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# Species Co-Occurrence Patterns among Lyme Borreliosis Pathogens in the Tick Vector *Ixodes ricinus*

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**Mixed infections have important consequences for the ecology and evolution of host-parasite interactions. In vector-borne diseases, interactions between pathogens occur in both the vertebrate host and the arthropod vector. Spirochete bacteria belonging to the *Borrelia burgdorferi sensu lato* genospecies complex are transmitted by *Ixodes* ticks and cause Lyme borreliosis in humans. In Europe, there is a high diversity of *Borrelia* pathogens, and the main tick vector, *Ixodes ricinus*, is often infected with multiple *Borrelia* genospecies. In the present study, we characterized the pairwise interactions between five *B. burgdorferi sensu lato* genospecies in a large data set of *I. ricinus* ticks collected from the same field site in Switzerland. We measured two types of pairwise interactions: (i) co-occurrence, whether double infections occurred more or less often than expected, and (ii) spirochete load additivity, whether the total spirochete load in double infections was greater or less than the sum of the single infections. Mixed infections of *Borrelia* genospecies specialized on different vertebrate reservoir hosts occurred less frequently than expected (negative co-occurrence) and had joint spirochete loads that were lower than the additive expectation (inhibition). In contrast, mixed infections of genospecies that share the same reservoir hosts were more common than expected (positive co-occurrence) and had joint spirochete loads that were similar to or greater than the additive expectation (facilitation). Our study suggests that the vertebrate host plays an important role in structuring the community of *B. burgdorferi sensu lato* genospecies inside the tick vector.**

Most hosts are infected with multiple parasite species or parasite strains (1–3). Interactions among parasite taxa infecting the same host can take a variety of forms (4). Competition for limited host resources can result in the elimination of the less competitive parasite; for example, *Wolbachia* bacteria prevent dengue viruses, chikungunya viruses, and malaria parasites from infecting *Aedes aegypti* mosquitoes (5). Alternatively, the presence of one pathogen may facilitate opportunistic infection by another pathogen; for example, HIV and fungal/bacterial infections (6–8), intestinal helminth and malaria infections (9), or influenza virus and pneumococcal infections (10). Multiple infections involving genetically distinct clones of the same parasite species are of particular interest, because their interactions are important in shaping the evolution of parasite virulence and disease severity (11–13), as shown in malaria (14, 15) and *Pasteuria* infections (16). Characterizing the interactions among parasite taxa in multiply infected hosts represents a major challenge for understanding the epidemiology of infectious diseases. In the case of vector-borne diseases, this task is further complicated because the parasites interact in both the arthropod vector and the vertebrate host.

The *Borrelia burgdorferi sensu lato* complex is a group of tick-borne spirochete bacteria that cause Lyme borreliosis, the most common tick-borne disease in the Northern Hemisphere (17). In Europe, there are at least 10 different *B. burgdorferi sensu lato* genospecies (18–21). All of these spirochetes are vectored by the hard-bodied tick *Ixodes ricinus* and maintained in a variety of vertebrate reservoir hosts (mostly birds and small mammals) (22). Previous field surveys have repeatedly shown that questing *I. ricinus* ticks often carry multiple spirochete infections (21, 23), providing ample opportunity for interactions among *Borrelia* genospecies. While many studies report the incidence of single and multiple *Borrelia* infections in *I. ricinus* (reviewed in reference 21), to our knowledge only one study (24) has tested whether the in-

cidence of multiple infections deviates from the random expectation. To better understand patterns of co-occurrence and abundance of these tick-borne pathogens, it is crucial that we compare their observed distributions to the random or neutral expectation (25, 26).

The specificity of *Borrelia* genospecies for their vertebrate reservoir hosts plays a key role in shaping the ecology of mixed *Borrelia* infections in both the host and the tick vector. Previous work has shown that *B. burgdorferi sensu stricto*, *B. afzelii*, and *B. bavariensis* are specific for rodents (27–30), whereas other genospecies, such as *B. garinii* and *B. valaisiana*, are specific for birds (30, 31). This host specificity appears to be mediated by the complement system of the vertebrate host (32, 33). Thus, the vertebrate complement system reduces the probability of encounter between rodent- and bird-adapted *Borrelia* genospecies. When encounters between maladapted coinfection partners do occur, the vertebrate immune system likely plays a key role in shaping the joint spirochete load in the tick vector. Spirochete growth rates may also influence the outcome of the mixed infection inside the tick vector. For example, the total spirochete population first expands following the larval blood meal and then declines during the larval premolting period (34), providing further opportunities for competitive interactions among *Borrelia* genospecies.

