# ORIGINAL ARTICLE

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# Host dispersal shapes the population structure of a tick-borne bacterial pathogen

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LEY-MOLECULAR ECOLOGY

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### Abstract

Birds are hosts for several zoonotic pathogens. Because of their high mobility, especially of longdistance migrants, birds can disperse these pathogens, affecting their distribution and phylogeography. We focused on Borrelia burgdorferi sensu lato, which includes the causative agents of Lyme borreliosis, as an example for tick-borne pathogens, to address the role of birds as propagation hosts of zoonotic agents at a large geographical scale. We collected ticks from passerine birds in 11 European countries. B. burgdorferi s.l. prevalence in Ixodes spp. was 37% and increased with latitude. The fieldfare Turdus pilaris and the blackbird T. merula carried ticks with the highest Borrelia prevalence (92 and 58%, respectively), whereas robin Erithacus rubecula ticks were the least infected (3.8%). Borrelia garinii was the most prevalent genospecies (61%), followed by B. valaisiana (24%), B. afzelii (9%), B. turdi (5%) and B. lusitaniae (0.5%). A novel Borrelia genospecies "Candidatus Borrelia aligera" was also detected. Multilocus sequence typing (MLST) analysis of *B. garinii* isolates together with the global collection of B. garinii genotypes obtained from the Borrelia MLST public database revealed that: (a) there was little overlap among genotypes from different continents, (b) there was no geographical structuring within Europe, and (c) there was no evident association pattern detectable among *B. garinii* genotypes from ticks feeding on birds, questing ticks or human isolates. These findings strengthen the hypothesis that the population structure and evolutionary biology of tick-borne pathogens are shaped by their host associations and the movement patterns of these hosts.

#### KEYWORDS

birds, Borrelia garinii, host-parasite interactions, Lyme borreliosis, migration, ticks

# 1 | INTRODUCTION

Wild birds are relevant for public health because of their role in the spread of emerging zoonotic pathogens that cause newly recognised diseases or diseases which are rapidly increasing in incidence or geographical range (Reed, Meece, Henkel, & Shukla, 2003). Some birds act as reservoirs of pathogens such as *Borrelia burgdorferi* sensu lato (s.l.), enterobacteria, flavivirus and influenza A virus, being

significantly involved in the direct infection of humans or arthropod vectors that transmit the disease agents to humans (Thomas, Hunter, & Atkinson, 2007). Wild birds, especially migratory species, may also carry the arthropod vectors (e.g., ticks) to different geographic areas creating new foci of disease (Reed et al., 2003). Studies that monitored tick infestation of birds during migration estimated that birds are responsible for the transport of 6.8–175 million ticks each spring between wintering and breeding areas (Ogden et al., 2008; Olsen,

of movements of its avian hosts and potential for dispersal and consequent spatial mixing of strains. To achieve this, we collected ticks feeding on common passerine species with different ranges of migratory movements in a coordinated study covering a broad geographical area in Europe (11 countries), and assessed the infection status of these ticks with *B. burgdorferi* s.l. The diversity of the common avian-associated *B. garinii* genospecies and potential phylogeographical patterns were determined using a multilocus sequence typing scheme (MLST) of eight house-keeping genes (Margos et al., 2008). Analyses based on these conserved genetic markers have been previously used to estimate and describe the degree of population genetic structure over the largest geographical range studied so far, and have the potential to reveal signatures of demographic processes, dispersal and migration (Hoen et al., 2009; Margos et al., 2012; Vollmer et al., 2013).

# 2 | MATERIALS AND METHODS

#### 2.1 | Birds and ticks

Birds were captured in collaboration with ornithologists and ringers in 11 European countries (Czech Republic, Estonia, Finland, Germany, Greece, Hungary, Netherlands, Portugal, Slovenia, Spain and Sweden) covering an area from 8°23'W to 24°57'E and from 40°35'N to 62°14'N. Capturing effort directed to tick collection was concentrated during the year of 2015, but additional data (including data from the years 2005-2008, 2013-2014 and 2016) was included for five of the 21 study sites (Table S1). These collection sites correspond to ringing stations or sites where bird populations of particular species have been studied in a long-term perspective and with which we could establish a collaborative protocol. Therefore, some European areas are missing from this study due to sampling limitations. Birds were captured using mist-nets or, when breeding in nest boxes: incubating females were captured by hand (in Estonia, in Gotland, Sweden and in Harjavalta, Finland), and both parents were caught when feeding the nestlings using spring or wire traps. Nestlings were sampled in the nest between eight and 15 days of age. Countries, collection sites, range of capturing dates and bird species sampled are detailed in Figure 1 and Table S1. Although classification of collection sites according to country has no biological meaning, for convenience, and because countries are related to geographical positioning, we refer to country of collection when reporting some results for an easy identification of sample origin, for comparability with previous localised studies in different countries, and for an integration with sequence data available in public databases. Birds were carefully inspected for attached ticks with special attention to the head (around the eyes, beak, ears, chin and crown) and neck, where ticks are most often attached. We removed infesting ticks with fine forceps and collected them into tubes containing 70%-99% ethanol according to each individual host. Because we were interested in B. burgdorferi s.l. infection prevalence in ticks feeding on birds, our analyses use ticks as sampling units. Therefore, we did not collect

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data on noninfested birds nor on tick infestation intensity. Our statistical analyses take into account sampling site, bird life cycle stage and month, to account for differences in sampling effort and uneven sample distribution across sites and time of year. The ticks were identified morphologically using identification keys (Estrada-Peña, Bouattour, Camicas, & Walker, 2004; Estrada-Peña, Nava, & Petney, 2014; Pérez-Eid, 2007).

## 2.2 | Molecular analysis

We extracted tick DNA in a subset of *lxodes* spp. ticks (n = 656 ticks; mean ticks ± SE = 65.6 ± 11.5 per country, 38.6 ± 14.28 per bird species, 1.54 ± 0.03 per bird), using a column DNA extraction kit (DNeasy, Qiagen, Hilden, Germany). Tick (nymph and adult) exoskeleton was broken by piercing followed by incubation with proteinase K, for 24 hr. We tested a subsample of these Ixodes spp. (n = 58, mean ticks  $\pm SE = 5.8 \pm 0.79$  randomly selected per country) using a conventional PCR targeting the mitochondrial 16S rRNA gene of ticks using the primers described by (Mangold, Bargues, & Mas-Coma, 1998) and an annealing temperature of 56°C, to confirm morphological identification of ticks by BLASTn search (https ://blast.ncbi.nlm.nih.gov/). For specimens (n = 4) for which the % of identity in BLASTn search was less than 98% we built a Maximum Likelihood phylogenetic tree, together with reference sequences (Chitimia-Dobler et al., 2018; Estrada-Peña et al., 2014) retrieved from Genbank, and confirmed the species identification with the obtained clustering patterns. We assessed infection of ticks by a nested PCR targeting the flaB gene of B. burgdorferi s.l. using the primers described in Johnson, Happ, Mayer, and Piesman (1992), with an annealing temperature of 52°C. We used the Invitrogen PCR Reagent System mix (Life Technologies), according to manufacturer's instructions, and a positive and a negative control were used in all PCR runs. Borrelia genospecies were identified by sequencing. The procedures listed above were performed at the Portuguese National Institute of Health Doutor Ricardo Jorge, Portugal.

