Heterogeneity in the abundance and distribution of *Ixodes ricinus* and *Borrelia burgdorferi* (*sensu lato*) in Scotland: implications for risk prediction

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**Abstract**

**Background:** Cases of Lyme borreliosis, a vector-borne zoonosis caused by bacteria in the *Borrelia burgdorferi* (*sensu lato*) species group, have increased in recent years in Europe. Knowledge of environmental factors associated with abundance of the tick vector *Ixodes ricinus* and the pathogen *B. burgdorferi* (*s.l.*) is of interest to understand responses to environmental changes, predict variation in risk and to inform management interventions.

**Methods:** Nineteen woodland sites across Scotland were surveyed in 2012 for *B. burgdorferi* (*s.l.*) infection in questing *I. ricinus* nymphs (*n* = 200 per site), deer abundance and vegetation. Climatic factors were extracted for each site. Six additional sites were surveyed for questing nymphs in both 2012 and 2013 (*n* = 200 per site and year) to test for variation in *B. burgdorferi* (*s.l.*) prevalence between years.

**Results:** The mean prevalence of *B. burgdorferi* (*s.l.*) across 19 sites was 1.7% (95% CI: 1.4–2.2%; range 0–6%), all four genospecies known to be present in the UK were detected: *B. garinii*, *B. afzelii*, *B. burgdorferi* (*sensu stricto*) and *B. valaisiana*. A higher prevalence of *B. burgdorferi* (*s.l.*), higher densities of nymphs and higher densities of infected nymphs were found at sites with warmer climates, estimated with growing degree-days. No association between infection prevalence in nymphs and woodland type (semi-natural mixed vs coniferous) or deer density was found. At six sites sampled in 2012 and 2013, there was a significant increase in *B. afzelii* prevalence at two sites and a decrease in *B. garinii* prevalence at one site.

**Conclusions:** This study highlights challenges for the prediction of risk of Lyme borreliosis, reflecting the sensitivity of both pathogen and vector ecology to habitat, host and climatic factors. Significant changes in the prevalence of individual genospecies at sites monitored across time are likely to be due to variability in the host community composition between years. Our results indicate the importance of monitoring dynamic variables such as reservoir host populations as well as climate and habitat factors over multiple years, to identify environmental factors associated with Lyme borreliosis risk.

**Keywords:** *Borrelia burgdorferi*, *Ixodes ricinus*, Risk prediction, Spatial heterogeneity, Host community

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Background

Vector-borne diseases represent a global public health burden and cause millions of cases of disease annually [1]. A number of vector-borne diseases are emerging, either from introductions into new regions or due to changing patterns of endemic disease [2]. Lyme borreliosis is an example of an emerging, endemic vector-borne disease, with an increasing geographical distribution in recent years and increased numbers of cases in many parts of Europe including Scotland [3–5]. Between 2004 and 2014, there has been a six-fold increase in the number of Lyme borreliosis cases recorded in Scotland [5]. Therefore, identifying environmental risk factors for the tick vector *Ixodes ricinus* and the bacterial pathogen *Borrelia burgdorferi* (sensu lato) is of importance.

Lyme borreliosis is caused by infection with spirochaete bacteria of the *B. burgdorferi* (s.l.) species complex. This pathogen complex is maintained amongst a large number of vertebrate hosts and ixodid ticks. There are at least 19 known genospecies of *B. burgdorferi* (s.l.) [6, 7]. In Scotland, four genospecies have been detected in questing ticks [8, 9]: *B. afzelii*, *B. garinii*, *B. burgdorferi* (sensu stricto) and *B. valaisiana*. All of these genospecies have been associated with cases of disease in humans [10, 11] although *B. afzelii* and *B. garinii* are most frequently associated with clinical disease in Europe [12]. Different genospecies of *B. burgdorferi* (s.l.) are associated with different clinical presentations in people [13, 14], for example *B. afzelii* is more frequently associated with dermal manifestations of disease, while *B. garinii* is more frequently detected in cases of neuroborreliosis [14]. Therefore, finer scale knowledge of the geographical distribution and abundance of each genospecies is potentially useful for public health messaging and clinical diagnosis.