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**TABLE 1** Description of the seven sampling occasions (A to G) which provided the data on *Borrelia* genospecies infections in *I. ricinus* nymphs

Study	Season of collection	Yr of collection	Sample size	Treatment	Publication
A	Spring	2009	500	Survival under hot conditions	35
B	Spring	2010	1,500	Humidity attraction	36
C	Spring	2010	2,250	Survival under hot conditions	Unpublished
D	Fall	2010	450	Survival under cold conditions	Unpublished
E	Spring	2011	900	Fat content quantification	38
F	Spring	2011	800	Survival under cold conditions	37
G	Fall	2011	1,000	Survival under cold conditions	37

The purpose of this paper was to characterize the pattern of interactions among *B. burgdorferi sensu lato* genospecies in *I. ricinus* ticks. We sampled ticks from the same sampling site in Switzerland multiple times during a 3-year period and used this data set (7,400 nymphal ticks) to test whether double infections occurred more or less often than the random expectation. We limited our study to genospecies interactions in questing *I. ricinus* nymphs, because this stage is responsible for infecting the next generation of reservoir hosts and is therefore the most important stage from an epidemiological point of view. We predicted that mixed infections of rodent- and bird-adapted *Borrelia* genospecies (e.g., *B. afzelii*-*B. garinii* and *B. afzelii*-*B. valaisiana*) in *I. ricinus* nymphs would be relatively rare compared to the random expectation. We also predicted that the joint spirochete load of such mixed infections would be lower than the additive expectation, because we expected the host complement system to suppress the density of the maladapted coinfection partner. We predicted that mixed infections of *B. burgdorferi sensu lato* genospecies that use the same vertebrate reservoir hosts (e.g., *B. garinii*-*B. valaisiana*) would occur more often than expected by chance. For these mixed infections, we did not have a clear prediction for the joint spirochete load. However, as the tick vector presumably sets some upper limit on the spirochete load, we expected negative interactions between spirochete loads to be the norm. Detecting positive and negative patterns of species co-occurrence is an important first step toward understanding species interactions in general and competitive and facilitative interactions in particular. Although such comparative data cannot decipher the underlying causal mechanisms, they are critical for generating new hypotheses that can be tested further experimentally.

## MATERIALS AND METHODS

**Meta-analysis of *Borrelia* infection data from the same population of *I. ricinus* ticks.** The present study is a meta-analysis of seven independent sampling occasions that were conducted by Coralie Herrmann over the course of her Ph.D. thesis (35–38). The ticks collected on these sampling occasions were used in studies to quantify fat content in ticks and to test how temperature, humidity, and *Borrelia* infection influenced the physiology, behavior, and survival of *I. ricinus* nymphs (Table 1). Over a period of 3 years (2009 to 2011), 7,400 questing *I. ricinus* nymphs were collected from the same sampling site in Switzerland, and all of these ticks were processed in the same way with respect to quantification of spirochete load and *Borrelia* genospecies identification. Therefore, this data set pre-

sented a unique opportunity to test how *B. burgdorferi sensu lato* genospecies interact in the tick vector. After pooling the data from the sampling occasions, the large sample size gave us sufficient statistical power to test whether the frequency and spirochete load of mixed infections was different from the random expectation.

**Tick collection, spirochete load, and *Borrelia* genospecies identification.** The sampling site was a mixed forest situated 600 m above sea level on the south-facing slope of Chaumont Mountain, Neuchâtel, Switzerland (47°00' N, 6°57' E). Field sampling of questing ticks and subsequent molecular methods have been described in detail elsewhere (35, 36–38). Briefly, our molecular protocol consisted of two successive steps: (i) quantitative PCR (qPCR) to identify *Borrelia*-infected nymphs and to estimate the spirochete load, and (ii) reverse line blot (RLB) of the *Borrelia*-infected nymphs to identify the *Borrelia* genospecies. Thus, the sensitivity of our protocol depended on the ability of the qPCR to identify the *Borrelia*-infected ticks, whereas the ability to discriminate among the various *Borrelia* genospecies depended on the RLB. As the qPCR protocol estimated the total spirochete load, we do not have separate spirochete loads for each genospecies in the case of doubly infected nymphs.

We used the qPCR protocol from Schwaiger et al. (39), which targets the flagellin gene. For the RLB, we used the primers of Alekseev et al. (40) to amplify the variable spacer region between 2 repeated copies of the 23S and 5S ribosomal genes. The RLB protocol contains three general probes for *B. burgdorferi sensu lato* in addition to specific probes that allow us to detect each of the five *B. burgdorferi sensu lato* genospecies present at our study site: *B. burgdorferi sensu stricto*, *B. afzelii*, *B. garinii*, *B. valaisiana*, and *B. bavariensis* (23). Our RLB protocol also contains a probe that allows us to detect the relapsing fever-like spirochete *B. miyamotoi* (23). However, in the present paper, we only consider the pairwise interactions between *Borrelia* genospecies belonging to the *B. burgdorferi sensu lato* complex.

**Statistical methods. (i) Positive and negative co-occurrence of *B. burgdorferi sensu lato* genospecies.** We focused on interactions between pairs of *B. burgdorferi sensu lato* genospecies because higher-order interactions (i.e., triple infections) were exceedingly rare. Our study area contained five different *B. burgdorferi sensu lato* genospecies (*B. burgdorferi sensu stricto*, *B. afzelii*, *B. garinii*, *B. valaisiana*, and *B. bavariensis*), resulting in 10 possible genospecies pairs. We used log-linear analysis to characterize the co-occurrence patterns between pairs of *B. burgdorferi sensu lato* genospecies in the tick vector. For each genospecies pair, the data consist of the counts of four groups of ticks: (i) uninfected, (ii) infected with genospecies A, (iii) infected with genospecies B, and (iv) infected with genospecies A and B. A log-linear analysis essentially consists of modeling these count data as a function of genospecies A (presence/absence), genospecies B (presence/absence), and the genospecies A-genospecies B interaction using a generalized linear model with a Poisson error distribution. The sign (positive or negative) and statistical significance of the interaction term test whether the two genospecies co-occurred more or less often than expected by chance. The advantage of log-linear analyses over chi-square tests is that the sign and magnitude of the interaction term provide a quantitative estimate of the genospecies co-occurrence pattern.

