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Effects of Parasitism and Morphology on Squirrerpox Virus Seroprevalence in Grey Squirrels (Sciurus carolinensis)

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Abstract

Invasive species have been cited as major causes of population extinctions in several animal and plant classes worldwide. The North American grey squirrel (Sciurus carolinensis) has a major detrimental effect on native red squirrel (Sciurus vulgaris) populations across Britain and Ireland, in part because it can be a reservoir host for the deadly squirrelpox virus (SQPV). Whilst various researchers have investigated the epizootiology of SQPV disease in grey squirrels and have modelled the consequent effects on red squirrel populations, less work has examined morphological and physiological characteristics that might make individual grey squirrels more susceptible to contracting SQPV. The current study investigated the putative relationships between morphology, parasitism, and SQPV exposure in grey squirrels. We found geographical, sex, and morphological differences in SQPV seroprevalence. In particular, larger animals, those with wide zygomatic arch widths (ZAW), males with large testes, and individuals with concurrent nematode and/or coccidial infections had an increased seroprevalence of SQPV. In addition, males with larger spleens, particularly those with narrow ZAW, were more likely to be exposed to SQPV. Overall these results show that there is variation in SQPV seroprevalence in grey squirrels and that, consequently, certain individual, or populations of, grey squirrels might be more responsible for transmitting SQPV to native red squirrel populations.

Introduction

There is a strong link between infectious disease and biodiversity because individual success, survival, and, consequently, the viability of entire host populations depend upon interactions with external pathogens [1]. The effects of infection are related to a number of factors including pathogen virulence [2], the nutritional and reproductive status of the host [3,4], as well as various biotic and abiotic factors such as ambient temperature and intraspecific competition [5,6]. Virulent pathogens may result in the rapid death of their host and initiate subsequent host population decline [7], whereas others can persist as low-grade chronic infections that are mediated immunologically [8]. The pathology associated with chronic disease may be important, being manifest as reductions in host growth, longevity and/or fecundity [1]. In addition, chronically diseased animals can be “carriers” within an ecosystem, potentially transmitting infection to other individuals [9]. The arrival of novel pathogens and/or infected hosts/ vectors into a geographical area where the resident species were not previously exposed could facilitate disease transmission [10]. Diseases in resident individuals may then present as acute rather than chronic [11].

Many impacts of disease on host behaviour and life history are mediated through modifications in host energy expenditure [12]. Investment in one aspect of somatic function, such as fighting an infection, inevitably means that those resources cannot be invested in others such as reproduction or growth [13]. Conversely, the success of a host population depends upon an adequate supply of food from the environment such that reductions in food availability can increase the probability of infection [1]. For example, sub-lethal effects of a viral pathogen infecting Indian meal moths, Plodia interpunctella, were more apparent when food was limited [4]. As food availability within an environment may fluctuate over time, e.g. seasonally, the prevalence and impacts of infectious disease may also coincide with such fluctuations. Finally, the effects of infection may be exacerbated by the presence of multiple pathogens, such that animals harbouring one pathogen may be more susceptible to, or suffer greater effects from a secondary infection. This may be a consequence of a diversion of the host’s resources towards the primary infection, thereby allowing the secondary pathogen to evade the immune system [14,15]. For example, myxoma virus correlates with nematode and cestode burden in the European rabbit (Oryctolagus cuniculus), which was suggested to occur as a result of the animal’s reduced immune response following the viral infection [16].

In addition to the effects of extrinsic factors on disease, intrinsic (host-specific) factors may correlate with disease prevalence. Host body size is often positively related to parasitic infection, perhaps...
Materials and Methods

Ethics statement
Squirrel carcasses were obtained from the Northern Ireland (N.I.) Forest Service, who granted permission for their use. Animals were culled according to methods outlined in “Control of Grey Squirrels for Red Squirrel Conservation” [36]. No individuals were obtained specifically for the purposes of this study.

