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Host tropism determination by convergent evolution of immunological evasion in the Lyme disease system

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Abstract

Pathogens possess the ability to adapt and survive in some host species but not in othersan ecological trait known as host tropism. Transmitted through ticks and carried mainly by mammals and birds, the Lyme disease (LD) bacterium is a well-suited model to study such tropism. Three main causative agents of LD, Borrelia burgdorferi, B. afzelii, and B. garinii, vary in host ranges through mechanisms eluding characterization. By feeding ticks infected with different Borrelia species, utilizing feeding chambers and live mice and quail, we found species-level differences in bacterial transmission. These differences localize on the tick blood meal, and specifically complement, a defense in vertebrate blood, and a polymorphic bacterial protein, CspA, which inactivates complement by binding to a host complement inhibitor, Factor H (FH). CspA selectively confers bacterial transmission to vertebrates that produce FH capable of allele-specific recognition. CspA is the only member of the Pfam54 gene family to exhibit host-specific FH-binding. Phylogenetic analyses revealed convergent evolution as the driver of such uniqueness, and that FH-binding likely emerged during the last glacial maximum. Our results identify a determinant of host tropism in Lyme disease infection, thus defining an evolutionary mechanism that shapes host-pathogen associations.

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Author summary

The evolution of pathogens in response to a host environment often leads to a specificity of these pathogens to survive in particular hosts, known as host tropism. However, the mechanisms of host tropism remain unclear. The Lyme disease (LD) bacterium is an ideal model to study such mechanisms because the causative agent of this disease (bacteria species) are transmitted by ticks and carried by multiple vertebrates, including mammals and birds. We found that different LD bacteria species vary in their ability to survive in mice and quail, and in ticks fed on human or quail blood after transmission. We also showed that such a variation is dependent on whether tick blood meals have functional complement, an innate defense in vertebrate blood. We attributed such species-to-species differences of transmission to the polymorphisms of a bacterial complement inactivating protein, CspA: CspA, by inactivating vertebrate complement in tick blood meals in a species-specific manner, allows bacterial transmission to selective vertebrates. We further demonstrated that this CspA-mediated bacterium species-specific complement evasion arose through convergent evolution, as an evolutionary mechanism to permit such vertebrate-bacteria interaction. The mechanisms defined in this multi-disciplinary work provide new insights into the host-pathogen interactions.

Introduction

The interactions between hosts and pathogens (e.g. bacteria, viruses, and protozoa) have arisen through numerous evolutionary events [1,2], often resulting in generalist pathogens, which adapt to most host environments, or specialists, which selectively survive in particular host species. The association between pathogens and their respective hosts is defined as "host tropism (or host specialization)" [3]. For vector-borne pathogens, such a host tropism can be dictated by not only host factors but also host constituents in the vectors (e.g. blood meals) [4]. Because many bacteria varying in host specificity are involved in the infection cycle, the Lyme disease bacterium is one of the models regularly applied to investigate the host-pathogen interactions [4]. Carried by Ixodes ticks, this disease is the most common vector-borne disease in the northern hemisphere [5]. The causative agent of Lyme disease is a genospecies complex of the spirochete Borrelia burgdorferi sensu lato (also known as Borreliella burgdorferi sensu lato (NCBI taxid: 139), Lyme borreliae) [6]. Among these genospecies, the most frequently isolated spirochetes from both ticks and vertebrate hosts are B. afzelii, B. garinii, and B. burgdorferi sensu stricto (hereafter referred to as B. burgdorferi) [7]. Following the tick bite, spirochetes need to survive in tick blood meals, which permit transmission to the bite site of the host skin. Survival in the host bloodstream then is a prerequisite for hematogenous dissemination and colonization of distant tissues, resulting in varied disease manifestations involving different organs in incidental hosts such as humans [8]. In nature, Lyme borreliae can invade vertebrate reservoirs (mainly birds and rodents) but varying in their ability to infect these animals in a host-specific manner [9,10]. For example, some Lyme borreliae species, such as B. burgdorferi, can invade a wide range of hosts whereas others are selectively infectious in few host taxa (e.g. B. afzelii for rodents and B. garinii for birds). However, the molecular basis for such host tropism is largely unclear [9,10].

The elimination of pathogens by host immune responses is a major bottleneck of infectivity, limiting the breadth of host competence [9,11,12]. Complement is one of the first lines of host defenses in vertebrate animals and can be activated through three canonical routes: the classical, lectin, and alternative pathways [13]. The activation of complement on the pathogen

surface results in the formation of a protein complex called C3 convertase. C3 convertase is essential for complement function because it serves as a protease to cleave C3 protein to its active fragments. Further complement activation leads to the lysis of pathogens due to the formation of membrane attack complex pores (C5b-9) on pathogen surfaces. C3 convertase formed by alternative pathway activation, called C3bBb (composed of C3b and Bb proteins), activates more C3 molecules and results in a positive feedback of the C3 amplification loop for the complement activation. To avoid tissue damage from unwanted complement activation in the absence of pathogens, hosts possess complement inhibitors [13,14]. One of these inhibitors is factor H (FH), which binds to C3b and blocks further activation of complement [13,15]. Like other pathogens, Lyme borreliae exploit such host self-regulatory mechanisms by recruiting complement inhibitors on their surface [10,16-18], which allows spirochetes to evade complement-mediated killing in the host bloodstream or tick blood meals [10,16–18]. In fact, spirochetes bind to FH through the production of several bacterial FH-binding proteins, called Complement Regulators Acquiring Surface Proteins (CRASPs): CspA, CspZ, and OspErelated proteins [19,20]. Among these CRASPs, CspA is uniquely required for bacterial transmission from nymphal ticks to vertebrate animals by binding to FH, resulting in complement evasion in those feeding ticks [21-23].

Interestingly, the ability of CspA to bind to FH from different vertebrate hosts varies by alleles and dictates the specificity of a Lyme borreliae species to survive in the sera of various animals [21,24,25]. Specifically, the ability of CspA variants to bind to mammalian FH and survive in homologous sera correlates with the ability of these variants to promote tick-to-mouse transmission [21]. These findings raise a possibility that CspA-mediated, FH binding-dependent complement evasion drives Lyme borreliae host tropism [9,10]. However, that possibility could not be fully demonstrated until we could elucidate the roles of CspA variants that do not promote tick-to-mouse transmission to confer tickborne transmission to other hosts. Additionally, if CspA is one of the spirochete determinants of host tropism, what evolutionary mechanisms give rise to the host-spirochete associations mediated by this protein?

In this study, we used Lyme disease spirochetes, avian and mammalian hosts, and the blood of mammals and birds as models to examine the role of complement in driving host tropism of pathogens. We further identified CspA as a molecular determinant for such tropism and elucidated the evolutionary mechanisms resulting in the allelically specific roles of this protein in conferring maintenance of pathogens in diverse hosts during the infection cycle.

Results

Lyme borreliae genospecies differ in their levels of transmission to wildtype but not complement-deficient mice and quail

To examine tick-to-host transmission among Lyme borreliae species, we intradermally injected wild-type BALB/c (wild-type; WT) mice with *B. burgdorferi* B31-5A4, *B. garinii* ZQ1, or *B. afzelii* CB43. The tissues from B31-5A4- or CB43-infected mice had significantly greater spirochete burdens than those from uninfected mice (S1A–S1D Fig). In contrast, except for the bladders from two mice, bacterial burdens in the tissues of ZQ1-infected mice were below detection limits (10 bacteria per 100ng total DNA) (S1A–S1D Fig). To generate ticks harboring equal loads of spirochetes, we intradermally injected each of these strains into C3-deficient BALB/c mice (C3^{-/-}) mice, which do not have functional complement. After allowing *I. scapularis* larvae to feed on these mice, we found similar burdens of these strains in all tested tissues, fed larvae, and post molting flat nymphs (S1E–S1J Fig).

We then permitted the nymphs carrying each of these strains to feed on WT mice and determined the spirochete burdens in the replete nymphs, the skin at the tick bite site, and

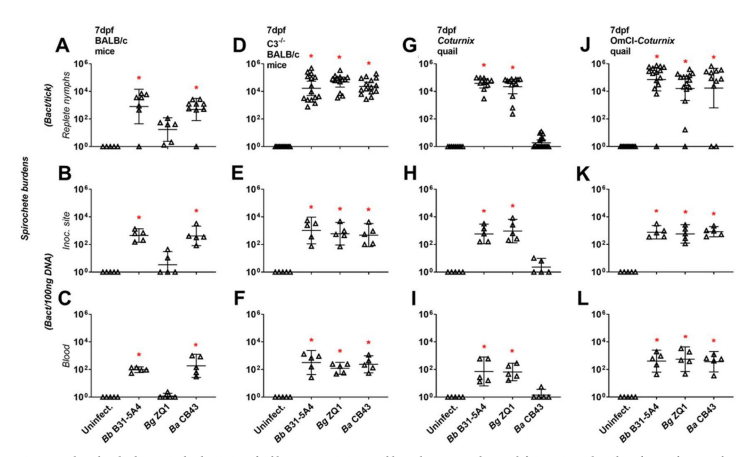


Fig 1. Lyme borreliae display species-level variation of tickborne transmission to wild-type but not complement-deficient mice and quail. *Ixodes scapularis* nymphs infected with *B. burgdorferi* B31-5A4 (*"Bb* B31-5A4"), *B. garinii* ZQ1 (*"Bg* ZQ1"), or *B. afzelii* CB43 (*"Ba* CB43") fed on (A-C) BALB/c mice, (D-F) C3^{-/-} BALB/c mice, (G-I) *Coturnix* quail, or (J-L) OmCI-treated *Coturnix* quail. Uninfected nymphs and mouse and quail tissues were included as control ("Uninfect."). Fed nymphs were collected upon repletion, and blood and the tick bite sites of skin were collected at 7 days post nymph feeding ("dpf"). Spirochete burdens in (A, D, G, and J) replete nymphs, (B, E, H, and K) tick bite site of skin ("Inoc. site"), and (C, F, I, and L) blood were determined by qPCR. For the burdens in tissue and blood samples, the resulting values were normalized to 100 ng total DNA. Shown are the geometric means of bacterial loads ± 95% confidence interval of bacterial burdens in tissues and blood from 5 mice or quail per group or nymphs feeding on mice (7 nymphs carrying *Bb* B31-5A4, 6 nymphs carrying *Bb* B31-5A4, 0r *Bg* ZQ1, or 17 nymphs carrying *Ba* CB43), or 0 mCI-treated quail (15 nymphs carrying *Bb* B31-5A4, 11 nymphs carrying *Bg* ZQ1, or 15 nymphs carrying *Ba* CB43). Significant differences (p < 0.05, Kruskal-Wallis test with the two-stage step-up method of Benjamini, Krieger, and Yekutieli) in the spirochete burdens relative to uninfected ticks or tissues are indicated with an asterisk.

