Investigating BB0405 as a novel *Borrelia afzelii* vaccination candidate in Lyme borreliosis

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BB0405 is a surface exposed *Borrelia burgdorferi* protein and its vaccination protected mice against *B. burgdorferi* infection. As BB0405 is highly conserved across different *B. burgdorferi* sensu lato species, we investigated whether vaccination with recombinant BB0405 or through intradermal bb0405 DNA tattoo vaccination could provide protection against different *Borrelia* species, specifically against *Borrelia afzelii*, the predominant *B. burgdorferi* sensu lato genospecies causing Lyme borreliosis across Eurasia. We immunized C3H/HeN mice with recombinant BB0405 or with a codon-optimized bb0405 DNA vaccine using the pVAC plasmid and immunized corresponding control groups mice with only adjuvant or empty vectors. We subsequently subjected these immunized mice to a tick challenge with *B. afzelii* CB43-infected *Ixodes ricinus* nymphs. Upon vaccination, recombinant BB0405 induced a high total IgG response, but bb0405 DNA vaccination did not elicit antibody responses. Both vaccine formulations did not provide protection against *Borrelia afzelii* strain CB43 after tick challenge. In an attempt to understand the lack of protection of the recombinant vaccine, we determined expression of BB0405 and showed that *B. afzelii* CB43 spirochetes significantly and drastically downregulate the expression of BB0405 protein at 37 °C compared to 33 °C, where as in *B. burgdorferi* B31 spirochetes expression levels remain unaltered. Vaccination with recombinant BB0405 was previously shown to protect against *B. burgdorferi* sensu strico. Here we show that vaccination with either recombinant BB0405 (or non-immunogenic bb0405 DNA), despite being highly conserved among *B. burgdorferi* sl genospecies, does not provide cross-protection against *B. afzelii*, mostly likely due to downregulation of this protein in *B. afzelii* in the mammalian host.

Lyme borreliosis is the most common vector-borne disease in the Northern hemisphere and is caused by spirochetes belonging to the *Borrelia burgdorferi* sensu lato (sl) group. They are transmitted by *Ixodes* ticks and although humans get infected by *B. burgdorferi* sl, they are accidental hosts and do not play a role in the spirochete’s enzootic life cycle. Because *B. burgdorferi* sl are extracellular pathogens, the outer membrane of the spirochete, containing multiple surface exposed lipoproteins, is continuously exposed to the immune system of the host. Many studies have focused on identifying new *B. burgdorferi* outer surface proteins (Osps), because these are key targets for the host’s humoral immune response and thus may be potential new vaccinogens. Indeed, multiple protective Osps have been identified and OspA formed the basis of the only anti-Lyme vaccine that was publically available. There is however a wide genetic diversity among *B. burgdorferi* sl genospecies and the spirochetes change surface proteins throughout their life cycle, which makes it challenging to identify protective antigens.

Originally, Brooks et al. identified several surface-exposed *B. burgdorferi* sensu strico (ss) outer membrane proteins to which specific anti-*B. burgdorferi* antibodies were shown to be bactericidal. Among these proteins was BB0405, an outer membrane protein unique for *B. burgdorferi* sl species. With 78 to 90% identity between BB0405 orthologues, its sequence is highly conserved among *B. burgdorferi* ss, *B. afzelii* and *B. garinii*, the three major genospecies causing Lyme borreliosis. Although initially thought not to be immunogenic during natural
infection it was shown by Brooks et al. that BB0405 is immunogenic and actively expressed during nonhuman primate infection by *B. burgdorferi* by detecting the protein with sera from infected baboons. Multiple studies show that BB0405 is necessary for establishing infection in mice, since *bb0405*-deletion mutants are unable to be transmitted from ticks and establish infection in mammalian hosts. Of importance, vaccination with recombinant *BB0405* also protected mice from *B. burgdorferi* infection by *B. burgdorferi*-infected ticks. Thus, *bb0405* is a highly conserved antigen with the potential to form the basis for a vaccine protecting against multiple *B. burgdorferi* sl genospecies.