Specimen collection
Grey squirrel specimens were collected as part of a government forestry culling programme, from various sites across N.I. (February 2008–February 2009 inclusive). Samples were available during all four seasons (spring, summer, autumn, and winter). The locations, sample sizes, and years that culling of grey squirrels began were: Belvoir (n = 10) (2001); Larchfield (n = 12) (no data); Lissan (n = 11) (2008); Tollymore (n = 1) (1998); Drum Manor (n = 12) (2000); Portglenone (n = 15) (2002); Loughgall (n = 119) (1992); Gosford (n = 43) (1988); Drumlanagher (n = 19) (1993); and Derrynoyd (n = 51) (2001) (Fig. 1). Specimens were stored at −20°C until processed.

Determination of SQPV exposure
The presence of antibodies against SQPV was determined from blood samples for all specimens using an ELISA [35]. The presence of antibodies was used as a measure of SQPV exposure as it is difficult to replicate the virus in high titres. This provides a measure of viral exposure at the time of culling. An optical density reading of >0.2 at 492 nm was taken as the cut-off to differentiate specific antibody positive serum samples from negative serum samples.

Statistical analyses
Data analyses were carried out in “R” version 2.15.2 [38]. Data distributions for all variables were significantly different from normal (Kolmogorov-Smirnov test, p<0.001). Season, sex, and location differences in SQPV seroprevalence were compared using Fisher’s exact tests. Variation in SQPV exposure was examined according to the length of time that grey squirrel colonies were established in each location to determine whether seropositivity within a location changed with time [39]. The date that culling was first carried out by N.I. Forest Service was used as a proxy for the establishment of grey squirrels in a given area.

Generalized Linear Models with a binomial distribution and log link function were used to assess the potential relationships between SQPV seroprevalence, parasite load, and morphological features. Minimal models were selected based on comparisons of
Akaike Information Criteria (AIC) [40]. First, any collinearity between morphological features was examined using Spearman’s rank correlations to determine if any organ masses could be omitted from models. No collinearity was observed (p > 0.05 in all cases) so the associations between each morphological feature (body mass, ZAW, mass of the testes, and spleen mass) and SQPV exposure were examined by entering each feature singly as an explanatory variable in individual models with SQPV exposure as a response variable. In the event of a variable holding an explanatory power of p < 0.1, that variable was then entered into a single model comprising the main effects and two-way interactions between morphological features (except mass of the testes) on SQPV exposure (Model A, Table 1). This model was then expanded to include the main effects and two-way interactions between morphological features and various forms of parasitism (nematode, coccidia, and ectoparasites) on SQPV exposure in a single model (Model B, Table 1). The final models examined the combined effects of morphology, parasitism, and sex on exposure to SQPV, these were: The main effects and two-way interactions between morphology and sex on SQPV exposure (Model C1, Table 1) and the main effects and two-way interactions between parasitism and sex on SQPV exposure (Model C2, Table 1).

The inclusion of mass of the testes as an explanatory variable excluded females from the analysis. Therefore, further models examined male specimens only. Initial analysis revealed that there was a significant seasonal variation in mass of the testes (Kruskal-Wallis: $\chi^2 = 9.13$, df = 3, p = 0.028). Hence the main effects of season, mass of the testes, and body mass, and the interactions between season and mass of the testes, as well as mass of the testes and body mass on SQPV exposure were included in a combined model. In this case, none of the terms showed a significant explanatory power (p > 0.05). Therefore, the factor “season” was omitted from further models. The first model featuring data for just males included the main effects and two-way interactions of all morphological features (body mass, ZAW, mass of the testes, and spleen mass) on SQPV exposure (Model D, Table 1). Thereafter, additional models examined the interactions between morphology and the various types of parasitism (coccidia, nematodes, and ectoparasites) (Models E-G, Table 1).

Results

Of the 236 grey squirrel specimens examined, 113 were female and 123 were male. Of these, 33.6% of females and 32.5% of males tested positive for the presence antibodies against SQPV. There was no significant difference between the proportion of males and females exposed to SQPV ($\chi^2 = 0.002$, df = 1, p = 0.96).