Species of the *B. burgdorferi* (s.l.) complex infect a wide range of vertebrate reservoir hosts. Some genospecies are host generalists, while others specialise on particular taxonomic groups [15]. For example, *B. afzelii* is transmitted mainly by rodent reservoir hosts [16] and *B. garinii* and *B. valaisiana* are transmitted mainly by birds [17, 18] while *B. burgdorferi* (s.s.) is a generalist genospecies able to infect both birds and mammals [19, 20]. Trans-stadial infection of immature stages of ixodid ticks feeding on competent reservoir hosts is considered to be the most significant mode of transmission [21]. Co-feeding transmission between an uninfected tick feeding close to an infected tick is thought to be less significant epidemiologically [22–24]. Deer are generally considered not to transmit *B. burgdorferi* (s.l.) [25, 26] and deer sera has been found to be borreliacidal [27]; however some studies suggest potential for co-feeding transmission [28, 29]. Deer provide blood meals to ticks and are important in many areas as tick reproduction hosts, feeding reproductively active adult female ticks [30].

In western Europe, the main tick vector of *B. burgdorferi* (s.l.) is the exophilic, three-host tick *Ixodes ricinus*. Temperature, humidity and a combination of these (the saturation deficit) are the primary factors in determining conditions suitable for host-seeking behaviour [31, 32], while the community composition and abundance of hosts for blood meals are strong determinants of vector abundance and *B. burgdorferi* (s.l.) infection rates [33, 34]. Each of the three tick life-stages, larva, nymph and adult, takes a blood meal. In a diverse host community, different proportions of each life-stage feed on different types of host. Larvae tend to feed on all members of the host community including small mammals and birds, nymphs tend to feed mainly on medium-sized vertebrates and larger hosts, and adult ticks tend to feed on large mammal hosts such as deer and sheep (also termed ‘tick reproduction hosts’) [30].

A small number of surveys of *I. ricinus* populations and *B. burgdorferi* (s.l.) in the UK have indicated broad scale patterns in genospecies distribution. Bird-associated genospecies *B. garinii* and *B. valaisiana* appear to dominate in England [6, 35, 36], with some reports of rodent-associated *B. afzelii* [37, 38]. In Scotland, *B. afzelii* appears to be the most common genospecies [8]. There is some evidence for spatial restriction in the distribution of *B. burgdorferi* (s.s.) in Scotland, as this genospecies has been found mainly in the northeast region of the country, based on one previous large-scale study of 25 sites [8]. This survey which selected sites associated with cases of Lyme borreliosis, also found a lower prevalence of *B. burgdorferi* (s.l.) in *I. ricinus* nymphs collected at higher altitudes [8] and a higher prevalence of infection associated with semi-natural mixed woodlands compared to coniferous forest [8, 9]. A study from northern England also found a higher prevalence of infection in *I. ricinus* nymphs sampled from semi-natural mixed woodlands compared to coniferous forests [36]. It has been suggested that the higher prevalence of infection detected in semi-natural mixed woodlands may be due to higher densities of competent reservoir hosts. In particular, there may be higher densities of small mammal hosts such as wood mice (*Apodemus sylvaticus*), bank voles (*Myodes glareolus*) and shrews (*Sorex spp.*), as well as ground-foraging birds.

The prevalence of *B. burgdorferi* (s.l.) and the risk of Lyme borreliosis to humans (defined as the density of infected ticks) are expected to be affected by the density of competent reservoir hosts such as rodents and birds, as well as incompetent hosts such as deer which host ticks but do not infect them with the pathogen. In comparison to data on climate and habitat, it is more challenging to collect data on the vertebrate host community, particularly during cross-sectional studies of
several sites. As described above, habitat proxies have been used in previous studies to assume differences in the relative abundance of reservoir hosts (rodents and birds) between semi-natural mixed woodlands and conifer plantations. However, reservoir host populations can vary between years within habitat types, particularly rodent populations which can vary significantly in abundance between years depending on environmental conditions. Correlative studies using such proxies may have limited use if the number of infected ticks and/or the genospecies composition changes significantly over time. Therefore, studies which measure the variation in B. burgdorferi (s.l.) prevalence across time are needed.

In this study we aimed to: (i) Identify environmental variables associated with the prevalence of B. burgdorferi (s.l.) in questing I. ricinus nymphs and the density of B. burgdorferi (s.l.) infected nymphs at sites across Scotland. As individual genospecies may have different environmental associations (due to different reservoir hosts) we also tested for environmental associations with the most commonly detected genospecies B. afzelii and B. garinii; (ii) Compare these results to a previous study from Scotland which selected sites associated with known cases of Lyme borreliosis [8]; (iii) Test for significant variation in the prevalence of genospecies of B. burgdorferi (s.l.) between years at a subset of sites; and (iv) Identify environmental variables associated with the abundance of questing I. ricinus nymphs at sites across Scotland.

