

# Structural evolution of an immune evasion determinant shapes pathogen host tropism

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Modern infectious disease outbreaks often involve changes in host tropism, the preferential adaptation of pathogens to specific hosts. The Lyme disease-causing bacterium Borrelia burgdorferi (Bb) is an ideal model to investigate the molecular mechanisms of host tropism, because different variants of these tick-transmitted bacteria are distinctly maintained in rodents or bird reservoir hosts. To survive in hosts and escape complement-mediated immune clearance, Bb produces the outer surface protein CspZ that binds the complement inhibitor factor H (FH) to facilitate bacterial dissemination in vertebrates. Despite high sequence conservation, CspZ variants differ in human FH-binding ability. Together with the FH polymorphisms between vertebrate hosts, these findings suggest that minor sequence variation in this bacterial outer surface protein may confer dramatic differences in host-specific, FH-binding-mediated infectivity. We tested this hypothesis by determining the crystal structure of the CspZ-human FH complex, and identifying minor variation localized in the FH-binding interface yielding bird and rodent FH-specific binding activity that impacts infectivity. Swapping the divergent region in the FH-binding interface between rodent- and bird-associated CspZ variants alters the ability to promote rodent- and bird-specific early-onset dissemination. We further linked these loops and respective host-specific, complement-dependent phenotypes with distinct CspZ phylogenetic lineages, elucidating evolutionary mechanisms driving host tropism emergence. Our multidisciplinary work provides a novel molecular basis for how a single, short protein motif could greatly modulate pathogen host tropism.

Lyme disease | CspZ | factor H | complement | host tropism

The emergence of infectious disease outbreaks often involves changes in host tropism, the preferential adaptation of pathogens to selectively invade and persist in hosts (1). Such host tropism is often the result of ongoing host-pathogen interactions (2). Evolution theoretically favors the emergence of host-specializing pathogens, but host-generalist strategies can be advantageous in environments where pathogens have the potential to regularly interact with multiple hosts (3). Specialism vs. generalism can be attributed to polymorphisms within pathogen proteins that differentially interact with host ligands (2). Such polymorphism-mediated host ranges are often thought to require complex host-specific adaptive mechanisms (4 and 5), but actually can be conferred by a few variable amino acids (AAs) (6). Understanding how such minor differences impact diverse host-adapted phenotypes can elucidate the mechanistic insights into the emergence of modern infectious diseases, allowing for the development of earlier and more efficient public health interventions.

Species within the *Borrelia* (*Borreliella*) *burgdorferi* sensu lato genospecies complex cause Lyme disease (LD). These spirochetal bacteria are transmitted by ticks and maintained by several reservoir hosts, primarily rodents and birds (7). Lyme borreliae have been genotyped using different polymorphic loci, such as *ospA*, *ospC*, the 16S-23S rRNA intergenic spacer (RST [typing]), and multilocus sequence typing (8, 9). Laboratory and field studies have shown not only that Lyme borreliae species differ in the host species they infect, but that individual genotypes within single species display distinct preferential host associations, particularly within *B. burgdorferi* sensu stricto (hereafter *B. burgdorferi*, "*Bb*"), the primary LD agent in North America (10, 11). This species- and genotype-specific host selectivity distinguishes Lyme borreliae as a model system for studying the molecular basis and evolutionary history of host tropism (10, 11).

Lyme borreliae host associations require the spirochetes to transmit from infected ticks to hosts, establish infection at tick-biting sites, and disseminate hematogenously to persist in distal tissues (10, 11). The survival of Lyme borreliae during these discrete events

#### Significance

Changes to pathogens' ability to infect particular hosts ("host tropism") often dictate modern infectious disease outbreaks. Lyme disease bacteria present a suitable model to study the molecular drivers of host tropism, as variants selectively survive in different hosts. To escape from killing by complement, part of the immune system, these bacteria produce the outer surface protein CspZ, which binds the complement inhibitor factor H (FH) to promote infection. Here, we applied multidisciplinary approaches to identify a small variable AA motif within the FH-binding site that drives host-specific FH-binding activity and infectivity. We further elucidated the evolutionary history of the recent CspZ divergence to provide new insights into the emergence of pathogen host tropism.

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The authors declare no competing interest.

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necessitates evasion of multiple host immune responses, including complement, a first-line innate immune defense mechanism (12–14). Complement can be activated by the classical, lectin, and/or alternative pathways (15). The activation of these pathways results in the cascading cleavage and recombination of multiple complement components, ultimately resulting in phagocytic clearance, inflammation, and lytic pore formation on the pathogen surface by the C5b-9 protein complex. Complement is downregulated by diverse regulatory proteins that inhibit activation, preventing native cell damage in the absence of pathogens. For example, the alternative pathway is inhibited by factor H (FH), which is comprised of 20 individual short consensus repeat (SCR) domains (16).