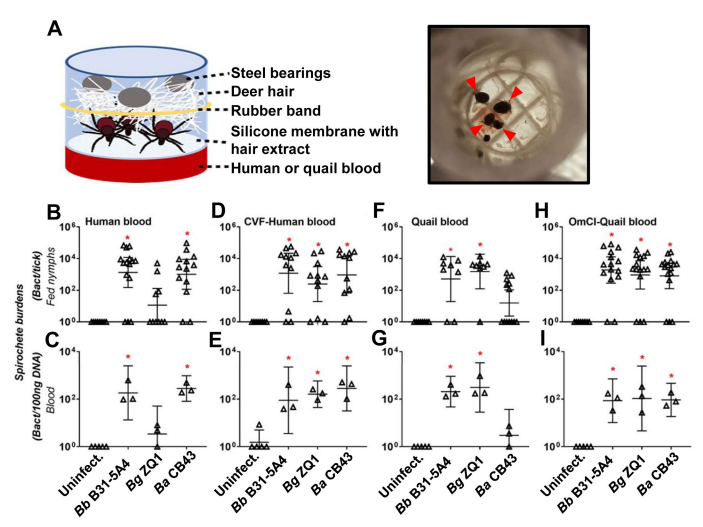
blood from these animals at 7 days post feeding (dpf), and uninfected nymphs were included as control. Strains B31-5A4 and CB43 survived at these sites ($\sim 10^3$ spirochetes per tick or 10^2 to 10^3 spirochetes per 100ng DNA of tissues, Fig 1A–1C). Two out of six fed nymphs had undetectable loads of ZQ1 whereas the other four ticks have bacterial loads ranging from 39 to 111 spirochetes per tick (Fig 1A). These low, variable values were not significantly different compared to uninfected nymphs (Fig 1A, p = 0.52). Strain ZQ1 was also largely undetectable in tick bite sites and blood (three and five out of five bite sites and blood samples, respectively; Fig 1B and 1C). Additionally, we fed nymphs carrying B31-5A4, CB43, or ZQ1 on C3^{-/-} mice and found that these strains were detected in fed nymphs, bite sites, and blood at similar levels (Fig 1D–1F). These results indicate that ZQ1 is less capable of surviving in fed nymphs and establishing infection than B31-5A4 and CB43 after ticks fed only on WT mice, but not on the mice lacking C3, suggesting that mouse complement dictates spirochete transmission. We further studied the tickborne transmission of these strains in a similar fashion using *Coturnix* quail, the avian model of Lyme disease [26,27]. We detected B31-5A4 and ZQ1 in fed nymphs, tick bite sites, and bloodstream (~10⁴ spirochetes per tick or 10^2 to 10^3 Lyme borreliae per 100ng DNA of tissues, Fig 1G–1I). Conversely, strain CB43 was not detected above the detection limit in these ticks and tissue samples (Fig 1G–1I). The nymphs carrying each of these strains were then allowed to feed on quail treated with *O. moubata* complement inhibitor (OmCI), which is the only tick salivary protein reported to block quail and other avian complement at the level of C5 activation (S2 Fig)[28]. We found similar levels of spirochetes in fed nymphs, tick bite sites and blood (Fig 1J–1L). These data showed less efficient tick-to-quail transmission of CB43 than that of B31-5A4 and ZQ1, and as was the case in mice, quail complement dictates transmission efficiency among those spirochete strains.

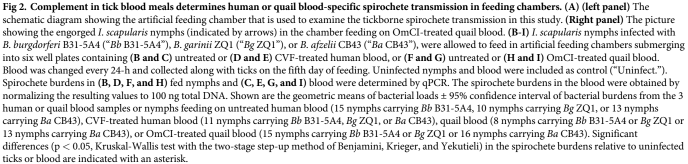
The different abilities of Lyme borreliae genospecies to evade complement in tick blood meals determine mammalian- or avian-specific spirochete transmission

To examine the role that the source of the blood meals plays in determining spirochete transmission, nymphs infected with B31-5A4, ZQ1, or CB43 were allowed to feed on artificial feeding chambers with human blood, the mammalian blood representative [29](Fig 2A). Uninfected nymphs were included as controls. We did not detect statistical difference of the number of attached ticks feeding on quail blood, compared to those on human blood, consistent with our previous study [29]. We found that B31-5A4 and CB43 survive in fed nymphs and blood ($\sim 10^3$ spirochetes per tick and 10^2 spirochetes per 100ng DNA of blood, Fig 2B and 2C). In contrast, ZQ1 was undetectable in the majority of human blood-fed nymphs (6 out of 10 ticks) (Fig 2B). Similarly, none of the blood samples fed on by ZQ1-infected nymphs had spirochete burdens significantly greater than uninfected blood (Fig 2C). We also allowed nymphs carrying B31-5A4, ZQ1, or CB43 to feed on human blood treated with Cobra Venom Factor (CVF), which depletes human complement cascade from the level of C3 [30]. We found similar burdens of these strains in the fed nymphs and the blood samples ($\sim 10^3$ spirochetes per tick and $\sim 10^2$ spirochetes per 100ng DNA of blood, Fig 2D and 2E). These results indicate that ZQ1 is less competent than B31-5A4 or CB43 to survive in the tick blood meals from humans during transmission, and active human complement in the blood meals drives survival differences. We also performed similar work using quail blood and detected the strains B31-5A4 and ZQ1 in fed nymphs and blood ($\sim 10^3$ spirochetes per tick and $\sim 10^2$ spirochetes per 100ng total DNA of blood, Fig 2F and 2G). However, the strain CB43 was not detected in the majority of fed nymphs (7 out of 13 nymphs, Fig 2F; P > 0.05 compared to uninfected nymphs). Further, spirochete burden values in blood samples fed on by nymphs carrying CB43 were indistinguishable from those of uninfected blood samples (Fig 2G). Similar levels of each strain were observed when nymphs carrying each of these strains were fed on OmCI-treated quail blood ($\sim 10^3$ spirochetes per tick and 10^2 spirochetes per 100ng DNA of blood, Fig 2H and 2I). These findings indicate that quail complement limits CB43 survival when ticks feed on quail blood.

CspA-mediated quail FH-binding activity promotes tick-to-quail transmission of spirochetes by complement evasion

We previously showed that a *cspA*-deficient mutant *B. burgdorferi* producing a spirochete outer surface protein, CspA, from the *B. burgdorferi* B31 (CspA_{B31}) or *B. afzelii* PKo (CspA_{PKo}) but not from *B. garinii* ZQ1 (CspA_{ZQ1}), facilitates tick-to-mouse transmission by surviving in fed nymphs [21]. This isogenic strain-specific transmission is dependent on the mouse FH-binding activity of these CspA variants to evade complement [21] (S1 Table) and recapitulates





the mouse-specific tickborne transmissibility of *B. burgdorferi* B31, *B. afzelii* CB43, and *B. garinii* ZQ1 (Fig 1). Note that CspA from *B. afzelii* CB43 (CspA_{CB43}) shares 99% amino acid identity with CspA_{PKo}, making these CspA variants likely confer similar FH-binding and transmission phenotypes (S3A Fig). To extend that isogenic strain-specific phenotype to other small mammals, we allowed nymphs carrying the *cspA*-deficient *B. burgdorferi* producing CspA_{B31}, CspA_{PKo}, or CspA_{ZQ1} to feed on a rodent reservoir of Lyme borreliae, *Peromyscus leucopus*. We found that expression of CspA_{B31} and CspA_{PKo}, but not CspA_{ZQ1}, permitted spirochete transmission to that rodent species (S4 Fig). Further, our previous findings of CspA_{ZQ1} (and $CspA_{B31}$) binding to quail FH and promoting survival in quail serum raised the hypothesis that CspA variants that bound FH drives tick-to-quail transmission [21] (S1 Table). To test this hypothesis, quail were fed on by the nymphs carrying WT B. burgdorferi B31-5A15 (5A15), B. burgdorferi B31-5A4NP1ΔcspA harboring an empty vector (ΔcspA/Vector), or the *cspA*-deficient strain carrying plasmids to express CspA_{B31}, CspA_{ZQ1}, or CspA_{PK0} in the background of B31-5A4NP1 $\Delta cspA$. We also included an isogenic strain producing CspA_{B31}-L246D. This mutant protein is derived from CspA_{B31} and selectively devoid of quail FH-binding activity but retaining other previously reported ligand-binding activities of CspA and thus can be used as negative control as a FH-binding deficient variant [21,31] (S1 Table). Uninfected ticks were included as control. The strain 5A15 but not $\Delta cspA/Vector \text{ or } cspA_{B31}$ -L246D-complemented strain, had burdens above detection limits in the fed nymphs, tick bite sites and blood (Fig 3A-3C). Strains producing CspA_{B31} or CspA_{ZQ1} but not CspA_{PKo}, had detectable burdens in the fed nymphs or tick bite sites and blood ($\sim 10^4$ spirochetes per tick and $\sim 10^2$ spirochetes per 100ng DNA of tissues or blood, Fig 3A-3C). In contrast, when nymphs carrying each of these spirochete strains were permitted to feed on OmCI-treated quail, all strains showed comparable burdens in fed nymphs, tick bite sites, and blood (Fig 3D-3F). These findings suggest that the CspA_{ZO1} and CspA_{B31} as quail FH binders promote tick-to-quail transmission by evading complement, and CspA-mediated quail FH-binding activity dictates such a transmission.