Most research on new Lyme vaccines focuses on recombinant proteins, but DNA vaccination constitutes an alternative vaccination platform. For instance, a previous study by Wagemakers et al. has shown that DNA vaccination by tattoo with *B. afzelii* strain PKo Outer surface protein C (OspC) was fully protective against *B. afzelii* challenge in mice and induced favorable humoral immune responses compared to recombinant protein vaccination. In line with this, we were able to show protection against *B. burgdorferi* strain N40 in a similar set-up, in which OspC from *B. burgdorferi* strain N40 was used both as recombinant as well as DNA vaccine (Klouwens et al. manuscript in preparation).

In the current study we aimed to investigate the role of BB0405 in providing protection across *B. burgdorferi* sl genospecies. To this end, mice were immunized with *B. burgdorferi* B31-derived recombinant BB0405 or *bb0405* DNA vaccine and subsequently challenged with *B. afzelii* CB43-infected ticks, after which immunogenicity and host protection of the two different vaccination approaches were determined using established methods.

### Results

#### Immunogenicity of *bb0405* antigens.

As described previously, BB0405 is a highly conserved *B. burgdorferi* sl surface protein and alignment of the protein *BB0405* of *B. burgdorferi* B31 and *B. afzelii* CB43 showed an identity of 88% and similarity of 96% at the amino acid sequence level (Fig. 1). To determine whether antibodies against BB0405 would protect across different *B. burgdorferi* sl genospecies, we performed a vaccination study in mice. Recombinant BB0405 and a DNA vaccine for *bb0405*, both generated from *B. burgdorferi* B31, were constructed as well as an empty DNA vaccine, functioning as the negative control. From our previous published and unpublished studies it is known that an empty DNA vaccine, i.e. a pVAX vector without inserted target sequences, does not affect *B. burgdorferi* sl infection (10 and Klouwens et al. manuscript in preparation).

![Figure 1](https://example.com/figure1.png)

**Figure 1.** Alignment between the BB0405 protein sequence between *Borrelia burgdorferi* B31 and *Borrelia afzelii* PKO. *Borrelia burgdorferi* B31 corresponds to NCBI Reference Sequence: NP_212539. *Borrelia afzelii* PKO is identical to CB43 (GenBank: CP002933.1; translated in ExPASy translate tool). Identity between sequences is 88.2%. Additionally there are 17 similar positions bringing the similarity to 96.5%. Alignment was performed with Clustal Omega. * = identical, : = similar.

In the current study we aimed to investigate the role of BB0405 in providing protection across *B. burgdorferi* sl genospecies. To this end, mice were immunized with *B. burgdorferi* B31-derived recombinant BB0405 or *bb0405* DNA vaccine and subsequently challenged with *B. afzelii* CB43-infected ticks, after which immunogenicity and host protection of the two different vaccination approaches were determined using established methods.

#### Host-protection via BB0405 immunization.

Next, we assessed whether both vaccines were able to provide protection against *B. afzelii* infection by performing qPCR and culture of several tissues, obtained by sacrificing the mice 21 days after challenge with *B. afzelii* strain CB43-infected ticks. *B. afzelii* loads in ear, skin (tick feeding site), heart, bladder and ankle were quantified and normalized to mouse β-actin. In addition, bladder and skin tissues from the feeding site of the ticks were cultured and checked weekly for the growth of spirochetes. Despite the high antibody titers, results of qPCR and culture indicate that vaccination with BB0405, either recombinant or as DNA vaccine, did not protect against heterologous *Borrelia* challenge in mice (Table 1).
Indeed, there were no significant differences between the separate groups in the different tissues, nor between experimental groups at a cumulative level.

**BB0405 surface expression B. burgdorferi sensu stricto and B. afzelii.** To study the location of the protein and also the potential accessibility of the protein to antibodies we performed a proteinase K assay with viable *B. burgdorferi* ss and *B. afzelii* spirochetes followed by Western Blot analysis. We show that BB0405 is partly surface exposed as the protein is partly digested upon proteinase K treatment for both 30 and 60 min both in *B. burgdorferi* ss as well as in *B. afzelii*. Flagellin B, a periplasmic control protein, was, as expected, not affected by proteinase K treatment (Fig. 3).