Similar to many bloodborne pathogens that evolved mechanisms to circumvent complement-mediated clearance, Lyme borreliae produce several outer surface proteins that bind and recruit complement components and/or regulatory proteins on the spirochete surface to inactivate complement (12–14). One of these proteins is an FH-binding protein, CspA (also known as Complement Regulator Acquiring Surface Protein 1, CRASP-1, or BBA68 of pFam54-60 in *Bb* strain B31) that confers spirochete survival via complement evasion in the tick bloodmeal (17). However, this protein is downregulated after spirochetes infect hosts, suggesting other functionally redundant proteins may facilitate spirochete survival at this infection stage. In fact, CspZ (also known as CRASP-2 or BBH06 in *Bb* strain B31), which binds to the 6th and 7th domains of the SCR (SCR6-7), is upregulated when spirochetes reside in hosts (18, 19). This protein is highly conserved both between and within different Lyme borreliae species [>85% AA identity between species and 98% between Bb strains] (20–23). We and others have previously reported that the FH-binding activity of one CspZ variant promotes spirochete survival in vertebrate sera, and consequently promotes host infectivity using non-physiologically relevant infection routes (i.e., not tick-transmitted) (24, 25). However, CspZ variants differ in their human FH-binding activity (20, 21), and vertebrate complement components or regulatory proteins vary between host taxa (e.g., ~40% AA identity between mammalian and avian FH) (26). These findings raise several intriguing questions: Could this minor divergence among CspZ variants confer host-specific differences in FH-binding activity, resulting in varying host infectivity phenotypes? Further, how did such divergences evolve to impact Lyme borreliae host associations?

In this study, we solved the high-resolution structure of the CspZ-human FH complex, identified a polymorphic CspZ motif within the FH-binding interface, and defined its contribution to host-specific FH-binding activity. We then examined the role of this motif in dictating strain-specific, host-dependent dissemination, using mice and quail as rodent and avian models, respectively. Paired with evolutionary analyses of cspZ, we further elucidated how the evolutionary history behind a minor divergence in an immune evasion determinant can impact pathogen host tropism.

## Results

A Polymorphic CspZ Loop Structure Defines Human FH-Binding Activity. We set to identify the regions driving the differences in human FH-binding activity among CspZ variants. We first pinpointed the AAs that are involved in FH-binding by cocrystalizing SCR6-7 of human FH and a recombinant CspZ variant with human FH-binding activity, from *Bb* strain B408 ("CspZ<sub>B408</sub>") (21). The resulting crystal structure (2.59 Å) showed an extensive binding interface between CspZ<sub>B408</sub> and both SCR6 and SCR7 (Fig. 1*A*). Specifically, Asp47, Tyr50, Asn51, Thr54, Asn58, and Thr62 in helix-B, Arg142 in helix-F, Asn183 from helix-G, and Tyr214 in helix-I of CspZ<sub>B408</sub> interacted with SCR7 (Fig. 1 *B* and *C* and *SI Appendix*, Table S1). Additionally, Asp71 from the loop between helix-B and -C, and Asp73 and Ser75 from the same region, bound to SCR7 and SCR6, respectively (Fig. 1 *B* and *C* and *SI Appendix*, Table S1).