We used a two-step approach to hypothesis testing. In the first step, we used three-way log-linear analysis to test whether we were justified in pooling the data from the seven sampling occasions for each of the 10 *B. burgdorferi sensu lato* genospecies pairs. Specifically, we tested whether the genospecies co-occurrence pattern was the same among the seven sampling occasions by evaluating the three-way interaction term for sampling occasion, genospecies A, and genospecies B. In the second step, we used two-way log-linear analysis to test the co-occurrence pattern for each of the 10 genospecies pairs depending on the results of the first step. If the three-way interaction (sampling occasion-genospecies A-genospecies B) was not statistically significant, we pooled the data from the different sampling occasions and estimated the global two-way interaction. If the three-way interaction was statistically significant, we did not pool the data and analyzed the two-way interaction separately for each of the seven sampling occasions. We used the glm() function in R with a Poisson error

**TABLE 2** Data on single and double infections

Genospecies <sup>a</sup>	Genospecies <sup>b</sup>					
	af	bav	ga	miy	ss	vs
af	788	1	13	20	35	15
	12,277					
bav	6,040	92	0	1	0	10
		84,841				
ga	9,574	NA	290	3	3	106
			41,895			
miy	19,778	45	33,888	8	3	0
				259		
ss	85,034	NA	4,987	7,251	79	1
					9,697	
vs	1,084	47,967	36,970	NA	732	263
						7,969

<sup>a</sup> af, *B. afzelii*; bav, *B. bavariensis*; ga, *B. garinii*; miy, *B. miyamotoi*; ss, *B. burgdorferi sensu stricto*; vs, *B. valaisiana*.

<sup>b</sup> Number of single and double infections (upper right, unshaded area) and the corresponding mean spirochete load (lower left, shaded area) for the six *Borrelia* genospecies (on the diagonal) and the 15 genospecies pairs (off the diagonal) in questing *I. ricinus* nymphs ( $n = 7,400$ ) sampled in Neuchâtel, Switzerland. Some genospecies associations did not occur ( $n = 0$ ), so the mean spirochete load was not available (NA).

distribution to run the log-linear analyses. All statistics were calculated with R for Mac OS X (41).

**(ii) Spirochete load in doubly infected ticks (inhibition and facilitation).** We analyzed the subset of *B. burgdorferi sensu lato*-infected ticks ( $n = 1,731$  infected nymphs – 35 *B. miyamotoi*-infected nymphs = 1,696 *B. burgdorferi sensu lato*-infected nymphs [Table 2]) to test whether the spirochete load of mixed infections deviated from the neutral (additive) expectation. For this analysis, we pooled the results of the seven sampling occasions to maximize our sample size and statistical power. The naive or null hypothesis of additivity assumes that the expected spirochete load in a doubly infected tick ( $\bar{X}_{A \cap B}$ ) is simply the sum of the mean spirochete loads of genospecies A ( $\bar{X}_A$ ) and genospecies B ( $\bar{X}_B$ ) in singly infected ticks (i.e.,  $\bar{X}_{A \cap B} = \bar{X}_A + \bar{X}_B$ ). The two alternative hypotheses are inhibition and facilitation where the spirochete load of doubly infected ticks is less than ( $\bar{X}_{A \cap B} < \bar{X}_A + \bar{X}_B$ ) or greater than ( $\bar{X}_{A \cap B} > \bar{X}_A + \bar{X}_B$ ) the additive expectation.

For each of the 10 *B. burgdorferi sensu lato* genospecies pairs, we calculated the observed mean spirochete load ( $\bar{X}_{A \cap B}$ ) for the sample of coinfecting ticks ( $n_{AB}$ ) (Table 2). To generate the null distribution of additivity, we randomly sampled (with replacement)  $n_{AB}$  pairs of spirochete loads from the set of singly infected ticks for each genospecies in the pair. We summed each randomly sampled pair of spirochete loads to form the sample of joint spirochete loads and then calculated the expected mean spirochete load. We repeated this random sampling protocol 100,000 times to create a null distribution with sufficient precision for calculating *P* values. Thus, for each genospecies pair, the mean and variance of the null distribution were  $\bar{X}_A + \bar{X}_B$  and  $\sigma_A^2 + \sigma_B^2$ , respectively. We used the 2.5th and 97.5th percentiles from the null distribution as the 95% confidence limits (CL) of the mean expected spirochete load. To facilitate comparison among genospecies pairs, we divided the mean expected spirochete load and the 95% CL by the corresponding observed mean spirochete load. Genospecies pairs where the 95% CL of the expected mean/observed mean ratio overlap 1.0 exhibit additivity. Genospecies pairs where the 95% CL of the ratio are above or below 1.0 exhibit inhibition and facilitation, respectively.

