed twice with PBS containing

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**Table 1. Primers Used in PCR Amplification of dbpAB, dbpA or dbpB.**

<table>
<thead>
<tr>
<th>Primer</th>
<th>Vector</th>
<th>Sequence (5’-3’)</th>
</tr>
</thead>
<tbody>
<tr>
<td>dbpA/BbssREV</td>
<td>pET-30</td>
<td>GAC GAC GAC AAG GGA TTA AAA GGA AAA ACA AAA ATC</td>
</tr>
<tr>
<td>dbpA/BbssREV</td>
<td>pET-30</td>
<td>GAG GAG AAG CCC GGT TTA GTT ATT TTT GCA TTT TTC ATC</td>
</tr>
<tr>
<td>dbpA/BbssREV</td>
<td>pET-30</td>
<td>GAC GAC GAC AAG GGA TTA GTA GAA AGA ACA AAT GC</td>
</tr>
<tr>
<td>dbpA/BbssREV</td>
<td>pET-30</td>
<td>GAG GAG AAG CCC GGT TTA TTT CTT TTT GCT TTT ATT AT</td>
</tr>
<tr>
<td>dbpAB/BbssREV</td>
<td>pBSV2</td>
<td>AAAA GGA TGC ACA AGC CAG ATT GCA TAG C</td>
</tr>
<tr>
<td>dbpAB/BbssREV</td>
<td>pBSV2</td>
<td>AAAA GGA TGC ACA TTA TTT GGC AAA CTG GC</td>
</tr>
<tr>
<td>dbpAB/BbssREV</td>
<td>pBSV2</td>
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<td>pBSV2</td>
<td>AAAA GGA TGC CCC CTG GCA AAA TAA AAT TC</td>
</tr>
<tr>
<td>dbpAB/BbssREV</td>
<td>pBSV2</td>
<td>AAAA GGA TGC TTA TTT TTG ATT TTT AGT TTG TTT TTC</td>
</tr>
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**NOTE.** Restriction enzyme sites in the primers are bold type.
5 mM MgCl₂ and were diluted to 1 × 10⁸ bacteria/mL. Proteinase-K was added to the cell suspension at a final concentration of 100 μg/mL, incubated at room temperature for 30 min, and washed twice with the aforementioned buffer before preparing the samples for Western blot analyses.

**Western Blot Assay with Biotinylated Decorin**

Decorin (from bovine cartilage; Sigma) was biotinylated with EZ-Link NHS-LC-Biotin (Pierce) according to the manufacturer’s instructions. Biotinylated decorin (1 μg/mL) was used as the primary reagent, and HRP-conjugated streptavidin (0.1 μg/mL; Pierce) was used as the secondary reagent in the Western blot assay.

**Surface Plasmon Resonance Analysis**

Flow cell of the CM5 chip (GE Healthcare) was coated with the recombinant Dbp (50 μg/mL) in coating buffer (10 mM sodium acetate; pH 4.5 or 5.0) with use of the amine coupling kit, according to the manufacturer’s instructions, until ~8000 resonance units (RUs) were reached. A control cell lacking the adhesin was prepared under identical conditions. The changes in RUs caused by binding of decorin to Dbps on the chip were measured using Biacore X (Biacore AB). All binding experiments were run at room temperature. Decorin was dissolved in HBS-P running buffer (10 mM HEPES [pH, 7.4]; 0.15 M sodium chloride; 0.005% (v/v) Surfactant P20), and concentrations of 10 μg/mL, 50 μg/mL, and 100 μg/mL were used. Flow rate of 10 μL/min was applied in all measurements. The background signal from the buffer-treated reference cell was subtracted from the Dbp signal. The BIAevaluation program (version 3.0; Biacore) was used in the affinity curve calculations.