A subsample of B. garinii-positive specimens were tested using MLST targeting eight housekeeping genes (clpA, clpX, nifS, pepX, pyrG, recG, rplB, uvrA) according to Margos et al. (2008) at the German National Reference Centre for Borrelia, Bavarian Health and Food Safety Authority, Germany. Those isolates were selected from common bird species (Turdus spp. and hole-nesting birds) that covered the geographical range in which *B. garinii* was detected in our study. All samples were analysed following the same protocol (see https:// pubmlst.org/borrelia/sequencing for protocol and primer information). We compared the obtained sequences with those available in the Borrelia MLST database (https://pubmlst.org/borrelia/) located at the University of Oxford, to obtain allele and sequence type (ST) numbers for each isolate (Jolley & Maiden, 2010). Novel alleles or STs were given novel numbers. Samples that contained mixed infections were discarded from further MLST/MLSA analyses because allele numbers and sequence types could not be identified for these samples. We submitted all data to the Borrelia MLST database.



**FIGURE 1** Map of countries and sampling locations where birds were screened for infesting ticks. Sampling locations closer than 40 km apart are represented under the same pin and are numbered according to study site listed in Table S1. Light grey pins, sites where birds were screened for ticks but no ticks were found; dark grey pins, sites where ticks were collected feeding on birds. Details on sampling locations, range of bird capturing dates, bird species, number of bird individuals infested and number of *lxodes* spp. ticks collected at each location are detailed in Table S1. Adapted from the Cartographic Research Laboratory of the University of Alabama

A novel *Borrelia* genospecies was detected and characterised by PCR amplification of the 5S-23S rRNA intergenic spacer (Rijpkema, Molkenboer, Schouls, Jongejan, & Schellekens, 1995), and subsequent sequencing of PCR amplicons of the 16S rRNA locus (Radulović, Milutinovic, Tomanovic, & Mulenga, 2010) and of the *clpX* gene (Margos et al., 2008).

# 2.3 | Statistical analyses

Factors affecting the prevalence of *B. burgdorferi* s.l. in ticks from birds (each tick feeding on a bird was used as a sampling unit) were tested using a Generalized Linear Mixed Model (GLMM) with a binomial error distribution (logit function). The full model included bird species (10 levels), latitude, longitude, tick stage (larvae or other), tick species (four levels), bird life cycle stage (breeding or

nonbreeding) and month (12 levels) as fixed effects. Bird identity (bird\_ID, to control for ticks tested from the same individual), nested within collection site, was included as a random effect. We ran a set of models with different fixed effects structures and used the Akaike Information Criterion (AIC) to select the best model (best model selection table is presented in Table S2). After ranking the models using AIC, we used those with a  $\Delta$ AIC < 2 with respect to the top model to calculate conditional model-averaged parameter estimates. Significance level was defined at *p* = .05.

In these models, only bird species for which we had information on *Borrelia* infection for at least eight ticks (from different individuals) were used. Tick stage was considered in statistical analyses and was divided into larvae and other tick stages because nymphs represented the majority whilst adults were too infrequently found to justify their own group for statistical analysis. This procedure was adopted because transovarial transmission (i.e., the acquisition of Borrelia by larvae via vertical transmission from the parent) is considered to be very low in *Ixodes* spp. (Eisen & Lane, 2002). For this reason, an infection of a larva is highly unlikely if they were collected from an uninfected or incompetent reservoir bird host. On the other hand, nymphs/adults may have acquired an infection during a previous blood meal on an infected host. In such case Borrelia DNA may be detected in the tick independent of the bird hosts they were collected from. Tick species was also included in the models because within the genus Ixodes, different species differ in their vector competence for B. burgdorferi s.l. (Eisen & Lane, 2002; Heylen, Krawczyk, et al., 2017; Heylen, Sprong, et al., 2014). We controlled for the effects of timing of tick collection throughout the year including month of collection as explanatory variable and also by grouping those ticks collected during the birds' breeding season (April-July) and those collected outside the breeding season (August-March), because breeding is one of the most stressful periods in the birds' life cycle and the stress associated with breeding duties may suppress their immune system potentially leading to spirochetaemia. Therefore, this could affect the probability of infection of the ticks feeding on the birds. We used the same statistical approach to test factors affecting prevalence of infection by the most prevalent Borrelia genospecies in our study (B. garinii). The models were run in R (R Core Team, 2013) using the packages LME4, LMERTEST, BBMLE, MUMIN and ARM (Bates, Mächler, Bolker, & Walker, 2015; Kuznetsova, Brockhoff, & Christensen, 2017).

# 2.4 | Multilocus sequence typing and multilocus sequence analysis (MLST/MLSA)

In order to study the population structure and the phylogenetic relationships of *B. garinii* in a global context, we selected a subset of 82 *B. garinii*- positive samples, as to include isolates from all European countries sampled in this study and represent all bird species sampled with *B. garinii*- infected ticks ( $n \ge 5$  infected ticks), and tested those by MLST.

Complete allelic profiles of *B. garinii*- positive specimens obtained in our study were analysed with goeBURST analysis using Phyloviz (Francisco, Bugalho, Ramirez, & Carriço, 2009) together with other *B. garinii* genotypes detected worldwide (identification of the isolates used for MLST/MLSA is given in Table S3; we included each ST only once for each country). Relationships among STs were evaluated through triple locus variants (TLV), and founder clonal complexes were identified to infer patterns of descent.

We estimated nucleotide diversity ( $\pi$ ; Nei, 1987) and Tajima *D* (Tajima, 1989) for each gene using R packages PEGAS v. 0.9 (Paradis, 2010) and APE v. 3.5 (Paradis, Claude, & Strimmer, 2004) on each continent for a sample set including 304 isolate sequences (198 from Europe, 85 from Asia and 21 from North America), and on each country (n = 11) for which more than five isolates were available (see Table S3 for identification of the isolates included in this analysis). The sequences for gene *clpX* were realigned using MAFFT v7.205

(Katoh & Standley, 2013) as there was a deletion of three bases in some isolates.