Allelically variable, CspA-mediated FH-binding activity confers spirochete complement evasion in tick blood meals and transmissibility in a mammalian and avian blood-specific manner

We sought to examine whether CspA-mediated FH-binding activity facilitates spirochete evasion of complement in tick blood meals, and if that ability determines tickborne transmission in a host-specific manner. Human blood was allowed to be ingested by the nymphs carrying 5A15, *\(\Delta\)*cspA/Vector, or this strain producing CspA_{B31}, CspA_{ZQ1}, CspA_{PK0}, or CspA_{B31}-L246D using feeding chambers. We detected 5A15, but not $\Delta cspA/Vector$ or CspA_{B31}-L246D-producing strains, in fed nymphs and blood (Fig 4A and 4B). The CspA_{B31}- or CspA_{PKo}-producing strains were found in fed nymphs and in human blood ($\sim 10^4$ spirochetes per tick (Fig 4A) and more than 10 spirochetes per 100ng total DNA of blood (Fig 4B)). Conversely, the CspA_{ZQ1}producing strain was not detectable in these samples (Fig 4A and 4B). When we allowed nymphs carrying the same strains to feed on CVF-treated human blood, all strains were detected at similar burdens in fed nymphs and blood (Fig 4C and 4D). These results suggest that CspA_{B31} and CspA_{PKo}, but not CspA_{ZO1}, permitted transmission to human blood by facilitating human FH-binding mediated complement evasion in tick blood meals. We also permitted the nymphs carrying the above-mentioned strains to feed on quail blood in the same fashion and found B31-5A15, but not $\Delta cspA/Vector \text{ or } CspA_{B31}-L246D$, in the fed nymphs and blood had detectable burdens in both nymphs and blood (Fig 4E and 4F). The CspA_{B31}or CspA_{ZO1}-producing strain was readily detected in fed nymphs and blood (Fig 4E and 4F). Though three and two ticks carrying the CspA_{PKo} and CspA_{B31}-L246D-producing strain, respectively, had burdens greater than detection limits, we did not detect any spirochetes in the remaining 10 nymphs (Fig 4E). Additionally, the burdens of these strains in the blood were statistically indistinguishable from uninfected blood samples (Fig 4F). When we performed similar experiments using OmCI-treated quail blood, all strains were found in nymphs and blood at comparable levels (Fig 4G and 4H). These results show the contribution of quail FHbinding dependent complement evasion in tick blood meals in promoting transmission to quail blood, and CspA_{B31} and CspA_{ZQ1}, but not CspA_{PK0}, conferred these activities.

CspA homologs showed discontinuous sequence variation and genospeciesspecific polymorphisms

Given the finding that homologous CspA proteins from single strains of *B. burgdorferi*, *B. afzelii*, or *B. garinii* confer distinct host tropism, we examined CspA variation in publicly available

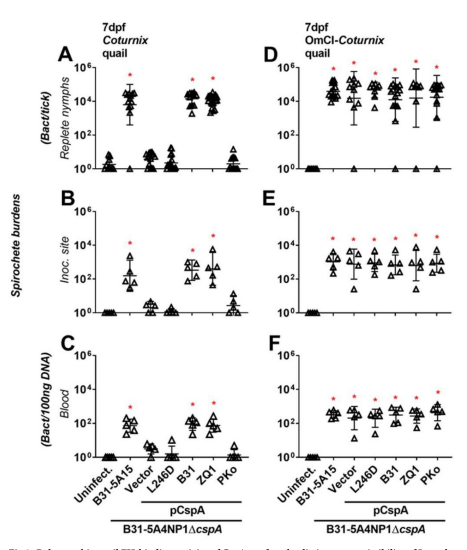


Fig 3. Polymorphic quail FH-binding activity of CspA confers the distinct transmissibility of Lyme borreliae to quail. Ixodes scapularis nymphs infected with WT B. burgdorferi B31-5A15 ("B31-5A15"), B. burgdorferi B31-5A4NP1/cspA transformed with an empty shuttle vector ("Vector"), or this deletion strain producing a mutated variant of CspA from B. burgdorferi B31 selectively devoid of quail FH binding activity ("L246D"), WT B. burgdorferi B31 ("B31"), B. garinii ZQ1 ("ZQ1"), or B. afzelii PKo ("PKo") were allowed to feed on (A-C) untreated or (D-F) OmCI-treated quail. Uninfected nymphs and quail tissues were included as control ("Uninfect."). Fed nymphs were collected upon repletion, and blood and tissues were collected at 7 days post nymph feeding ("dpf"). Spirochete burdens in (A and D) replete nymphs, (B and E) tick bite sites of skin ("Inoc. site"), and (C and F) blood were determined by qPCR. For the burdens in tissue samples, the resulting values were normalized to 100ng total DNA. Shown are the geometric means of bacterial loads ± 95% confidence interval of bacterial burdens from 5 quail tissues and blood or nymphs feeding on untreated quail (13 nymphs carrying the strains B31-5A15 or pCspA-PKo, 15 nymphs carrying the strains "Vector", 12 nymphs carrying the strain pCspA-B31, 21 nymphs carrying the strain pCspA-ZQ1, or 17 nymphs carrying the strain pCspA-L246D), or the nymphs from OmCI-treated quail (15 nymphs carrying the strains B31-5A15, pCspA-B31, or pCspA-PKo, 8 nymphs carrying the strains "Vector", 9 nymphs carrying the strain pCspA-L246D, 8 nymphs carrying the strain pCspA-ZQ1). Significant differences (p < 0.05, Kruskal-Wallis test with the two-stage step-up method of Benjamini, Krieger, and Yekutieli) in the spirochete burdens relative to uninfected ticks or quail tissues are indicated with an asterisk.

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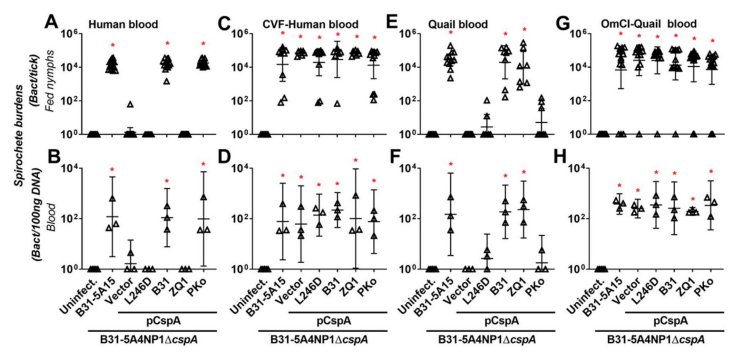


Fig 4. Allelically variable FH-binding activity of CspA dictates human- and quail blood-specific transmission in feeding chambers by evading complement in tick blood meals. *I. scapularis* nymphs infected with WT *B. burgdorferi* B31-5A15 ("B31-5A15"), *B. burgdorferi* B31-5A4NP1*AcspA* transformed with an empty shuttle vector ("Vector"), or this deletion strain transformed to produce a mutated variant of CspA from *B. burgdorferi* B31 selectively devoid of FH binding activity ("L246D"), WT *B. burgdorferi* B31 ("B31"), *B. garinii* ZQ1 ("ZQ1"), or *B. afzelii* PKo ("PKo") were allowed to feed in feeding chambers submerged into 6-well plates containing (**A and B**) untreated or (**C and D**) CVF-treated human blood, or (**E and F**) untreated or (**G and H**) OmCI-treated quail blood. Blood was changed every 24-h and was collected along with ticks on the fifth day of feeding. Uninfected nymphs and blood were included as control ("Uninfect."). Spirochete burdens from (**A, C, E and G**) fed nymphs and (**B, D, F, and H**) blood were determined by qPCR. For the burdens in tissue samples, the resulting values were normalized to 100ng total DNA. Shown are the geometric means of bacterial loads and 95% confidence interval of bacterial burdens from 3 human and quail blood samples, or nymphs feeding on untreated human blood (15 nymphs carrying the strains B31-5A15 or pCspA-B31, 14 nymphs carrying the strains "Vector" or pCspA-PKo, 17 nymphs carrying the strains "Vector", pCspA-ZQ1, or 11 nymphs carrying the strains pCspA-L246D or pCspA-PKo), untreated quail blood (11 nymphs carrying the strains B31-5A15, pCspA-PKo, or 7 nymphs carrying the strains B31-5A15, pCspA-PKo, or pCspA-PKo), or pCspA-B31, or pCspA-B31, or pCspA-PKo, or 7 pCspA-L246D) or OmCI-treated quail blood (15 nymphs carrying the strains pCspA-L246D or pCspA-PKo), untreated quail blood (11 nymphs carrying the strains B31-5A15 or "Vector", PCspA-PKO, or 7 nymphs carrying the strains B31-5A15 or "Vector", 8 nymphs carrying the strains pCspA-PKO, or 7 pCspA-PKO, or p