**Expression of BB0405 or homologous proteins in B. burgdorferi sensu stricto and B. afzelii.** Finally, in an attempt to understand why, in contrast to previous observations for *B. burgdorferi* ss, BB0405 did not protect against *B. afzelii*, we next precisely compared the expression levels of BB0405 between these two different *B. burgdorferi* sl genospecies under different experimental conditions. We did not find differences in expression of *bb0405* at the RNA level between *B. burgdorferi* ss and *B. afzelii* grown at 33 °C or 37 °C (data not shown). However, we also assessed BB0405 expression at the protein level. To this end, equal amounts of whole lysates of spirochetes, cultivated in vitro at different temperatures, were run on a Western blot and incubated with BB0405 specific antibodies. As can be appreciated from Fig. 4A, *afzelii* CB43 produces considerably lower amounts of BB0405 at 37 °C, the condition similar to the conditions in the mammalian host. We also quantified the protein bands on the Western Blots (Image J software) and measured the relative density of the bands compared to a corresponding Flagellin B loading control (Fig. 4B). Indeed, the relative density of BB0405 in *B. afzelii* CB43 cultured at 37 °C was significantly lower compared to BB0405 expression at 33 °C and to *B. burgdorferi* B31 at 37 °C, offering a possible explanation for our findings.
Discussion

An ideal Lyme vaccine candidate would need to provide protection against the most dominant \textit{B. burgdorferi} sensu stricto and \textit{B. afzelii}. The main \textit{B. burgdorferi} sensu stricto genospecies causing Lyme borreliosis in North-America is \textit{B. burgdorferi ssp}, whereas in Europe this is \textit{B. afzelii}. To explore the potential of BB0405, in the current study we investigated whether vaccination with BB0405—a surface-exposed \textit{B. burgdorferi} ss protein that is well-conserved among different \textit{B. burgdorferi} sensu stricto genospecies including \textit{B. afzelii}—protected against heterologous challenge with \textit{B. afzelii} through tick-bite in an experimental mouse model. Moreover, two different methods of vaccination with BB0405 were used, recombinant protein vaccination and DNA vaccination by tattoo. Finally, to provide further insights into our findings we assessed the differential BB0405 expression of \textit{B. burgdorferi} ss and \textit{B. afzelii} under varying conditions.