Methods

Field site selection

To estimate the prevalence of B. burgdorferi (s.l.) and the abundance of I. ricinus, 19 woodland sites were chosen across Scotland (Fig. 1, Table 1). Sites were selected to broaden the geographical area sampled from a previous survey in Scotland in 2007–2008 [8] in order to test for spatial trends in B. burgdorferi (s.l.) prevalence and genospecies composition. Sites were chosen to be from the same basic habitat types and altitudes as a previous survey in Scotland [8], and were composed of semi-natural mixed (n = 11) and coniferous woodland (n = 8). Sites were sampled between April and July in 2012.

To investigate variability in B. burgdorferi (s.l.) prevalence and genospecies composition across time, a further six sites were selected, which were a subset of sites sampled in the previous survey [8]. These sites had been sampled between March and October one or more times in 2007 and/or 2008 and were chosen to include three sites with relatively high prevalence (mean 9.7%, range 7–14%) and three sites with relatively low prevalence (mean 1.7%, range 1–2%). For the purposes of this study, we classified these as “high prevalence” and “low prevalence” sites. These sites were re-sampled in the current study in May and June in 2012 and 2013.

Questing tick sampling

Sampling involved slowly dragging a 1 m² white blanket across the surface of the vegetation for 10 m [39]. This method samples a portion of the questing tick population, and can be used to obtain an index of questing nymph relative abundance [39]. Nymphs were recorded for statistical analysis as this life-stage is the most important for transmitting B. burgdorferi (s.l.) to humans [40]. Twenty standardized 10 m long blanket drags were carried out at each site, each drag separated by at least 50 m. Sampling was carried out between 10:00 and 16:00 h during dry conditions or at least two hours after heavy rainfall so as not to wet the blanket. Sites were sampled once per year.

Host and environmental predictors of tick abundance and B. burgdorferi (s.l.) prevalence

Site and drag specific environmental data were recorded. Briefly, at the start of each standardized drag, the GPS location was recorded with a handheld Garmin etrex vista Hcx unit (Garmin, Schaffhausen, Switzerland) and the ground temperature and humidity measured with a RS 1360A Humidity and Temperature Meter (RS Components Ltd, Corby, UK). The mean saturation deficit (a measure of the drying power of the environment) was calculated for each site [41]. The presence or absence of deer dung was recorded in a 1 m² area at the beginning and end of each 10 m blanket drag [39] and averaged by the total number of drags to create an index of deer abundance for each site [39]. The ground vegetation type and height were recorded at the start of each drag. The dominant vegetation on each drag was recorded and categorised as (i) grasses and herbaceous species; (ii) ericaceous and vaccinium species; (iii) moss species; and (iv) bracken and ferns [8]. After carrying out the 20 standardised drags at each site, additional blanket drags were added to collect a minimum of 200 nymphs per site to estimate the prevalence of B. burgdorferi (s.l.)

A number of climatic variables that could affect tick development rates or survival were extracted from GIS datasets for each site. These variables included growing degree-days (day by day sum of the number of degrees the mean temperature is greater than 4 °C, over the summer months, from long term average climate data, 1971–2000), the number of ground frost days and average annual precipitation. Climate data were derived from UK Met Office Long Term Average climate data [42]. The site locations were uploaded to ArcGIS 9.3.1 (ESRI, California, USA, 2009) and raster and polygon data files
sampled at each point location using the Intersect Point tool in Hawth’s Tools [43].

**DNA extraction, B. burgdorferi (s.l.) detection and genospecies determination**

The DNA from individual nymphs was extracted using an ammonia hydroxide technique [44]. B. burgdorferi (s.l.) was detected using a real time PCR as previously described [45]. Genospecies determination was by reverse line blotting [46, 47] and/or sequencing of the 5S-23S rRNA intergenic spacer region (IGS) [48]. Each trimmed IGS sequence was subjected to a basic local alignment search tool analysis (BLAST) against the National Centre for Biotechnology (NCBI) Nucleotide database. Then, each sequence was examined for polymorphisms within the IGS region which discriminate between genospecies [49].

**Statistical analysis**

**Prevalence of B. burgdorferi (s.l.) in questing nymphs and the density of infected nymphs**

Using data collected from the cross sectional survey of 18 sites in 2012 (Fig. 1, Table 1; site MF was excluded as no climatic data were available), B. burgdorferi (s.l.) infection in individual nymphs (infected or uninfected) at each site was modelled using a logit link and binomial distributed errors as a function of the following site level host and environmental factors: woodland type (seminatural mixed or coniferous), growing degree-days, nymph abundance (mean nymphs/10 m²) and the index of deer abundance based on deer dung, with an observation level random effect [50, 51]. An observation level random effect was incorporated to account for overdispersion; variation which is not partitioned into explanatory variables or other random effects is incorporated.
into this variable. As individual genospecies may have different environmental associations (due to different reservoir host associations), the described analysis was repeated individually for the two dominant genospecies *B. afzelii* and *B. garinii*.