To locate the residues that impact the variable human FH-binding activity, we aligned the sequences of CspZ<sub>B408</sub> with one CspZ variant that binds to human FH (CspZ from *Bb* B31, "CspZ<sub>B31</sub>"), and one that lacks human FH-binding ability from *Bb* strain B379 ("Csp $Z_{B379}$ ") (21). All the aforementioned residues involved in human FH-binding are conserved among these variants, except Asn51 and Asp71 in CspZ<sub>B408</sub> (*SI Appendix*, Fig. S1A). We also determined the crystal structures of CspZ<sub>B379</sub> and CspZ<sub>B408</sub> (2.10 and 2.45 Å, respectively). The previously crystalized  $CspZ_{B31}$  superimposes very well with both  $CspZ_{B379}$  (RMSD: 0.88 Å over 190 aligned C $\alpha$  atoms) and CspZ<sub>B408</sub> (RMSD: 0.90 Å over 195 aligned C $\alpha$  atoms), reflecting the high conservation between the proteins (23) (Fig. 1A). Moreover, the crystal structure of CspZ<sub>B408</sub> shows very high similarity to CspZ<sub>B408</sub> from the complex structure (RMSD: 0.93 Å over 201 aligned C $\alpha$  atoms). For the two aforementioned nonconserved AAs involved in human FH-binding: in helix-B, Asn51 in CspZ<sub>B408</sub> is substituted by a Ser51 in CspZ<sub>B379</sub>. The superimposed structure suggests that the relatively small side chain of serine would not inhibit human FH-binding in CspZ<sub>B379</sub> because the position of helix-B is identical in both proteins (RMSD: 0.15 Å over 16 aligned C $\alpha$  atoms) (Fig. 1 B and C). The second nonconserved AA, Asp71 in  $CspZ_{B408}$ , is part of an insertion that is not present in  $CspZ_{B31}$ and CspZ<sub>B379</sub> (Fig. 1*B*), and CspZ<sub>B31</sub> binds human FH. Therefore, these two non-conserved residues alone cannot explain the differential human FH-binding ability of these CspZ variants. However, the C-terminus of helix-B and the subsequent loop region between helix-B and helix-C harbor polymorphisms (Fig. 1A): CspZ<sub>B379</sub> has a unique duplication of the last residues in helix-B (Ile60, Met61, Thr62, and Tyr63) (SI Appendix, Fig. S1A), resulting in an extended helix-B in this variant (Fig. 1B). This structural extension leads to steric hindrance between CspZ<sub>B379</sub> and human FH to selectively prevent human FH-binding activity by CspZ<sub>B379</sub>. The abovementioned regions are well defined in the electron density map and are not influenced by crystal packing (SI Appendix, Fig. S2). Thus, our results suggest variant-specific CspZ structural differences in a loop and the adjacent helix-B (hereafter, "loop structures"), providing mechanistic insights into the CspZ-mediated, polymorphic human FH-binding activity.

CspZ Loop Structures Determine Host-Specific FH-Binding Activity. FH SCR6-7 is divergent among vertebrate species (*SI Appendix*, Fig. S1*B*), suggesting the CspZ loop structures may dictate host-specific FH-binding activity. We found that CspZ<sub>B31</sub> and CspZ<sub>B408</sub>, but not CspZ<sub>B379</sub>, bound to mouse FH. CspZ<sub>B31</sub> and CspZ<sub>B379</sub>, but not CspZ<sub>B408</sub>, bound to quail FH (Fig. 2A and SI Appendix, Fig. S3 and Table S2). We then searched for structural evidence of this FH-binding activity by superimposing the complex structure of  $CspZ_{B408}$ -human FH with  $CspZ_{B31}$ , CspZ<sub>B379</sub>, the previously resolved structures of mouse SCR6-7, and the AlphaFold-predicted structure of quail SCR6-7 (pLDDT > 70) (Fig. 2*C*). There are similar tertiary structures between mouse and human SCR6-7 (RMSD: 2.47 Å over 108 aligned C $\alpha$  atoms), including segments facing the protein complex interface which is consistent with the ability of both FH variants to selectively bind CspZ<sub>B31</sub> and CspZ<sub>B408</sub> (Fig. 2A and SI Appendix, Fig. S3 and Table S2) (21). The CspZ<sub>B379</sub>-specific duplication showed



**Fig. 1.** Superimposed crystal structures of the CspZ<sub>B408</sub>-SCR6-7 complex, CspZ<sub>B31</sub>, and CspZ<sub>B379</sub> reveal the FH-binding mechanisms of CspZ variants. (*A*) The crystal structure of human FH domains SCR6 (light gray) and SCR7 (dark gray) in complex with CspZ<sub>B408</sub> (orange) was superimposed with that of CspZ<sub>B31</sub> (purple) and CspZ<sub>B379</sub> (green). (*B*) The FH-binding interface with the location of the CspZ<sub>B408</sub> residues involved in human SCR6-7 binding, and the equivalent residues in CspZ<sub>B379</sub>. The numbering of the residues is given for CspZ<sub>B408</sub>.  $\alpha$ -helices in CspZ are labeled from A to I starting from the N terminus. The polymorphic loop region between  $\alpha$ -helices B and C in CspZ<sub>B408</sub> (residues 60-IMTYSEVNNVTD-71), CspZ<sub>B31</sub> (residues 60-IMTYSEGT-67) and CspZ<sub>B379</sub> (residues 60-IMTYIMTYSEGT-71) are indicated. A more detailed illustration of the CspZ<sub>B408</sub>-SCR6-7 complex interface involving the CspZ<sub>B408</sub> residues (*C*, *Top*) Tyr50, Thr54, Asn58 and Arg142 and (*C*, *Bottom*) the residues Asp47, Asn51, Asn183 and Tyr214 is shown.