**RESULTS**

**Prevalence of *Borrelia* genospecies and spirochete loads.** Among the 7,400 questing nymphs sampled, there were 1,520 single, 211

**TABLE 3** Three-way log-linear analyses testing whether the pairwise co-occurrence between *B. burgdorferi sensu lato* genospecies differed among the seven sampling occasions for each of the 10 genospecies pairs<sup>c</sup>

Genospecies <sup>a</sup>		df	Deviance	<i>P</i> value
A	B			
af	bav	6	4.938	0.5518
af	ga	6	5.486	0.4832
af	ss	6	9.125	0.1667
af	vs	6	20.232	0.0025 <sup>b</sup>
bav	ga	6	<0.001	1.0000
bav	ss	6	<0.001	1.0000
bav	vs	6	1.849	0.9330
ga	ss	6	9.444	0.1501
ga	vs	6	7.750	0.2570
ss	vs	6	2.847	0.8278

<sup>a</sup> af, *B. afzelii*; bav, *B. bavariensis*; ga, *B. garinii*; ss, *B. burgdorferi sensu stricto*; vs, *B. valaisiana*.

<sup>b</sup> Only the af-vs genospecies pair had a significant three-way interaction after Bonferroni correction ( $P = 0.05/10 = 0.005$ ).

<sup>c</sup> Shown are the degrees of freedom, the deviance, and the *P* value testing whether the three-way interaction between sampling occasion, genospecies A, and genospecies B was statistically significant. After Bonferroni correction,  $\alpha/n = 0.05/10 = 0.005$  (where *n* is the number of pairwise comparisons).

double, and 10 triple infections. The triple infections were excluded from our analyses. *B. afzelii* was the most common genospecies, representing 50.4% of single and double infections (872/1,731) (Table 2), followed by *B. garinii* (24.0%; 415/1,731) and *B. valaisiana* (22.8%; 395/1,731). *B. burgdorferi sensu stricto* (7.0%; 121/1,731), *B. bavariensis* (5.4%; 94/1,731), and the relapsing fever-like spirochete *B. miyamotoi* (2.0%; 35/1,731) were less common. Among nymphs infected with a single *Borrelia* genospecies, the rank order of median spirochete load per tick was *B. bavariensis* (23,050), *B. garinii* (5,080), *B. burgdorferi sensu stricto* (3,410), *B. afzelii* (3,140), *B. valaisiana* (1,830), and *B. miyamotoi* (1,160).

**Positive and negative co-occurrence of *B. burgdorferi sensu lato* genospecies.** The three-way log-linear analysis showed that the co-occurrence between *B. burgdorferi sensu lato* genospecies was the same across the seven sampling occasions for 9 of the 10 genospecies pairs (Table 3). This result means that we were justified in pooling the data of the seven sampling occasions and interpreting the global two-way interaction for these genospecies pairs. Of the nine genospecies pairs where pooling was justified, three showed positive co-occurrence and six showed negative co-occurrence (Table 4). Two of the three positive co-occurrences (*B. afzelii*-*B. burgdorferi sensu stricto* and *B. garinii*-*B. valaisiana*) and three of the six negative co-occurrences (*B. afzelii*-*B. bavariensis*, *B. afzelii*-*B. garinii*, and *B. bavariensis*-*B. garinii*) were statistically significant. Pooling was not justified for the *B. afzelii*-*B. valaisiana* genospecies pair (degrees of freedom [df] = 6; deviance = 20.232;  $P = 0.0025$ ) (Table 3), and examination of the seven sampling occasions found six cases of negative co-occurrence (four were statistically significant) and one case of positive co-occurrence (not statistically significant) (data not shown).

**Spirochete load in doubly infected ticks (inhibition and facilitation).** We used simulations to describe the spirochete load of nymphal ticks infected with two *B. burgdorferi sensu lato* genospecies. Specifically, for each genospecies pair we set out to test

**TABLE 4** Two-way log-linear analyses testing whether the pairwise associations between *B. burgdorferi sensu lato* genospecies are positive or negative when the seven independent sampling occasions are combined into a single data set<sup>d</sup>

Genospecies <sup>a</sup>		Deviance	P value	Significance <sup>b</sup>	Interaction term		Co-occurrence type	No. (%) of co-occurrences by study <sup>c</sup>
A	B				Estimate	SE		
af	bav	19.390	0.00001	***	-2.656	1.006	Negative	7 (100)
af	ga	43.425	<0.00001	***	-1.483	0.285	Negative	7 (100)
af	ss	26.730	<0.00001	***	1.158	0.205	Positive	6 (86)
af	vs	31.567	<0.00001	***	-1.238	0.266	Negative	6 (86)
bav	ga	13.254	0.0003	**	-24.78	55500	Negative	7 (100)
bav	ss	3.314	0.0687	NS	-22.640	38540	Negative	7 (100)
bav	vs	2.776	0.0957	NS	0.607	0.338	Positive	5 (71)
ga	ss	2.601	0.1068	NS	-0.829	0.588	Negative	5 (71)
ga	vs	198.560	<0.00001	***	2.049	0.129	Positive	7 (100)
ss	vs	7.395	0.0065	NS	-1.917	1.006	Negative	7 (100)

<sup>a</sup> af, *B. afzelii*; bav, *B. bavariensis*; ga, *B. garinii*; ss, *B. burgdorferi sensu stricto*; vs, *B. valaisiana*.

<sup>b</sup> Level of significance after Bonferroni correction: no significance (NS),  $P > 0.05/10 = 0.005$ ; low significance (\*),  $P = 0.05/10 = 0.005$ ; intermediate significance (\*\*),  $P = 0.01/10 = 0.001$ ; high significance (\*\*\*),  $P = 0.001/10 = 0.0001$ .