An ancestry recombination graph for the 85 STs present in Europe was reconstructed with BEAST2 software v. 2.5 (Bouckaert et al., 2019) and package BACTER v. 2.2 (Didelot, Lawson, Darling, & Falush, 2010; Vaughan et al., 2017) using sequences of the eight housekeeping genes. BEAST2 was run three times with a unique tree and substitution model for the eight loci but with a lognormal-relaxed clock model for each locus. We used the following priors: HKY substitution model (Hasegawa, Kishino, & Yano, 1985), Gamma site heterogeneity model with four gamma categories, Tree prior: Coalescent Constant Populations. Due to the high number of STs and the complex model including recombination, the chain was slow to converge (as was shown by Tracer v. 1.6 (Rambaut, Suchard, Xie, & Drummond, 2014) and we thus extended the original 10M states to 17M for one run and 19M for the other two. A consensus tree was reconstructed with the tool ACGAnnotator present in the BACTER package after removing 40%-70% burnin depending on the run.

### 3 | RESULTS

#### 3.1 | Ticks collected from birds

In total, 2,308 ticks were collected from 843 infested birds, belonging to 28 bird species (Table S1). Ticks collected from these birds belonged to three genera: Haemaphysalis (n = 3), Hyalomma (n = 48) and *lxodes* (n = 2,255). Two ticks could not be identified to genus by morphological criteria as they were damaged. We identified four species of Ixodes: I. ricinus (n = 1,779), I. arboricola (n = 214), I. frontalis (n = 164), and I. ventalloi (n = 24) but 74 Ixodes ticks could not be identified to species because they lacked critical body structures needed for morphological identification (Table S4). The vast majority of collected ticks were immatures (2,175 out of 2,255 Ixodes spp.), and from these, 63% were nymphs. Adults belonged to I. arboricola (n = 63), I. frontalis (n = 11), I. ricinus (n = 1) and I. ventalloi (n = 4). Amplification and sequencing of the ribosomal 16S rRNA gene of ticks (Mangold et al., 1998) confirmed tick morphological identification in 84% of the cases, corresponding to a misidentification rate of 16%.

The blackbird Turdus merula, the song thrush T. philomelos, the redwing T. iliacus, the great tit Parus major, the collared-flycatcher Ficedula albicollis, and the Eurasian jay Garrulus glandarius presented coinfestations by ticks of different species (Table S4).

# 3.2 | Prevalence of *Borrelia burgdorferi* s.l. in ticks collected from birds

Out of 656 *Ixodes* ticks collected from birds and analysed for *B. burg-dorferi* s.l. infection, 244 (37.2%) were positive. Of these, 22 were larvae (prevalence of *B. burgdorferi* s.l. in larvae = 20%, 22/110), and

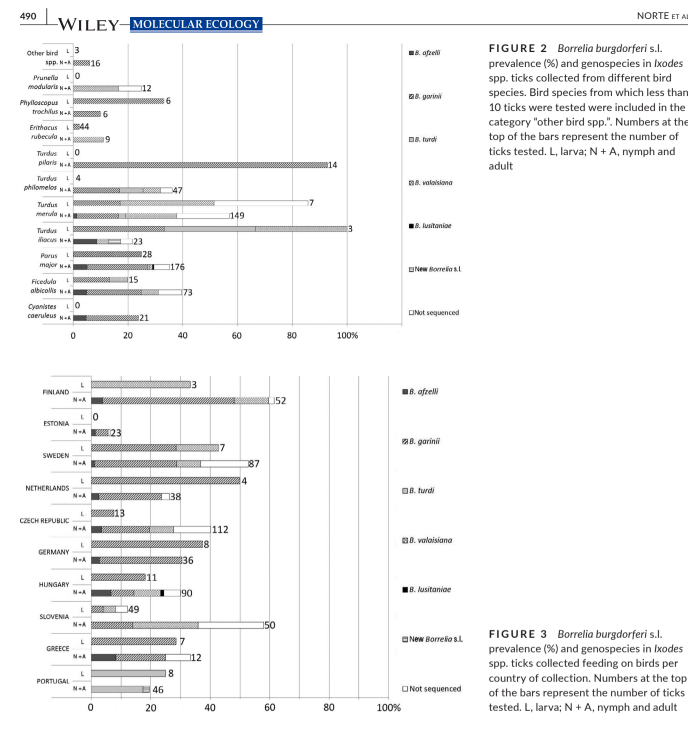


FIGURE 2 Borrelia burgdorferi s.l. prevalence (%) and genospecies in Ixodes spp. ticks collected from different bird species. Bird species from which less than 10 ticks were tested were included in the category "other bird spp.". Numbers at the top of the bars represent the number of ticks tested. L, larva; N + A, nymph and adult

214 were nymphs (prevalence of *B. burgdorferi* s.l. in nymphs = 41%, 214/521). Ixodes ricinus was the most infected tick species (40.2%, 210/522), followed by I. ventalloi (31.3%, 5/16), I. arboricola (29.7%, 14/47) and I. frontalis (20.5%, 9/44).

Borrelia burgdorferi s.l. prevalence differed significantly between ticks collected from different bird species and was affected by latitude ( $\chi_{10.616}$  = 90.10, p < .0001; Figures 2 and 3). Longitude, tick stage, tick species, month and the birds' life cycle stage did not affect B. burgdorferi s.l. prevalence. The model selection table is presented in Table S2 and the conditional model averaged coefficient parameters obtained from the generalized linear mixed models (GLMMs) that best explained the prevalence of B. burgdorferi s.l. in ticks collected from birds are presented in Table 1. In comparison with the

reference species (the blue tit Cyanistes caeruleus), ticks collected from T. merula (estimate  $\pm$  SE = 2.50  $\pm$  0.91, z = 2.76, p = .006) and *Turdus pilaris* (estimate  $\pm$  SE = 4.29  $\pm$  1.51, z = 2.84, p = .005) showed higher infection rates, whereas those collected from the robin E. rubecula had the lowest infection rates (estimate  $\pm$  SE = -2.41  $\pm$  1.25, z = -1.93, p = .054). Borrelia burgdorferi s.l. prevalence increased with latitude (estimate  $\pm$  SE = 0.08  $\pm$  0.03, z = 2.28, p = .022). The fieldfare T. pilaris was the bird species that carried ticks with the highest Borrelia prevalence (92%), followed by the blackbird T. merula (58%). Only two out of 53 (3.8%) ticks feeding on the robin E. rubecula were positive for B. burgdorferi s.l. (Figure 2).

