

1 Genome-wide association and genome partitioning reveal novel
2 genomic regions underlying variation in gastrointestinal nematode
3 burden in a wild bird

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12 partitioning; missing heritability

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14 **Abstract**

15 Identifying the genetic architecture underlying complex phenotypes is a notoriously difficult prob-
16 lem that often impedes progress in understanding adaptive eco-evolutionary processes in natural
17 populations. Host-parasite interactions are fundamentally important drivers of evolutionary pro-
18 cesses, but a lack of understanding of the genes involved in the host's response to chronic parasite
19 insult makes it particularly difficult to understand the mechanisms of host life-history trade-offs and
20 adaptive dynamics involved. Here we examine the genetic basis of gastrointestinal nematode (*Tri-*
21 *chostrongylus tenuis*) burden in 695 red grouse (*Lagopus lagopus scotica*) individuals genotyped at
22 384 genome-wide SNPs. We first use genome-wide association to identify individual SNPs associated
23 with nematode burden. We then partition genome-wide heritability to identify chromosomes with
24 greater heritability than expected from gene content, due to harbouring a multitude of additive SNPs
25 with individually undetectable effects. We identified five SNPs on five chromosomes that accounted
26 for differences of up to 556 worms per bird, but together explained at best 4.9 % of the phenotypic
27 variance. These SNPs were closely linked to genes representing a range of physiological processes
28 including the immune system, protein degradation and energy metabolism. Genome partitioning

29 indicated genome-wide heritability of up to 29 % and three chromosomes with excess heritability of
30 up to 4.3 % (total 8.9 %). These results implicate SNPs and novel genes underlying nematode burden
31 in this system and suggest that this phenotype is somewhere between being based on few large-effect
32 genes (oligogenic) and based on a large number of genes with small individual but large combined
33 effects (polygenic).

34 Introduction

35 Host-parasite interactions are widely recognised as fundamentally important drivers of evolutionary and
36 ecological processes in natural populations. The Red-Queen dynamics of the host-parasite co-evolutionary
37 arms race dictate that the ability of a host to cope with parasite insult is a major component of individual
38 fitness (van Valen, 1973; Brockhurst, 2011). This selective pressure on the host can influence how energetic
39 resources are allocated to immune defence over other vital cellular functions and life-history components,
40 and thus how parasite-driven life-history trade-offs are resolved (Sheldon & Verhulst, 1996; Zuk & Stoehr,
41 2002; Schmid-Hempel, 2003). Such decisions affect individual mating behaviour and sexual selection
42 processes (Hamilton & Zuk, 1982; Hill & Farmer, 2005) as well as population demography and dynamics
43 (Lochmiller, 1996; Hudson *et al.*, 2006).

44 A proper understanding of host-parasite eco-evolutionary dynamics requires knowledge of the genetic
45 basis of individual host resistance or susceptibility (Paterson & Piertney, 2011). Traditionally, and
46 perhaps intuitively, most focus has been placed on the immune system (Lochmiller & Deerenberg, 2000;
47 Schmid-Hempel, 2003; Sadd & Schmid-Hempel, 2009; Owen *et al.*, 2010). As such, a range of classic
48 immunological candidate genes have been implicated, including the Major Histocompatibility Complex
49 MHC (Piertney & Oliver, 2006; Tobler *et al.*, 2014), interferon gamma IFNG (Coltman *et al.*, 2001;
50 Stear *et al.*, 2007), Toll-like receptors TLR (Downing *et al.*, 2010; Gavan *et al.*, in press) and cytokines
51 (Downing *et al.*, 2010; Turner *et al.*, 2012). However, a focus on the immune system alone may be
52 misleading in systems where hosts do not attain complete parasite resistance and instead suffer chronic
53 infection (Schmid-Hempel, 2003; Sadd & Schmid-Hempel, 2009). These cases might necessitate a complex
54 constitutive response by the host, which may involve a multitude of genes across broad physiological
55 processes rather than one or few particular key immune genes. Indeed, parasites in general and nematodes
56 in particular often evade or manipulate the host's immune response and have extensive multidimensional
57 effects on host behaviour and physiology (Thomas *et al.*, 2010; Biron & Loxdale, 2013; Poulin, 2013). If
58 the host response is then mediated through changes in the allocation of resources among the different
59 life-history categories and cellular processes (Sheldon & Verhulst, 1996; Lochmiller & Deerenberg, 2000),
60 this will inevitably involve a broad suite of genotypic mechanisms with a range of physiological functions
61 divorced from the immune system (Hill, 2011).

62 In order to identify novel genes that affect a host's response to parasites, functional transcriptomic

63 assays can provide extensive suites of gene targets, which can then be screened for genotypic variation
64 (Hoffmann & Willi, 2008; Piertney & Webster, 2010; Orsini *et al.*, 2011; Pemberton *et al.*, 2011; Gossner
65 *et al.*, 2013; Wenzel & Piertney, 2015). However, this strategy does not necessarily provide a robust
66 understanding of the genetic basis of a phenotype because transcriptomic responses are context dependent
67 and are not always linked to genotypic variation in homologues of different species (Orsini *et al.*, 2012;
68 Brown *et al.*, 2013; Wenzel & Piertney, 2015). A more robust genome-scale understanding of genetic
69 architecture can be gained by quantitative-trait-loci (QTL) mapping or genome-wide association (GWA)
70 analysis directly in the target species (Slate, 2005; Hill, 2012). These approaches are based on associations
71 between genome-wide genetic markers and phenotypic traits, usually without *a priori* marker selection
72 based on putative function. QTL mapping and GWA analysis have been undertaken extensively for
73 gastrointestinal nematode burden in sheep or cattle breeds and have highlighted that nematode burden is
74 associated with small numbers of predominantly anonymous loci, some of which are located near classic
75 immunological candidate genes (e.g., Beh *et al.*, 2002; Davies *et al.*, 2006; Beraldi *et al.*, 2007; Silva *et al.*,
76 2012; Riggio *et al.*, 2013). These results are consistent with insights from QTL studies in other host-
77 parasite systems and suggest that parasite burden may after all be primarily based on small numbers of
78 large-effect immune genes (Wilfert & Schmid-Hempel, 2008).