potential structural hindrance in the mouse FH-binding interface, similar to human FH (Fig. 2*D*). However, the equivalent region in quail SCR7 (median pLDDT: 92) is positioned away from the CspZ-FH-binding interface, allowing sufficient space for the extended helix-B of CspZ<sub>B379</sub> to interact with quail FH (Fig. 2*D*). Further, the crystal structure of CspZ<sub>B408</sub>-SCR6-7 superimposed with the predicted structure of quail SCR6-7 showed that Asp71 of CspZ<sub>B408</sub> may collide with quail FH, while the equivalent region in mouse SCR7 would not interfere with the FH-binding of CspZ<sub>B408</sub> (Fig. 2*D*).

These structural differences in mouse/human vs. quail FH, paired with the respective impactful AAs from CspZ variants, suggest the loop structures of CspZ may cause host-specific FH-binding activity. To test this, we swapped the loop regions of the two CspZ variants with distinct host-specific FH-binding ability to generate two chimeric proteins: CspZ<sub>B379</sub>L<sub>B408</sub> has the backbone of the strictly quail FH-binder (CspZ<sub>B379</sub>), and the loop structures of the strictly mouse FH-binder (CspZ<sub>B408</sub>), whereas CspZ<sub>B408</sub>L<sub>B379</sub> has the backbone of CspZ<sub>B408</sub> and the loop structures of CspZ<sub>B379</sub> (*SI Appendix*, Fig. S1A). We found that CspZ<sub>B379</sub>L<sub>B408</sub> and CspZ<sub>B408</sub>L<sub>B379</sub> selectively bound to mouse and quail FH, respectively, demonstrating the CspZ loop structures are a determinant of host-specific FH-binding activity (Fig. 2*B* and *SI Appendix*, Fig. S3 and Table S2).

The CspZ Loop Structures Dictate Spirochete Strain- and Host-Specific Complement Inactivation. We next examined the hostdependent complement inactivation of these CspZ variants produced on the spirochete surface. We thus obtained a wild-type Bb strain B31-A3 (referred to hereafter as "WT B31-A3") and a cspZ-deficient mutant strain in this background harboring an empty vector (" $\Delta cspZ$ /Vector"). We complemented this mutant with a plasmid encoding  $cspZ_{B31}$  ("pCspZ<sub>B31</sub>") (25),  $cspZ_{B379}$  ("pCspZ<sub>B379</sub>"),  $cspZ_{B408}$  ("pCspZ<sub>B408</sub>"), or the loop-swapped mutants ("pCspZ<sub>B379</sub>L<sub>B408</sub>" and "pCspZ<sub>B408</sub>L<sub>B379</sub>"). We verified that these strains had similar generation times (SI Appendix, Table S3), no differences in the CspZ surface production level (SI Appendix, Fig. S4), and that surface-produced CspZ bound FH in the same manner as the recombinant proteins (SI Appendix, Fig. S5). We then used flow cytometry to determine the ability of these CspZ variants and mutants to inactivate mouse and quail complement by measuring the deposition levels of mouse C5b-9 and quail C8, respectively, after incubation with sera from these animals. All strains had levels of mouse C5b-9 or quail C8 deposition significantly lower than the control strain: high passage, noninfectious, and mouse and quail complement-susceptible *Bb* strain B313 (Fig. 3 A and B and SI Appendix, Fig. S6 A and B) (19).  $\Delta cspZ$ /Vector had significantly greater levels of mouse C5b-9 and quail C8 deposition than WT B31-A3 or pCspZ<sub>B31</sub> (Fig. 3



**Fig. 2.** The polymorphic loop region in CspZ promotes host-specific FH-binding ability. (*A* and *B*) Untagged CspZ variants or mutants were flowed over the chip surface, conjugated with indicated FH variants. Binding was measured in response units (R.U.) by SPR. The  $k_{on}$ ,  $k_{off}$ , and  $K_D$  values were determined from the average of these three experiments (*SI Appendix*, Table S2). (*C*) The crystal structure of CspZ<sub>B408</sub>-SCR6-7 where human SCR6-7 (gray) is superimposed with mouse SCR6-7 (gold, PDB: 2YBY), and the predicted structure of quail SCR6-7 (brown), while CspZ<sub>B408</sub> (orange) is superimposed with CspZ<sub>B379</sub> (green). (*D*) The loop region in CspZ proteins located between  $\alpha$ -helices B and C. The Asp71 in CspZ<sub>B408</sub>, and the residues 60-IMTYIMTYSEGT-71 in CspZ<sub>B379</sub> are highlighted. The polymorphic loop region between  $\alpha$ -helices B and C in CspZ variants, as well as the loop region in human, mouse and quail SCR6-7 located at the interface with helix-B in CspZ, is showed with a striped pattern. The potential structural hindrance region between CspZ<sub>B408</sub> and quail FH, as well as the structural hindrance between CspZ<sub>B408</sub> and both mouse and human FH, are indicated with the starbursts.