The genospecies of 193 positive samples was identified by sequencing the flaB gene. The most prevalent genospecies was B. garinii **TABLE 1** Conditional model averagedcoefficient parameters from thegeneralized linear mixed models (GLMMs)that best explained (lowest AICc, $\Delta$ AICc < 2, see Table S2) the prevalence</td>of Borrelia burgdorferi s.l. in ticks collectedfrom birds

Conditional model averaged coefficients Parameter Estimate SE Adjusted SE Z value p-value -5.72 2.07 2.07 2.76 .0058 Intercept 1.25 -2.41 1.25 1.93 .054 Bird\_species\_E. rubecula Bird\_species\_F. albicollis 1.27 0.88 0.88 1.44 .15 Bird species P. major 1.12 0.82 0.83 1.35 .18 1.24 Bird\_species\_P. trochilus 1.24 0.09 .93 0.11 Bird species P. modularis -0.30 1.27 1.27 0.24 .81 Bird\_species\_T. iliacus 0.89 1.05 1.05 0.84 40 Bird\_species\_T. merula 2.50 0.91 0.91 2.76 .006 0.95 Bird\_species\_T. philomelos 1.11 0.95 1.17 .24 4.29 1.51 1.51 2.84 .0046 Bird\_species\_T. pilaris Tick species I. frontalis 0.47 1.03 1.03 0.45 .65 Tick\_species\_I. ricinus 0.70 0.71 .096 1.18 1.66 0.78 Tick\_species\_I. ventalloi 0.96 1.22 1.22 .44 0.079 0.03 0.03 2.28 .022 Latitude Latitude \* Longitude 0.0008 0.0005 0.0005 1.56 .12

UK (n = 33) Germany (n = 26)Russia (n = 26) Latvia (n = 21)Japan (n = 16) Canada (n = 12)Finland (n = 12)France (n = 11)Norway (n = 11)China (n = 7)Sweden (n = 6)Czech Republic (n = 4)Italy (n = 4)Slovenia (n = 3)Netherlands (n = 2)Serbia (n = 2)Austria (n = 1)Estonia (n = 1)Hungary (n = 1)Mongolia (n = 1)Yugoslavia (n = 1)ST detected in tick from a terrestrial bird V ST detected in a tick from a marine bird

**FIGURE 4** Borrelia garinii STs distribution between countries in Europe (blue/green), Asia (red) and North America (purple). GoeBURST analysis included 137 STs, using TLV as parameter. Sixteen major clonal complexes and four minor clonal complexes (consisting of only two associated STs) were found. A total of 33 isolates formed singletons. A clonal founder was identified in five out of the 16 major clonal complexes (ST86, ST88, ST184, ST244 and ST251), in red. N refers to the number of *B. garinii* isolates used in the analyses

(60.7%, 116/191), followed by *B. valaisiana* (23.6%, 45/191), *B. afzelii* (9.4%, 18/191) and *B. turdi* (5.2%, 10/191). *B. lusitaniae* (0.5%, 1/191) and a novel genospecies (0.5%, 1/191) were also detected (Figures 2 and 3; Table S5).

The most abundant genospecies associated with *T. pilaris*, *T. philomelos*, *P. major* and *F. albicollis* was *B. garinii* with a prevalence varying between 100% (13/13) in *T. pilaris* and 53% (8/15) in *T. philomelos* (Figure 2), whereas *B. valaisiana* was the most prevalent genospecies

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detected in ticks collected from *T. merula* (50%, 30/60; Figure 2). The model explaining the variation in *B. garinii* prevalence, the most common genospecies detected in ticks feeding on birds, was identical to the one explaining *B. burgdorferi* s.l. prevalence, with the exception that the prevalence of *B. garinii* in ticks feeding on *T. merula* and *E. rubecula* was not significantly different from the reference bird species. Under the assumption that there was no cofeeding transmission (i.e., when larvae acquire the infection due to feeding in close proximity to other infected tick stages; Randolph, Gern, & Nuttall, 1996), data on larval infection suggested that *F. albicollis* and *T. merula* may act as reservoirs for *B. garinii* and *B. valaisiana*, *T. iliacus* for *B. garinii*, *B. valaisiana* and *B. turdi*, *P. major* for *B. garinii*, and the willow warbler *P. trochilus* and *E. rubecula* for *B. garinii* (Figure 2; Table S5).

Borrelia afzelii DNA was detected only in nymphs, mostly feeding on *P. major*, and to a lesser extent on other bird species (*C. caeruleus*, *F. albicollis*, *T. iliacus* and *T. merula*). Borrelia lusitaniae DNA was detected in one *I. ricinus* nymph feeding on *P. major*. Borrelia turdi DNA was detected in all stages of two tick species, *I. frontalis* (66.7%; six out of nine positive ticks), and *I. ventalloi* (80%; four out of five positive ticks), feeding on *T. iliacus*, *T. merula*, *T. philomelos* and *P. major*.

DNA of the new *Borrelia* genospecies (*Candidatus* Borrelia aligera) was detected in an *I. ventalloi* nymph feeding on a *T. iliacus* in Portugal. Its *flaB* sequence was 100% identical to a *flaB* sequence previously detected in one *I. ricinus* nymph feeding on a Sardinian warbler *Sylvia melanocephala* in Portugal (isolate T794A; accession number KT207789; Norte et al., 2015). The PCR targeting the 5S-23S rRNA intergenic spacer was positive showing that this genospecies belongs to the *B. burgdorferi* s.l. group. Its 16S rRNA sequence showed only 97% similarity to several *B. burgdorferi* genospecies, including *B. bissettiae* and *B. mayonii*. The sequence of the housekeeping gene *clpX* showed 36 nucleotide differences from all previously detected alleles available at the MLST database (http://pubmlst.org/borrelia). Detailed information on specimens from which different *Borrelia* genospecies were detected in this study is presented in Table S5.

# 3.3 | Multilocus sequence typing/multilocus sequence analysis (MLST/MLSA)

Twenty-nine complete allelic profiles with sequences for all eight genes were obtained from a subset of 82 *B. garinii*- positive samples selected as to include isolates from all European countries sampled in this study. Some samples (n = 25) were excluded because they represented *B. garinii* mixed infections, and, therefore, allelic profiles could not be determined. These complete 29 profiles represented all countries from which *B. garinii* was detected in ticks from birds in this study, apart from Greece, for which we did not obtain any complete profiles. Comparison of alleles from an incomplete ST (i.e., not obtaining sequences for all alleles) from a tick feeding on a bird in Greece showed that they were identical to samples previously reported from the UK.

The 29 *B. garinii* allelic profiles were resolved into 20 STs, nine of which were new. These sequence data were supplemented with

sequences of *B. garinii* isolates available at the MLST database (see Table S3 for identification of isolates included in this analysis) and used for goeBURST (n = 172; Figure 4) and phylogenetic analyses (n = 110; Figure 5).

At a global scale, out of the 201 *B. garinii* isolates (137 STs) analysed (downloaded from the MLST database and our own data), 2% (three STs: ST244, ST86 and ST575) were found in more than one continent, 21% (29/137) were found in more than one country and 9.5% (13/137) were found in three or more countries. When a ST was detected in more than one country, those countries were generally distant (i.e., did not border each other, 96.6%). Eleven STs (out of the 20) found in ticks feeding on birds, and typed during the course of this study, were detected in more than one continent and four widespread STs (detected in five to nine countries; Figure 4). Among these 20 STs described as part of this study, two were shared between migrant bird species and species with both resident and mixed populations.