An adequate humoral response is essential in clearing \textit{Borrelia} spirochetes and specific immunoglobulins play a key role in affording protective host immunity. Since BB0405 is an outer surface protein, it is surface exposed and therefore likely to be antibody-accessible. Surprisingly, Kung et al. described the interesting phenomenon that no BB0405 specific antibodies are generated during natural infection in mice; however Brooks et al. showed that recombinant BB0405 is immunogenic in rats and antibodies raised against BB0405 could kill \textit{B. burgdorferi ssp} in vitro in the presence of complement. In addition, in mice immunized with recombinant BB0405 a strong and long-lasting antibody response was induced that provided protective immunity against tick-transmitted infection with \textit{B. burgdorferi}. In the current study we also show a robust humoral immune response in mice vaccinated with recombinant BB0405 compared to control mice, corroborating the observation that recombinant BB0405 is immunogenic. In contrast, we did not observe antibody responses against BB0405 in the mice vaccinated with the \textit{bb0405} DNA vaccine. Interestingly, the same DNA vaccination approach has been successful in the past for other \textit{Borrelia} outer surface proteins. In our study this could indicate that the mice cells were not able to transcribe, translate or translocate \textit{bb0405} and that the utility of DNA vaccination against \textit{B. burgdorferi} sensu stricto is highly dependent on the target. However, as murine infection with \textit{B. burgdorferi ssp}, either by syringe or ticks, does not result in BB0405-specific antibody responses, it could also be that \textit{bb0405} needs the adjuvants used in recombinant protein vaccinations and is not immunogenic by itself.
In the study described here, we have challenged the BB0405 (B. burgdorferi B31 derived)-immunized mice with B. afzelii infected ticks. We did not observe any protection of BB0405, neither as recombinant vaccine nor as DNA tattoo. Based on our observations, BB0405 does not seem to be a suitable vaccine candidate for the European situation, as it does not provide cross-protection between B. burgdorferi ss and B. afzelii—despite high antibody titers after vaccination with recombinant BB0405. Although we have performed a heterologous challenge, the in silico analysis showed high sequence homology between these proteins. Indeed in Figs. 3 and 4 we show that the raised antibodies recognize and bind to BB0405 expressed by B. afzelii. It seems therefore unlikely that antigen recognition can explain the observed lack of vaccine efficacy. Future studies could assess whether using different species-specific BB0405 homologues from different B. burgdorferi ss genospecies could protect against heterologous challenge.
As stated above, antigen specificity of the generated antibodies is unlikely to explain the observed lack of efficacy. Therefore we firstly wanted to determine whether BB0405 was indeed surface expressed in *B. afzelii* strain CB43, as has been shown *B. burgdorferi* ss strain B31. We have shown that BB0405 is indeed partly expressed at the cell surface by performing a Western Blot analysis after treating viable *B. afzelii* B43 spirochetes with proteinase K. We used *B. burgdorferi* B31 as a control. The reason that there is only partial surface expression in both *B. burgdorferi* ss strains can be explained by BB0405 being a transmembrane protein and thus partly intracellular or membrane bound, explaining the lower protein products on the Western Blots of Fig. 3. Secondly, there are over 20 different species within the *B. burgdorferi* sl complex and these are capable of molecular adaptations in order to ensure efficient transmission by their vector and to survive in different environments of their host. Temperature is a key environmental factor known to affect *B. burgdorferi* sl gene expression. More specifically, it has been described by Ojaimi et al. that BB0405 is upregulated by temperature suggesting upregulation in conditions that mimic the situation in the mammalian host. We therefore wondered whether a difference in protein expression of BB0405 between American and European *B. burgdorferi* sl genospecies could explain the lack of protection in our vaccination experiment. At the protein level we observed a distinct and significantly lower expression of BB0405 in *B. afzelii* CB43 spirochetes grown at 37 °C, as compared to the expression in the same spirochetes grown at 33 °C. More importantly, expression of BB0405 in *B. afzelii* CB43 spirochetes grown at 37 °C was also significantly lower compared to *B. burgdorferi* B31 spirochetes grown at the same temperatures. This difference was not apparent at the RNA level. However, it is well-known that RNA expression levels do not necessarily correspond to the protein expression levels. Thus, the BB0405 protein seems to be evidently less expressed by *B. afzelii* CB43 in conditions reflecting the mammalian host and this might explain the observed lack of protection against *B. afzelii* strain CB43, despite the presence of specific antibodies upon vaccination with recombinant BB0405. An alternative explanation for the lack of protection could be an indirect effect of failure of antibody recognition. *B. burgdorferi* sl is able to alter surface protein in vivo via differential gene expression and via VlsE recombination which might impact spirochete recognition by host-generated antibodies. There are multiple examples of surface-exposed Borrelia antigens that are masked by neighboring (abundant) proteins. Although BB0405 is conserved across *B. burgdorferi* ss and *B. afzelii*, which may account for different surface topology and outcome of antibody-mediated protections.

In conclusion, being a surface-exposed, immunogenic and well-conserved *B. burgdorferi* sl protein, BB0405 was shown to be an interesting vaccine candidate to protect against Lyme borreliosis caused by *B. burgdorferi* ss. However, we here show that vaccination with *B. burgdorferi* ss-derived BB0405 does not protect against heterologous challenge with *B. afzelii* through tick-bite, and our data suggest that this could be due to lower expression of the BB0405 homologue at the protein level in *B. afzelii* in the mammalian host. Nevertheless, experiences with the OspA vaccine have shown that a vaccine does not have to protect against all *B. burgdorferi* sl genospecies to be commercially viable. Future experiments should investigate whether multivalent BB0405 vaccines are able to protect against multiple *B. burgdorferi* sl genospecies.

**Methods**

**Ethics statement.** All experiments were reviewed and approved by the Animal Research Ethics Committee of the Academic Medical Center, Amsterdam, The Netherlands (protocol 208AK-1 and 271AA-1). Experiments have been conducted according to European and national guidelines http://eur-lex.europa.eu/legal-content/EN/TXT/?uri=celex:32010L0063 and in compliance with the ARRIVE guidelines (http://www.nc3rs.org.uk/page.asp?id=1357).

**Recombinant BB0405 protein.** The *bb0405* gene was cloned without the N-terminal leader sequence into pET28a (Invitrogen), produced in *E. coli*, and purified using Ni–NTA resin as detailed elsewhere. Sequence alignment between BB0405 from different strains was performed using Clustal Omega.