The number of *B. burgdorferi* (s.l.) infected nymphs at each site was modelled with a log link and Poisson distributed errors and the following explanatory variables: woodland type (semi-natural mixed or coniferous), growing degree-days, an index of deer abundance based on dung counts and an observation level random effect. In addition, an offset variable was included in the model to account for the different area blanket dragged at each site which converted the outcome variable to the number of infected nymphs per m² (density of infected nymphs). The total area blanket dragged to collect 200 nymphs was estimated by calculating the mean density of nymphs per m² from the 20 standardized 10 m drags and using this to estimate the total area dragged at each site.

Data collected in the survey of six sites which were sampled in 2007/8, 2012 and 2013 were used to test whether there was a significant difference in *B. burgdorferi* (s.l.) prevalence between years. As the total number of nymphs tested in 2007/8 was not available, we tested for differences in *B. burgdorferi* (s.l.) prevalence at sites sampled in 2012 and 2013. *Borrelia burgdorferi* (s.l.) prevalence (infection in individual nymphs, infected or uninfected) was modelled using a logit link and binomially distributed errors with the year of sampling as an explanatory variable, and random effects of sampling site and observation. The model was re-run to test for differences in *B. afzelii* prevalence and *B. garinii* prevalence between 2012 and 2013. Other genospecies and mixed infections were present too infrequently to model separately.

The prevalence of each genospecies at individual sites and the 95% confidence interval were compared between 2012 and 2013 to see if there were significant genospecies differences at the site level between years.

### Table 1

| Site | Year | Woodland type | Mean altitude (m) | Growing degree-days | Deer dung index 2012 (2013) |
|------|------|---------------|------------------|---------------------|-----------------------------|
| SW   | 2012 | Coniferous    | 70               | 134,134             | 0.45                        |
| GL   | 2012 | Deciduous     | 85               | 89,549              | 0.15                        |
| MCP  | 2012 | Deciduous     | 118              | 125,045             | 0.05                        |
| LK   | 2012 | Deciduous     | 300              | 91,168              | 0.05                        |
| HG   | 2012 | Coniferous    | 133              | 130,502             | 0.00                        |
| BR   | 2012 | Coniferous    | 89               | 104,415             | 0.10                        |
| CA   | 2012 | Deciduous     | 117              | 122,919             | 0.00                        |
| CG   | 2012 | Deciduous     | 262              | 100,233             | 0.00                        |
| CL   | 2012 | Coniferous    | 293              | 105,822             | 0.05                        |
| FF   | 2012 | Deciduous     | 47               | 141,376             | 0.00                        |
| GA   | 2012 | Coniferous    | 243              | 95,511              | 0.00                        |
| GF   | 2012 | Coniferous    | 130              | 146,246             | 0.00                        |
| IW   | 2012 | Deciduous     | 92               | 135,780             | 0.00                        |
| KN   | 2012 | Coniferous    | 75               | 115,821             | 0.10                        |
| LS   | 2012 | Deciduous     | 15               | 142,322             | 0.00                        |
| PC   | 2012 | Coniferous    | 250              | 76,429              | 0.05                        |
| PV   | 2012 | Deciduous     | 60               | 122,324             | 0.05                        |
| CB   | 2012 | Deciduous     | 30               | 122,197             | 0.20                        |
| MF   | 2012 | Deciduous     | 125              | na                  | 0.00                        |
| COM  | 2012/3| Coniferous    | 154              | 112,953             | 0.00 (0.00)                 |
| CR   | 2012/3| Deciduous    | 231              | 96,798              | 0.05 (0.10)                 |
| DR   | 2012/3| Deciduous    | 82               | 94,397              | 0.00 (0.00)                 |
| TB   | 2012/3| Deciduous    | 50               | 127,111             | 0.00 (0.08)                 |
| WB   | 2012/3| Coniferous    | 200              | 105,889             | 0.00 (0.13)                 |
| FZ   | 2012/3| Coniferous    | 170              | 89,892              | 0.05 (0.00)                 |
**Questing nymph abundance**

Using data collected from the cross sectional survey of 18 sites in 2012 (Fig. 1, Table 1; site MF was excluded as no climatic data were available), nymph abundance (counts of nymphs per 10 m drag) was modelled using a log link and Poisson distributed errors, as a function of the following explanatory variables: ground vegetation type (at the drag level), the mean saturation deficit on the day of sampling, an index of deer abundance based on deer dung, woodland type (seminatural mixed or coniferous), growing degree-days, average annual precipitation, average number of days of ground frost (all at the site level). The model also included an interaction term between latitude and longitude of the site to test for spatial trends in questing nymph abundance. Random effects of site and at the observation level (individual drag) were included to control for inter-site variation in abundance and overdispersion, respectively [50, 51]. The ratio of residual deviance to residual degrees of freedom was calculated in the R package RVAideMemoire [52].