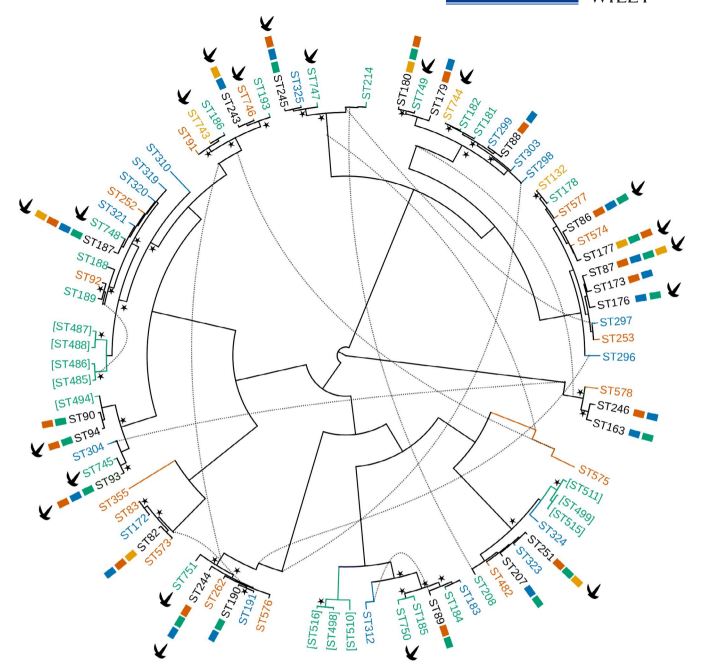
In the goeBURST analysis of the global collection (137 STs), using TLV as parameter, 16 major clonal complexes and four minor clonal complexes (consisting of only two associated STs) were found. A total of 33 isolates formed singletons (Figure 4). In five out of the 16 major clonal complexes, a clonal founder could be identified: those were ST86, ST88, ST184, ST244 and ST251. The goe-BURST analysis further revealed that STs from different continents belonged to different clonal complexes, with only a few exceptions: seven out of the 137 STs were shared between continents, or clustered together (e.g., ST364, ST694; Figure 4). Two of the STs found in more than one continent were also detected in ticks from European birds investigated during the course of this study.

Focusing on European STs, the pattern of clonal complexes was not related to geographical distribution and there was no evidence that STs from different countries formed separate clonal complexes, except for four STs detected only in Norway (ST487, ST488, ST498 and ST516; Tveten, 2013) (Figure S6, marked with an \*). There was also no indication of clustering according to bird species (Figure S6).

STs detected in ticks from birds worldwide did not cluster tightly as clonal complex but were distributed amongst clonal complexes, including those from migrant and birds which have both resident and mixed populations (Figure S7).

The averaged nucleotide diversity of *B. garinii* for all eight genes together was of the same order for the three continents and all countries (ranging from 0.007 to 0.010), except Norway and Sweden which presented the lowest nucleotide diversity ( $\pi$  = 0.005; Table 2). Tajima's *D* was close to zero for most countries, showing that there is no specific sign of selection or expansion in these genes. However, the population in Norway showed a comparatively high Tajima's *D* of 1.085, which could be a sign of a bottleneck, being in agreement with the low genetic diversity observed in this population.

At the European scale, *B. garinii* showed no spatial structuring in goeBURST analysis (Figure S6): STs for which more than one isolate has been obtained (e.g., ST86, ST187, ST251, ST94, ST82) were not



**FIGURE 5** Ancestry Recombination Graph of 85 European sequence types reconstructed with BEAST2 v. 2.5 and package BACTER v. 2.2. Labels are coloured by geographic origin: green, Northern Europe (Estonia, Finland, Latvia, Norway and Sweden); orange, Eastern Europe (Czech Republic, Hungary, Serbia, Slovenia and Yugoslavia); red, Central Europe (France, Germany, Italy, Netherlands); blue, British islands (United Kingdom). Branches leading to taxa found in one geographic region only show the corresponding colour. Black labels refer to STs present in several geographic regions indicated in coloured rectangles and corresponding in coloration to the regions defined (North, East, Central Europe and British islands). STs isolated from birds (this study) are marked with a bird next to the label name. Dashed lines show recombination events present in at least 50% of all posterior graphs and stars mark high confidence nodes (present in at least 80% of all posterior graphs)

regionally restricted, but originated from distant countries such as UK, Latvia, Slovenia, Hungary or Austria.

This is also what we observed on the ancestry recombination graph reconstructed using sequences of the 85 European STs and generated in BEAST2 (Bouckaert et al., 2019; Figure 5). This method was used because we suspected recombination between housekeeping loci. Indeed, 12 occurrences of recombination (dashed lines) were identified in at least 50% of the sampled graphs in the phylogeny showing that recombination does occur but that there is a global clonal frame. These recombination events concerned six out of the eight loci. The ancestry recombination graph shows no evident geographical clustering for four main European regions (Northern Europe - Estonia, Finland, Latvia, Norway and Sweden; Eastern Europe - Czech Republic, Hungary, Serbia, Slovenia and II FY-MOLECULAR ECOLOGY

**TABLE 2** Nucleotide diversity ( $\pi$ ) and Tajima's *D* averaged over the eight MLST genes (*clpA*, *clpX*, *pepX*, *pyrG*, *nifS*, *recG*, *rplB*, *uvrA*) of *Borrelia garinii* strains (strain IDs included in this analysis are available in Table S3)

Continent   Europe 227 0.008 -0.222   Asia 85 0.009 -1.184   North America 21 0.008 0.617   Country Canada 21 0.008 0.617   China 8 0.009 0.022   Finland 12 0.007 0.157	N strains	Population	ins Mean $\pi$	Mean Tajima's D
Asia 85 0.009 -1.184   North America 21 0.008 0.617   Country Canada 21 0.008 0.617   China 8 0.009 0.022   Finland 12 0.007 0.157		Continent		
North America 21 0.008 0.617   Country Canada 21 0.008 0.617   China 8 0.009 0.022   Finland 12 0.007 0.157	227	Europe	0.008	-0.222
Country Canada 21 0.008 0.617   China 8 0.009 0.022   Finland 12 0.007 0.157	85	Asia	0.009	-1.184
Canada210.0080.617China80.0090.022Finland120.0070.157	21	North America	0.008	0.617
China 8 0.009 0.022   Finland 12 0.007 0.157		Country		
Finland 12 0.007 0.157	21	Canada	0.008	0.617
	8	China	0.009	0.022
F 40 0.007 0.000	12	Finland	0.007	0.157
France 18 0.007 -0.090	18	France	0.007	-0.090
Germany 55 0.007 -0.326	55	Germany	0.007	-0.326
Japan 21 0.010 -0.865	21	Japan	0.010	-0.865
Latvia 30 0.010 -0.192	30	Latvia	0.010	-0.192
Norway 16 0.005 1.085	16	Norway	0.005	1.085
Russia 54 0.008 -1.026	54	Russia	0.008	-1.026
Sweden 6 0.005 -0.205	6	Sweden	0.005	-0.205
UK 70 0.009 0.219	70	UK	0.009	0.219

Yugoslavia; Central Europe -France, Germany, Italy, Netherlands; and British islands - United Kingdom) except for the 11 STs present in Norway (marked with square brackets on Figure 5) that cluster into three monophyletic groups and one isolated ST. Out of the 85 STs present in Europe, 25% (n = 22) were present in at least two of the geographical regions defined (Northern, Eastern, Central Europe and the British islands). *Borrelia garinii* STs can thus disperse very far at the continental scale. STs detected in birds (this study) were dispersed among other STs isolated from ticks or humans.