*A* and *B* and *SI Appendix*, Fig. S6 *A* and *B*) (25, 27). Compared to  $\Delta cspZ/Vector$ , pCspZ<sub>B379</sub> had indistinguishable levels of mouse C5b-9 deposition but significantly lower levels of quail C8, whereas pCspZ<sub>B408</sub> harbored significantly lower levels of mouse C5b-9 but indistinguishable levels of quail C8 (Fig. 3 *A* and *B* and *SI Appendix*, Fig. S6 *A* and *B*). Further, pCspZ<sub>B379</sub>L<sub>B408</sub> recruited significantly lower levels of mouse C5b-9 but indistinguishable levels of mouse C5b-9 but in

bound indistinguishable levels of mouse C5b-9 but significantly lower levels of quail C8 than  $\Delta cspZ/Vector$  (Fig. 3 *A* and *B* and *SI Appendix*, Fig. S6 *A* and *B*). These results correspond with these variants' host-specific FH-binding activity, suggesting the loop structure-driven host-specific FH-binding activity confers host-specific complement inactivation.

We also measured the survival of these strains in rodent and quail sera by counting the number of live bacteria using both live/



**Fig. 3.** The CspZ loop-driven, variable FH-binding activity confers host-specific complement inactivation and early dissemination. (*A* and *B*) Indicated *Bb* strains were incubated with mouse or quail sera (20%) followed by straining with antibodies against mouse C5b-9 or quail C8. The deposition levels of (*A*) mouse C5b-9 or (*B*) quail C8 on the surface of the indicated strains were measured by flow cytometry. Each bar represents the mean of three independent experiments ±SEM. Significant differences in the deposition levels relative to B313 (" $\Phi$ "), *ΔcspZ*/Vector ("\*"), or between two strains relative to each other ("#"), are indicated. (*C* and *D*) the *B*. *burgdorferi* strains were incubated with (*C*) white-footed mouse sera or (*D*) quail sera (40%). The number of live spirochetes was assessed microscopically using live/dead staining. The percent survival of the strains was calculated using the number of live spirochetes at 4-h post incubation normalized to that prior to incubation with sera. Each bar represents the mean of three independent experiments (n = 3) ±SEM. Significant differences in the percent survival of spirochetes relative to each other ("#") are indicated. (*E-M*) *I*. *scapularis* nymphs carrying indicated *Bb* strains were allowed to feed until repletion on indicated animals. Shown are the geometric mean of *recA* copies in animal tissues, normalized to 10<sup>4</sup> copies of (*E-I*) mouse nidogen or (*J-M*) quail β-actin ±SEM of five animals per group, except for the BALB/c blood from the B31-A3 and *ΔcspZ*/Vector groups, which have six and 10 mice per group, respectively. Significant differences in the spirochete survival relative to each other ("#") are indicated.

dead staining and the number of colony-forming units (CFUs) after plating. Note that sera from white-footed mice (*Peromyscus leucopus*) rather than house mice (*Mus musculus*) were used to represent rodent sera. House mouse complement is highly labile in vitro, which leads to inconsistent results (28). We found that WT B31-A3 survived in white-footed mouse and quail sera more efficiently than  $\Delta cspZ$ /Vector (Fig. 3 C and D and SI Appendix, Fig. S7). Because the percent survival yielded from live/dead staining and CFU counting were indistinguishable (*SI Appendix*,

Fig. S7), counting the number of live bacteria via live/dead staining was used to determine the survival of the rest of the strains. We found that pCspZ<sub>B31</sub> also survived in white-footed mouse and quail sera at greater levels than  $\Delta cspZ$ /Vector (Fig. 3 *C* and *D*) (25). Compared to  $\Delta cspZ$ /Vector, pCspZ<sub>B379</sub> survived at significantly greater levels in quail but not white-footed mouse sera, whereas pCspZ<sub>B408</sub> survived at significantly higher levels in white-footed mouse but not quail sera (Fig. 3 *C* and *D*). Relative to  $\Delta cspZ$ /Vector, pCspZ<sub>B379</sub>L<sub>B408</sub> survived significantly higher in white-footed mouse but not in quail sera, whereas  $pCspZ_{B408}L_{B379}$ survived significantly higher in quail but not in white-footed mouse sera (Fig. 3 *C* and *D*). However, these differences were not observed in the presence of sera treated with Cobra Venom factor or *Ornithodorus moubata* complement inhibitor (OmCI) that deplete functional rodent and quail complement, respectively (*SI Appendix*, Fig. S8) (29, 30). This suggests that CspZ loop structures determine spirochete host-specific serum survival.