sequences. CspA is nested in the fourth clade of a protein family encoded on the linear plasmid 54, lp54 (PFam54-IV) (41–81% nucleotide identity; Fig 5A) [32,33]. However, the homology of all PFam54-IV proteins makes it difficult to identify CspA variants, leading to inaccurate annotations and misidentification [32]. We thus compiled publicly available gene sequences encoding PFam54-IV available in GenBank from *B. burgdorferi*, *B. afzelii*, and *B. garinii*, and compared the pairwise nucleotide identities of codon alignments for these genes from each species. Within PFam54-IV-encoding genes of any one particular *B. burgdorferi* strain, we identified one-to-one orthologous genes based on sequence conservation (>95% identity, green in S5 Fig). In contrast, we found moderate conservation among PFam54-IV homologs lacking such one-to-one orthology (<81% identity, red and yellow in S5 Fig). Sequence divergence patterns (inlets in S5–S7 Figs) allowed us to identify genes encoding CspA_{B31}, CspA_{PKo}, and CspA_{ZQ1} as CspA orthologs in *B. burgdorferi*, *B. afzelii*, and *B. garinii*, respectively. Among these CspA orthologs, intraspecific diversity (i.e. within genospecies) exceeded 93% identity, while interspecific diversity (i.e. between genospecies) ranged from 67 to 72% (Fig 5B and 5C). These results suggest a genospecies-specific polymorphism among CspA variants,

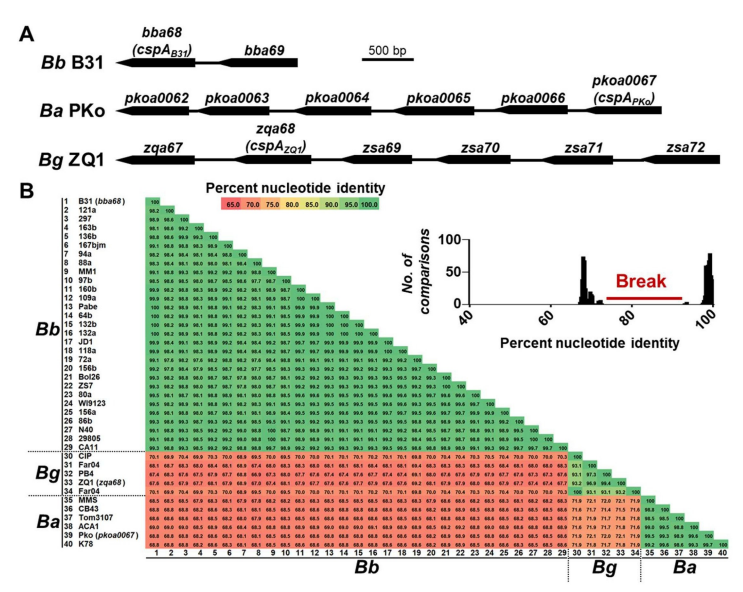


Fig 5. Genetic divergence in CspA variants across *B. burgdorferi* **B31**, *B. afzelii* **PKo**, and *B. garinii* **ZQ1**. (A) Synteny of Pfam54-IV loci in *B. burgdorferi* B31, *B. afzelii* PKo, and *B. garinii* ZQ1 (B) Pairwise nucleotide identity across the three spirochete species. (inset) Frequency distribution of pairwise genetic distances. The pairwise identity numbers are coded by color gradually from identical (100% pairwise identity; green) to divergent sequences (65% pairwise identity; red). The clear break in the frequency distribution separates the highly similar (> 95% pairwise identity) from moderately divergent comparisons (< 80%).

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whereby variants of the same genospecies share notably high identity, and variants of different genospecies share relatively lower identities.

Host-specific FH-binding activity of CspA variants arose through convergent evolution

An average identity of 74% among the genes encoding CspA and other Pfam54-IV proteins raises the possibility that non-CspA members of Pfam54-IV share FH-binding functions. We thus examined the mouse (*Mus musculus*) and quail FH-binding ability of Pfam54-IV from *B. burgdorferi* B31-5A4, *B. afzelii* MMS, and *B. garinii* ZQ1 using ELISA. Note that PFam54-IV members from *B. afzelii* MMS, PKo, and CB43 are nearly identical (>99% identity) (S3A and

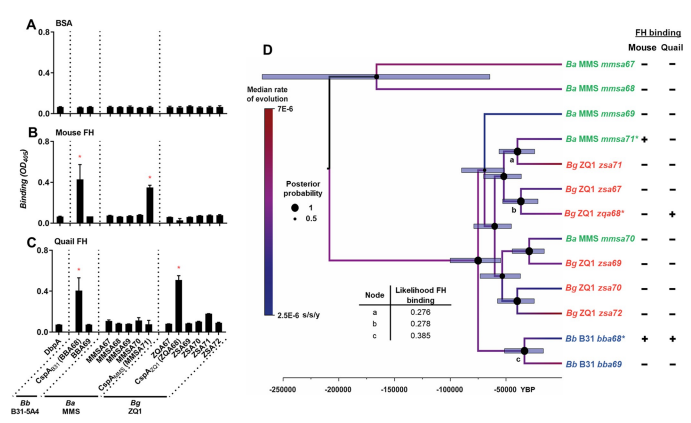
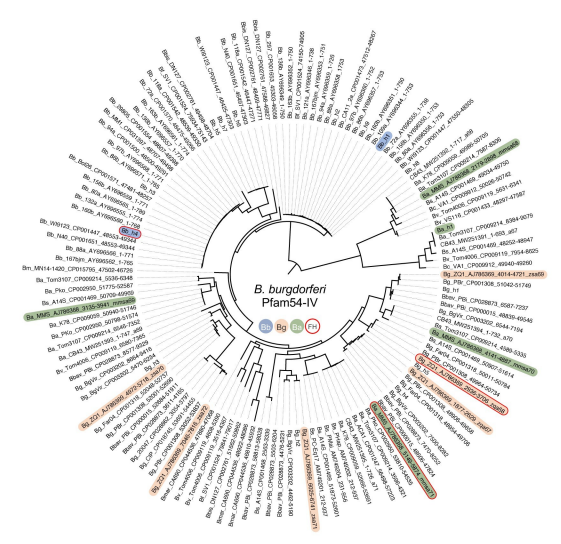
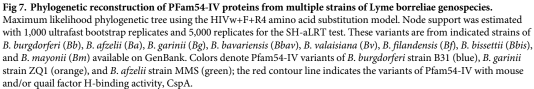


Fig 6. Allelically variable, host-specific FH-binding activity of CspA has emerged from convergent evolution. (A to C) Two μ M of histidine-tagged Pfam54-IV proteins from *B. burgdorferi* B31-5A4, *B. afzelii* PKo, or *B. garinii* ZQ1 or recombinant histidine-tagged DbpA from *B. burgdorferi* B31-5A4 (negative control) were added in triplicate to wells coated with of purified (A) BSA (negative control) or (B) mouse or (C) quail FH. Protein binding was measured by ELISA in three independent experiments. Each bar represents the geometric mean \pm 95% confidence interval of three replicates in one representative experiment. Significant differences (p < 0.05, Kruskal-Wallis test with the two-stage step-up method of Benjamini, Krieger, and Yekutieli) in the levels of FH binding of indicated proteins relative to DbpA are indicated (*). The ability of each protein in binding to factor H is summarized in S1 Table. (D) Maximum clade credibility tree is based on Pfam54-IV genes from *B. burgdorferi* B31 (blue), *B. afzelii* MMS (green), *B. garinii* ZQ1 (red). The scale bar represents an approximate timeline of evolution, in years before present ("YBP"), using the estimated substitution rate of 4.75×10⁻⁶ substitutions/site/year. Node bars represent the 95% highest posterior density of the node age. Node circles represent the posterior probability support. Branches are colored based on estimate FH-binding activities at ancestral nodes.

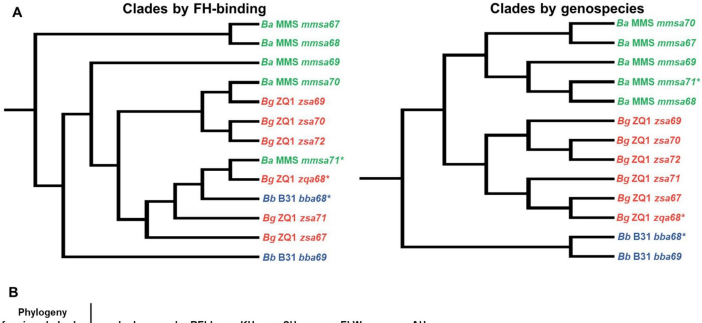
S6 Figs). We used Pfam54-IV of MMS to represent these proteins of *B. afzelii* given that the recombinant version of these proteins from MMS had been generated in our previous work [24]. As expected, PFam54-IV from *B. burgdorferi* B31-5A4, *B. afzelii* MMS, and *B. garinii* ZQ1 did not bind to BSA (Fig 6A). We found that CspA_{B31} and CspA_{MMS} bound to mouse FH at levels greater than a negative control spirochete protein, DbpA [21] (Fig 6B). Despite a high concentration (2 μ M) of other recombinant Pfam54-IV used, none bound to mouse FH over baseline levels seen with DbpA (Fig 6B). Furthermore, we observed that CspA_{B31} and CspA_{ZQ1}, but none of other tested Pfam54-IV, bound to quail FH (Fig 6C). These results indicate that the host-specific and allelically variable FH-binding activity of Pfam54-IV is CspA-dependent.