79 Notwithstanding, a major issue with these methods is that those genotypic variants that are identi-
80 fied through QTL or GWA usually explain only a small fraction of the phenotypic variance even if the
81 phenotype is highly heritable (Manolio *et al.*, 2009). This “missing heritability” issue fuels an ongoing
82 controversy over the power of candidate gene or QTL/GWA approaches to detect small-effect polymor-
83 phisms and whether the endeavour to identify individual genotypic variants linked to complex phenotypes
84 is inherently misguided (Amos *et al.*, 2011; Rockman, 2012; Robinson *et al.*, 2013). The central tenet
85 of the argument is that polygenic phenotypes that are underpinned by the joint effect of a multitude of
86 loci each of small effect, corresponding to Fisher’s infinitesimal model (Fisher, 1919), are intractable with
87 SNP-by-SNP association approaches because only loci with individually large effects can be detected. As
88 such, these loci may be misleading about the distribution and effect size of the genetic variants that truly
89 underpin phenotypic evolution (Rockman, 2012). The high incidence of missing heritability suggests that
90 polygenic architectures may be common and thus reinforces the argument to abandon classic methods
91 involving candidate genes (Yang *et al.*, 2011b; Hill, 2012; Rockman, 2012). Parasite susceptibility may
92 well be such a polygenic phenotype if parasites have extensive physiological and behavioural effects on
93 their host (Thomas *et al.*, 2010; Biron & Loxdale, 2013; Poulin, 2013). Although heritability of parasite
94 burden is well established, for example in sheep, cattle and poultry (e.g., Stear *et al.*, 2007; Kaufmann
95 *et al.*, 2011), and previous candidate gene and QTL/GWA studies have indeed identified some large ef-
96 fect genes, these approaches may well have failed to identify genes with subtle individual, but large joint
97 effects.

98 One way forward in addressing this issue is to compare results obtained from GWA with emerging
99 “genome partitioning” methods based on quantitative genetics. These approaches do not identify in-
100 dividual SNPs or haplotypes, rather than linkage groups of SNPs (usually chromosomes) that together
101 explain more phenotypic variance than expected from a polygenic null model where all markers contribute
102 equally (Yang *et al.*, 2011b,a; Hill, 2012). Genome partitioning has proven extremely useful for retrieving
103 some of the missing heritability unaccounted for by genotypic variants identified through GWA. This has
104 been well illustrated in complex phenotypes, for example in human height (Yang *et al.*, 2010, 2011b) and
105 disorders such as schizophrenia, Tourette syndrome and obsessive-compulsive disorder (Lee *et al.*, 2012;
106 Davis *et al.*, 2013). Genome partitioning can also be useful in non-model systems with limited genomic
107 resources. Gastrointestinal nematode burden in sheep has recently been shown to be primarily polygenic,
108 but five chromosomes explained disproportionate amounts of phenotypic variance, implying particular
109 genomic regions as targets for fine-mapping causal variants (Al-Kalaldeh *et al.*, 2013). The utility of
110 comparing across QTL/GWA and genomic partitioning approaches is further illustrated by contrasting
111 results in recent studies that examined the genetic basis of wing length in birds. In spite of multiple QTLs
112 identified in zebra finches (Schielzeth *et al.*, 2012) and reed warblers (Tarka *et al.*, 2010), genome parti-
113 tioning in great tits with substantial numbers of samples and markers instead suggests that wing length
114 may be a purely polygenic additive phenotype (Robinson *et al.*, 2013). This highlights that genomic
115 architectures may be idiosyncratic among different species and that genome partitioning may be a useful
116 method for reconciling differences in the numbers and identities of large-effect genetic polymorphisms.

117 Here, we examine the genetic basis of chronic nematode infection in a wild bird species using GWA
118 and genome partitioning approaches to identify genotypic variants that explain detectable amounts of
119 phenotypic variance and genomic regions that harbour missing heritability. We focus on red grouse
120 (*Lagopus lagopus scotica*) and its primary gastrointestinal nematode parasite *Trichostrongylus tenuis*.
121 This host-parasite system has become an important model for understanding parasite-driven demographic
122 and ecological dynamics in the absence of a classic immune response (Hudson *et al.*, 1998; Redpath *et al.*,
123 2006a; Webster *et al.*, 2011b). Red grouse are an economically important game bird endemic to the upland
124 heather moors of Scotland and northern England (Martínez-Padilla *et al.*, 2014). *T. tenuis* displays a
125 direct life cycle where infective larvae are ingested with heather shoots, adults establish in the caeca and
126 eggs are shed with faeces (Hudson & Dobson, 1989). Adult worms burrow into the caecal walls where
127 they cause haemorrhaging and necrosis with substantial impacts on grouse fitness (Hudson, 1986; Watson
128 *et al.*, 1987; Hudson *et al.*, 1992; Delahay *et al.*, 1995; Delahay & Moss, 1996). Prevalence of infection with
129 *T. tenuis* in grouse populations is typically greater than 90 % and grouse bear chronic nematode burdens
130 due to an inability to purge the infection despite mounting directed responses (Wilson, 1983; Shaw &
131 Moss, 1989; Webster *et al.*, 2011a). This long-term exposure to nematode insult has marked effects
132 on grouse life-history trade-offs and population dynamics. Interactions between nematode burden and

133 testosterone-mediated territorial contest behaviour and sexual selection by females have been identified
134 as key drivers of life-history trade-offs in male grouse in particular (MacColl *et al.*, 2000; Mougeot *et al.*,
135 2003, 2004, 2005c, 2009; Piertney *et al.*, 2008; Bortolotti *et al.*, 2009; Wenzel *et al.*, 2013). Further,
136 the direct impact of nematode infection on grouse fecundity and survival together with intrinsic density-
137 dependent aggression and stress-mediated immuno-suppression implicates *T. tenuis* as a major component
138 in regulating grouse population dynamics (Hudson *et al.*, 1998; Mougeot *et al.*, 2005b; Seivwright *et al.*,
139 2005; Redpath *et al.*, 2006a; Webster *et al.*, 2011b).

140 Motivated by this importance of chronic nematode infection for red grouse ecology, substantial effort
141 has gone into characterising the physiological categories involved in the molecular response to infection
142 and into identifying genomic regions associated with variation in nematode burden among individual
143 grouse. Transcriptomic assays following experimental nematode infection have highlighted a broad range
144 of physiological categories beyond the immune system and have identified testosterone-dependent tran-
145 scription dynamics that are consistent with trade-offs involving depression of parasite defence mechanisms
146 (Webster *et al.*, 2011a,b; Wenzel *et al.*, 2013). These insights have been channelled into developing novel
147 candidate genes for nematode susceptibility (Wenzel *et al.*, 2015), which have been confirmed to explain
148 variation in nematode burden among grouse individuals in a network of moors in north-east Scotland
149 (Wenzel & Piertney, 2015). In that same study system, DNA methylation patterns at specific genomic
150 regions have been linked to nematode burden, suggesting that parasites may affect epigenetic mechanisms
151 impacting the regulations of specific genes (Wenzel & Piertney, 2014). In concert, this previous work
152 has established that the host-parasite interactions in this system may be linked to multiple large-effect
153 candidate genes involved in immune system, oxidative stress, energy metabolism and cell cycle processes.
154 However, it is unknown whether previously undetected genome-wide large-effect genotypic variants ex-
155 ist, and to what extent linkage groups of variants with individually undetectable effects contribute to
156 variation in nematode burden.

157 We extend and expand on these issues by carrying out genome-wide association and genome parti-
158 tioning of variation in nematode burden in a sample of 695 red grouse individuals from five locations
159 in Scotland and England across twelve years and genotyped at 384 genome-wide SNPs. First, we fit
160 custom generalized linear models to identify individual SNPs that are associated with individual nema-
161 tode burden. Second, we ascertain putative functions of these identified SNPs via linked genes identified
162 through homology with the chicken genome. Finally, we estimate genome-wide and chromosome-specific
163 heritabilities of nematode burden, and identify chromosomes that contribute disproportionately to heri-
164 tability through multiple additive genotypic variants of individually small effects.