## 4 | DISCUSSION

In this study *I. ricinus* was the most abundant tick collected from common passerine birds across a large geographical area in Europe. Overall, *B. burgdorferi* s.l prevalence in ticks collected from birds was 37%. Thrushes (*Turdus* spp.) were the most important carriers of infected *Ixodes* spp., supporting the notion that different bird species contribute differently to *B. burgdorferi* s.l. genospecies complex maintenance and dispersal. Our genetic characterization of the most prevalent genospecies detected in ticks feeding on birds, *B. garinii*, showed that this tick-borne pathogen presents little overlap of STs among continents, but no geographical population structuring was detected in Europe, or according to isolation source (bird-derived ticks or questing ticks/human isolates). Taken together this provides supporting evidence that birds act as important reservoirs for *B. garinii* and are a main source of infection of this genospecies to ticks and ultimately humans (through the bite of an infected tick). Given the

importance of birds as main hosts of this tick-borne pathogen, they have the potential to modulate its phylogeography by homogenising the distribution of STs within the European continental range through dispersal and migratory movements. Studying the different factors in action driving this complex host-vector-parasite system is important for a full understanding of *B. burgdorferi* s.l. enzootic cycle and potentially other (not only tick- borne) bird-associated zoonotic pathogens.

Ixodes ricinus is a generalist tick and birds are known to be important hosts for its immature stages (Norte et al., 2012; Santos-Silva et al., 2011). The other tick species (I. frontalis, I. arboricola and I. ventalloi) and genera (Haemaphysalis and Hyalomma) collected from birds in this study have also been previously reported on birds (Diakou et al., 2016; Norte et al., 2012; Pérez-Eid, 2007) and differ in vector competence for Borrelia (Eisen & Lane, 2002; Heylen, Krawczyk, et al., 2017; Heylen, Sprong, et al., 2014). Some bird species such as hole-nesting birds (P. major and F. albicollis), T. merula and T. philomelos were hosts for different tick species; however, the opportunities for cofeeding transmission of B. burgdorferi s.l. between different tick species are limited by spatial and temporal tick species distribution. The misidentification rate of ticks based on morphological features in our study was similar to that reported for the genus Ixodes (14%; Estrada-Peña et al., 2017). Although I. persulcatus occurs in part of the geographic range included in our study (e.g., Finland and Estonia; ECDC, 2018; Laaksonen et al., 2017), and its morphological distinction from I. ricinus in immature stages is difficult, none of the tested samples were identified as I. persulcatus by 16 rRNA sequencing.

Overall, the prevalence of B. burgdorferi s.l. (37%) was within the range reported for ticks collected from hosts in The Netherlands and Belgium (34%; Heylen, Fonville, et al., 2017). However, it was higher than in ticks collected from birds migrating through Italy (30.7%; Toma et al., 2014) and Sweden (26.6%; Olsen et al., 1995), birds from central Europe (25.8%-28%; Dubska, Literak, Kocianova, Taragelova, & Sychra, 2009; Hanincova et al., 2003; Taragel'ova et al., 2008), Germany (25%; Kipp, Goedecke, Dorn, Wilske, & Fingerle, 2006), Poland (13.3%; Michalik, Wodecka, Skoracki, Sikora, & Stanczak, 2008), Switzerland (19.6%-22.5%; Lommano, Bertaiola, Dupasquier, & Gern, 2012; Poupon et al., 2006), Spain (9.2%; Palomar et al., 2016) and Portugal (7.3%; Norte et al., 2015). We cannot exclude the possibility that this may be related to different methodologies used for B. burgdorferi s.l. detection in different studies (real-time PCR vs.conventional PCR and target genes). Our results, using the same detection method across samples from different geographical origins revealed that prevalence of B. burgdorferi s.l. in ticks from birds varied significantly according to latitude. The fact that prevalence increased with latitude is in accordance with Scandinavian countries such as Finland and Sweden showing relatively high prevalence when compared with other European countries, as reported in previous studies on Lyme borreliosis incidence and infection loads of questing ticks (Hubalek, 2009; Rauter & Hartung, 2005; Wilhelmsson et al., 2013). Meta-analyses for Europe revealed an overall prevalence of 12.3%-13.7% in questing ticks, higher in central Europe and Sweden (Rauter & Hartung, 2005; Strnad, Hönig, Růžek, Grubhoffer, & Rego,

rinii, B. afzelii and B. burgdorferi s.s.

2017), but with a significant increase with longitude, rather than latitude, as in our study (Rauter & Hartung, 2005). Nonetheless, it is also known that Lyme borreliosis presents a focal pattern of distribution, determined by the heterogeneous spatial distribution of vector ticks, and also that the north-south gradient has a greater influence on disease incidence at its distributional range limits (Hubalek, 2009). A heterogeneous geographical distribution of *Borrelia* genospecies was also reported by Olsen et al. (1995). These authors showed that *Borrelia* infections in ticks collected from birds arriving to Sweden from the south or southeast in spring were mainly caused by *B. garinii*, whereas the genospecies distribution was more heterogeneous in ticks from birds coming from the southwest, and included *B. ga* 