**Model selection**

All statistical analyses were carried out in R version 3.1 (R Development Core Team, Vienna, Austria) using the lme4 package [53] for generalised linear mixed models (GLMMs). Confidence intervals were calculated using the exact binomial confidence interval in R. For all models, a maximal global model was initially fitted, which included all potential explanatory variables and specified interactions. A backward stepwise model selection was carried out based on the model’s Akaike's Information Criterion (AIC) and variables were dropped sequentially based on their effect on the model AIC [54]. The model with the lowest AIC was selected. Initial data exploration revealed that altitude was positively correlated with growing degree-days and negatively correlated with the average number of days of ground frost. To avoid collinearity problems [55], each of these variables was included in the full model separately, in turn and the parameter associated with the best-fit model was selected for inclusion, based on the AIC as described above. For all models the AIC was corrected for small sample size (AICc) in the R package AICcmodavg [56].

**Results**

The overall prevalence of *B. burgdorferi* (s.l) from the 19 sites sampled in 2012 was 1.7% (95% CI: 1.3–2.2%; range 0–6%; 66 infected ticks out of 3800 tested). Five sites had no *B. burgdorferi* (s.l) infected nymphs detected (Fig. 2, Additional file 1: Table S1). Of the infected nymphs across all 19 sites 45.5% of infections were *B. afzelii*, 28.8% were *B. garinii*, 7.6% were *B. valaisiana*, 9.1% were *B. burgdorferi* (s.s) and 6.1% were mixed *Borrelia* genospecies infections where two or more genospecies were detected in the same nymph. Mixed genospecies infections were composed of three *B. garinii* and *B. afzelii* infections and one combined *B. afzelii* and *B. burgdorferi* (s.s) infection. The prevalence and genospecies composition of infected ticks at sites sampled in this study in 2012 and the previous study from Scotland [8] are shown in Fig. 3.

The best-fit model for *B. burgdorferi* (s.l) prevalence included a positive effect of growing degree-days (delta AICc = 15.1) and a negative effect of nymph abundance (delta AICc = 7.1) (Table 2). The best-fit model for the density of infected nymphs also included a positive effect of growing degree-days (delta AICc = 11.9) as did the best-fit model for *B. afzelii* prevalence (delta AICc = 3.3). The intercept model was the best-fit model for *B. garinii* prevalence. Woodland type (seminatural mixed or coniferous) and the index of deer abundance based on deer dung were not supported as significant predictors in any of the best-fit models (Table 2).
At the six sites sampled in multiple years, (1200 nymphs tested in each year), the prevalence at ‘low’ and ‘high’ prevalence sites varied through time (Fig. 4, Additional file 2: Table S2). The best-fit model for *B. burgdorferi* (s.l.), *B. afzelii* and *B. garinii* prevalence at the six sites sampled in 2012 and 2013 was the intercept model, year of sampling was not supported in any of these models (Additional file 3: Table S3). There was a non-significant tendency for an increased prevalence of *B. afzelii* in 2013 compared to 2012 (delta AICc = 1.6, *P* = 0.06, Additional file 3: Table S3). At the site level, there were significant increases in *B. afzelii* prevalence at two of the six sites, ‘COM’ and ‘CR’ and a significant decrease in the prevalence of *B. garinii* at site DR between 2012 and 2013 (Fig. 4).

The mean number of nymphs per 10 m² blanket drag at each site ranged between 0.6–11.5 nymphs (Additional file 1: Table S1). The best-fit model for nymph abundance included growing degree-days, and the ground vegetation type. Inclusion of an observation level random effect improved the model fit and absorbed overdispersion in the data [50, 51].