165 **Materials and Methods**

166 **Phenotypic data**

167 A total of 695 red grouse individuals were sampled at four sites near Deeside in north-east Scotland and
168 one site in Catterick, northern England from 1995–2012 (Table 1). The core sampling effort was carried
169 out at Invermark and Invercauld in 2011–2012 and was supplemented with birds that were sampled as
170 part of previous studies investigating grouse population ecology (e.g., MacColl *et al.*, 2000; Redpath *et al.*,
171 2006b), behavioural ecology (e.g., Mougeot *et al.*, 2003, 2009) and physiological responses to nematode
172 infection (e.g., Webster *et al.*, 2011b; Wenzel *et al.*, 2013; Wenzel & Piertney, 2014, 2015). Birds were
173 either captured at night using lamping and netting techniques or collected following driven or walked-up
174 sporting shoots. *T. tenuis* burdens were estimated from faecal egg counts using the standard McMaster
175 chamber slide method and empirical prediction functions for worm burden (Moss *et al.*, 1990; Seivwright
176 *et al.*, 2004). For shot birds, caecum samples were stored cold immediately after sampling and processed
177 within one week to ensure reliable estimates (Seivwright *et al.*, 2004). Sex and age were determined
178 morphologically using plumage and supra-orbital comb size. Birds that fledged in the year of sampling
179 were classed as “young”, and all other birds were classed as “old” (>1 year), allowing for statistical control
180 of typically lower nematode burden in young birds (Shaw & Moss, 1989; Hudson & Dobson, 1997). Body
181 mass was measured to the nearest 10 g using a spring balance and wing length was measured to the
182 nearest mm. As a measure of physiological condition, the scaled mass condition index (Peig & Green,
183 2009) was then calculated for each individual (i) as:

$$CI_i = m_i \left(\frac{w_0}{w_i} \right)^b$$

184 where m_i is body mass, w_i is wing length, w_0 is mean wing length and b is the slope of standardised
185 major axis regression of $\ln m$ on $\ln w$ across the data set.

186 **Genetic data**

187 DNA was extracted from feather calamus or liver tissue samples using a standard salting out method
188 (Hogan *et al.*, 2008). Individuals were genotyped at 384 genome-wide SNPs using the Illumina Gold-
189 enGate BeadXpress platform (NBAF-S, University of Sheffield). Based on orthology with the chicken
190 genome (*galGal4* assembly) following BLAT searches (Kent, 2002), these SNPs were designed such that
191 the number of SNPs per chromosome was approximately proportional to chromosome size. As such,
192 chromosomes 1–15, 17–26, 28 and Z were covered by 1–82 SNPs (median 6) with a median distance
193 between consecutive SNPs of 1.9 Mbp (supplementary materials S1).

194 Quality filtering and calculation of summary statistics were carried out in PLINK 1.9 (Chang *et al.*,

214). SNPs with calling rate below 0.90 and minor allele frequency (MAF) below 0.05 were removed. The inbreeding coefficient (F_{IS}) for each SNP and individual was computed from observed and expected heterozygosities, and Hardy-Weinberg equilibrium exact tests were carried out to identify SNPs with genotyping errors. Quality filtering resulted in a remaining total of 271 SNPs. All 695 individuals had a genotyping rate larger than 80 %, with an overall genotype coverage of 99.6 %. Linkage disequilibrium (r^2) was estimated among SNP pairs within each chromosome. The degree of relatedness among individuals was ascertained from the realized relationship matrix based on genetic identity-by-state (Yang *et al.*, 2011a). Genetic structure was ascertained by computing global F_{ST} among sampling sites and graphically examining the first 20 principal components of the genetic relationship matrix (Johnston *et al.*, 2014). The minimum number of principal components that capture most of the genetic structure was then identified by examining the eigenvalues and selecting a principal component cut-off such that the differences in eigenvalues across subsequent principal components were minimal and relatively stable. To aid the identification of this cut-off, an *ad hoc* statistic ΔE was calculated that relates the changes in eigenvalues from each principal component to the following and the preceding component ($\Delta E_{pc} = \frac{|E_{pc} - E_{pc+1}|}{|E_{pc} - E_{pc-1}|}$), and peaks in ΔE were examined.

210 Genome-wide association analysis

211 GWA analysis was carried out by implementing custom generalized linear models with negative binomial error structure in the statistical software R 3.0.3 (R Core Team, 2014) and the package *MASS* (Venables & Ripley, 2002). The discrete worm counts were not normally distributed and therefore inappropriate for analysis with common GWA analysis packages. Exploratory modelling including all sampling-related and phenotypic variables and applying a Poisson error structure indicated substantial over-dispersion ($\phi > 1000$), and diagnostic residuals plots confirmed that a negative-binomial error structure provided the best fit for these data.

218 Relationships among nematode burden and sampling-related or phenotypic variables were ascertained by graphical exploration and linear modelling to identify confounding variables. Nematode burden varied considerably among sampling sites, years and months, but sample sizes were very low for several years and months (supplementary materials S2). To avoid statistical issues with small sample sizes, these two variables were simplified by combining factor levels. Years were binned into three time periods with substantially different nematode burdens (1995–1999; 2000–2006; 2010–2012; supplementary materials S2). Similarly, sampling activity was predominantly limited to spring (April) and autumn (August, September, October), reflecting breeding and territory establishment seasons respectively (Mougeot *et al.*, 2005a; Redpath *et al.*, 2006b). In consequence, months were binned into two seasons (“spring”: January to June; “autumn”: July to December) that captured seasonal differences in nematode burden (supplementary materials S2). Nematode burden was higher in males ($\beta_1 = 0.29 \pm 0.16$; $z_{693} = 1.86$, $P = 0.06$) and old

229 birds ($\beta_1 = 0.61 \pm 0.15$; $z_{693} = 4.22$, $P < 0.001$), confirming expectations from previous studies (Mougeot
230 *et al.*, 2004, 2005d, 2009; Martínez-Padilla *et al.*, 2010; Vergara *et al.*, 2012) and justifying inclusion of
231 sex and age as explanatory variables. Similarly, significant variation in nematode burden was explained
232 by body mass ($\beta_1 = 0.003 \pm 0.001$; $z_{691} = 2.553$, $P = 0.011$), wing length ($\beta_1 = 0.035 \pm 0.008$; $z_{648} =$
233 4.679 , $P < 0.001$) and condition index ($\beta_1 = -0.003 \pm 0.001$; $z_{647} = -3.149$, $p = 0.001$) individually,
234 though all of these variables were stratified by sex and age (supplementary materials S2).