<sup>c</sup> Number of times the co-occurrence type was observed out of the seven sampling occasions.

<sup>d</sup> Shown are the deviance, the P value, and the estimate and standard errors (SE) of the two-way interaction term describing the pairwise associations between *B. burgdorferi sensu lato* genospecies. The co-occurrence type column indicates whether the genospecies pair exhibits positive co-occurrence (two-way interaction is positive) or negative co-occurrence (two-way interaction is negative). The by-study column indicates the number of studies where the sign of the two-way interaction was in the same direction (positive or negative) as the overall estimate (estimate column). The three-way log-linear analysis (Table 3) found a significant effect of study on the pairwise association for the *B. afzelii*-*B. valaisiana* genospecies pair (shaded gray).

whether the joint spirochete load observed in doubly infected ticks was unusual relative to null distribution. We generated the null distribution by randomly sampling and then summing the spirochete loads of the singly infected ticks belonging to each member of the genospecies pair. We calculated a null distribution for 8 of the 10 genospecies pairs (Fig. 1). Two genospecies pairs were excluded (*B. bavariensis*-*B. burgdorferi sensu stricto* and *B. bavariensis*-*B. garinii*) because they had no doubly infected ticks. Of the eight remaining genospecies pairs, two pairs (*B. afzelii*-*B. garinii* and *B. afzelii*-*B. valaisiana*) showed inhibition (95% CL of the ratio > 1.0), one pair (*B. afzelii*-*B. burgdorferi sensu stricto*) showed facilitation (95% CL of the ratio < 1.0), and the remaining five pairs showed additivity (Fig. 1 and Table 5). After Bonferroni correction, three of the eight spirochete load interactions remained statistically significant (Table 5). Spirochete loads in nymphs with double infections of *B. afzelii*-*B. garinii* or *B. afzelii*-*B. valaisiana* were 6 to 19 times lower than the additive expectation. In contrast, spirochete loads of double infections of *B. afzelii*-*B. burgdorferi sensu stricto* were four times greater than the additive expectation.

**Relationship between co-occurrence and spirochete load inhibition in doubly infected ticks.** *B. burgdorferi sensu lato* genospecies pairs fell into two broad categories when jointly considering the two types of interactions measured in this study (Fig. 2). Genospecies pairs that had negative co-occurrence had combined spirochete loads that were lower than expected (stronger inhibition), whereas genospecies pairs with positive co-occurrence had spirochete loads that met or surpassed the additive expectation (weaker inhibition or facilitation). Thus, broadly speaking, *B. burgdorferi sensu lato* genospecies pairs either fell in the negative co-occurrence/inhibition or the positive co-occurrence/facilitation quadrant (Fig. 2). Across the set of eight genospecies pairs, there was a significant negative correlation between the two types of interactions (Pearson  $r = -0.851$ ;  $t = -3.97$ ;  $df = 6$ ;  $P = 0.0074$ ).

## DISCUSSION

To our knowledge, this is the first study to test the frequency and spirochete load of mixed *B. burgdorferi sensu lato* infections in the epidemiologically relevant nymphal stage of *I. ricinus* against the background null hypotheses of random species co-occurrence patterns and additive spirochete loads in double infections. The value of our statistical approach was that it allowed us to quantify the degree of co-occurrence and the degree of spirochete load inhibition between co-infection partners. Plotting these measures of co-occurrence and spirochete load inhibition revealed broad patterns of interactions between *B. burgdorferi sensu lato* genospecies pairs (Fig. 2). Common coinfections have higher-than-expected joint spirochete loads inside the nymphal tick (Fig. 2). *B. burgdorferi sensu lato* genospecies that share the same vertebrate reservoir hosts (*B. afzelii*-*B. burgdorferi sensu stricto* and *B. garinii*-*B. valaisiana*) frequently occur together and exhibit weak inhibition and even facilitation with respect to the spirochete load inside the nymphal tick. Conversely, *B. burgdorferi sensu lato* genospecies pairs that are specialized on different vertebrate reservoir hosts (*B. afzelii*-*B. garinii*, *B. afzelii*-*B. valaisiana*, *B. garinii*-*B. burgdorferi sensu stricto*, and *B. burgdorferi sensu stricto*-*B. valaisiana*) rarely occur together and exhibit strong inhibition with respect to spirochete load. This negative association between occurrence and spirochete load inhibition is likely driven by the vertebrate immune system. Specifically, the vertebrate complement system, which is present in the tick midgut (42), is capable of lysing *B. burgdorferi sensu lato* spirochetes that are not adapted to that particular vertebrate host (32, 33). In cases where the complement system fails to eliminate the maladapted coinfection partner, the spirochete load of the latter likely would be much reduced, resulting in the observed pattern of inhibition in the nymphal tick. For example, when the same larva takes multiple blood meals from different vertebrate hosts after interrupted attachments (43, 44), the complement system of the second host could reduce the spirochete