### Table 2

Results from the final selected generalised linear mixed model to explain variation in *B. burgdorferi* (s.l.) prevalence in questing nymphs, the density of *B. burgdorferi* (s.l.) infected nymphs and *B. afzelii* and *B. garinii* prevalence in questing nymphs

| Model description | Fixed effects | Mean (estimated) | SE | *P*-value | Delta AICc |
|-------------------|---------------|------------------|----|-----------|------------|
| *B. burgdorferi* (s.l.) | (Intercept) | -9.7 | 1.5 | 9.9 × 10^{-11} | – |
|  | Growing degree-days | 0.5 | 0.1 | 1.8 × 10^{-5} | 15.1 |
|  | Mean nymph abundance/10 m² | -0.2 | 0.1 | 0.0010 | 7.1 |
| Density of *B. burgdorferi* (s.l.)-infected ticks | (Intercept) | -12.4 | 1.8 | 9.6 × 10^{-12} | – |
| *B. afzelii* | Growing degree-days | 0.5 | 0.1 | 0.00015 | 11.9 |
|  | (Intercept) | -12.1 | 3.3 | 0.00021 | – |
| *B. garinii* | Growing degree-days | 0.5 | 0.3 | 0.031 | 3.3 |
|  | (Intercept) | -5.8 | 0.4 | <2.0 × 10^{-16} | – |

Delta AICc indicates the change in AICc after removing each variable from the best-fit model.

*UK Met Office Long Term Average climate data [35]*

*Abbreviation: SE standard error*
of the residual deviance to the residual degrees of freedom was 0.6 with the observation level random effect, compared to 3.2 without it. Increased numbers of nymphs were found with higher growing degree-days (delta AICc = 2.5) and in areas where the forest floor was dominated by grasses compared to all other ground vegetation types (delta AICc = 2.7) (Additional file 4: Table S4). The index of deer abundance based on deer dung, the saturation deficit, woodland type (seminatural mixed or coniferous) and an interaction term between latitude and longitude of the site were not supported as significant predictors in the best-fit model.

Discussion

The overall genospecies composition in this study was consistent with previous data from Scotland [9]. *Borrelia afzelii* was the most common genospecies detected in questing nymphs, followed by *B. garinii* and both *B. valaisiana* and *B. burgdorferi* (s.s.) were found at lower prevalences. Consistent with previous reports [9, 36], *B. valaisiana* was detected only in seminatural mixed woodland and *B. burgdorferi* (s.s.) was detected only in coniferous woodland. Our study also showed a wider distribution for *B. burgdorferi* (s.s.) than previously described, with detection of this genospecies in central and western Scotland as well as the northeast Highlands (Fig. 3). *Borrelia valaisiana* is associated with bird hosts, and there are many potential competent reservoir hosts among bird communities in semi-natural mixed woodlands [57]. *Borrelia burgdorferi* (s.s.) is a generalist genospecies, able to infect both birds and mammals. Several studies have suggested an association between red squirrels (a conifer specialist) and *B. burgdorferi* (s.s.) [9, 58, 59] but their contribution in proportion to other hosts in maintaining this genospecies remains unknown. The significance of red squirrels as a reservoir host for *B. burgdorferi* (s.s.) warrants further investigation as this could be useful in predicting the distribution of this genospecies.

The overall prevalence of *B. burgdorferi* (s.l.) in this study (mean: 1.7%; 95% CI: 1.3–2.2%; range: 0–6%) is significantly lower than previously reported in a study of 25 woodland sites around Scotland (mean: 5.6%; 95% CI: 4.4–7.0%; range: 0.8–13.9%) [8], and within the lower quartile of the range of prevalence reported across Europe [60]. The previous survey of ticks and *B. burgdorferi* (s.l.) in Scotland selected woodland sites thought to be associated with cases of Lyme borreliosis from questionnaire surveys [8, 61]. The difference in prevalence between these studies may be due to the method of site section, the geographic area which was sampled, differences in PCR testing methodology or reflect real differences in *B. burgdorferi* (s.l.) circulation between years (e.g. due to fluctuations in reservoir host populations). A further difference from the previous study [8] was that we did not detect *B. burgdorferi* (s.l.) at five of our study sites, based on a sample size of 200 nymphs per site. This suggests absence of *B. burgdorferi* (s.l.) or persistence at very low levels (mean: 0%, 95% CI: 0–1.8%). This is an unusual finding; in a meta-analysis of European studies, only two out of 110 studies found a prevalence less than 1.8% [62]. Another study from the UK carried out in northern England also did not detect *B. burgdorferi* (s.l.) in ticks tested from four of 11 sites [36].