The seroprevalence of SQPV varied with location (Fisher’s exact test: p = 0.007) (Fig. 1A). At one site, 63.2% of the individuals tested positive for SQPV antibodies, whereas all other sites had seroprevalences ≤50% and three populations showed no evidence of exposure. There was no significant relationship between the percentage of seropositive individuals and the time since first culling in that location (least squares regression, $F_{1,7} = 2.10$, $r^2 = 0.231$, p = 0.191). However, there was a significant difference between seroprevalence in winter (40.3% of individuals seropositive) compared to summer (21.7% of individuals seropositive) ($\chi^2 = 4.12$, df = 1, p = 0.042).

Specimens were infected with a variety of ecto- and endoparasites (Table 2). Ectoparasites found included the flea, *Ochropaenus hircudii*, the louse, *Neoehematophilus sciurinus*, the tick, *Ixodes ricinus*, and the mite, *Androlaelaps fahrenholzi*. The overall prevalence of ectoparasites was 70.3%. Endoparasite species identified included coccidia (*Eimeria* sp.) oocysts and nematodes, *Typanoxyuris* (Rodentoxyuris) *sciuri*, *Trichostrongylus retortaeformis*, and *Trichuris* sp. The prevalence of coccidial parasites and nematodes was 70.3% and 53.7% respectively.
<table>
<thead>
<tr>
<th>Model</th>
<th>Cohort</th>
<th>Maximal model</th>
<th>Minimal model</th>
<th>Sig. effects</th>
<th>z</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>All</td>
<td>BM+ZAW+SPL+BM<em>ZAW+BM</em>SPL+ZAW*SPL</td>
<td>BM+ZAW+SPL+BM<em>ZAW+ZAW</em>SPL</td>
<td>BM</td>
<td>2.47</td>
<td>0.014</td>
</tr>
<tr>
<td>B</td>
<td>All</td>
<td>BM+ZAW+SPL+COX+NEM+ECT+BM<em>ZAW+BM</em>SPL+BM<em>COX+BM</em>NEM+BM<em>ECT+ZAW</em>SPL+ZAW<em>COX+ZAW</em>NEM+ZAW<em>ECT+SPL</em>COX+SPL<em>NEM+SPL</em>ECT+NEM<em>COX+NEM</em>ECT+NEM*COX</td>
<td>BM+ZAW+NE+COX+ZAW+SPL+ZAW+NEM+ZAW*COX</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
</tr>
<tr>
<td>C1</td>
<td>All</td>
<td>BM+SPL+ZAW+SEX+BM<em>SPL+BM</em>ZAW+BM<em>SEX+SPL</em>ZAW+SPL<em>SEX+ZAW</em>SEX</td>
<td>BM+SPL</td>
<td>BM</td>
<td>2.98</td>
<td>0.003</td>
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<tr>
<td>C2</td>
<td>All</td>
<td>COX+NEM+ECT+SEX+COX<em>NEM+COX</em>ECT+COX<em>SEX+NEM</em>ECT+NEM<em>SEX+ECT</em>SEX</td>
<td>NEM</td>
<td>NS</td>
<td>NS</td>
<td>NS</td>
</tr>
<tr>
<td>D</td>
<td>Males</td>
<td>BM+TES+ZAW+SPL+BM<em>TES+BM</em>ZAW+BM<em>SPL+TES</em>ZAW+TES<em>SPL+ZAW</em>SPL</td>
<td>BM+SPL+TES+BM*TES</td>
<td>SPL; TES</td>
<td>1.97</td>
<td>3.38</td>
</tr>
<tr>
<td>E</td>
<td>Males</td>
<td>BM+SPL+TES+ZAW+COX+BM<em>TES+BM</em>ZAW+BM<em>COX+BM</em>TES+SPL<em>ZAW+SPL</em>COX+TES<em>ZAW+TES</em>COX+ZAW*COX</td>
<td>BM+SPL+TES+ZAW+COX+BM+SPL+BM<em>TES+BM</em>ZAW+BM<em>COX+ZAW</em>COX</td>
<td>TE<em>S; COX; BM</em>TE<em>S; BM</em>COX</td>
<td>2.81</td>
<td>–2.31</td>
</tr>
<tr>
<td>F</td>
<td>Males</td>
<td>BM+SPL+ZAW+NEM+BM<em>SPL+BM</em>ZAW+BM<em>NEM+BM</em>TES+BM<em>ZAW+SPL</em>ZAW+SPL<em>NEM+BM</em>ZAW+NEM</td>
<td>BM+SPL+ZAW+NEM+BM<em>ZAW+SPL</em>ZAW+SPL<em>NEM+BM</em>ZAW+NEM</td>
<td>ZAW; BM<em>ZAW; SPL</em>ZAW; ZAW*NEM</td>
<td>–2.67</td>
<td>–2.18</td>
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<tr>
<td>G</td>
<td>Males</td>
<td>BM+SPL+TES+ZAW+ECT+BM<em>TES+BM</em>ZAW+BM<em>SPL+BM</em>TES+BM<em>ZAW+BM</em>SPL<em>ECT+BM</em>TES+ZAW*SPL</td>
<td>BM+SPL+ZAW+TES+BM<em>SPL+BM</em>ZAW+SPL<em>ECT+BM</em>TES+ZAW*SPL</td>
<td>SPL*ZAW</td>
<td>–2.01</td>
<td>0.045</td>
</tr>
</tbody>
</table>