235 Five GWA models were constructed with different biological and statistical complexity to balance out
236 biological consistency and statistical power. The base-line model contained *site*, *period*, *season*, *sex* and
237 *age* variables (model 1; $n = 695$). Two additional models were designed to account for condition-specific
238 covariance in nematode burden. First, by including *body mass* as an additional variable, omitting two
239 observations with missing data (model 2; $n = 693$). Second, by substituting the *condition index* for body
240 mass, omitting 44 further observations with missing data (model 3; $n = 649$). These two models introduce
241 some collinearity among sex, age and body mass or condition index. Although variance inflation factors
242 for these variables were low (< 2), an alternative model was constructed to resolve collinearity. Mixed
243 correspondence analysis (MCA) was applied on the *sex*, *age* and *condition index* variables using the R
244 package *ade4* (Dray & Dufour, 2007) and all three independent principal components were then fitted
245 instead of the original variables (model 4; $n = 649$). This model was also implemented in an alternative
246 version with full sample size (model 5; $n = 695$), where missing condition indices were imputed before
247 MCA, using the multiple imputation method implemented in the R package *missMDA* (Josse & Husson,
248 2012).

249 Associations between individual SNPs and nematode burden were then examined by extending these
250 five models to include an additional categorical explanatory variable that represented the three possible
251 genotypes. To check for allele-specific effects, all models were also implemented for each individual
252 allele independently, by including a single binary explanatory variable representing allele presence. To
253 account for genetic relatedness and population structure, the first eight principal components of the
254 genetic relationship matrix were included as explanatory variables. Although multiple elbow points were
255 apparent when plotting eigenvalues, the greatest change in eigenvalues followed by comparatively steady
256 decrease was after the eighth component (Figure 1). Including more components into models provided
257 qualitatively similar results but incurred a disproportionate power penalty through overfitting.

258 Coefficient estimates and P -values for the genetic term were extracted from each model, and P -values
259 were corrected for multiple testing using the Benjamini-Hochberg false-discovery-rate (FDR) method
260 (Benjamini & Yekutieli, 2005) genome-wide within each model. Results were visualised using Manhattan
261 plots and associations were deemed significant at FDR-corrected $q \leq 0.1$ and highly significant at $q \leq 0.05$.
262 Predicted nematode burdens for each SNP genotype were calculated for significant SNPs as least-square
263 population means from GWA model 2 when keeping all other variables constant at mean values, using the

264 R package *doBy* (Højsgaard & Halekoh, 2013). Effect size was then calculated as the absolute difference
265 between the largest and the smallest estimate, and standard errors were propagated. Finally, to estimate
266 the proportion of phenotypic variance explained by each individual SNP, the SNP genotypes were fitted
267 as the sole predictor of \log_{10} -transformed nematode burden in a simple linear model. The OLS coefficient
268 of determination (r^2) then gives a liberal estimate of the heritability accounted for by a single SNP.

269 Identified SNPs were mapped to the chicken genome to ascertain genomic context and putative physi-
270 ological functions of genes linked to the identified polymorphism. The SNP probe sequences were mapped
271 to the *galGal4* assembly using BLAT (Kent, 2002) and genomic locations were visualised using the UCSC
272 genome browser (Kent *et al.*, 2002). Proximal characterised ENSEMBL chicken genes were then explored
273 in ENSEMBL BIOMARTS (Kinsella *et al.*, 2011), identifying orthologues and extracting GENEONTOLOGY
274 (The Gene Ontology Consortium, 2000) annotations.

275 **Genome partitioning**

276 Genome-wide and chromosome-specific narrow-sense heritabilities (h^2) of nematode burden were esti-
277 mated using genome partitioning analysis as implemented in the software GCTA (Yang *et al.*, 2011a).
278 This method is based on partitioning phenotypic variance across multiple additive genetic variance com-
279 ponents to test whether these components explain significant amounts of phenotypic variance. Each
280 genetic component represents an arbitrary group of genetic markers, usually chromosomes or smaller
281 linkage groups. Additive genetic variance components are estimated from identity-by-state genetic rela-
282 tionship matrices (GRMs) rather than from pedigree data (Yang *et al.*, 2011b; Hill, 2012). In order to
283 satisfy the requirement for a Gaussian quantitative phenotype in GCTA analysis, nematode burden was
284 \log_{10} -transformed (Okada *et al.*, 2010; Brown *et al.*, 2013).

285 Genome-wide heritability was estimated by generating a single GRM in PLINK using all SNPs and
286 fitting the GRM as a single genetic variance component in linear mixed models in GCTA. Statistical
287 significance of the genetic variance was obtained from a log-likelihood ratio test in comparison to a
288 null model without a genetic variance component. Two types of sensitivity analysis were undertaken.
289 First, to ascertain sensitivity to the number of explanatory variables included, three nested analyses were
290 carried out and compared: 1) including sex and age explanatory variables and eight PCA components to
291 account for genetic stratification; 2) adding sampling site and period explanatory variables; and 3) adding
292 sampling season and body mass explanatory variables. This final model contained the same explanatory
293 variables as model 2 in the GWA analysis. Second, the sensitivity to SNP density was examined by
294 generating 500 replicate GRMs from random genome-wide draws of 5–95 % (in 5 % increments) of all
295 SNPs and estimating heritability from each GRM independently. This was carried out for the base-line
296 model without control variables and for the three models with nested control variables.

297 Chromosome-specific heritability was estimated by generating one GRM for each chromosome using

298 only SNPs on that chromosome and fitting this GRM alongside a second GRM based on all genome-wide
299 SNPs apart from those on that chromosome. A likelihood-ratio test was then carried out between this full
300 model and a model excluding the chromosome-specific component. Given a null-model of polygenic ar-
301 chitecture, chromosome-specific heritability should scale linearly with chromosomal gene content (Davis
302 *et al.*, 2013; Robinson *et al.*, 2013). To test this hypothesis, heritability estimates were regressed on
303 numbers of RefSeq genes in chicken chromosomes (*galGal4* assembly), retrieved from the UCSC Table
304 Browser (Karolchik *et al.*, 2004). Given the imbalanced SNP design across chromosomes, heritability
305 was also regressed on numbers of genotyped SNPs and an interaction between both predictors in inde-
306 pendent models. Finally, to test whether the heritability of some chromosomes is larger than expected
307 given chromosomal gene content, another GCTA analysis was carried out where the chromosome-specific
308 component was fitted alongside a GRM using all genome-wide SNPs (Robinson *et al.*, 2013). All analyses
309 were carried out for the four nested models as detailed above for genome-wide heritability estimation.

310 Results

311 Summary statistics

312 Observed and expected heterozygosities at each of 271 SNPs ranged from 0.086–0.534 (median 0.348)
313 and 0.107–0.500 (median 0.349) respectively. A total of 20 SNPs on 10 chromosomes displayed significant
314 departures from Hardy-Weinberg equilibrium ($P \leq 0.05$). However, only three SNPs on chromosome 5
315 remained significant after FDR-correction for multiple testing ($q \leq 0.05$) and none of these displayed
316 extreme genotype distributions or P -values that would suggest genotyping errors. All SNPs were in
317 effective linkage equilibrium, with r^2 ranging from 0.000 to 0.088 (median: 0.001; 99th percentile: 0.014)
318 (supplementary materials S3).