To further study the evolutionary mechanisms leading to host-specific and allelically variable FH-binding activity of CspA, we estimated the phylogenetic relationships among gene sequences encoding PFam54-IV from *B. burgdorferi* B31, *B. afzelii* MMS, and *B. garinii* ZQ1. We found that those sequences of the same genospecies do not form monophyletic assemblages, but CspA variants grouped in separate clades with moderate to high internode branch





support (Bayesian posterior probabilities (PP) of 0.81 at CspA_{B31} and CspA_{MMS} nodes and 0.86 at CspA_{MMS} and CspA_{ZQ1} nodes) (Fig 6D). Similar branching patterns were seen when phylogenetic relationships were estimated among homologs encoding PFam54-IV available on GenBank from *B. burgdorferi*, *B. afzelii*, and *B. garinii* and other Lyme borreliae genospecies with species forming paraphyletic assemblages (Fig 7). We then tested the plausibility of this evolutionary scenario by placing CspA variants in the same clades, grouped by FH-binding functions (Fig 8A, left) or by grouping PFam54-IV variants from the same genospecies together (Fig 8A, right). The sequence data did not support either alternative evolutionary scenarios based on phylogenetic tree topology testing (Fig 8B), in favor of the convergent evolution scenario (better tree log-likelihood value, P = 1.0). Further, the supported phylogeny raises the possibility that CspA-mediated FH-binding activities arose from 1) a common



forming clades by	logL	bp-RELL	p-KH	p-SH	c-ELW	p-AU
Convergent evolution	-7669.526515	1	1	1	1	1
FH-binding activity	-7814.374806	0	0	0	1.81×10 ⁻³⁶	4.95×10 ⁻⁶
Genospecies	-7761.994918	0	0	2×10 ⁻⁴	1.45×10 ⁻¹⁵	1.56×10 ⁻⁴

Fig 8. Competing evolutionary scenario evaluation of PFam54-IV gene divergence rejects alternative tree topologies of PFam54-IV proteins forming clades by FH-binding activity or spirochete genospecies. (A) Phylogenetic scenarios showing a single emergence of FH-binding or PFam54-IV genes forming clades by genospecies. (B) Log-likelihood value for each tree (logL), RELL bootstrap proportion (bp-RELL; 1.0 is full support for that tree), Kishino-Hasegawa test, Shimodaira-Hasegawa test, Expected Likelihood Weights (c-ELW; 1.0 is full support for that tree), and the Approximately Unbiased test (p-AU). Significant p-values indicate comparisons where the best-known ML tree explains the sequence data better than the alternative scenario.

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FH-binding ancestor or 2) the convergent evolution of PFam54-IV (Fig 6D). However, our results from maximum likelihood and parsimony-based ancestral state reconstruction methods, rejected the FH-binding ancestor scenario (Fig 6D), indicating that the allelically-variable, host-specific FH-binding activity of CspA is a result of convergent evolution within PFa-m54-IV. Based on a chromosome mutation rate estimated in a previous study [34], this evolutionary event likely occurred approximately 15,000–55,000 years before present, coinciding with the end of the last glacial maximum (Fig 6D).

Discussion

Frequent interactions of pathogens with one or a few hosts may drive them to adapt and become specialized in such host(s), thus maximizing fitness [4,35]. However, the fact that generalists are present in nature suggests that broad host ranges can also confer fitness advantages [4,35]. For vector-borne pathogens, the process leading to host tropism can be driven by host-derived components (i.e. immune molecules or nutrients) either in the hosts or acquired by vectors [4]. The molecular determinants and evolutionary mechanisms by which pathogens specialize or generalize to be associated with hosts are largely unclear. Reflected by the variable hosts carrying different spirochete genospecies transmitted through *Ixodes* ticks, the Lyme disease bacterium is a well-suited model to study host tropism [9,10]. One long-held concept that

have not been definitively tested is that evading host complement immune responses by spirochetes determines Lyme borreliae host tropism [36]. A hurdle for such an investigation is the inability to easily maintain non-mammalian hosts, such as birds, in laboratory and/or persistently infect them [37-47]. Though some wild-birds have been brought into laboratories to study spirochete infectivity [37-47], molecular mechanisms have not been elucidated because of the lack of avian-specific reagents. We and others have used *Coturnix* quail as avian Lyme disease model as this species was found to be fed on by *Ixodes* ticks [48,49] and can sustain detectable spirochete burdens for more than eight weeks [26,27]. In this study, we allowed ticks carrying spirochetes to feed on quail, similar to previous work performed in this species and other domestic aves [50-52]. We found that B. garinii ZQ1 and B. burgdorferi B31-5A4 survive in fed ticks and are transmitted to quail whereas B. afzelii CB43 did not. These results demonstrate that the genospecies variation of spirochete transmissibility to birds, in agreement with prior studies (reviewed in [9]), supporting the use of quail as an avian host representative. In contrast, when nymph feeding was performed on wild-type mice, B. afzelii CB43 and B. burgdorferi B31-5A4 survived in fed ticks and migrated to these animals while B. garinii ZQ1 did not. All three species survived in fed ticks and were transmitted to complement-deficient quail or mice. In support of previous *in vitro* evidence [36,53], this study establishes complement evasion by spirochetes as a driver of Lyme borreliae host tropism. As complement is enriched in vertebrate blood, our findings raise a possibility that spirochetes must evade host complement specifically in tick blood meals.

Tick feeding on live animals used in our study may introduce confounding factors of blood meal-independent, complement-mediated clearance [54]. To address this potential confounder, we use different sources of blood in "artificial feeding chambers" without the involvement of animals. This chamber assay allows us to demonstrate that spirochete evasion to complement in tick blood meals dictates Lyme borreliae host tropism [29]. Nonetheless, it should be noted that heparin as an anti-coagulant required to be used in the blood derived from this feeding chamber assay was reported to impact the complement activation [55–57]. Thus, such a caveat emphasizes the importance of considering the combined evidence from both feeding chamber assays and tick infection animal models.

A potential limitation of the experimental design is that we used *I. scapularis* ticks for all experiments. B. afzelii and B. garinii are not endemic to North America where I. scapularis are commonly found, and have been only isolated from other Ixodes ticks (i.e. I. ricinus and I. persulcatus) in the field. B. burgdorferi is the only Lyme borreliae genospecies in this study that is circulated in I. scapularis in nature [5]. Specifically, vector competence to a particular pathogen is determined by not only pathogen persistence in molting ticks and the following tick-to-host transmission but also the levels of initial acquisition for a pathogen acquired by uninfected ticks, which was not measured in this study. Thus, utilizing I. scapularis as a vector representative may not completely address the role of vector competence in impacting host tropism of spirochetes [58–59]. However, using a single tick species carrying each of the tested spirochete species allows for attribution of the observations solely to host and/or pathogen determinants, the emphasis of this study. Additionally, I. scapularis ticks were shown in laboratory infections to acquire and transmit B. burgdorferi, B. afzelii, and B. garinii at similar levels [60-61], suggesting the feasibility to use this tick in representing the Ixodes vectors of Lyme disease. Additionally, Lyme borreliae-infected mice have been frequently used to generate nymphs harboring spirochetes through blood feeding by naïve larvae [62]. However, rearing nymphs carrying similar burdens of each spirochete species in this fashion is difficult because wild-type mice do not maintain equal loads of these spirochetes [63–64]. Using complement-deficient mice (C3^{-/-} mice), we found similar burdens of B. burgdorferi B31-5A4, B. afzelii CB43, and B. garinii ZQ1 in fed larvae, post-molting flat nymphs, and the tissues derived from spirochete-infected mice.

Furthermore, it should be noted that C3 not only plays a role as a complement component to confer complement activation but also cross-talks with adaptive immune responses [65], raising the possibility of the adaptive immune response as a confounding factor to influence transmissibility. However, the infection stage tested in this work is very early (7 days post tick feeding), suggesting that this possibility is unlikely. Taken together, our results provide a strategy to overcome the difficulty in tick-rearing and infection, and support the concept that complement controls the spirochete infectivity during infection [21,63,66].