Statistical models used to determine the effects of various morphological features as well as parasitism and sex on SQPV exposure. All models used a binomial distribution and log link function. Abbreviations are denoted as follows: BM = Body mass, SPL = Spleen mass, TES = Mass of testes, ZAW = Zygomatic arch width, SEA = Season, COX = Coccidial burden, NEM = Nematode burden, ECT = Ectoparasite burden, SEX = Sex, * = interaction, NS = No significance (p > 0.05 for all effects in minimal model).

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Effects of morphology and parasitism on SQPV exposure in all individuals

There were no sex-differences in body mass, spleen mass, or ZAW (Table 2). When the main effects and two-way interactions between morphological features (body mass, ZAW, and spleen mass) were considered in a single model, only body mass had a significant association with SQPV exposure, with larger individuals experiencing exposure more often than smaller individuals (Model A, Table 1). There were no significant interactions between any features in the minimal model. ZAW and spleen mass were not significantly related to SQPV exposure either (p > 0.05). Similarly, when parasitism, morphology, and their interactions were considered, there were no significant relationships with SQPV exposure (p > 0.05 in all cases) (Model B, Table 1). Body mass was the only variable which remained significant with the combined effects of morphology and sex on SQPV exposure (p = 0.028). Examination of the associations between morphological features (body mass, ZAW, and spleen mass) were considered in a single model, only body mass had a significant association with SQPV exposure, with larger individuals experiencing exposure more often than smaller individuals (Model D, Table 1). There were no significant interactions between any features in the minimal model. ZAW and spleen mass were not significantly related to SQPV exposure either (p > 0.05). Similarly, when parasitism, morphology, and their interactions were considered, there were no significant relationships with SQPV exposure (p > 0.05 in all cases) (Model E, Table 1). Body mass was the only variable which remained significant with the combined effects of morphology and sex on SQPV exposure (p = 0.036). Mean, standard error of the mean (S.E.) (* denotes median and standard error of the median), and sample size (N) of various internal and external morphometrics (measured in grams (g) and millimetres (mm)), and ecto- and endo-parasite burdens in males and females. Also shown are the p-values (P) from Mann-Whitney U tests examining sex differences for each parameter. No sex differences were evident in any of these tests (p > 0.05) except with tick burden where males exhibited higher burdens than females (p = 0.036).