319 Inbreeding coefficients (F_{IS}) of individual samples were normally distributed, ranging from -0.251 to
320 0.247 with a mean of 0.011 ± 0.085 SD (supplementary materials S3). Relatedness among individuals
321 was approximately centred on zero (median: -0.003 ; 1st percentile: -0.144 ; 99th percentile: 0.156)
322 (supplementary materials S3). This distribution is consistent with a simulated distribution of genetic
323 relatedness estimates for unrelated red grouse from neutral polymorphisms (Piertney *et al.*, 1999), and as
324 such suggests that the genotyped individuals are effectively unrelated. Population genetic differentiation
325 among the five sampling sites was very low ($F_{ST} = 0.005$), though the English site was more strongly
326 differentiated from the Scottish sites ($F_{ST} = 0.020$), as would be expected under isolation-by-distance.
327 Accordingly, little genetic clustering was apparent in PCA eigenvector plots (Figure 1).

Genome-wide association analysis

GWA analysis yielded support for associations between nematode burden and genotypes or alleles of five SNPs on five chromosomes (Figure 2). Congruence among models incorporating sex and age (model 1) or sex, age and body mass (model 2) variables was very high, highlighting genotypes of all five SNPs and alleles of three SNPs (Table 2). However, models incorporating sex, age and condition index or equivalent principal components of mixed correspondence analysis (models 3 and 4) highlighted weaker associations ($q \leq 0.1$) for alleles or genotypes in only three of these five SNPs (Table 2). These models probably suffered from decreased power due to reduced sample sizes, given that the equivalent model 5 with imputed principal components (and hence full sample size) yielded almost identical results to models 1 and 2 (Table 2). Notwithstanding, each of these five SNPs is highlighted by at least three models, indicating that these are robust results despite the potential for reduced power in some models.

Of these, SNPs X1407, X5104 and X1375 were best supported, displaying consistently strong associations ($q \leq 0.05$) of genotypes or alleles in the three full models, and weaker associations ($q \leq 0.1$) of genotypes or alleles in the two reduced models (Table 2). Genotypes in two further SNPs X2277 and X2298 were moderately supported across models, but there was no support for alleles in any model. Some models additionally highlighted weak associations ($q \leq 0.1$) for two further SNPs on chromosome 1 and 2 (not shown), but these were inconsistent and as such were disregarded. Predicted nematode burdens among genotypes of the five highlighted SNPs indicated additive allelic effects in most cases, and absolute effect sizes ranged from 313–556 worms per bird (Figure 3). None of these SNPs deviated from Hardy-Weinberg equilibrium and hence there was no evidence for heterozygosity advantage or disadvantage. The proportion of phenotypic variation explained by these SNPs individually in simple linear models ranged from 0.1–2.8 % (sum: 4.9 %), which should be taken as liberal best-case estimates (Table 2). Full association statistics for all models and SNPs alongside tests for Hardy-Weinberg equilibrium are available in supplementary materials S4–S5.

Homology with the chicken genome indicated that SNPs X1407 and X1375 were located in introns of genes *MAPKBP1* (mitogen-activated protein kinase binding protein 1) and *KLHL34* (kelch-like family member 34) on chromosomes 5 and 1 respectively (Table 2). All other SNPs were located in non-coding regions on chromosomes 10, 13 and 20, approximately 6–540 kbp (equivalent to 0.02–1.51 cM assuming 2.8 cM/Mbp, Hillier *et al.*, 2004) remote from the following upstream and downstream genes: *RAPGEF6* (Rap guanine nucleotide exchange factor 6), *FNIP1* (folliculin interacting protein 1), *CEBPB* (CCAAT/enhancer binding protein β), *PTPN1* (protein tyrosine phosphatase, non-receptor type 1), *NR2F2* (nuclear receptor subfamily 2, group F, member 2) and *MCTP2* (multiple C2 domains, transmembrane 2). The putative functions of these linked genes involve innate and adaptive immune system processes, protein degradation and energy metabolism (Table 2; supplementary materials S6).

Genome partitioning

Estimated genome-wide heritability of nematode burden based on all SNPs together was $h^2 = 0.294$. This estimate decreased rapidly when increasing numbers of explanatory variables were added to the model (Table 3). Sensitivity analysis by randomly sampling subsets of genome-wide SNPs and repeating heritability estimation identified an asymptotic relationship between heritability and SNP density, though this was much weaker when additional explanatory variables were included (supplementary materials S7). Although these heritability estimates are likely to increase at higher SNP densities, these SNPs capture a statistically significant amount of heritability that is consistently larger than the heritability estimates of the five SNPs identified using GWA.

Partitioning genome-wide heritability across chromosomes was limited by variation in SNP density and power reduction when multiple explanatory variables were included in the models. Chromosomes 21, 22, 24, 26 and 28 could not be analysed because of insufficient SNP density. For other chromosomes, some models failed to converge for particular combinations of variables. Estimates of chromosome-specific heritability across all models ranged from 10^{-6} to 0.099, of which 1–11 estimates were significantly different from zero (Figure 4). The summed chromosome-specific heritabilities within models were very similar to genome-wide heritabilities, despite the reduced number of chromosomes analysed (Table 3), suggesting some covariance across chromosomes that may inflate individual estimates. Heritability estimates were significantly positively associated with chromosomal gene content and number of genotyped SNPs in simple models only, presumably due to reduced power in more complex models (Table 3). Although these two predictors were highly correlated as a consequence of the SNP design (Spearman's $\rho = 0.93$; $P < 0.001$), there was a significant interaction among them (Table 3), indicating an effect beyond the mere number of genotyped SNPs per chromosome. All these relationships were primarily driven by chromosomes 1 and 2, which had particularly large heritability, size and SNP density (Figure 4). When these two chromosomes were removed, the relationships remained significant for the model without control variables only (not shown). Chromosomes 5 and 17 displayed significantly ($p < 0.05$) greater heritability than expected from their gene contents in some models (excess $h^2 = 0.029 \pm 0.020$ and $h^2 = 0.017 \pm 0.017$ respectively; Figure 4). Some weaker evidence for excess heritability was also apparent for chromosomes 1 ($h^2 = 0.043 \pm 0.034$), 13 and 14 ($h^2 = 0.010 \pm 0.012$ in both cases). Total excess heritability across all chromosomes was $h^2 = 0.066 - 0.089$.

These results were partially consistent with the GWA results. Chromosomes 1 and 5 contained the best supported SNPs X1407 and X1375, and chromosome 1 also contained other suggestive SNPs (Figure 2). SNPs on chromosomes 10, 13 and 20 were highlighted in the GWA analysis, but these chromosomes showed no significant excess heritability. In contrast, chromosome 17 displayed evidence for excess heritability, but no SNP on this chromosome was highlighted in the GWA analysis.