The CspZ Loop Structures Define Host-Specific, Complement-Dependent, Early-Onset Dissemination. We further evaluated whether the CspZ loop structure variants determine host infectivity by allowing *Ixodes scapularis* nymphs carrying similar burdens of each of the aforementioned Bb strains to feed on mice and quail (*SI Appendix*, Fig. S9 A and D). The burdens of  $\Delta cspZ$ /Vector were significantly lower than WT B31-A3 at 7- and 10-days postfeeding (dpf) in mouse blood and at 10 dpf in distal tissues (joints, heart, and bladder), and at 9 dpf in quail blood and distal tissues (brain and heart) (SI Appendix, Fig. S10). There were no differences in burdens by a later time point (i.e., 14 dpf, *SI Appendix*, Fig. S10). We thus performed subsequent experiments using our entire Bb strain cohort at 10 dpf and 9 dpf in mouse and quail, respectively. In mice, we found indistinguishable spirochete burdens at initial infection sites (tick bite sites) between all seven strains tested at 10 dpf (Fig. 3*E*). pCspZ<sub>B31</sub>, pCspZ<sub>B408</sub>, and pCspZ<sub>B379</sub>L<sub>B408</sub>, but neither  $pCspZ_{B379}$  nor  $pCspZ_{B408}L_{B379}$ , colonized mouse blood and distal tissues at significantly higher levels than  $\Delta cspZ/$ Vector (Fig. 3 *F–I*). At 9 dpf in quail, we found no significantly different spirochete burdens at tick bite sites among strains. However, pCspZ<sub>B31</sub>, pCspZ<sub>B379</sub>, and pCspZ<sub>B408</sub>L<sub>B379</sub>, but neither  $pCspZ_{B408}$  nor  $pCspZ_{B379}L_{B408}\text{, colonized quail blood and}$ distal tissues at significantly greater levels than  $\Delta cspZ/Vector$ (Fig. 3 J-M). There were no significantly different burdens of any strains in any tissue from complement-deficient mice and

quail (C3<sup>-/-</sup> mice or OmCI-treated quail) (*SI Appendix*, Fig. S11). These results demonstrate that the CspZ loop structures define complement-dependent, spirochete strain- and host-specific early onset dissemination.

The Population-Wide CspZ Loop Structures Evolved from a Variant with Versatile FH-Binding Ability. To investigate the evolutionary history behind the CspZ loop structures, we mined *Bb cspZ* from NCBI GenBank and Sequence Read Archive (totaling 174 high-quality cspZ isolates originating from ticks, reservoir hosts, and patients, across North America, Europe, and Asia). A haplotype-based phylogenetic network distinctly separated three individual lineages, each of which contained one of the abovetested alleles:  $cspZ_{B31}$ ,  $cspZ_{B379}$ , or  $cspZ_{B408}$  (SI Appendix, Fig. S12). Examining the entire dataset, isolates within the same lineage had over 99% overall sequence identity and the same loop structure as  $cspZ_{B408}$ ,  $cspZ_{B379}$ , or  $cspZ_{B31}$  (defined as the unique insertion, duplication, or neither, respectively) (Fig. 4). We pinpointed single AA polymorphisms (SAPs) in the isolates and found none of the SAPs within each lineage were in the FH-binding interface, except for one SAP in one isolate in the CspZ<sub>B379</sub> lineage (SI Appendix, Tables S1 and S4). The majority of the SAPs were between lineages, located in the loop structures driving the FH-binding abilities. While this present study has been the only one to investigate the abilities of CspZ variants to bind house mouse and quail FH, the human FH-binding abilities of 13% of these 174 isolates have been evaluated (20, 21, 25). Incorporating these findings with the phylogeny, we found a 100% correlation of known host-specific FH-binding ability with lineages (Fig. 4). These results suggest each CspZ lineage with distinct loop structures can be linked to CspZ-specific, host-dependent FH-binding ability.