The higher prevalence of B. burgdorferi s.l. in ticks removed from birds than that reported from questing ticks is in accordance with birds acting as reservoirs for some Borrelia genospecies and transmitting the infection to feeding ticks (Heylen, Matthysen, Fonville, & Sprong, 2014; Humair, Postic, Wallich, & Gern, 1998; Kurtenbach, Carey, Hoodless, Nuttall, & Randolph, 1998; Norte, Lopes de Carvalho, Núncio, Ramos, & Gern, 2013). Additionally, Borrelia starts dividing in feeding ticks and may be more readily detected by PCR (Schwan & Piesman, 2002). Our study revealed a nonhomogeneous distribution of Borrelia among bird species in bird-derived ticks. Thus, our data corroborate previous reports that not all bird species contribute equally to the Borrelia enzootic cycle, as it is also known for different mammal species (LoGiudice, Ostfeld, Schmidt, & Keesing, 2003; Talleklint & Jaenson, 1994), and suggested by studies including different lizard species (Norte et al., 2014; Szekeres, Majláthová, Majláth, & Földvári, 2016). Contributing factors may not only be different infestation rates with vector ticks, which may be related to foraging behaviour and consequent probability of exposure (Norte et al., 2012), but also to other intrinsic factors related to host competence, including the host's adaptive and innate immune system (Kurtenbach et al., 2006). Turdus species have been identified as main reservoirs for B. garinii and B. valaisana in Europe (Dubska et al., 2011; Mannelli et al., 2005; Michalik et al., 2008) and Asia (Miyamoto & Masuzawa, 2002). In addition, T. merula has also been proven as competent reservoir for B. turdi through xenodiagnosis (Heylen, Krawczyk, et al., 2017; Humair et al., 1998; Norte, Lopes de Carvalho, et al., 2013). In an experimental setup in which migratory restlessness was induced, latent B. garinii infections were re-activated in T. iliacus (Gylfe, Bergstrom, Lunstrom, & Olsen, 2000). Although B. burgdorferi s.l. infected ticks, including larvae, have occasionally been collected from E. rubecula (this study; Poupon et al., 2006), our results suggest that this bird species, although often infested by vector ticks (Norte et al., 2012), plays a minor role in B. burgdorferi s.l. enzootic cycle because of the very low prevalence of B. burgdorferi s.l. in its ticks.

Ticks associated with hole-nesting birds such as *P. major* and *F. albicollis* presented infection rates of 33.8%-36.4%. The most prevalent genospecies was also *B. garinii*, which made up 64%-77.9% of infections in these bird species. *Parus major* has been

– MOLECULAR ECOLOGY – WIL

shown experimentally to selectively amplify B. garinii and B. valaisiana, whereas B. afzelii prevalence in moulted adult ticks that fed as nymphs on this bird species tended to decrease in successive infestations of the birds with wild questing nymphs (Heylen, Matthysen, et al., 2014). The finding of this mammal-associated genospecies in attached ticks derived from birds has been suggested to result from previously acquired infection from another (mammal) host because these spirochetes were found to be unviable by culturing (Heylen, Sprong, et al., 2017). In our study, all B. afzelii- positive ticks were nymphs and we cannot comment on birds' reservoir competence for B. afzelii with these findings because PCR does not allow distinguishing between viable and nonviable bacteria. We cannot rule out that nymphs acquired the infection during a previous blood meal as larvae from a mammalian host, or that larvae were infected via transovarial transmission (Bellet-Edimo, Betschart, & Gern, 2005), because larvae of I, ricinus have been shown to transmit B. afzelii and B. miyamotoi to vertebrate hosts (van Duijvendijk et al., 2016). In Europe, the role of transovarial transmission for different tick and Borrelia species has not been resolved (Bellet-Edimo et al., 2005; Eisen & Lane, 2002; Humair & Gern, 2000; van Duijvendijk et al., 2016). Thus, the role of birds in B. afzelii transmission needs to be further scrutinized.

Borrelia turdi, originally described in Japan (Fukunaga et al., 1996), has been increasingly detected in Europe, often in association with the ornithophilic tick I. frontalis and its bird hosts (Heylen, Tijsse, Fonville, Matthysen, & Sprong, 2013; Norte et al., 2015). In this study, it was detected only in Portugal, although it is known to be present in Spain, Belgium and Norway (Hasle, Bjune, Midthjell, Røed, & Leinaas, 2011; Heylen et al., 2013; Palomar et al., 2016). In our study it has been detected in I. frontalis and I. ventalloi only, which are host- specialised tick species (to birds and rabbits, respectively; Hillyard, 1996). Ixodes frontalis has been proven to be a competent vector for B. turdi (Heylen, Krawczyk, et al., 2017) but vector competence of I. ventalloi remains unknown. This Borrelia genospecies may have been overlooked in the past in questing ticks such as I. ricinus probably because of its low prevalence (Heylen, Krawczyk, et al., 2017). Because B. turdi prevalence in our study was relatively low, we were unable to evaluate statistically its bird host and vector species' associations. Furthermore, the small sample size for tick species other than I. ricinus, may have hampered the detection of significant associations between Borrelia and tick species, to infer tick vector competence. Such relationships may be better evaluated in experimental transmission studies (Heylen, Fonville, et al., 2017; Heylen, Sprong, et al., 2014).

Besides the avian-associated genospecies *B. valaisiana*, *B. garinii* and *B. turdi*, we also detected DNA of a new *Borrelia* genospecies that has not been previously described. Although PCR amplification and sequencing of 16S rRNA, *flaB* and *clpX* was possible and clearly indicated the genetic distinction of the isolate from other *Borrelia* species, it is conceivable that its genetic dissimilarity precluded a deeper characterisation involving other house-keeping genes (which could not be amplified). This finding adds to the growing evidence of the diversity of genospecies in circulation in cryptic cycles in bird WII FY-MOLECULAR ECOLOGY

hosts. Specificities of reservoir host and/or vector competence may explain why this novel *Borrelia* sp. genospecies was not detected before.

Borrelia lusitaniae, a genospecies whose main reservoirs are lizards (De Sousa et al., 2012; Dsouli et al., 2006; Norte et al., 2014), has been occasionally detected in ticks feeding on birds, including larvae (Poupon et al., 2006). In our study, only one tick feeding on a bird was positive for *B. lusitaniae*. However, the paucity of these findings suggests that birds, at most, have a minor role as reservoirs for this genospecies. These infections could be the result of a previous incomplete blood meal on a lizard, transovarial or cofeeding transmission. Surveys in endemic areas in Italy and Portugal in which hundreds of bird-derived ticks were tested revealed no *B. lusitaniae* positive specimens and thus, did not provide evidence that birds may serve as reservoir hosts for *B. lusitaniae* (Amore et al., 2007; Norte et al., 2014; Norte, Ramos, Gern, Núncio, & Lopes de Carvalho, 2013).