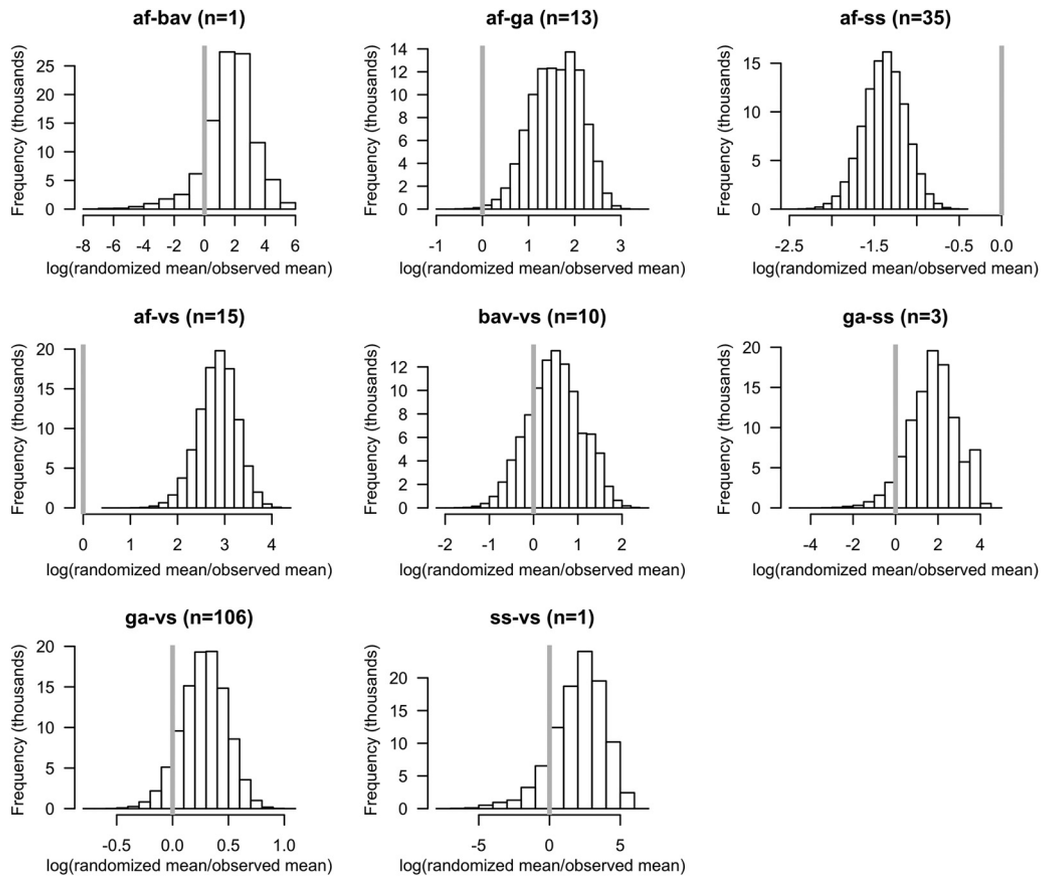


FIG 1 Distribution of the expected *B. burgdorferi sensu lato* spirochete load under the null hypothesis of additivity for eight genospecies pairs. To facilitate visualization of the null distribution, expected values were divided by the observed value and subsequently log transformed. Applying this transformation sets the observed value to zero (vertical gray line). Null distributions centered on zero indicate additivity. Null distributions centered on positive values (right-shifted) indicate inhibition among genospecies pairs. Null distributions centered on negative values (left-shifted) indicate facilitation. Abbreviations: af, *B. afzelii*; bav, *B. bavariensis*; ga, *B. garinii*; ss, *B. burgdorferi sensu stricto*; vs, *B. valaisiana*.

load of the *Borrelia* pathogen from the first host. Thus, the vertebrate immune system is the most likely explanation as to why coinfections involving rodent- and bird-adapted *B. burgdorferi sensu lato* genospecies exhibit lower-than-expected spirochete loads. Our results for the more common genospecies associations correspond to what has previously been reported in the literature (17, 21, 24). Rauter and Hartung (21) reported

that *B. garinii*-*B. valaisiana* was the most common mixed infection of *I. ricinus* ticks, but they did not test whether mixed infections occurred more or less often than the random expectation. Kurtenbach et al. (24) used a simple chi-square test to show that *B. garinii*-*B. valaisiana* and *B. afzelii*-*B. garinii* mixed infections were more and less common than the random expectation in a variety of European countries. However, their

TABLE 5 Results from the randomization protocol testing the null hypothesis of additivity of *B. burgdorferi sensu lato* spirochete load<sup>b</sup>

Genospecies <sup>a</sup>	Genospecies				
	af	bav	ga	ss	vs
af		16.137 (0.08–82.59)	<b>5.668 (1.71–12.47)</b>	<b>0.259 (0.15–0.40)</b>	<b>18.732 (7.39–36.25)</b>
bav	0.2428		NA	NA	1.938 (0.50–4.92)
ga	<b>0.0040</b>	NA		10.350 (0.55–42.63)	1.349 (0.88–1.90)
ss	<b>&lt;0.0002</b>	NA	0.1206		24.117 (0.10–134.43)
vs	<b>&lt;0.0002</b>	0.4444	0.1660	0.2512	

<sup>a</sup> af, *B. afzelii*; bav, *B. bavariensis*; ga, *B. garinii*; ss, *B. burgdorferi sensu stricto*; vs, *B. valaisiana*.