We explored the evolutionary mechanisms that could have led to this minor loop structure-dependent FH-binding specificity. No evidence for recombination was found (GARD method: 38



**Fig. 4.** Phylogeny and sequence comparisons support polymorphic CspZ loop arising from host-specific adaptation of Lyme borreliae. Unrooted likelihood phylogenetic tree of 174 *Bb cspZ* isolates generated with IQ-tree and visualized in iTOL. The names of the isolates are highlighted based on their CspZ loop types. The percent sequence identity of each isolate relative to CspZ<sub>B31</sub> (purple), CspZ<sub>B408</sub> (orange), or CspZ<sub>B379</sub> (green) is indicated on the outer ring. The known FH-binding activity is marked: black or gray stick figures, mice, and quail indicate FH binding or lack thereof, respectively.

potential and one inferred breakpoints) or past changes in effective population size (Bayesian skyline analysis: all predicted mean population sizes are within the lowest and highest HPD95% of all timepoints). The evidence for episodic diversifying selection along cspZ was weak (BUSTED method including site-to-site synonymous rate variation and double/triple nucleotide substitutions: p =0.025) while excluding terminal branches on the gene phylogeny. Several codons appear to deviate from expectations of neutrality (SI Appendix, Fig. S1A). We reconstructed the possible last common ancestor states ("LCAS") of the entire Bb CspZ dataset using the previously estimated chromosome mutation rate (31). The diversification between the three lineages was dated at 261 to 784 years B.P. (YBP) (SI Appendix, Table S5), whereas the diversification of the Bb CspZ LCAS as a whole likely occurred 2166 YBP (HPD95% 896 to 5,718). The LCAS variants possess a loop region resembling that of CspZ<sub>B31</sub> both in primary and tertiary structure (SI Appendix, Figs. S1A and S13), and bound to human, mouse, and quail FH, similar to CspZ<sub>B31</sub> (SI Appendix, Fig. S14 and Table S6). These results suggest that modern CspZ variants arose relatively recently from a generalist variant with versatile FH-binding features.

#### Discussion

Host tropism is an outcome of host-pathogen interactions (11). In vector-borne zoonotic pathogens, such interactions include vector-to-host transmission, pathogen dissemination and persistence, and host-to-vector acquisition (32). For Lyme borreliae to survive these steps, spirochetes need to overcome numerous host immunological mechanisms, including complement-mediated killing. In fact, Lyme borreliae inactivate complement in the tick blood meal for tick-to-host transmission, and in the host bloodstream for dissemination and persistence (25, 33, 34). In this study, we examined this inactivation conferred by the FH-binding protein, CspZ. We showed reduced burdens of a tick-introduced, cspZ-deficient mutant at the early stages of bloodstream survival and distal tissue colonization, defining this polymorphic protein as a contributor to survival at these early stages. Superimposing the crystal structures of the three polymorphic CspZ variants revealed a short, variable AA motif consisting of duplication and insertion/deletion events (23). Our newly-resolved and softwarepredicted complex structures of CspZ with human, mouse, and quail SCR6-7 further linked these variable residues to distinct host-specific FH-binding activity, in a unique manner compared previously reported pathogenic FH-binding proteins (SI Appendix, Fig. S15) (35). These results are congruent with the ability of CspZ variants and their associated mutants with swapped loop structures to promote host-specific FH-binding activity, complement evasion, and dissemination, defining this CspZ loop structure as a determinant of host tropism. Our findings thus demonstrate the concept of how minor variation functionally impacts host-adapted phenotypes, providing a new mechanism for host tropism modulation by pathogens (36-38).

We integrated an evolutionary approach to investigate the emergence of such host tropism determinants, an approach whose importance has been recently demonstrated to understand and track modern disease emergence (39). Our findings demonstrate that modern CspZ variants emerged from a last common ancestor that could bind human, mouse, and quail FH. This diversification occurred relatively recently compared to the origin of *Bb* 60,000 YBP, and the diversification of another FH-binding protein, CspA, about 25,000 YBP (31, 33). The lack of CspZ orthologs also supports the predicted young age of this protein, and as such it may be undergoing widespread neutral evolution (23, 40). We detected diversifying selection using multiple models on three codons, all of which are located in helices. No evidence for positive selection was found in the loop structures driving FH-binding phenotypes, despite those loops harboring most of the variation and insertions/deletions across CspZ. Alignment columns with gaps are excluded from most selection scans, rendering the detection of selective pressure more challenging and less trustworthy.