396 Discussion

397 This study examined the broad genome-wide basis of variation in gastrointestinal nematode burden
398 among red grouse individuals. Genome-wide association identified five SNPs that were closely linked to
399 novel genes putatively involved in multiple physiological processes beyond the immune system, consistent
400 with the broad functions of previously identified genes using candidate gene approaches (Wenzel *et al.*,
401 2015; Wenzel & Piertney, 2015) and epigenetic DNA methylation analysis (Wenzel & Piertney, 2014).
402 Genome partitioning indicated moderate genome-wide heritability of nematode burden, and highlighted
403 that some chromosomes contain additive genotypic variants of individually small effects that represent
404 disproportionate amounts of heritability. These findings contribute to elucidating the broad genomic
405 basis of parasite susceptibility (Wilfert & Schmid-Hempel, 2008) and suggest that nematode burden in
406 grouse may be somewhere between a purely polygenic phenotype corresponding to Fisher's infinitesimal
407 model (Fisher, 1919) and a purely oligogenic phenotype where few loci explain a large proportion of
408 phenotypic variance (Rockman, 2012).

409 *T. tenuis* infection is localised to the caecum where adult worms cause haemorrhaging and tissue
410 necrosis with substantial impact on grouse condition and fitness (Watson *et al.*, 1987; Hudson *et al.*,
411 1992; Delahay *et al.*, 1995; Delahay & Moss, 1996). The majority of genes closely linked to the five SNPs
412 identified through GWA are putatively involved in immune system processes that may be associated with
413 chronic intestinal *T. tenuis* infection: MAPKBP1 is involved in signal transduction during inflammation
414 and in intestinal homeostasis (Lecat *et al.*, 2012). In humans, MAPKBP1 (alias JNKBP1) is an antagonist
415 to the NOD2 receptor which recognises bacterial cell wall components and helps orchestrate an innate and
416 adaptive immune response (Lecat *et al.*, 2012). Failure to regulate expression of NOD2 in intestinal tissue
417 by MAPKBP1 disrupts cytokine signal transduction and is implicated in Crohn's disease, a degenerative
418 disorder of the intestine (Hugot *et al.*, 2001; Lecat *et al.*, 2012). Similarly, CEBPB is involved in signal
419 transduction following an inflammatory immune response (Ramji & Foka, 2002), and may specifically
420 regulate antibacterial activity of macrophages and repair of necrotic tissue (Ruffell *et al.*, 2009). Other
421 genes represent broader immunoregulatory factors: RAPGEF6 is a guanine nucleotide exchange factor
422 for RAP GTPases (Kuiperij *et al.*, 2003), which are involved in regulating signalling interactions between
423 antigen-presenting cells and T-cells and in regulating leukocyte integrin activation (Katagiri *et al.*, 2002;
424 Scheele *et al.*, 2007). FNIP1 is an adapter protein that interacts with a range of factors that regulate
425 cellular energy metabolism (Park *et al.*, 2012). Disruption of FNIP1 function arrests development of B
426 lymphocytes and invariant natural killer T-cells due to an inability to regulate metabolic homeostasis
427 during cell proliferation, particularly under metabolic stress (Park *et al.*, 2012, 2014). MCTP2 is involved
428 in intercellular signal transduction and has been implicated in various neurological disorders such as
429 schizophrenia and autism (Shin *et al.*, 2005; Djurovic *et al.*, 2009). Its link to parasite infection is unclear,
430 but transcriptomic studies suggest that it may be cryptically involved in immune system function (Verner

431 *et al.*, 2012), possibly through regulation of T-helper cell differentiation and function (Äijö *et al.*, 2012).

432 The remaining identified genes represent a range of other physiological pathways that may play a
433 key role in underpinning host-parasite interactions in red grouse. KLHL34 is part of the evolutionarily
434 highly conserved kelch-like protein family whose functions are not well understood, though they are
435 associated with a range of human diseases (Dhanao *et al.*, 2013). The best studied function is a role
436 in protein ubiquitination (Gupta & Beggs, 2014), which links kelch-like proteins to a broad range of
437 cellular processes including inflammatory immune response and protein degradation, potentially as part
438 of a xenobiotics detoxification response (Ben-Neriah, 2002; von Mikecz, 2006). In infected red grouse,
439 such xenobiotics may originate from primary *T. tenuis* infection or from secondary pathogen infection
440 following caecal haemorrhaging (Watson *et al.*, 1987). Further, both NR2F2 (alias COUP-TFII) and
441 PTPN1 are involved in regulating glucose homeostasis, energy expenditure and adipogenesis (Li *et al.*,
442 2009; Tsou *et al.*, 2012), which could plausibly impact physiological condition with broad consequences
443 for the ability of red grouse to cope with parasite infection (Sheldon & Verhulst, 1996; Svensson *et al.*,
444 1998).

445 Although GWA provides no direct evidence for a mechanistic involvement of these genes, their putative
446 physiological functions are broadly consistent with novel candidate genes previously identified through
447 comparative transcriptomics between infected and nematode-free grouse (Wenzel *et al.*, 2015; Wenzel
448 & Piertney, 2015). These candidate genes represent innate and adaptive immune system processes,
449 in particular eosinophil-mediated parasite expulsion and antimicrobial peptides, alongside key enzymes
450 in the xenobiotics detoxification and oxidative stress pathways, and broad cell proliferation regulators
451 (Wenzel & Piertney, 2015). Together, these insights reinforce the notion that host-parasite interactions
452 in red grouse are not a primarily immunological matter. This provides an intriguing perspective on
453 the classic immunological paradigm of host life-history trade-offs (Sheldon & Verhulst, 1996; Lochmiller
454 & Deerenberg, 2000; Zuk & Stoehr, 2002; Schmid-Hempel, 2003; Owen *et al.*, 2010). In spite of well-
455 established immunological links in red grouse (Mougeot & Redpath, 2004; Webster *et al.*, 2011a; Wenzel
456 *et al.*, 2013; Wenzel & Piertney, 2015) and sheep or cattle species (e.g., Beh *et al.*, 2002; Davies *et al.*,
457 2006; Beraldi *et al.*, 2007; Silva *et al.*, 2012; Riggio *et al.*, 2013), the cost of chronic parasite infection,
458 particularly in the case of nematodes, may well extend to other physiological and perhaps even behavioural
459 categories (Thomas *et al.*, 2010; Biron & Loxdale, 2013; Poulin, 2013). In red grouse, for example, DNA
460 methylation states at specific loci linked to genes involved in immune system, metabolism, cell cycle
461 regulation and epigenetic mechanisms have been associated with parasite burden in the field (Wenzel
462 & Piertney, 2014), suggesting that chronic parasite infection may have a broad epigenetic component
463 (Poulin & Thomas, 2008). This perspective is still consistent with the classic idea of life-history trade-
464 offs (Hamilton & Zuk, 1982; Sheldon & Verhulst, 1996; Lochmiller & Deerenberg, 2000), but suggests
465 that physiological condition is an amalgam of a multitude of cellular functions that may either directly

466 contribute to parasite defence or affect its efficacy indirectly by broad effects on energy metabolism or
467 molecular signalling cascades (Hill, 2011).