**Adhesion of Recombinant Dbps to Decorin in Microtiter Plate Assay**

Microtiter plates (Thermo Fisher Scientific) were coated with respective Dbp (10 μg/mL) in PBS and blocked with 1% BSA/PBS. The wells were incubated with biotinylated decorin (1 μg/mL), washed 3 times with PBS-T (PBS; 0.05% Tween 20), and incubated with 1 μg/mL alkaline phosphatase-conjugated streptavidin (Pierce). After washings, p-NPP-Na₂ substrate (1 mg/mL; Reagena) was added for 25 min before the reaction was stopped with 1M sodium hydroxide, and absorbances

Figure 1. A, Binding of decorin to recombinant Dbps in a Western blot assay. The upper panel represents SimplyBlue staining of the SDS-PAGE gel. The lower panel shows the adhesion of biotinylated decorin (1 μg/mL) to Dbps. B, Binding of decorin to recombinant Dbps in a microtiter plate assay. Recombinant Dbps (10 μg/mL) were attached on microtiter plates, and unspecific binding was blocked with BSA. The binding of biotinylated decorin (1 μg/mL) was detected with alkaline-phosphatase streptavidin and p-NPP-Na₂ substrate. Results are expressed as geometric mean of OD₄₅₀ values, subtracted with background, ± standard deviation of quadruplicate samples. Columns with same letter do not differ at 5% level of probability (Tukey HSD test).

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Figure 2. Surface plasmon resonance assay of decorin-Dbp interaction. Recombinant Dbps were attached covalently on CM5 chips (GE Healthcare), and unlabelled decorin was allowed to adhere to the chip. The binding was detected with Biacore 2000 equipment. A, Dose response of decorin (10, 50, and 100 μg/mL) binding to DbpB/Bg. B, Binding of decorin (100 μg/mL) to Dbps. The results are expressed as resonance units (RUs).
were measured using a Multiskan EX spectrophotometer (Thermo Fisher Scientific). All incubations were at 37 °C for 1 h, except for the substrate. Results are expressed as OD405 values subtracted with the background (a well without biotinylated decorin) absorbance. His-tag specific antibody (Penta-His biotin conjugate; Qiagen) was used to demonstrate the presence of Dbps on the wells (data not shown).

In inhibition experiments, the inhibitors were preincubated on the plates or with biotinylated decorin at 37°C for 1 h, and the assay was otherwise done as described above. Peptide LALREAKVQAIVETG (based on the DbpB sequence of Bg; 10 µg/mL; EAKVQA-peptide; Haartman Institute), dermatan sulphate (50 µg/mL; from shark cartilage; Sigma), chondroitin-6-sulphate (50 µg/mL; from shark cartilage; Sigma), and chondroitin-4-sulphate (50 µg/mL; from bovine trachea; Sigma) were used as inhibitors. Decorin (10 µg/mL) and BSA fraction V (100 µg/mL; Serological Proteins) were used as a positive and a negative control, respectively.

**Dot Adhesion Assay**

Wild-type and recombinant borrelia strains were washed and resuspended in PBS. One µL (5 × 10⁵ or 1 × 10⁶ bacteria) of the bacterial suspension was applied on a nitrocellulose membrane (Protran BA 85/20 0.45 µm; Schleicher & Schuell). The air-dried membrane was blocked with 3% nonfat dry milk in TBST (50 mM Tris-HCl [pH, 7.4]; 150 mM sodium chloride; 0.1% Tween 20) and incubated with biotinylated decorin (1 µg/mL) in TBST for 1 h at room temperature. The binding of biotinylated decorin was detected with streptavidin-HRP (0.1 µg/mL) in TBST. Detection was done as in the Western blot assay. The intensities of the dots were quantified with UVP BioSpectrum AC Imaging System. The unspecific binding of decorin to B313 was given a value of 1.

**Cell Adhesion Assay**

Human foreskin fibroblasts (HFFs) were cultured in Dulbecco’s modified Eagle medium (Invitrogen) containing 7% fetal calf serum (Hyclone) and gentamycin (Nalgene) in Lab-Tek 8-well chamber slide (Nunc) in a carbon dioxide incubator at 37°C for 2 days. Cells were fixed with −20°C acetone for 15 min and washed twice with PBS-T.

Recombinant borrelia strains were stained with carboxyl-fluorescein diacetate succinimidyl ester (CFSE; Molecular probes) as described elsewhere [28]. B313 and B313/pBSV2 were used as negative controls. CFSE-stained borrelia (10⁷) were allowed to adhere to the cells in slide chambers at 37°C for 1 h. The slides were washed twice with PBS, and the results were imaged with confocal microscope (Zeiss LSM510 META). Pictures of 10 random fields per chamber were analyzed with Image J analyzing program, and the number of individual bacteria was counted from every field.