<sup>b</sup> Entries above the diagonal show the ratio of the mean expected spirochete load to the mean observed spirochete load; the 95% confidence limits are in parentheses (the 2.5th and the 97.5th quantiles from the null distribution of expected spirochete loads divided by the observed spirochete load). Ratios with 95% confidence limits that overlap 1.0 indicate additivity, ratios with 95% confidence limits of less than 1.0 indicate facilitation, and ratios with 95% confidence limits greater than 1.0 indicate inhibition. Entries below the diagonal show the proportion of randomized values that were more extreme than the observed mean spirochete load multiplied by 2.0 (two-tailed test). The randomization protocol was not possible for two genospecies pairs, and these are described as not available (NA). For the remaining eight genospecies pairs, we used a Bonferroni-corrected  $\alpha$  level of  $0.05/8 = 0.00625$  to determine the statistical significance (shown in boldface).

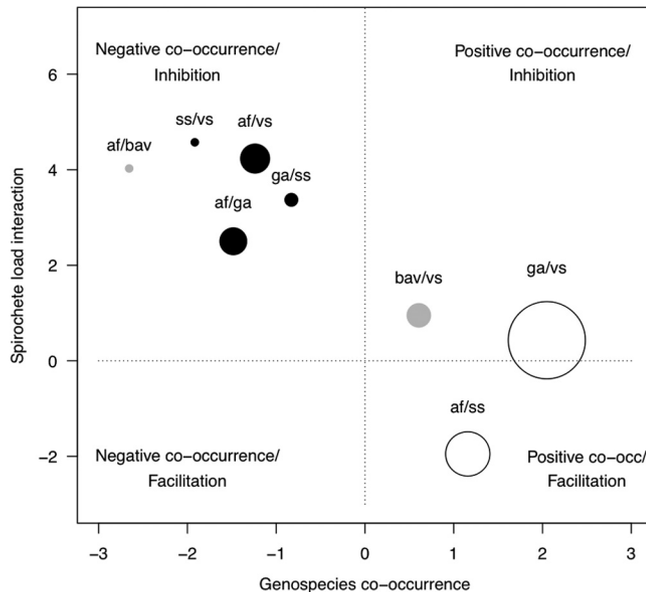


FIG 2 Relationship between our two measures of interaction for the eight pairs of *B. burgdorferi sensu lato* genospecies. Genospecies co-occurrence refers to whether double infections occurred more or less often than expected (positive or negative). The spirochete load interaction refers to whether the total spirochete load in the doubly infected ticks was greater or less than the sum of the single infections (facilitation or inhibition). The circle size refers to the sample size of double infections for each genospecies pair (Table 2). Solid black dots and open dots refer to genospecies pairs for which we had *a priori* predictions of negative and positive co-occurrence, respectively. Solid gray dots refer to genospecies pairs for which we had no *a priori* predictions for the co-occurrence pattern. The dotted lines (0 on *x* and *y* axes) refer to the null hypotheses of independence (i.e., the frequency of double infections equals the product of the frequencies of single infections) and additivity (i.e., spirochete load of double infection equals the sum of the single infections). To facilitate the graphing of the spirochete interaction, we log transformed the ratios shown in Table 5 (negative values indicate ratios of  $<1$  [representing facilitation], whereas positive values indicate ratios of  $>1$  [inhibition]). Abbreviations: af, *B. afzelii*; bav, *B. bavariensis*; ga, *B. garinii*; ss, *B. burgdorferi sensu stricto*; vs, *B. valaisiana*.

study focused on adult ticks, which mostly feed on incompetent hosts such as deer and do not contribute significantly to the epidemiology of Lyme borreliosis.

The specificity of rodent and avian-adapted *Borrelia* genospecies for their respective reservoir hosts is not perfect. Our study found that double infections between rodent and avian-adapted *Borrelia* genospecies do occur (Table 2), and we suggest four mechanisms. First, some *B. burgdorferi sensu lato* strains might be generalists that are capable of infecting both rodent and bird reservoir hosts. For example, a recent field study on the Siberian chipmunk found one double infection with *B. afzelii* and *B. garinii* (45). Second, larvae may acquire a double infection via a combination of systemic and co-feeding transmission (46). Systemic transmission refers to the standard mode where ticks acquire spirochetes that have established a widespread and long-term infection in the host. Co-feeding transmission refers to the process where hosts are not systemically infected but instead function as a temporary bridge (47) that facilitates transmission between infected and uninfected ticks feeding in close proximity to each other (48). Third, double infections may result from a combination of vertical (transovarial) and horizontal (blood meal) transmission of *Borrelia* genospecies, although the former is believed to

be rare or nonexistent in *Ixodes* ticks (49–51). Fourth, larval ticks taking multiple blood meals from different hosts (interrupted blood meals) (44) could also produce double infections of rodent- and bird-specific genospecies (43). These four mechanisms illustrate the diversity of transmission pathways that can produce ticks doubly infected with rodent- and bird-adapted *Borrelia* genospecies. These double infections are of interest because they connect the avian and rodent Lyme borreliosis systems and provide opportunities for genetic exchange between *Borrelia* genospecies.