468 The genome partitioning analysis broadly supported this perspective. In spite of low heritability of the
469 five individual SNPs, genome-wide heritability using all SNPs together was moderate ($h^2 = 0.07 - 0.29$)
470 and within the range of reported heritability estimates for nematode burden in sheep (Dominik, 2005;
471 Stear *et al.*, 2007) and chicken (Kaufmann *et al.*, 2011; Wongrak *et al.*, 2014). This could imply that
472 parasite susceptibility may to some extent be a polygenic phenotype involving large numbers of genes
473 with individually small, but large joint effects (Yang *et al.*, 2011b; Hill, 2012; Rockman, 2012). However,
474 chromosomes 1, 5 and 17 contributed disproportionate amounts to total heritability despite different
475 SNP densities, suggesting that nematode burden may not be primarily based on a large number of
476 genome-wide polymorphisms with small effects, rather than on moderate numbers of small effect genes
477 in these three chromosomes. In fact, chromosomes 1 and 5 contain two SNPs highlighted by the present
478 GWA analysis and seven previously identified novel candidate genes for response to parasite infection
479 (Wenzel *et al.*, 2015), most of which have been confirmed to explain variation in parasite load in the
480 field (Wenzel & Piertney, 2015). The two identified SNPs are not in the vicinity of any of these genes
481 (40–80 Mbp and 7–8 Mbp distance on chromosomes 1 and 5 respectively), highlighting their novelty. This
482 chromosomal congruence across candidate genes, GWA and genome partitioning provides support for a
483 primarily oligogenic basis of parasite susceptibility in chromosomes 1 and 5, making them priority targets
484 for fine-mapping novel genomic candidate regions in addition to those already discovered. Chromosome 17
485 may also be a good target on the quest for the “missing heritability”, though it has not been implicated
486 in GWA or previous candidate genes studies to contain large-effect genes. The genome partitioning
487 results are consistent with a study in sheep, where five chromosomes explained disproportionate amounts
488 of variance in gastrointestinal nematode burden, despite an overall relationship between chromosomal
489 heritability and gene content (Al-Kalaldeh *et al.*, 2013).

490 Despite some congruence in numbers and function of identified genes across red grouse studies with
491 different approaches, there are some inconsistencies that deserve consideration. First, some SNPs high-
492 lighted by the GWA analysis are on chromosomes that do not display excess heritability or contain
493 any previously identified candidate genes. Similarly, many previously identified candidate genes are on
494 chromosomes that are not highlighted by GWA or genome partitioning (Wenzel *et al.*, 2015; Wenzel &
495 Piertney, 2015). Each of these three techniques highlights a different set of genomic regions overall, reflect-
496 ing different ways of statistical inference with different power and potential for false positives (Ellegren
497 & Sheldon, 2008; Amos *et al.*, 2011). In the present study, one caveat to consider with respect to genome
498 partitioning in particular is that SNP density was low and the study design was based on comparatively
499 few samples. Although few chromosomes displayed significant total heritability, we were able to detect
500 three chromosomes with more heritability than expected, suggesting that power was sufficient to detect

501 large heritability effects but may have been insufficient to detect more subtle effects. A large-scale study
502 with thousands of samples and markers will be required to fully reconcile these differences and resolve
503 the underlying genomic architecture (Lee *et al.*, 2012).

504 Second, in spite of the putative functional importance of the linked genes identified from orthology
505 with chicken, the actual molecular effects of the identified SNPs are unclear. None of the identified
506 SNPs were in exonic regions, suggesting that the association of alleles with parasite load may not be due
507 to amino acid substitution in the gene product, though these SNPs may be linked to unknown exonic
508 SNPs with such effect. Instead, intronic SNPs may affect transcript splicing or downstream mRNA
509 processing with effect on translation dynamics (Pagani & Baralle, 2004; Hunt *et al.*, 2014). Similarly,
510 non-coding SNPs may affect gene expression or chromatin organisation through epigenetic mechanisms
511 such as miRNAs or transcription factors (Pagani & Baralle, 2004; Chamary *et al.*, 2006). Epigenetic
512 processes may play a particularly important role in the red grouse system because DNA methylation at a
513 range of non-coding genomic regions is associated with parasite load (Wenzel & Piertney, 2014), SNPs in
514 previously identified candidate genes associated with parasite load were predominantly non-synonymous,
515 intronic or untranslated (Wenzel & Piertney, 2015) and some of these genes are themselves involved in
516 regulating gene expression (Wenzel & Piertney, 2015).

517 In summary, genome-wide association and genome partitioning have added new insight to our previous
518 work on red grouse that highlighted a range of genetic polymorphisms in or close to candidate genes
519 that are linked to nematode burden among grouse individuals, accounting for differences of as much
520 as 666 worms per bird (Wenzel & Piertney, 2015). By combining a range of strategies for examining
521 the genetic basis of a complex phenotype, we have reinforced the idea that parasite susceptibility may
522 involve a considerable number of genes involved in a range of physiological categories beyond the immune
523 system. We further identified particular chromosomes as priority targets for fine-mapping polymorphisms
524 of small effect in the quest for the missing heritability of nematode burden. These findings have painted
525 a consistent picture of the genetic basis of nematode burden in red grouse and illustrate that genome
526 partitioning is a powerful addition to the classic strategies for studying the genetic basis of complex
527 phenotypes.

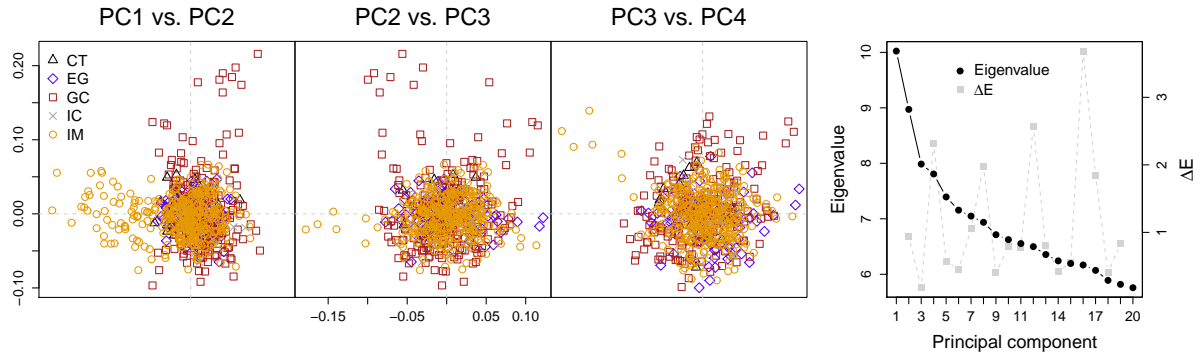


Figure 1: Principal components analysis of the genome-wide genetic relationship matrix based on 271 SNPs. The first three panels are scatterplots of sequential combinations of the first four principal components (eigenvectors), highlighting sampling sites (CT = Catterick; EG = Edinglassie; GC = Glas Choille; IC = Invercauld; IM = Invermark). The last panel plots the eigenvalues of the first 20 principal components and an *ad hoc* statistic ΔE (quotient of the eigenvalue change to the following versus to the preceding principal component) whose peaks aid identification of elbow points.