**Generation of the BB0405 DNA vaccine.** The pVAXhTPA-BB0405 DNA vaccine was designed as described before in Wagemakers et al. In the BB0405 gene sequence of *B. burgdorferi* B31 (NCBI Reference Sequence: NC_001318.1) the 18aa signal sequence (predicted by SignalP 4.0 web-based software, CBS, Lyngby, Denmark) was replaced with the human tissue plasminogen activator (hTPA) signal sequence (genbank AA61213.1). The resulting sequence was codon-optimized to mouse tRNA usage and via VlsE recombination which might impact spirochete recognition by host-generated antibodies. There are multiple examples of surface-exposed Borrelia antigens that are masked by neighboring (abundant) proteins. Although BB0405 is conserved across *B. burgdorferi* ss and *B. afzelii*, which may account for different surface topology and outcome of antibody-mediated protections.

**Vaccination experiments.** Six to eight weeks old female C3H/HeN mice were purchased from Charles River and the experiment was designed with 3 groups of eight mice each. The vaccination experiment was carried out as described in a previous publication of our group. The first group was vaccinated with a recombinant BB0405 vaccine (recBB0405), the second group with a BB0405 DNA vaccine, and the third group consisted of an empty vector DNA vaccination group as a negative control. Mice were vaccinated at *t* = 0, *t* = 14 and *t* = 28 days and sera were collected at each time point for use in ELISA experiments. For the recombinant BB0405 vaccine 10 μg protein was emulsified with complete Freund's adjuvant at *t* = 0 and 5 μg in incomplete Freund's adjuvant at *t* = 14 and *t* = 28 days (Fig. 3). All vaccinations were administered subcutaneously. For the pVAX-hTPA-BB0405 DNA vaccine and the negative control hair was removed from the mice abdomens using hair removal
Borrelia control sample in which OspA is still detectable + 3xSD. replaced by the value of the OspA detection limit. The detection limit is defined as the highest dilution of the B. afzelii known to be infected with B. afzelii. In case of a negative value for OspA in a sample, the OspA value was 20 s and 72 °C 20 s. Reactions were performed in triplicate. Results were analyzed using LinRegPCR software using SYBR green dye (Roche) using the following PCR protocol: 95 °C 6 min, and 60 cycles of 95 °C 10 s, 60 °C C-3.

β-actin forward 5′-AAA AAT ATT TAT TGG GAA TAG GTC T-3′ and reverse 5′-AGC GGG AAA TCG TGC GTG -3′ were used for normalization. The qPCRs were performed on the LightCycler480 (Roche, Nutley, NJ, USA) ELISA plate reader at 450–655 nm.

B. afzelii DNA in mouse tissues and gen, Venlo, The Netherlands). Quantitative (q)PCR was used to quantify was performed according to previously described protocol. OspA primers were used for quantification; for -ward 5′-TGC AGC GAA ATT TTT TGA, mouse Beta-actin forward 5′-AGCGGGAATCTGCGGTG-3′ and reverse primer 5′-CAGGGTACATGGTGTTGC G-3′ were used for normalization. The qPCRs were performed on the LightCycler480 (Roche, Nutley, NJ, USA) using SYBR green dye (Roche) using the following PCR protocol: 95 °C 6 min, and 60 cycles of 95 °C 10 s, 60 °C 20 s and 72 °C 20 s. Reactions were performed in triplicate. Results were analyzed using LinRegPCR software (Amsterdam, The Netherlands). Negative and positive controls were included in each qPCR run. A positive Borrelia load was determined by at least one melting curve exactly matching the positive control (skin tissue known to be infected with B. afzelii). In case of a negative value for OspA in a sample, the OspA value was replaced by the value of the OspA detection limit. The detection limit is defined as the highest dilution of the control sample in which OspA is still detectable + 3xSD.

ELISA. To measure IgG directed against BB0405, ELISAs were performed according to previous described protocol. High-binding 96-well ELISA plates (Greiner Bio-one, Kremsmünster, Austria) were coated overnight at 4 °C with 1 μg/ml recBB0405, washed with PBS–Tween (phosphate-buffered saline–0.05% Tween) and incubated with blocking buffer (1% BSA in PBS) for 2 h at room temperature. Mouse sera (collected at day 42 before tick challenge) were diluted in blocking buffer, added to the wells and incubated for 1 h at room temperature. Plates were washed and incubated for 1 h with horseradish peroxidase (HRP)-linked anti-mouse IgG (Cell Signaling, Beverly, MA, USA) diluted 1:1000 in blocking buffer. The plates were washed again and developed using TMB substrate (50 μl TMB chromogene in 5 ml TMB substrate buffer (8.2 g NaAc and 21 gr citric acid monohydrate dissolved in 1 L H2O + 10 μl 3% H2O2) and optical density was measured in a Biotek (Winooski, VT, USA) ELISA plate reader at 450–655 nm.