A number of possible mechanisms could contribute to a lower prevalence and an apparent absence of *B. burgdorferi* (s.l.) transmission at some sites in this study. These include a high density of incompetent hosts such as deer, low densities of reservoir hosts such as rodents and birds, and host vector interactions such as lower numbers of nymphs on reservoir hosts [15]. We did not
find a relationship between the index of deer abundance and *B. burgdorferi* (s.l.) prevalence, unlike a previous survey from Scotland [8] which reported a positive relationship between red deer dung and *B. burgdorferi* (s.l.) prevalence. The relationship between the density of deer and *B. burgdorferi* (s.l.) prevalence can be variable based on theoretical [15, 63] and empirical studies, with positive [8], negative [64, 65] and no effect of deer density on *B. burgdorferi* (s.l.) prevalence [66, 67] reported in field studies. Lower *B. burgdorferi* (s.l.) prevalence was associated with higher nymph abundance in this study. This may indirectly suggest a role for high densities of incompetent hosts such as deer which act as tick reproduction hosts and can ‘dilute’ the prevalence of *B. burgdorferi* (s.l.) if high numbers of immature ticks feed on them [15, 30, 63]. However, unlike previous studies [8, 39, 68, 69], an index of deer density based on deer dung was not supported in the nymph abundance model in this study. Although using a method to estimate deer abundance which had been used successfully in previous studies from the same area [8, 39], we suspect that index of relative deer abundance in this study is not well correlated with actual deer abundance. We counted low amounts of deer dung or an absence of dung at sites which other evidence of deer presence (e.g. tracks) were seen. To improve measures of deer abundance in future studies, it may be useful to examine a larger area at each site for deer dung.

Semi-natural mixed woodland has been associated with an increased prevalence of *B. burgdorferi* (s.l.) in nymphs compared to coniferous plantation in previous studies in the UK [8, 9, 36]; however we did not find any significant association between the prevalence of *B. burgdorferi* (s.l.) and woodland type. It is possible that the low prevalence in this study may have decreased our power to detect an effect of woodland type. A consequence of reduced power is decreased confidence in interpreting explanatory variables which are not maintained in the model due to an increased chance of type II errors. In contrast, when significant effects are detected, such as the association between growing degree-days and prevalence in this study, this shows strong support for these explanatory variables (Table 2). In this study, a higher prevalence of *B. burgdorferi* (s.l.), *B. afzelii* and a higher density of infected nymphs were found at sites with warmer climates, as these may be the areas which are more suitable for rodent reservoir hosts. A higher likelihood of finding infected ticks was reported at lower altitudes in previous studies in Scotland [8] and Switzerland [70]. Temperature is the main abiotic factor which varies with altitude [39], with warmer climates present at lower altitudes. So, these studies also suggest that climatic factors (either directly, or as a proxy of rodent host abundance) affect the prevalence of *B. burgdorferi* (s.l.) in questing ticks.

Whether local prevalence and the density of infected ticks remains stable over time is an important consideration for developing models to predict variation in risk-based on long-term averaged abiotic and habitat factors. By sampling woodland sites in multiple years, we found that the *B. burgdorferi* (s.l.) prevalence and genospecies composition at individual sites initially categorised as ‘high’ and ‘low’ prevalence can be highly variable between years (Fig. 4). Sampling variation could contribute to a proportion of the observed variation, however the sample size of nymphs collected at each site (n = 200) was chosen to be high to minimize this. Variation in prevalence between years has also been reported at sites sampled over several years in Europe [71, 72] and North America [73], although a recent study in the Netherlands found surprisingly little inter-year variation at all sites [74].

Between-year variation in the prevalence and genospecies composition of *B. burgdorferi* (s.l.) in questing ticks is most likely to be due to changes in the vertebrate host community composition. For example, a study from North America has shown that the densities of mice and chipmunks were found to be a significant predictor of the numbers of infected ticks the following year [73]. This suggests that in order to develop methods to predict areas of ‘high’ and ‘low’ environmental risk for humans, measures of both dynamic environmental factors such as host population densities will need to be measured as well as more static factors such as long term averaged climatic factors, habitat and vegetation. As *I. ricinus* life-stages generally take one blood meal per year, then emerge after developing into the next stage the following year, we would expect the prevalence of infection in nymphs to be affected by reservoir host densities in the previous year.

For the six sites sampled repeatedly in subsequent years, we found a non-significant trend towards higher numbers of nymphs infected with *B. afzelii* in 2013 compared to 2012 (Additional file 3: Table S3). At the individual site level, there was a significant increase in *B. afzelii* at two sites in 2013 compared to 2012 (Fig. 4). Another longitudinal study carried out in the Netherlands, in which *B. afzelii* was the dominant genospecies, found an alternating annual trend in prevalence at three out of four sites [71]. A decrease in *B. afzelii* prevalence between consecutive years was also noted at several sites in a study from Germany [72]. Rodent species are the most competent hosts for *B. afzelii* [16]. Parallel fluctuations in the prevalence of *B. afzelii* at several sites could be due to changes in rodent populations, perhaps in response to similar environmental conditions between these sites. None of these studies including the current study, monitored rodent abundance along with the prevalence of infection in questing ticks. Such studies are needed.
to test whether variability in small mammal density affects prevalence in questing ticks as has been described in North America [73].