Table 2. Sex differences in morphology and parasitism.

<table>
<thead>
<tr>
<th>Feature</th>
<th>Males</th>
<th>Females</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>N</td>
<td>Mean</td>
</tr>
<tr>
<td>Body mass (g)</td>
<td>151</td>
<td>528</td>
</tr>
<tr>
<td>ZAW (mm)</td>
<td>148</td>
<td>33.8</td>
</tr>
<tr>
<td>Spleen mass (g)</td>
<td>151</td>
<td>0.31</td>
</tr>
<tr>
<td>Mass of testes (g)</td>
<td>152</td>
<td>0.47</td>
</tr>
<tr>
<td>Ticks</td>
<td>127</td>
<td>0.06</td>
</tr>
<tr>
<td>Fleas</td>
<td>127</td>
<td>4.77</td>
</tr>
<tr>
<td>Mites</td>
<td>107</td>
<td>0.05</td>
</tr>
<tr>
<td>Lice</td>
<td>127</td>
<td>1.13</td>
</tr>
<tr>
<td>Total ectoparasites</td>
<td>127</td>
<td>5.99</td>
</tr>
<tr>
<td>Nematodes</td>
<td>151</td>
<td>3.83</td>
</tr>
<tr>
<td>Coccidial score</td>
<td>139</td>
<td>1*</td>
</tr>
</tbody>
</table>

Mean, standard error of the mean (S.E.) (* denotes median and standard error of the median), and sample size (N) of various internal and external morphometrics (measured in grams (g) and millimetres (mm)), and ecto- and endo-parasite burdens in males and females. Also shown are the p-values (P) from Mann-Whitney U tests examining sex differences for each parameter. No sex differences were evident in any of these tests (p > 0.05) except with tick burden where males exhibited higher burdens than females (p = 0.036).

doi:10.1371/journal.pone.0083106.t002

Discussion

Invasive species have been cited as major causes of population extinctions in several animal and plant classes worldwide [41]. The North American grey squirrel has a major detrimental effect on native red squirrel populations in Britain and Ireland, in part because they are reservoir hosts for SQPV, which is deadly to red squirrels. Although some researchers have investigated the epizootiology of SQPV disease [42–44], less work has examined the various morphological, physiological, and environmental characteristics that might make individual grey squirrels more susceptible to, and potentially better able to transmit the virus to red squirrels [33,45]. The current study investigated the putative relationships between morphology, parasite infection, and SQPV seroprevalence in grey squirrels, and documented any geographical and seasonal variation that was found in these characteristics. We found that larger individuals and males with larger testes were more likely to have been exposed to SQPV. This is consistent with predictions that larger, dominant individuals may experience increased exposure to infection, possibly by moving further distances and spending more time performing activities such as foraging which might expose them to pathogens [46].
with an increased likelihood of SQPV seroprevalence. It is unclear why this might be the case but coccidial infection is often present as a more serious, acute infection in young or juvenile animals, and more chronic in adults [47]. Thus, adults may appear to have an increased coccidial prevalence and perhaps this is then associated with an increased likelihood of SQPV seroprevalence.

There were sex differences in the relationships between morphology and parasitism, with many significant effects on SQPV exposure evident in males but not in the pooled data of males and females. For example, males with a wide ZAW and high nematode burdens were more likely to be SQPV seropositive, which suggests that older dominant males may be more likely to be exposed to SQPV. The fact that apparent correlates of dominance and reproductive activity are related to parasitism and exposure to SQPV suggests that there are costs to these traits, i.e. some individuals, often males, may increase infection risk in favour of reproductive success [19,20,48,49]. Male squirrels are also known to be more aggressive, have large home ranges, fight amongst each other, and chase females during oestrus [49,47]. These behaviors might increase exposure to SQPV. The fact that some forms of parasitism co-occur with SQPV exposure supports the contention that one pathogen may be exploiting the diversion of the host's immune response to the other [14], or that immunomodulation, either by the parasites or SQPV, can increase susceptibility to the other.