To confirm the presence of decorin on the fixed cells, they were stained with monoclonal anti-decorin antibody (Santa Cruz Biotechnology) in PBS at 37°C for 1 h and washed twice with PBS-T. The unspecific binding of decorin to B313 was given a value of 1.
with PBS-T and once with PBS. HRP-conjugated anti-mouse IgG was used as a secondary antibody and incubated at 37°C for 1 h with the cells. After the washes, HRP was allowed to react with a peroxidase substrate (1 mM 3-amino-9-ethylcarbazole, 5% (v/v) dimethylformamide, 0.1% (v/v) hydrogen peroxidase in acetate buffer [pH, 5.5]) at room temperature for 30 min, and the decorin-stained cell cultures were imaged with optical microscope.

Inhibition assays were performed with B313/DbpAB/Bg and B313/DbpAB/Bbss. The inhibitors were preincubated with the CFSE-stained bacteria or the cells for at 37°C for 1 h, and the assay was otherwise performed as described above. EAKVQA-peptide (10 µg/mL), dermatan sulphate (50 µg/mL), chondroitin-6-sulphate (50 µg/mL), and chondroitin-4-sulphate (50 µg/mL) were used as inhibitors. Decorin (100 µg/mL) was used as a positive control, and BSA (100 µg/mL) was used as a negative control.

**Statistical Analyses**

The statistical significance of differences in the microtiter plate assay, dot adhesion assay, and cell adhesion assay were determined using analysis of variance (PASW statistics, version 18). Response variables were log10-transformed in microtiter plates and cell adhesion assay to normalize error distributions. Post hoc comparisons between means were done using Dunnett t test when there was a clear control and with Tukey’s honestly significant difference test in other cases.

**RESULTS**

**DbpB of Bg and Bbss Bind Decorin in Western Blot Assay**

First, the binding properties of recombinant His-tagged Dbps were studied. The PCR-amplified genes were sequenced (data not shown), and proteins of correct size were identified on sodium dodecyl sulfate polyacrylamide gel electrophoresis (Figure 1A). The binding of decorin to Dbps was analyzed using a Western blot assay using biotinylated decorin as the probe. Under these conditions, only DbpB/Bg and DbpB/Bbss showed clear binding to decorin.

**DbpB of Bg and Bbss Are the Strongest Decorin Binders in a Microtiter Plate Assay**

To study interaction between the Dbps and decorin under native conditions and to quantitate the binding, a microtiter plate assay was performed. This assay also showed clear binding of DbpB/Bg and DbpB/Bbss to decorin, which was statistically significantly higher than the binding of other Dbps (Figure 1B). The signals of the wells coated with DbpA/Bg, DbpA/Ba, DbpB/Ba, or DbpA/Bbss remained at a low level, although the binding of DbpB/Ba differed statistically significantly from other Dbps.

**DbpA and B of Bg Have the Highest Affinities to Decorin in Surface Plasmon Resonance Assay**

To characterize in more detail the affinities of individual Dbps to decorin, a surface plasmon resonance assay was performed.
Decorin binding to Dbps was dose dependent (Figure 2A). DbpB/Bg had the highest affinity to decorin (Figure 2B). In contrast to Western blot and microtiter plate assays, DbpA/Bg also showed clear binding to decorin. DbpB/Bbss exhibited moderate affinity to decorin, whereas DbpA/Bbss and DbpB/Ba showed only modest binding. The binding of DbpA/Ba to decorin was negligible at the tested conditions.

Taken together, the above results suggest that, depending on the mode of analysis, DbpB/Bg, DbpA/Bg, and DbpB/Bbss are the strongest decorin adhesins among the studied Dbps.

Characterization of Recombinant Borrelia Strains

Next, recombinant borrelia strains in the surface protein–deficient Bbss B313 background strain were constructed to study the role of Dbps in the adhesion of borrelia spirochetes to decorin and decorin-expressing cells. Correctness of the cloned operons was confirmed by sequencing (data not shown).