Importantly, while the prevalence of *B. burgdorferi sensu lato* genospecies often fluctuates through time and space, the nature of the pairwise interaction appeared to be robust. For 6 of 10 genospecies pairs, the nature of the pairwise interaction was always in the same direction for each of the seven sampling occasions (Table 4), despite the fact that the questing nymphs had been collected in different years and seasons and had been exposed to different abiotic conditions prior to *B. burgdorferi sensu lato* genospecies identification (Table 1). Similarly, the fact that the three-way interaction was not statistically significant for 9 of the 10 genospecies pairs (Table 3) indicates that the different experimental conditions of the seven sampling occasions did not bias the co-occurrence patterns of the *B. burgdorferi sensu lato* genospecies. Thus, an important aspect of this study is our demonstration that the genospecies associations appear to be relatively constant over time at our site.

Most of the double infections had spirochete loads that were considerably lower than the additive expectation, suggesting that genospecies compete for limited resources in the tick (i.e., for seven of the eight genospecies pairs shown in Table 5, the average ratio of the mean expected spirochete load to the mean observed spirochete load is greater than 1, suggesting inhibition). The two statistically significant inhibition interactions include the *B. afzelii*-*B. garinii* and the *B. afzelii*-*B. valaisiana* genospecies pairs. As previously discussed, this inhibition was likely caused by the host complement system suppressing the spirochete load of the maladapted coinfection partner. In contrast, the only statistically significant facilitation interaction involved two genospecies, *B. afzelii*-*B. burgdorferi sensu stricto*, that are both adapted to rodent reservoir hosts (28–30, 52). One possible explanation for facilitation is that one genospecies suppresses the host immune system and thereby enhances the infection and transmission success of its coinfection partner.

Our approach of using qPCR to detect *Borrelia* infections followed by RLB to determine the community of *B. burgdorferi sensu lato* genospecies had its advantages and disadvantages. Advantages of our approach include low cost, relative simplicity, and time efficiency. Our qPCR protocol gives reliable estimates of the spirochete load, because the median spirochete load in the present study (3,200 spirochetes/tick across all *Borrelia* genospecies) was similar to that in another study (4,000 spirochetes/tick) on *B. burgdorferi sensu lato* infections in *I. ricinus* ticks (53). Our RLB protocol gives reliable identification of the *B. burgdorferi sensu lato* genospecies and *B. miyamotoi*, because there were only 3 out of 1,731 *Borrelia*-infected ticks (as detected by qPCR) where the PCR product failed to hybridize with a genospecies-specific probe (subsequent sequencing revealed two *B. afzelii* and one *B. garinii* infection). The reverse situation, where the RLB protocol identifies infections in ticks but the qPCR fails to detect them, may also occur (unpublished data). One major advantage of RLB is that the technique allows identification of mixed infections in samples. A disadvantage of our approach using qPCR is that we could not

estimate separate spirochete loads for each partner in the mixed infections. Thus, in the case of inhibition or facilitation, we do not know which of the two *B. burgdorferi sensu lato* genospecies reduced or increased their spirochete load in the mixed infection relative to the single infection. Future studies should use next-generation sequencing approaches that can identify all possible *B. burgdorferi sensu lato* genospecies and estimate genospecies-specific spirochete loads. In addition, experimental infections would greatly clarify the underlying mechanisms of the pairwise interactions observed in the present study.

In addition to *B. burgdorferi sensu lato* genospecies, the RLB identified a low prevalence of the relapsing fever-like spirochete *B. miyamotoi* (2.0%; 35/1,731) in our *I. ricinus* population (Table 2). The identity of this relapsing fever-like spirochete has been confirmed with DNA sequencing of the flagellin gene and the 16S rRNA genes of samples hybridizing with the *B. miyamotoi* RLB probe (23). Screening ticks for *Borrelia* infection using only the RLB protocol (i.e., no upstream qPCR identification of infected ticks) found that between 0.5% (7/1,324) (unpublished data) and 5.3% (5/94) (23) of nymphs collected in the same forest were infected with *B. miyamotoi*, showing that the prevalence rates of *B. miyamotoi* are similar regardless of the protocol. In the present study, double infections with *B. miyamotoi* and rodent-specialized genospecies such as *B. afzelii* ( $n = 20$ ) were more common than double infections with bird-specialized genospecies such as *B. garinii* ( $n = 3$ ) (Table 2). A recent study conducted on field-captured rodents in Switzerland (C. Burri, O. Schumann, C. Schumann, and L. Gern, unpublished data) showed that rodents are relevant reservoir hosts for *B. miyamotoi* and *B. afzelii*, which explains the high number of double infections involving these two genospecies. Alternatively, the co-occurrence of these two genospecies in nymphs may be due to a combination of transovarial transmission of *B. miyamotoi* (54) and horizontal transmission of *B. afzelii*.

In conclusion, the present study found that the co-occurrence and joint spirochete load of rodent- and bird-adapted *B. burgdorferi sensu lato* genospecies in *I. ricinus* nymphs both were lower than the neutral expectation. This observation is consistent with the theory that the vertebrate immune system plays an important role in structuring the *B. burgdorferi sensu lato* genospecies community in the *I. ricinus* tick vector. Conversely, *B. burgdorferi sensu lato* genospecies that are specialized on the same reservoir host co-occurred more often than expected from chance, and their joint spirochete loads followed or exceeded the additive expectation, suggesting facilitation. Future experimental infections will further elucidate the mechanisms shaping the community ecology of *B. burgdorferi sensu lato* genospecies inside the tick vector.

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