Spleen mass was an important indicator of SQPV exposure in males as those with larger spleens were more likely to have been exposed. When dominance was also considered, it was noted that males with a wide ZAW (potentially dominant individuals) with small spleens and males with a narrow ZAW (potentially subordinate individuals) with large spleens were more likely to have been exposed to SQPV (Fig. 2). It is unclear whether SQPV influences spleen size or vice versa but it would seem that there is a trade-off between being dominant and investing in immunity and that there is at least two modes of infection; one in which older, dominant individuals are exposed and another in which younger, subordinate animals are exposed. A potential explanation could be that individuals with small spleens are less well-equipped to deal with SQPV infection, especially if they have an increased risk of exposure to SQPV because they are reproductively active (as evidenced by wide ZAW and larger testes), or if they have elevated testosterone levels which may compromise immune system functionality (as indicated by larger testes) [21]. It is also possible that the relationship between mass of the testes and SQPV is a result of SQPV causing testicular swelling, which is characteristic of infection with other viruses such as mumps virus [50].

Concurrent with previous studies, the seroprevalence of SQPV varied according to location, with some areas apparently clear of the infection and other areas comprising large numbers of exposed animals [39]. While there was a possibility for the more established colonies from different areas to have greater percentage seroprevalence, this was not apparent. Geographical variation in SQPV seroprevalence might be a result of environmental factors such as the distribution of suitable habitats [51] as well as the presence of other organisms which could act as reservoir hosts or vectors for SQPV, however, at present no other hosts have been identified [39]. Although there was no overall seasonal effect on SQPV seroprevalence, there was a significant difference between winter and summer exposure, with almost twice as many individuals testing seropositive during the winter. As winter is the main breeding season [52], reproductive males may become exposed to SQPV when they patrol areas in search of receptive females [46]. Alternatively, increased SQPV seroprevalence during the winter may be a consequence of close social contact between individuals when communal nesting increases [53], or when individuals cluster around food resources [54].

Conclusions

Despite various conservation efforts, including culling grey squirrels [55], and reintroduction and supplementary feeding of red squirrels [54], the number of grey squirrels in the U.K. and Ireland continues to rise whilst the number of red squirrels falls. This study investigated the potential aspects of SQPV seroprevalence, which may contribute to this trend. While there may be potential biases in data collection (e.g. young or naïve individuals may be trapped more easily), the sample size was large enough to obtain representative animals spanning different cohorts within the population. Spatial and temporal variation in SQPV antibodies was evident, with notable increases during the winter. Reasons for this remain unclear but it may well be a result of individuals being in closer contact with each other either as a result of mating behaviours, feeding and/or nesting. Of note is that larger males with larger testes and those with concurrent nematode or coccidial infections were more likely to be exposed to SQPV. In addition, subordinate males with large spleens were also more likely to be exposed (Fig. 2). Hence, SQPV transmission may occur when individuals are large and dominant, and potentially already under stress from other infections and also when they are young and small. Finally, individuals with larger spleens were more likely to experience SQPV exposure, suggesting that spleen size may increase as a response to infection. It is hoped that natural immunity may develop in red squirrel populations, as has occurred in European rabbits presenting resistance to the myxoma virus [16]. It may therefore be prudent to encourage the expansion of those red squirrel populations which have already been exposed to SQPV and survived, perhaps by supplementary feeding, coupled with the culling programmes which are already in place for grey squirrels, to promote the development of natural immunity within the population.

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Author Contributions

Conceived and designed the experiments: NJM MS. Performed the experiments: NEM NJM CJM DD MS. Analyzed the data: NEM MS. Contributed reagents/materials/analysis tools: MS NJM CJM. Wrote the paper: NEM NJM CJM DD AM MS.

References


23. Webley GE, Pope GS, Johnson E (1985) Seasonal changes in the testes and
2. Wickings EJ, Dixson AF (1992) Testicular function, secondary sexual