Focusing on the genetic diversity and geographical distribution of the most prevalent genospecies detected in ticks feeding on birds, the avian-associated B. garinii, we found that its STs clustered according to continent showing some spatial structuring at this very wide geographical scale. However, there was one ST shared between Europe and Asia, one ST shared between Europe and North America, and one ST shared between Europe, Asia and North America providing evidence of overlap among distant areas at a global scale. One would expect that finding identical STs on continents separated by the Atlantic would be less likely than that between adjacent continents whose geographical barriers may be easily crossed by migrating birds. The movement of long-distance migratory birds, such as seabirds, which can travel thousands of miles and between hemispheres may be responsible for the spread of some B. garinii STs to distant geographical regions. Borrelia garinii is known to circulate in a marine cycle involving the ornithophilic tick I. uriae that infests seabirds at their colonies (Comstedt, Jakobsson, & Bergström, 2011; Gómez-Díaz et al., 2011). Migratory shorebirds such as the blacktailed godwit Limosa limosa, the common redshank Tringa totanus, and the little stint Calidris minuta were also reported to carry B. garinii (Lopes de Carvalho et al., 2012). To this point, B. garinii isolates sharing the same *flaB* sequence have been found in both Campbell Island (New Zealand), the Crozet Islands, and in the northern hemisphere (Egg and St. Lazaria Islands, USA; Comstedt et al., 2011). In our study, two of the transcontinental B. garinii STs were indeed found in ticks feeding on birds. One of these (ST244) was found in I. uriae on a Canadian island (Munro et al., 2017), in guesting I. ricinus in Europe and I. persulcatus in Russia (https://pubmlst.org/borrelia/), in human isolates in Germany (https://pubmlst.org/borrelia/), and in Ixodes spp. feeding in terrestrial birds in Finland and Germany (this study). Some passerine birds (e.g., the northern wheatear Oenanthe oenanthe) can also perform long distance migrations across the Atlantic (Bairlein et al., 2012). An overlap and exchange of strains between the marine and terrestrial cycles is, therefore, likely, as suggested by previous studies (Comstedt et al., 2011; Gómez-Díaz et al., 2011). However, Gómez-Díaz et al. (2011) reported a population division of B. garinii from seabirds between the Atlantic and Pacific basins. These researchers did not use the same MLST as employed in our study, thus, immediate comparison of the results is not possible.

When evaluating European B. garinii STs only, no pattern of geographical clustering was noticeable in our analysis, probably due to B. burgdorferi s.l./or ticks' dispersal promoted by the birds. Similarly, although seabird species show high fidelity to breeding colonies (Schreiber & Burger, 2001), and their main tick I. uriae occurs in seabird populations with strong host species associations (McCoy et al., 2005), no geographic structuring was observed in *B. garinii* within the Atlantic and Pacific Oceans (Gómez-Díaz et al., 2011). In contrast, the mammal-associated B. afzelii STs were shown to have much less geographical overlap in studies which compared geographical patterns and population structure of the avian-associated B. garinii and this mammal-associated genospecies (B. afzelii), using the same MLST scheme as this study (Vollmer et al., 2011, 2013). Their results illustrated that B. garinii showed higher spatial mixing than B. afzelii but that B. garinii presented population differentiation over a large geographical scale (Europe and China). However, Vollmer et al. (2013) included fewer strains from a smaller geographical range in Europe and China.

Although, in general, no overall apparent structure was found for European strains of *B. garinii*, some Norwegian samples were divergent. This could be due to a relative isolation of the study area in Norway, located in the northwest of the country (Tveten, 2013), or a recent invasion event, which would explain the lower diversity of these strains. Recent invasion would be consistent with the reported recent expansion of *I. ricinus* tick populations to northern latitudes in Norway (Gray, Dautel, Estrada-Peña, Kahl, & Lindgren, 2009), which could have caused a population bottleneck. This may also explain the evidence for selection and expansion on MLST genes revealed by the relatively high Tajima's *D* in this *B. garinii* population.

The uniform distribution of B. garinii STs among ticks collected from various bird species, and other sources (e.g., questing ticks), does not suggest specialization of certain B. garinii STs to certain hosts, contrary to the hypothesis of multiple niche polymorphism associated with OspC variation (Brisson, Drecktrah, Eggers, & Samuels, 2012) found for B. burgdorferi sensu stricto (Brisson & Dykhuizen, 2004; Vuong et al., 2014), but not for B. afzelii (Raberg et al., 2017). Our results are consistent with birds being the main reservoir hosts of B. garinii: they maintain its natural transmission cycle and are the source of infection for questing vector ticks. The lack of clustering of B. garinii STs regarding country of origin or isolation source at a finer scale (i.e., Europe), was also supported by the ancestry recombination graph. The clustering pattern between goeBURST (Figure S6) and that of the ancestry recombination graph was generally similar with only a few exceptions showing recombination among strains that could also be promoted by the avian-associated dispersal, which may increase chances of encounter between different strain types and mixing of strains.

We should acknowledge that for migrant bird species and those which have both resident and short-distance migrant populations, one cannot be completely confident that the *B. burgdorferi* s.l. infections which the bird-infesting ticks carried were acquired in the geographical area where the birds were captured. Birds (or their ticks) may have acquired the bacteria in a different area where they remained or stopped-over during migration. This may bias prevalence estimates and sequence type origin classification according to geographical location.

The results presented in this study demonstrate how B. burgdorferi s.l.- vector- host associations and the behaviour of hosts may shape and impact the spread and dispersal, and ultimately the evolutionary biology of B. burgdorferi s.l., used here as a model for tick-borne pathogens. Our data which includes B. garinii MLST characterization from bird-derived ticks from the largest geographical range investigated so far substantiates that bird migration and dispersal movements appear to be one of the main driving forces to shape B. garinii populations, one of the most genetically heterogeneous Lyme borreliosis- causing genospecies (Jacquot et al., 2014; Margos et al., 2008). Because birds are highly mobile and the main reservoir hosts not only for B. garinii, but also for other pathogens, they contribute to frequent, fast and long-range spatial mixing of strains and populations. Our study underlines that understanding pathogen variability and spatial distribution, and consequent modulation of transmission rates and evolution of new variants, is essential to understand disease risk.

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#### AUTHOR CONTRIBUTIONS

A.C.N., G.M., J.A.R., M.S.N., V.F. and I.L.C. designed the study; A.C.N., P.A., H.A., E.B., R.B., L.C., T.C., A.D., N.J.D., B.D., A.D., T.E., B.F., T.G., M.H., D.H., S.H., S.K., D.K., F.K., I.L., R.M., L.M., J.M., M.M., J.M.N., M.N., J.J.S., L.P.S., H.S., I.-S.T., J.T., T.T., Z.T. and M.E.V. carried out the fieldwork and provided samples for this study; A.C.N. and I.L.C. performed laboratory analyses; A.C.N., G.M., P.M.A. and N.S.B. carried out statistic and phylogenetic analyses; A.C.N., G.M., N.S.B., J.A.R. and I.L.C. wrote the manuscript.

#### DATA AVAILABILITY STATEMENT

Borrelia sp. "Candidatus Borrelia aligera" 16S rRNA and *clpX* gene partial sequences obtained in this study have been deposited in GenBank with the accession numbers MH068784 and MH157920, respectively. Borrelia garinii MLST sequences have been deposited in Borrelia MLST database (https://pubmlst.org/borrelia/) with the isolate id numbers 2451 to 2479.

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**II FY-MOLECULAR ECOLOGY** 

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# SUPPORTING INFORMATION

Additional supporting information may be found online in the Supporting Information section.

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