Western blot assay with specific antibodies of whole cell lysates showed that B313/DbpAB/Bg, B313/DbpAB/Ba, and B313/DbpAB/Bbss as well as wild-type Bg SBK40, Ba A91, and Bbss N40 express DbpA and B proteins (Figure 3A). Neither the background strain B313 (Figure 3A) nor B313 complemented with the pBSV2 vector only (data not shown) expressed the Dbps. Proteinase-K treatment showed the surface localization of Dbps in all recombinant strains (Figure 3B).

In line with the Western blot results using recombinant Dbps (Figure 1A) Western analysis of cell extracts from wild-type Bg SBK40 and Bbss N40 and recombinant B313/DbpAB/Bg and B313/DbpAB/Bbss identified one band reacting with decorin in each strain (Figure 3A). With Ba A91 or B313/DbpAB/Ba, no binding to decorin was detected.

Dbps of Bg and Bbss Support the Adhesion to Decorin

A dot adhesion assay was introduced to study the binding of intact borrelia to decorin (Figure 3C). The background level binding of B313 was given the value of 1. In agreement with aforementioned results, the B313/DbpAB/Bg and B313/DbpAB/Bbss recombinant strains bound decorin statistically

absorbances ± standard deviation of three experiments are shown. B and C, Inhibitions in the cell adhesion assay. The adhesion of B313/DbpAB/Bg and B313/DbpAB/Bbss to fibroblasts was inhibited with decorin, EAKVQA-peptide, dermatan sulphate, chondroitin-6-sulphate, and chondroitin-4-sulphate. BSA was used as a negative control. The average number of borrelia bound to fibroblasts in one confocal microscope field was counted from ten random fields in three independent experiments. Geometric mean ± standard deviations are shown. **P < .01, ***P < .001.
significantly, whereas the binding activity of B313/DbpAB/Ba was at the background level.

**Dbps of Bg and Bbss Mediate Adhesion to Decorin-Expressing Cells**

We next analyzed the ability of Dbps to support attachment of live borrelia to cells. HFF fibroblasts grown on chamber slides were shown to express decorin on their surface (Figure 4B). In the adhesion assay, CFSE-labelled borreliae were allowed to adhere to the cells, and the binding was quantitated using confocal microscopy. B313/DbpAB/Bg showed the highest adherence to the cells. B313/DbpAB/Bbss also bound well to the cells, whereas B313/DbpAB/Ba, B313/pBSV2, and B313 exhibited only background level of binding (Figure 4A and 4C). These results corroborate the unexpected finding that DbpA and DbpB of Ba, although expressed by this species, do not carry decorin binding activity.

**Molecular Dissection of the Interaction Between Dbps and Decorin**

It was previously shown with Bbss-derived Dbps that the decorin binding is inhibited by GAGs [19] or peptides containing the EAKVRA motif [17]. Therefore, we tested whether dermatan sulphate, chondroitin-6-sulphate, chondroitin-4-sulphate, or the EAKVQA-peptide could inhibit the interaction between decorin and DbpB/Bg. We also studied the role of these molecules in the cell adhesion assay by inhibiting the interaction between fibroblasts and B313/DbpAB/Bg or B313/DbpAB/Bbss.

In the microtiter plate assay, the binding of biotinylated decorin to DbpB/Bg was inhibited most efficiently with EAKVQA-peptide and dermatan sulphate, which both had effects similar to unlabelled decorin ($P \leq .001$). Chondroitin-6-sulphate and chondroitin-4-sulphate also inhibited the decorin binding significantly ($P \leq .001$), whereas BSA had no effect on the binding (Figure 5A).

In the cell adhesion assays, the adhesion of recombinant B313/DbpAB/Bg and B313/DbpAB/Bbss to fibroblasts was inhibited significantly with decorin, EAVKVA-peptide, and dermatan sulphate ($P \leq .001$). Chondroitin-6-sulphate and chondroitin-4-sulphate had only a minor, statistically non-significant effect on the binding. BSA, which was used as a negative control, did not influence the binding (Figure 5B and C).

In conclusion, the results of the inhibition assays confirm that the binding target of DbpB/Bg and DbpB/Bbss in decorin is the GAG side chain, especially dermatan sulphate and that the EAKVQA (or EAKVQA like) motif is the structure in the adhesins carrying the activity.