Both *Ixodes* ticks and Lyme borreliae produce complement-inactivating proteins to facilitate feeding and pathogen transmission [8,67-69]. Documented by the fact that many Lyme borreliae proteins are polymorphic, one attractive hypothesis is that polymorphisms in these proteins contribute to host tropism. Regrettably, using wild-type spirochete strains may not delineate the contribution of each of these proteins because these spirochetes have been known to generate multiple polymorphic complement-inactivating proteins [22,70–73]. Thus, identical spirochete background strains have been used to express genes or alleles (also known as "isogenic strains") to define bacterial determinants of particular phenotypes [21,74,75]. We have previously shown that isogenic strains in a *cspA*-deficient background producing CspA from B. burgdorferi B31 or B. afzelii PKo, but not B. garinii ZQ1, promote tick-to-mouse transmission. The ability of ticks to transmit Lyme borreliae is contingent on the ability of the spirochete to survive in fed ticks, consistent with the phenotypes observed using wild-type genospecies [21]. Such differential transmissibility depends on the ability of CspA to bind to mammal FH in the presence of mouse complement, leading to the possibility that allelicallyvariable, CspA-mediated FH-binding activity dictates host tropism [21,24,25]. Nonetheless, that concept requires identification of the animals that are susceptible to Lyme borreliae that express ZQ1-derived CspA. In agreement with our previous work showing CspA from ZQ1 and B31 but not PKo binds to quail FH [21], we found the ZQ1- and B31- (but not PKo-) derived CspA facilitates tick-to-quail transmission of spirochetes. We further showed that such an allele-dependent, quail-specific transmission is determined by the presence of quail complement and the ability of CspA to bind to this species' FH. Our results using feeding chambers, isogenic bacterial strains and complement-intact or -deficient human or quail blood indicated that CspA is a determinant of Lyme borreliae host tropism by promoting host-specific, FH-binding-dependent complement evasion in tick blood meals. Note that this study does not rule out the possibility of other anti-complement proteins in promoting hostspecific infectivity as *cspA* is downregulated after spirochete transmission to vertebrate animals [21,23]. Instead, these results demonstrated the role of CspA to facilitate spirochete in overcoming the initial barrier of tick-to-host transmission in determining host tropism.

CspA, like other Pfam54-IV proteins, experienced numerous duplication and loss events, resulting in moderate sequence identity (~40 to 80%) among variants in this protein family [32,33,76]. This is suggestive of rapid evolution, and potential neofunctionalization [32,33,77], supported by some important functions (e.g. complement evasion, cell adhesion, plasminogen binding, and tissue colonization) identified for CspA and other Pfam54 proteins [21,78–81]. However, the moderate sequence similarity from a limited number of strains in those studies renders the establishment of 1:1 orthology relationships difficult across Pfam54-IV [32,33], thus impacting the examination of evolutionary mechanisms giving rise to those functions. By comparing the sequences that encode PFam54-IV from different strains within each genospecies, we found a wide range of sequence diversity. These results allow us to define these highly similar sequences as paralogs. These findings also suggest that the phenotypes conferred by one Pfam54-IV protein (e.g. host-specific FH-binding activity/transmissibility) are likely shared by its orthologs from different strains within the same genospecies [82]. Furthermore, we observed notably divergence (pairwise identity <79%) between the genes encoding

Pfam54-IV among different spirochete genospecies, indicating that at least some genospeciesspecific polymorphisms have reached fixation. These results, combined with the fact that CspA variants confer variable complement evasion and host tropism [21,24,25], are in line with the findings of allelically-variable phenotypes in other Lyme borreliae polymorphic proteins [72,74,75]. Furthermore, these data suggest that any unidentified functions of CspA or other Pfam54-IV may vary among the strains from different genospecies.

Some phenotypes are shared among multiple Pfam54-IV proteins from the same Lyme borreliae genospecies (e.g. C7 and C9 binding-mediated complement evasion by Bga66 and Bga71 from B. bavariensis) [78]. This finding raises the possibility that FH-binding is a common feature for non-CspA Pfam54-IV proteins within the same genospecies. However, our finding showing no detectable mouse or quail FH-binding to non-CspA Pfam54-IV indicates that the FH-binding activity is unique to CspA, in agreement with the results of human FH-binding activity of Pfam54-IV [24]. Further, our data from phylogenetic reconstructions rejected the possibility of functional cladding of CspA or a common FH-binding ancestor, but supported convergent evolution as the mechanism leading to such an allelically variable, CspA-mediated FH-binding activity [83]. With the fact that such an activity dictates Lyme borreliae host tropism, allelically variable CspA-mediated FH-binding activity would effectively isolate different populations of Lyme borreliae in their respective hosts in nature, resulting in adaptive radiation of spirochetes. An intriguing question is whether the emergence of such host-specific FH binding is concurrent with allopatric ecological speciation of the Lyme borreliae whereby host switching events isolated groups of spirochetes. This would lead the spirochetes to evolve the ability, such as the host-specific FH binding activity of CspA, in response to disparate host environments and leading to adaptations [84-90]. However, the phylogenetic reconstruction of Pfam54-IV suggests that CspA of B. burgdorferi evolved approximately 15,000 to 55,000 years before present (BP), while the earliest common ancestor of B. burgdorferi in North America was dated at ~60,000 years BP [34]. Thus, the emergence of the CspA variants likely occurred after ancient Lyme borreliae speciation. Our timeline of CspA emergence coincides with the latter half of the last glacial maximum in North America and Europe, which was a period of major climactic change [91–92]. A plausible explanation is that this drastically changing environment resulted in novel selective pressures, such as changes to host body condition and immune response, or host and vector availability and ranges [93–95]. In response to such changes, the spirochetes may have evolved new strategies to be maintained in the enzootic cycle, such as allelically variable, CspA-mediated FH-binding activity. Using the Lyme disease bacterium as a model, this work is a pioneering study defining the mechanisms that dictate host tropism of pathogens, identifying the molecular determinants, and elucidating the evolutionary drivers of such host-pathogen associations. These findings will provide significant impacts into the origin of a vector-borne enzootic cycle and establish the groundwork for future studies to investigate the mechanisms in shaping host tropism.

Materials and methods

Ethics statement

All mouse and quail experiments were performed in strict accordance with all provisions of the Animal Welfare Act, the Guide for the Care and Use of Laboratory Animals, and the PHS Policy on Humane Care and Use of Laboratory Animals. The protocol was approved by the Institutional Animal Care and Use Committee (IACUC) of Wadsworth Center, New York State Department of Health (Protocol docket number 19–451) and Columbia University (Protocol docket number AC-AAAO4551). All efforts were made to minimize animal suffering.

Mouse, quail, tick, bacterial strains, OmCI, and FH

BALB/c mice were purchased from Taconic (Hudson, NY). C3^{-/-} mice in BALB/c background were generated from the C3^{-/-}(C57BL/6) purchased from Jackson Laboratory (Bar Harbor, ME) as described in our previous study [21]. *P. leucopus* mice were ordered from *Peromyscus* genetic stock center at University of South Carolina (Columbia, SC). *Coturnix* quail was purchased from Cavendish Game Birds Farm (Springfield, VT). *Ixodes scapularis* tick larvae were purchased from National Tick Research and Education Center, Oklahoma State University (Stillwater, OK) or obtained from BEI Resources (Manassas, VA). Lyme borreliae-infected nymphs were generated as described in the section "Generation of ticks carrying Lyme borreliae." The *Borrelia* and *Escherichia coli* strains used in this study are described in S2 Table. *E. coli* strains DH5α, M15, and derivatives were grown in Luria-Bertani (BD Bioscience) broth or agar, supplemented with kanamycin (50µg/ml), ampicillin (100µg/ml), or no antibiotics as appropriate. All *B. burgdorferi*, *B. afzelii*, and *B. garinii* strains were grown in BSK-II completed medium supplemented with kanamycin (200µg/mL), streptomycin (50µg/mL), gentamicin (50µg/mL), or no antibiotics (see S2 Table). Mouse FH was purchased from MyBiosource. Quail FH and recombinant OmCI proteins were generated as described previously [21,26,28].

Mouse infection using needle inoculation

Four-week-old female BALB/c or C3^{-/-} mice in BALB/c background were used for experiments involved in needle infection of Lyme borreliae strains. Mice were infected by intradermal injection as previously described [75] with 10⁶ of *B. afzelii* CB43, *B. garinii* ZQ1, or *B. burgdor-feri* B31-5A4. The plasmid profile of the strain B31-5A4 was verified prior to infection as described to ensure the stability of the vector and no loss of plasmids [96–97]. As the information of genome is not available for plasmid profiling, the strains CB43 and ZQ1 used in this study were less than ten passages. Mice were sacrificed at 21 days post-infection, the inoculation site of the skin, the ankle joints, ears, and bladder were collected to quantitatively evaluate levels of colonization during infection as described in the section "Determination of Lyme borreliae burdens in infected ticks, tissues and blood samples.".

Generation of ticks carrying Lyme borreliae

The procedure of the tick infection has been described previously [21,98]. Basically, fourweek-old male and female C3^{-/-} mice in BALB/c background were infected with 10⁶ of *B. afzelii* CB43, *B. garinii* ZQ1, or *B. burgdorferi* B31-5A4, B31-5A15, B31-5A4NP1 $\Delta cspA$ harboring a vector or this *cspA* mutant strain producing CspA_{B31}, CspA_{PKo}, CspA_{ZQ1}, or CspA_{B31}L246D by intradermal injection as described above. The ear punches from those mice were collected at 13 days post infection, and DNA was extracted to perform qPCR using *Borrelia* 16S rRNA primers as previously described [26] (S3 Table) to confirm the infection (See section "Determination of Lyme borreliae burdens in infected ticks, tissues and blood samples."). At 14 days post infection, the uninfected larvae were allowed to feed to repletion on those spirocheteinfected mice as described previously [21,98]. Approximately 100 to 200 larvae were allowed to feed on each mouse. The engorged larvae were collected and allowed to molt into nymphs in a desiccator at room temperature and 95% relative humidity in a room with light dark control (light to dark, 16:8 h). [21,96].