Borrelia afzelii detection and quantification. Murine bladder and skin samples were cultured in modified Kelly Pettenkofer (MKP) medium with rifampicin, 50 μg/ml and phosphomycin, 100 μg/ml) at 33 °C. The cultures were checked weekly (for a total of 8 weeks) for the presence of motile spirochetes with dark field microscopy as described before. For all samples DNA was extracted using Qiagen Blood and Tissue kit (Qiagen, Venlo, The Netherlands). Quantitative (q)PCR was used to quantify B. afzelii DNA in mouse tissues and was performed according to previously described protocol. OspA primers were used for quantification; forward 5′-AAAAATATTTATGGGAAATAGGTCT-3′ and reverse 5′-CACCAGGCAAAAATCTACTGAA-3′, mouse Beta-actin forward 5′-AGCGGGAATCTGCGGTG-3′ and reverse primer 5′-CAGGGTACATGGTGTTGC G-3′ were used for normalization. The qPCRs were performed on the LightCycler480 (Roche, Nutley, NJ, USA) using SYBR green dye (Roche) using the following PCR protocol: 95 °C 6 min, and 60 cycles of 95 °C 10 s, 60 °C 20 s and 72 °C 20 s. Reactions were performed in triplicate. Results were analyzed using LinRegPCR software (Amsterdam, The Netherlands). Negative and positive controls were included in each qPCR run. A positive Borrelia load was determined by at least one melting curve exactly matching the positive control (skin tissue known to be infected with B. afzelii). In case of a negative value for OspA in a sample, the OspA value was replaced by the value of the OspA detection limit. The detection limit is defined as the highest dilution of the control sample in which OspA is still detectable + 3xSD.

RNA and cDNA synthesis. B. burgdorferi B31 and B. afzelii CB43 were cultured in MKP medium at 33 °C or 37 °C to 1 × 10e6 spirochetes/ ml as assessed by a Petroff-Hausser counting chamber and dark-field microscopy. The cultures were centrifuged for 10 min at 10,000 rpm and the pellets were dissolved in 750 μl RNA Later (Qiagen) stored at − 80 °C until further use. All samples were thawed simultaneously and centrifuged for 5 min at 12,000 rpm. The pellets were then subsequently used for isolation of RNA using the Nucleospin RNA isolation kit (Macherey–Nagel), according to the manufacturer’s instructions. Subsequently, RNA samples were digested by DNase for a second time using the Qiagen RNase-Free DNase Set (#79254) and then cleaned up using the RNeasy MinElute Cleanup Kit (#74104, Qiagen). A total of 10 μl of each RNA sample was then used to generate cDNA and heated for 5 min at 85 °C and then cooled to 23 °C. To every sample, 10 μl of RT mix was added, consisting of 4 μl of M-MLV reverse transcriptase buffer (Promega), 0.5 μl of M-MLV reverse transcriptase enzyme 200U/ml (Promega), 2 μl dNTP mix (Invitrogen), 1 μl of random hexamers (), 0.5 μl DTT (Invitrogen), 0.5 μl RNaseOut (Invitrogen), as well as 1.5 μl of RNase free H2O. The PCR protocol for all samples was 23 °C 10 min, 42 °C 60 min and 95 °C 3 min (2720 Thermal Cycler, Applied Biosystems). To check the purity of the RNA, a PCR was performed using RNA and cDNA (100× diluted) of corresponding samples as a template with...
universal flaB primers (flaB forward 5'-GCTTCTGATGCTGCTGCTGCTG-3' and flaB reverse 5'-CGTCTGTAAGTGCCTGCCAT-3') and PCR Phusion High Fidelity Mastermix (#M0531S, NEB). As expected, there was no amplification in the samples when DNA was used as template, indicating there was no DNA contamination. Real-time quantitative polymerase chain reaction (qRT-PCR) was used to quantify the amount of BB0405 RNA in the two different B. burgdorferi sl strains, grown at different temperatures. The qRT-PCR was performed with BB0405 specific primers for B. afzelii CB43 (forward 5'-GCTTCTGATGCTGCTGCTGCTG-3' and reverse 5'-GGG TTTAAGTCCACGAACCC-3') and for B. burgdorferi B31 (forward 5'-GGCATATGTTTTGATTTGGG-3' and reverse 5'-CCATCACCAATAGGGCCAGG-3') and with Flagellin B universal primers as housekeeping genes (forward 5'-GCTTCTGATGCTGCTGCTGCTG-3' and reverse 5'-GCTGTAAGTGGCTGCTATTCC-3'). The cDNA from the two different B. burgdorferi sl strains grown under different conditions was diluted 100 times and 1 μl was used as template per reaction and qPCRs were performed using the LightCycler480 (Roche, Nutley, NJ, USA) and SYBR green dye (Roche) in triplicate. The PCR protocol was 95°C 60 min, and 60 cycles of 95°C 10 s, 60°C 20 s and 72°C 20 s. Result were analyzed using LinRegPCR software (Amsterdam, The Netherlands).