Other polymorphic anticomplement loci, particularly *ospC*, are undergoing balancing selection (41, 42). Negative frequencydependent selection (NFDS) and multiple-niche polymorphism (MNP) have been proposed as mechanisms driving ospC evolution. NFDS allows for high allele diversity to emerge through the negative correlation between pathogen fitness and the frequency of any alleles in the pathogen population (43). In Lyme borreliae, NFDS may explain the greater antibody-mediated clearance of spirochete strains harboring higher-frequency *ospC* alleles (44, 45). MNP maintains elevated levels of genetic diversity by varying pathogen fitness across reservoir hosts (42). As a systemically redundant anticomplement protein (46), CspZ may also be subjected to balancing selection. It is unlikely, but the lack of detectable selective pressure on the loop could also mean this locus is undergoing neutral selection. Balancing selection on one locus can increase diversity in another genetically-linked locus, and there is genetic linkage between cspZ and ospC types (20, 21, 27, 47). This raises the intriguing possibility that *ospC*-mediated balancing selection may promote diversification through drift in other functionally redundant genes such as *cspZ* without deleterious effects on the pathogen, warranting further investigations.

In this study, we used isogenic strains producing different CspZ variants under the control of an identical *cspZ* promoter to facilitate the equal production levels of each CspZ variant. While the strength of this approach can directly attribute phenotypes to the variation of coding sequences of CspZ (48), it does not consider the possibility that each *cspZ* variant is regulated differently in their parental strains. However, CspZ<sub>B379</sub> and  $CspZ_{B408}$  appear to be produced at similar levels in their parental strains (21), which may alleviate the abovementioned limitation. Additionally, it should be noted that other anticomplement proteins from ticks or spirochetes, or noncomplement-mediated mechanisms may (and likely do) contribute to the overall strain-specific, host-adapted phenotypes. This highlights the potential of polygenic host-specific phenotypes, which would need to be further considered in defining drivers of host tropism (49-51). Overall, the information and the platform established in this study would provide greater insights into pathogen-host interactions, facilitating the understanding of host tropism as a cause of infectious diseases.

## **Materials and Methods**

**Ethics Statement.** All animal experiments were performed in strict accordance with all provisions of guidelines detailed in *SI Appendix*.

**Bacteria**, **Mice**, **Quail**, **Ticks**, **Animal Sera**, **and Proteins**. The *Escherichia coli*, *Pichia pastoris*, and *Borrelia* strains, the sources of ticks, and animal sera, along with the FH and OmCl used in this study and their verification are detailed in the *SI Appendix*.

**Structure Determination by Crystallization and Prediction by AlphaFold.** CspZ<sub>B408</sub> (4 mg/mL) and human SCR6-7 (3 mg/mL) were mixed at a molar ratio of 1:2 and loaded on a pre-equilibrated HiLoad 16/600 Superdex 200 prep grade column (GE Healthcare, Chicago, IL, USA), followed by size exclusion chromatography and SDS PAGE. Crystals of this complex and those of CspZ<sub>B408</sub>, the diffraction data for these proteins and their complex, are detailed in Supplementary Information. AlphaFold v2.0 (52) was used to predict the 3D structure for quail FH SCR6-7 extrapolated from the sequences of *Coturnix japonica* complement FH (GenBank: XM\_015869474.2), detailed in the *SI Appendix*.

**ELISAs and SPR.** Enzyme-linked immunosorbent assay (ELISA) was used to determine FH-binding by CspZ proteins, detailed in the *SI Appendix*. For Surface Plasmon Resonance (SPR), interactions of CspZ proteins with FH were analyzed by SPR using a Biacore T200 (Cytiva, Marlborough, MA), detailed in *SI Appendix*.

**Flow Cytometry and Serum Resistance Assays.** CspZ production, FH-binding, complement deposition on spirochetes were determined using flow cytometry, detailed in *SI Appendix*. The serum resistance of *Bb* was measured as outlined in the *SI Appendix*.

**Mouse and Quail Infection.** The generation of flat, infected *I. scapularis* nymphs, the feeding of nymphs on mice and quail, and the determination of bacterial burdens at animal tissues and ticks are detailed in the *SI Appendix*.

**Genomic Analyses.** Sequences of both FH and CspZ were aligned and visualized, as detailed in *SI Appendix*. To generate the *cspZ* phylogenetic trees, we mined all publicly available *cspZ* sequences on NCBI as of September 2021. All resulting sequences were analyzed as outlined in the *SI Appendix*. The ancestor state of CspZ was reconstructed, described in the *SI Appendix*.

**Statistical Analysis.** Samples were compared using the Mann–Whitney *U* test or the Kruskal–Wallis test with the two-stage step-up method of Benjamini, Krieger, and Yekutieli (53).

**Data, Materials, and Software Availability.** The coordinates and the structure factors for CspZ<sub>B379</sub>, CspZ<sub>B408</sub>, and human SCR-CspZ<sub>B408</sub> have been deposited in the Protein Data Bank with accession codes 7ZJJ (54), 7ZJK (55), and 7ZJM (56), respectively. All study data are included in the article and/or *SI Appendix*.

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