Sites with a higher growing degree-day index generally had higher densities of questing nymphs, most likely due to effects on temperature-dependent inter-stadial development rates, survival rates, oviposition and egg development rates and levels of activity [75, 76]. Unlike other studies which have reported a positive effect of deer density on tick abundance [8, 68], an index of deer abundance (from dung counts) was not supported in models of tick abundance in this study. We suspect that the index of deer abundance used in this study may be poorly correlated with actual deer abundance as discussed above. The saturation deficit on the day of sampling was also not supported in this study in the best-fit model of tick abundance. A mean value for the saturation deficit over the previous 30 days prior to tick sampling may be a more useful predictor than sampling this parameter on one day [41]. We found that ground vegetation dominated by grass was associated with the highest density of questing nymphs compared to all other ground vegetation types. The category of ground vegetation associated with the highest tick densities varies between studies and geographic areas in Scotland using the same methodology. A previous study showed that bracken had lower tick counts on blanket drags compared to all other vegetation types [8]. A focal study around Loch Lomond in Scotland, found higher tick counts on blanket drags in ericaceous vegetation [77]. Thus, ground vegetation under woodland canopies appears to have limited predictive power for tick abundance over larger geographic areas.

Conclusions

We found that a higher prevalence of *B. burgdorferi* (s.l.), a higher prevalence of *B. afzelii*, higher nymph abundance and higher densities of ticks infected with *B. burgdorferi* (s.l.) (a measure of Lyme borreliosis risk) were associated with warmer sites in Scotland. In contrast to a previous study in Scotland [8], we did not find an association between *B. burgdorferi* (s.l.) prevalence and woodland type. We also did not find a relationship between an index of deer abundance (based on deer dung counts) and the abundance of *I. ricinus*, *B. burgdorferi* (s.l.) prevalence or the density of infected nymphs, as previously reported [8, 68]. However, we suspect that the index of deer abundance used in this study may not be well correlated with actual deer abundance and examining an increased area for the presence of deer dung may be beneficial in future studies. The prevalence of *B. afzelii* and *B. garinii* varied significantly at three of the six sites monitored over time, but no consistent trend in prevalence was seen across all sites. This suggests that there are changes in host communities and the proportion of competent reservoir hosts at sites over time, and that changes in host densities varies between locations. This emphasises the difficulty in forming generic rules of environmental associations with this suite of pathogens. Future challenges to predicting patterns of spatial and temporal variability in Lyme borreliosis risk depend on being able to directly or indirectly incorporate measures of the host community as well as other more static biotic and abiotic factors.

**Additional files**

Additional file 1: Table S1. Nymph abundance, and *B. burgdorferi* (s.l) prevalence and individual genospecies prevalence at 19 sites visited between April and July 2012 (n = 200 nymphs at each site). (DOCX 17 kb)

Additional file 2: Table S2. *B. burgdorferi* (s.l) prevalence and genospecies prevalence at each site sampled in, 2007/2008, 2012 and 2013. A full set of genospecies data and sample size to calculate the 95% CI were not available for all sites sampled in 2007/2008. (DOCX 17 kb)

Additional file 3: Table S3. Results for the final generalised linear mixed model to explain variation in the prevalence of infected nymphs collected at six sites sampled in 2012 and 2013. (DOCX 14 kb)

Additional file 4: Table S4. Results from the final selected generalised linear mixed model to explain variation in questing *Ixodes ricinus* nymph abundance. Delta AICc indicates the change in AICc after removing each variable from the best-fit model. (DOCX 14 kb)

**Abbreviations**

AIC: Akaike’s Information Criterion; BLAST: Basic local alignment search tool analysis; IGS: Intergenic spacer region; NCBi: National Centre for Biotechnology

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**Availability of data and material**

The data supporting the conclusions of this article are included within the article and its additional files.

**Authors’ contributions**

CM participated in the study design, field sampling, statistical analysis and drafted the manuscript. LG participated in the study design and field sampling and provided comments on drafts of the manuscript. PJ advised on the statistical analysis. MJ provided data from a previous study used for comparison with this study. EK participated in the DNA extractions and PCR testing of tick samples. RBs participated in PCR testing of tick samples. RBk participated in the study design and provided comments on drafts of the manuscript. All authors read and approved the final manuscript.

**Competing interests**

The authors declare that they have no competing interests.

**Consent for publication**

Not applicable.
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