**Binding of Decorin to Wild-Type Bbsl Strains**

Finally, to analyze whether the decorin-binding properties of Bg SBK40, Ba A91, and Bbss N40 could be generalized to other Bg, Ba, and Bbss strains, a dot adhesion assay was performed with an array of wild-type strains. In this experiment, none of the 5 tested Ba strains exhibited decorin binding activity, whereas 8 of 9 Bg and Bbss strains showed statistically significant binding to decorin, compared with B313 (Figure 6).

**DISCUSSION**

Decorin binding activity of Bbss has an important role in borrelia infection [21, 22, 29–31]. However, all the studies concerning the decorin binding activity of borrelia have been performed using Bbss strains and Dbps derived from Bbss. Of
importance, variation in the structure and also the potential variation in ligand specificity of Dbps, which may be factors that, at least in part, determine the different tissue tropism of the 3 genospecies, have drawn little attention. The results of the present study suggest that DbpA and B of different Lyme disease Borrelia genospecies have different binding properties to decorin. Dbps of Bg and Bbss clearly support the adhesion of the bacteria to decorin, whereas the Dbps of Ba, although expressed on the bacterial surface, do not carry this activity.

Our Western blot assays with Bbss N40 whole cell lysates identified only one protein (DbpB) binding to decorin, which is in contrast to the seminal results of Guo et al [4], who showed, in a similar assay, that N40 expresses 2 proteins, later identified as DbpA and DbpB, with decorin binding activity. However, our Western blot results with the recombinant Dbps are in line with Borrelia lysate results suggesting that, indeed, only DbpB of N40 carries significant decorin binding activity. The microtiter plate assay and SPR results further confirm this finding. The reason for this discrepancy could be random mutations in dbpA gene of the N40 strain that is in use in our laboratory, but no mutations were detected in dbpA and dbpB when they were sequenced after PCR and compared with sequences in databases. In addition, tick isolates are known to be heterogeneous, which may explain the disparate results obtained with N40 in different laboratories.

A third explanation might be the different source of decorin used in our assays (bovine cartilage) and in the assays of Guo et al (bovine skin) [4]. However, whether these 2 proteoglycans are differently glycosylated or have other structural dissimilarities remains unknown.

On the basis of the current understanding that Bbss and Ba are the main genospecies associated with manifestations of the connective tissue (arthritis and acrodermatitis chronica atrophicans, respectively), it could be assumed that they would also show avid binding to decorin. This was, however, not the case because in all assays, either DbpB/Bg or B313/DbpAB/Bg bound to purified decorin or decorin-expressing cells most efficiently. Of interest, decorin is also expressed in the central nervous system [32] and cerebral endothelial cells [33], which implies that the strong affinity of Bg to decorin might have a role in the colonization of the CNS by Bg in neuroborreliosis. Another interesting observation was that DbpA/Bg showed dramatic variation in the adherence to decorin under different experimental conditions. In Western blot and microtiter plate assays (ie, under stationary incubation), DbpA/Bg showed a weak or no adherence, whereas in SPR analysis, under flow, it exhibited even higher affinity to decorin than did DbpB/Bss, which was at least a moderate binder in all assays. The high affinity of both DbpA/Bg and DbpB/Bg to decorin under constant flow could, in fact, be essential for the adherence to cerebral endothelial cells and, thus, for the initiation of central nervous system colonization. It can be hypothesized that DbpA/Bg forms a “catch bond” with decorin and needs shear force to induce the binding, as has been shown for certain other bacterial adhesins [34].

In our studies, the individual recombinant Dbps of Ba showed only little or no adherence to decorin, and the Borrelia strains expressing Ba Dbps, both wild-type and recombinant (Figure 3), did not adhere to purified bovine decorin or to human cells expressing decorin (Figure 4), more than the Dbp deficient background strain B313. This was also the case with 4 other Ba wild-type strains (Figure 6). Thus, it is likely that the Dbps of Ba have biological functions and ligands other than the Dbps of Bg and Bbss. Experiments addressing this question and the role of decorin binding in central nervous system colonization by Bg have been launched.

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References