Serum resistance assays

Coturnix quail were subcutaneously injected with OmCI (1 mg/kg of quail) or PBS buffer, and the sera were collected at 6, 24, 48, 72, and 96 h post injection. A serum sensitive, high passaged

B. burgdorferi strain B313 was cultivated to mid-log phase, followed by being diluted to a final concentration of 5×10^6 bacteria/ml in BSKII medium without rabbit serum. These bacteria were then incubated with each of these quail serum samples (final concentration: 40% of serum). We also included the bacteria mixed with heat inactivated serum samples, which have been incubated at 56°C for 2h prior to being mixed with spirochetes. An aliquot was taken from each reaction at 0 and 4h post injection to determine the number of motile bacteria under a Nikon Eclipse E600 darkfield microscope, as previously described [21]. The survival percentage for those motile spirochetes was calculated using the number of mobile spirochetes at 4h post incubation normalized to that at the very beginning of incubation with serum.

Mouse, quail, and P. leucopus infection by ticks

The flat nymphs were placed in a chamber on four- to six-week old male and female BALB/c or C3^{-/-} mice in BALB/c background, and the engorged nymphs were collected from the chambers at seven days post nymph feeding as described [99]. For ticks feeding on quail, the feathers located on the back of quail's neck were plucked to expose approximately $2-3 \text{ cm}^2$ of skin, close to the back of its head. 1.2ml screw-top cryo-microcentrifuge vials (ThermoFisher Scientific) were cut to be used as mini chambers. The top of the caps from the chambers was pierced with a 25-gauge needle to create air holes, and sand papers were used to smooth any sharp edges along the cut surface edge of these chambers. Vetbond Tissue Adhesive (3M) was used to attach the chambers onto the exposed quail skin followed by manually restraining quail while the surgical glue dries (1-2min). Ten nymphs were placed into the chambers on mice or quail, which those ticks to feed on. For OmCI-treated quail, the quail were subcutaneously injected with OmCI (1mg/kg of quail) a day prior to the nymph feeding. The engorged nymphs were obtained from the chambers. The mice and quail were placed into a small cage, which then placed above the moat in a larger cage (for mice) or plastic bin (for quail). Ticks feeding on *Peromyscus leucopus* mice have been described previously [100–101]. Ten nymphs were placed in the ears of each mouse, five nymphs per ear and were allowed to feed until repletion. P. leucopus mice were separately placed in water cages which consisted of the cage being filled with approximately 2.5cm of water along the bottom, a wire rack to keep the mouse out of the water, and then being placed in a larger hamster cage with water to prevent ticks from escaping. The engorged nymphs were recovered from the water cage beginning five days post nymph feeding. Blood and tick placement site from quail, mice, and P. leucopus mice were collected at seven days post nymph feeding.

Feeding chamber assays by ticks

Artificial feeding chambers were prepared as described in our previous study [21]. In short, the silicone rubber-saturated rayon membrane was generated as described [21], with the exception of adhering fiberglass mesh (3-mm pore; Lowe's Inc.) to the membrane before attaching it to the rest of the chamber. Such membrane was attached to one side of a 2-cm length of polycarbonate tubing (hereafter called the chamber; inner diameter: 2.5 cm; outer diameter: 3.2 cm; (Amazon Inc.), as described, with the exception of using a rubber band to hold the chamber in place instead of a rubber O-ring [21]. Feeding stimuli including hair and hair extract from white-tailed deer (*Odocoileus virginianus*) and a plastic tile spacer (Lowe's Inc.) were added as described with the exception of using 3 stainless steel bearings (Amazon Inc.) instead of a nickel coin [21]. *I. scapularis* nymphs carrying *B. burgdorferi* B31-5A4, *B. afzelii* CB43, *B. garinii* ZQ1; or *B. burgdorferi* B31-5A4, B31-5A15, B31-5A4NP1 $\Delta cspA$ harboring a vector or this *cspA* mutant strain producing CspA_{B31}, CspA_{PKo}, CspA_{ZQ1}, or CspA_{B31}L246D were then added onto the chamber (5 to 8 ticks/chamber). Chamber feedings

were carried out as previously described using human (BioIVT, Westbury, NY) or quail blood (Canola Poultry Market, Brooklyn, NY) (Fig 2A), which were treated with heparin (final concentration 75 U/ml for quail blood) and pre-warmed at 37°C for 45 min. Chambers in blood were placed in a sealed Styrofoam cooler with wet paper towels to maintain humidity at approximately 87 to 95% in an incubator at 37°C. Depending on the experiments, blood was treated with CVF (ComTech) or OmCI to a final concentration of 17 μ g/ml. Blood was changed daily and was collected along with ticks only after 5 days of feeding. SYBR-based qPCR was used to determine bacterial burdens in the ticks and blood using *Borrelia* 16S rRNA gene primers (S3 Table).

Determination of Lyme borreliae burdens in infected ticks, tissues and blood samples

The ticks fed on quail, mice, *P. leucopus* mice, or feeding chambers were homogenized by hand in a 1.5 ml Eppendorf tube (Eppendorf) with a plastic pestle (ThermoFisher Scientific). The DNA from tissues or blood or homogenized ticks was extracted using the EZ-10 Genomic DNA kit (Biobasic). The quantity and quality of DNA was assessed using a Nanodrop 1000 UV/Vis spectrophotometer (ThermoFisher Scientific). The 280:260 ratio was between 1.75 and 1.85, indicating the lack of contaminating RNA or proteins. qPCR was then performed to quantitate bacterial loads. Spirochete genomic equivalents were calculated using an ABI 7500 Real-Time PCR System (ThermoFisher Scientific) in conjunction with PowerUp SYBR Green Master Mix (ThermoFisher Scientific), based on amplification of the Lyme borreliae 16S rRNA gene using primers 16SrRNAfp and 16SrRNArp (S3 Table), as described previously [23,75]. Cycling parameters for SYBR green-based reactions were 50°C for 2min, 95°C for 10min, and 45 cycles of 95°C for 15s, 55°C for 30s, and 60°C for 1min. The number of 16S rRNA gene extracted from *B. burgdorferi* strain B31-5A4, then comparing the Cq values of the experimental samples.

Sequence analysis of PFam54-IV

Nucleotide sequences of the PFam54-IV genes, including *bba68* (*cspA*_{B31}), and *bba69* from *B. burgdorferi* B31 (AE000790), *pkoa0062*, *pkoa0063*, *pkoa0064*, *pkoa0065*, *pkoa0066*, and *pkoa0067* (*cspA*_{PKo}), from *B. afzelii* PKo (CP002950), and *zqa67*, *zqa68* (*cspA*_{ZQ1}), *zsa69*, *zsa70*, *zsa71*, and *zsa72* from *B. garinii* ZQ1 (AJ786369), were used as queries against the NCBI Gen-Bank nr database using BLASTN [102]. Sequences from organisms other than *B. burgdorferi*, *B. afzelii*, and *B. garinii* and duplicate sequences were discarded. The remaining sequences were then adjusted to include the full open reading frames with removal of any sequences containing premature stop codons. Sequences were grouped by species, and the sequences of the genes encoding the third and fifth clade of PFam54 (PFam54-III and Pfam54-V, respectively) from *B. burgdorferi* B31, *B. afzelii* PKo, and *B. garinii* ZQ1 were added to the sequence set (GenBank accession codes are found on S5-S7 Figs). Codon alignments were generated using T-Coffee on the TranslatorX webserver [103–104], followed by calculation of pairwise sequence identity in Clustal Omega [105]. Only Pfam54-IV best hits were kept. The remaining sequences were then realigned by codon and pairwise identity was calculated as above. (S5–S7 Figs).

Generation of recombinant PFam54 proteins

The pQE30Xa vectors encoding the open reading frames lacking the putative signal sequences of *bba68* (*cspA*_{B31}), *bba69* from *B. burgdorferi* strain B31-5A4, *mmsa67*, *mmsa68*, *mmsa69*, or

mmsa70, and *mmsa71* (*cspA_{MMS}*) from *B. afzelii* strain MMS, or *zqa67*, *zqa68* (*cspA_{ZQ1}*), *zsa69*, *zsa70*, *zsa71*, or *zsa72* from *B. garinii* strain ZQ1 were obtained previously to generate recombinant histidine-tagged proteins [24]. The plasmids were transformed into *E. coli* strain M15, and the plasmid inserts were sequenced using Sanger sequencing on an ABI 3730xl DNA Analyzer (ThermoFisher Scientific) at the NYSDOH Wadsworth Center ATGC Core Facility. The resulting M15 derived strains were used to produce respective recombinant PFam54-IV (S2 Table). The histidine-tagged PFam54-IV were produced and purified by nickel affinity chromatography with Ni-NTA agarose according to the manufacturer's instructions (Qiagen).

FH binding assay by qualitative ELISA

Qualitative ELISA for FH binding by PFam54-IV was performed as described [21,106]. One microgram of BSA (negative control; Sigma-Aldrich) or FH from mouse or quail was coated onto microtiter plate wells by incubating the plate for overnight at 4°C. Then, 100 μ l 2 μ M of histidine-tagged DbpA from *B. burgdorferi* strain B31 (negative control) [107] or each of the PFam54-IV was added to the wells. Mouse anti-histidine tag 1:200× (Sigma-Aldrich) and HRP-conjugated goat anti-mouse IgG 1:1,000× (Seracare Life Sciences) were used as primary and secondary antibodies, respectively, to detect the binding of histidine-tagged proteins. The plates were washed three times with PBST (0.05% Tween 20 in PBS), and 100 μ l of tetramethyl benzidine (TMB) solution (ThermoFisher Scientific) was added to each well and incubated for 5 min. The reaction was stopped by adding 100 μ l of 0.5% hydrosulfuric acid to each well. Plates were read at 405nm using a Tecan Sunrise Microplate reader at 5 min post incubation (Tecan Life Science).