**Borrelia lysates and BB0405 western blots.** B. burgdorferi B31 and B. afzelii CB43 were cultured in MKP medium at 33 °C or 37 °C until log phase was reached, after which the cultures were centrifuged and the spirochete pellets were washed three times in PBS for 10 min at 10,000 g. Ultimately, the pellets were dissolved in 250 μl of PBS and were sonicated on ice (6 times 15 s, on 30 s off with 20% amplitude using Vibra-Cell High Intensity ultrasonic processor). Protein concentrations were measured using the Pierce BCA protein assay (#23225 from Thermo Scientific). Subsequently, lysates (2.5 μg protein per sample) were mixed in a 5:1 ratio with 0.0025% bromophenol blue) and subjected to sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS-PAGE). The membrane bound proteins were transferred onto a nitrocellulose membrane. After blotting, the membrane was cut in half just between the bands of Flagellin B (41 kDa) and proteinase K at a concentration of 200 μg/ml for either 30 or 60 min at room temperature. Phenylmethylsulfonyl fluoride (PMSF)(Sigma) was added (1.5 mM) to stop proteinase K activity. Spirochetes were again washed twice in PBS + 5 mM MgCl2 and were pelleted. Pellets were resuspended gently in PBS + 5 mM MgCl2 and split in equal volumes. They were incubated in PBS + 5 mM MgCl2 either in the absence or presence of proteinase K at a concentration of 200 μg/ml for either 30 or 60 min at room temperature. Phenylmethylsulfonyl fluoride (PMSF)(Sigma) was added (1.5 mM) to stop proteinase K activity. Spirochetes were again washed twice in PBS + 5 mM MgCl2 + 1 mM PMSF and pelleted. The pellets were resuspended in an appropriate volume of SDS-PAGE loading buffer for further Western Blot analysis. Western Blots were carried out as described above.

**Proteinase K assay.** Proteinase K assays were carried out as described elsewhere. Spirochetes in mid-phase growth were washed twice with PBS + 5 mM MgCl2 and were pelleted. Pellets were resuspended gently and split in equal volumes. They were incubated in PBS + 5 mM MgCl2 either in the absence or presence of proteinase K at a concentration of 200 μg/ml for either 30 or 60 min at room temperature. Phenylmethylsulfonyl fluoride (PMSF)(Sigma) was added (1.5 mM) to stop proteinase K activity. Spirochetes were again washed twice in PBS + 5 mM MgCl2 + 1 mM PMSF and pelleted. The pellets were resuspended in an appropriate volume of SDS-PAGE loading buffer for further Western Blot analysis. Western Blots were carried out as described above.

**Statistic methods.** Differences between experimental groups were compared using an unpaired t test (Graphpad Prism software version 5.0, San Diego, CA, USA). Differences in expression level between experimental groups on Western Blot were compared using LinRegPCR software (Amsterdam, The Netherlands).

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Author contributions
M.J.K. and J.J.T. designed the study. J.I.E., J.J.T. and M.J.K. performed the study procedures. F.N.M.P. assisted in the study procedures. R.S. and O.H. provided us with Exodes ricinus nymphs infected with Borrelia afzelii CBA3 and provided scientific input. U.P. and M.T. generated and provided the recombinant proteins and provided scientific input. J.W.H. supervised the study progress. All authors revised the draft manuscript and agreed to the final manuscript.

Competing interests
The authors declare no competing interests.

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