Phylogenetic reconstruction

The PFam54-IV codon alignment from B. burgdorferi B31, B. afzelii MMS, and B. garinii ZQ1 was used to estimate a Bayesian phylogenetic tree in BEAST v1.8.4 with a relaxed lognormal clock, an estimated mutation rate of 4.75×10^{-6} substitutions/site/year and a coalescent Bayesian skyline model [34,108]. A Markov chain Monte Carlo chain length of 10,000,000 steps was used with a 100,000-step thinning, resulting in effective sample sizes greater than 200, an indication of an adequate chain mixing. The resulting maximum clade credibility tree was visualized in FigTree v1.4.4 [109]. We evaluated alternative, competing evolutionary scenarios for PFam54-IV based on species-specific divergence, or clustering by FH-binding activity shown in Fig 8A [110]. A battery of statistical phylogenetic tests was deployed in IQ-TREE (Kishino-Hasegawa, Shimodaira-Hasegawa, Expected Likelihood Weights, and Approximately Unbiased tests)[111–114]. The more extensive dataset was created by aligning all relevant PFam54-IV protein sequences using M-Coffee [115] (http://www.tcoffee.org) and estimating phylogenetic relationships using maximum likelihood in IQ-TREE v1.6.14 [116]. The best fit substitution model HIVw+F+R4 among 344 candidate models, including empirical amino acid frequencies and among-site rate heterogeneity modeling, was selected based on the Bayesian Information Criterion (BIC-weight = 0.9583) using the ModelFinder [117] feature in IQ-TREE. Node robustness was assessed with 1000 ultrafast bootstrap replicates [118] and the SH-aLRT test with 5000 replicates [119].

Statistical analysis

Significant differences between samples were assessed using the Mann-Whitney *U* test or the Kruskal-Wallis test with the two-stage step-up method of Benjamini, Krieger, and Yekutieli. A P-value < 0.05 (*) or ([#]) was considered to be significant [120].

Supporting information

S1 Fig. Lyme borreliae vary in their ability to infect mice via intradermal injection of BALB/c but not C3^{-/-} BALB/c mice. (A to D) BALB/c or (E to H) C3^{-/-} BALB/c mice were injected with 10⁶ B. burgdorferi B31-5A4 ("Bb B31-5A4"), B. garinii ZQ1 ("Bg ZQ1"), or B. afzelii CB43 ("Ba CB43"). At 21 days post injection ("dpi"), spirochete burdens were determined in the (A and E) inoculation site ("Inoc. site"), (B and F) ears, (C and G) bladder, (D and H) ankles. Additionally, uninfected I. scapularis larvae were allowed to feed on those C3^{-/-} mice at 14 days post injection ("dpi") to repletion. Spirochete burdens in (I) replete larvae, and (J) post molting flat nymphs ("flat nymphs") were determined. The tissues from uninfected mice and uninfected nymphs were included as control ("Uninfect."). Shown are the geometric means of bacterial loads \pm 95% confidence interval of bacterial burdens from 6 replete larvae, flat nymphs or tissues from 5 BALB/c mice or indicated numbers of $C3^{-/-}$ BALB/c mice (9 Bb B31-5A4-infected inoculation sites and ears, 6 Bg ZQ1-infected inoculation sites and ears, or 5 all other tissues). Significant differences (p < 0.05, Kruskal-Wallis test with the two-stage stepup method of Benjamini, Krieger, and Yekutieli) in the spirochete burdens relative to uninfected ticks or tissues (*) are indicated. (TIF)

S2 Fig. The complement of quail inoculated with OmCI is inactivated. *Coturnix* quail were subcutaneously injected with OmCI (1 mg/kg of quail) or PBS buffer. Untreated (black bars) or heattreated (white bars) of sera collected from these quail at indicated time points after inoculation were incubated with a serum-sensitive, highly passaged *B. burgdorferi* strain B313 for 0-h and 4-h with a final concentration of 40%. The number of motile spirochetes was assessed through microscopy. The survival percentage of the spirochetes was calculated using the number of mobile spirochetes at 4 h post incubation normalized to that at 0 h of incubation with serum. Each bar represents the mean of three independent determinations \pm SEM from sera from five quail per group. Significant differences (p < 0.05, Kruskal-Wallis test with the two-stage step-up method of Benjamini, Krieger, and Yekutieli) in the percentage survival of spirochetes incubated with untreated sera from OmCIinoculated quail, compared to that in heat-inactivated sera from those quail (*). (TIF)

S3 Fig. Protein sequence alignment of CspA variants from *B. afzelii* **CB43, PKo, and MMS and the synteny from PFam54-IV proteins of these strains.** (A) CspA variants from *B. afzelii* CB43, PKo, and MMS aligned in M-Coffee. Blue shading indicates complete sequence conservation. (B) Synteny of Pfam54-IV genes. Colors indicate one-to-one orthologs. Locus *pkoa0066* is only found in strain PKo. Scale bar denotes 500 bp. (TIF)

S4 Fig. CspA variants confer differential transmissibility from nymphs to *P. leucopus* mice. *Ixodes scapularis* nymphs infected by WT *B. burgdorferi* B31-5A15 ("B31-5A15"), *B. burgdorferi* B31-5A4NP1 Δ cspA transformed with an empty shuttle vector ("Vector"), or this deletion strain producing a mutated variant of CspA from *B. burgdorferi* B31 selectively devoid of FH binding activity ("L246D"), WT *B. burgdorferi* B31 ("B31"), *B. garinii* ZQ1 ("ZQ1"), or *B. afzelii* PKo ("PKo") were allowed to feed to repletion on *P. leucopus* mice. Uninfected nymphs and *P. leucopus* mouse tissues and blood were included as control ("Uninfect."). Fed nymphs were collected upon repletion, and blood and tissues were collected at 7 days post nymph feeding ("dpf"). Spirochete burdens in (A) replete nymphs, (B) tick bite sites of skin ("Inoc. site"), and (C) blood were determined by qPCR. For spirochete burdens in tissue samples, the resulting values were normalized to 100 ng of total DNA. Shown are the geometric means of bacterial loads \pm 95% confidence interval of bacterial burdens in tissues from five *P*. *leucopus* mice per group or replete nymphs (8 nymphs carrying the strain B31-5A15, 12 nymphs carrying the strain "Vector", 13 nymphs carrying the strain pCspA-B31, 15 nymphs carrying the strain pCspA-PKo, or 13 nymphs carrying the strain pCspA-ZQ1). Significant differences (p < 0.05, Kruskal-Wallis test with the two-stage step-up method of Benjamini, Krieger, and Yekutieli) in the spirochete burdens relative to uninfected ticks or *P. leucopus* mouse tissues or blood are indicated (*). (TIF)

S5 Fig. Genetic divergence of Pfam54-IV alleles from *B. burgdorferi*. Nucleotide sequences encoding PFam54-IV proteins from *B. burgdorferi* B31 were used as queries to mine NCBI GenBank for orthologs. (**inset**) Frequency distribution of pairwise genetic distances. The pairwise identity numbers are coded by color gradually from identical (100% pairwise identity; green) to divergent sequences (70% pairwise identity; red). The clear break in the frequency distribution separates highly similar (> 95% pairwise identity) from moderately divergent comparisons (< 80%). The Pfam54-IV ortholog from a particular *B. burgdorferi* strain with > 95% identity to *cspA*_{B31} was defined as the *cspA* ortholog in that strain. (TIF)

S6 Fig. Genetic divergence of Pfam54-IV alleles from *B. afzelii*. Nucleotide sequences encoding PFam54-IV proteins from *B. afzelii* PKo were used as queries to mine NCBI Gen-Bank for orthologs. (**inset**) Frequency distribution of pairwise genetic distances. The pairwise identity numbers are coded by color gradually from identical (100% pairwise identity; green) to divergent sequences (40% pairwise identity; red). The clear break in the frequency distribution separates highly similar (> 98% pairwise identity) from moderately divergent comparisons (< 80%). The Pfam54-IV ortholog from a particular *B. afzelii* strain with > 98% identity to *cspA*_{B31} was defined as the *cspA* ortholog in that strain. (TIF)

S7 Fig. Genetic divergence of Pfam54-IV alleles from *B. garinii*. Nucleotide sequences encoding PFam54-IV proteins from *B. garinii* ZQ1 were used as queries to mine NCBI Gen-Bank for orthologs. (**inset**) Frequency distribution of pairwise genetic distances. The pairwise identity numbers are coded by color gradually from identical (100% pairwise identity; green) to divergent sequences (65% pairwise identity; red). The clear break in the frequency distribution separates highly similar (> 93% pairwise identity) from moderately divergent comparisons (< 80%). The Pfam54-IV ortholog from a particular *B. garinii* strain with > 93% identity to *cspA*_{B31} was defined as the CspA ortholog in that strain. (TIF)

S1 Table. Host-specific transmissibility and FH-binding activity of *B. burgdorferi*, *B. afze-lii*, and *B. garinii*, and their derived PFam54-IV proteins. (DOCX)

S2 Table. *Borrelia* sp. strains and plasmids used in this study. (DOCX)

S3 Table. PCR primers used in this study. (DOCX)

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