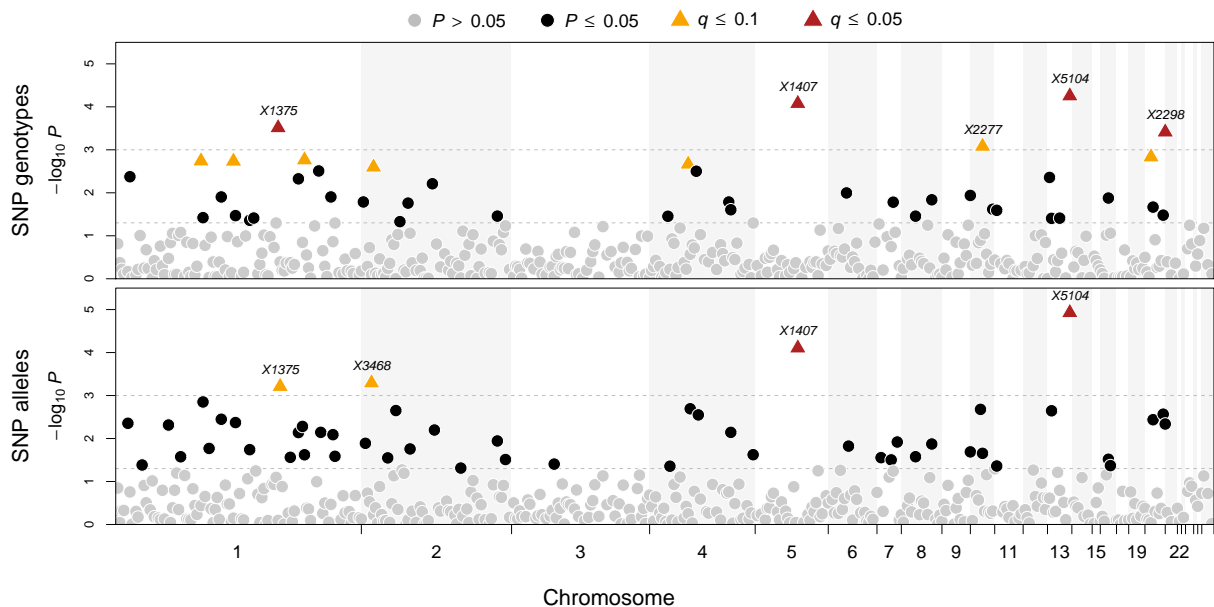


Figure 2: Genome-wide association results (model 2 with full set of control variables, Materials & Methods) for nematode burden using genotypes (top) or alleles (bottom) of 271 SNPs ordered along chromosomes based on orthology with the chicken genome. Each symbol represents statistical significance (P -value) of a single test (genotype comparison or allele presence). Statistical significance before and after correction for multiple testing (false discovery rate correction; q -values) is indicated by symbol shape and colours. Two common single-test significance thresholds ($P = 0.05$ and $P = 0.001$) are indicated by dashed lines for illustrative purposes.

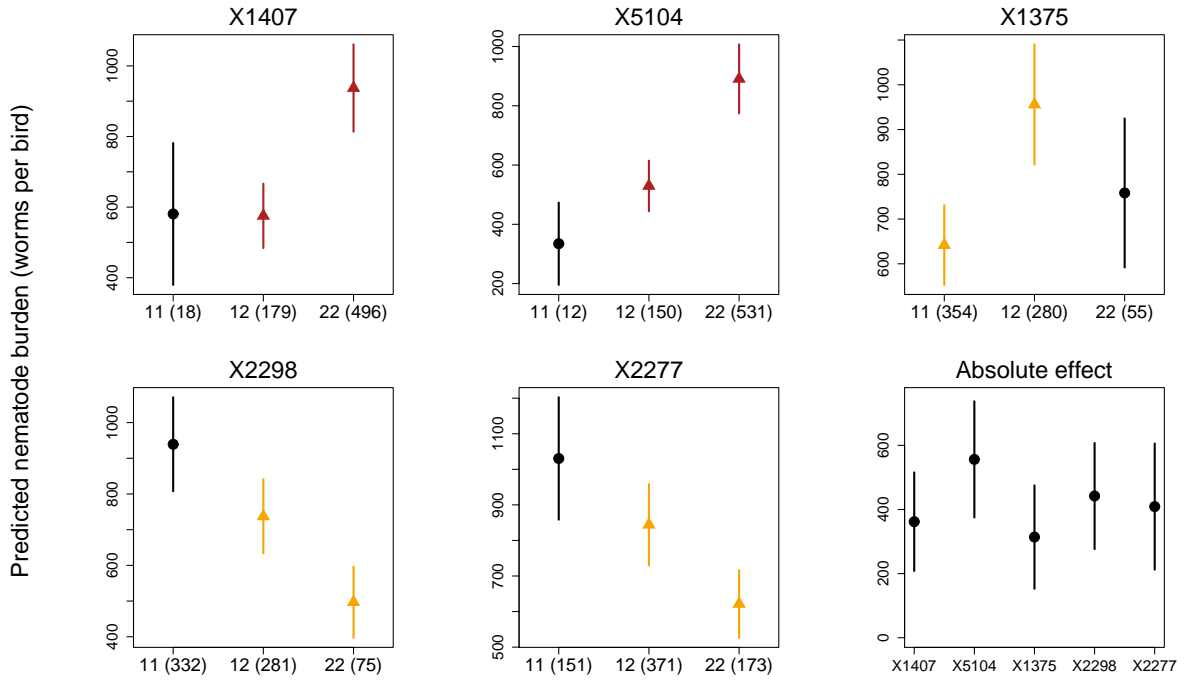


Figure 3: Predicted nematode burden according to three possible genotypes (11 and 22: homozygote for allele 1 or 2; 12: heterozygote) at five SNPs identified through genome-wide association. Least-square estimates (\pm SE) of model 2 (all other variables kept constant at mean values) are presented in the first five panels. Coloured triangles denote statistically significant ($q \leq 0.1$) genotype comparisons in the fitted model as in Figure 2. Numbers in brackets indicate sample sizes. The bottom right panel indicates overall effect size as the largest absolute difference in predicted estimates (\pm propagated SE).

Table 1: Sampling locations, sampling years and sample sizes by sex and age (young: hatched in same year; or old: >1 year) of 695 red grouse samples. Detailed sample information is presented in supplementary materials S2.

Site	Latitude	Longitude	Years	Sex		Age		Total
				F	M	Y	O	
Edinglassie (EG)	57.22 ^o	-3.19 ^o	2000–2012	13	86	31	68	99
Glas Choille (GC)	57.12 ^o	-3.32 ^o	1995–2003	63	136	131	68	199
Invercauld (IC)	57.08 ^o	-3.30 ^o	2012	15	16	10	21	31
Invermark (IM)	56.90 ^o	-2.88 ^o	2011–2012	109	238	76	271	347
Catterick (CT)	54.33 ^o	-1.87 ^o	2006	0	19	12	7	19
Total				200	495	260	435	695

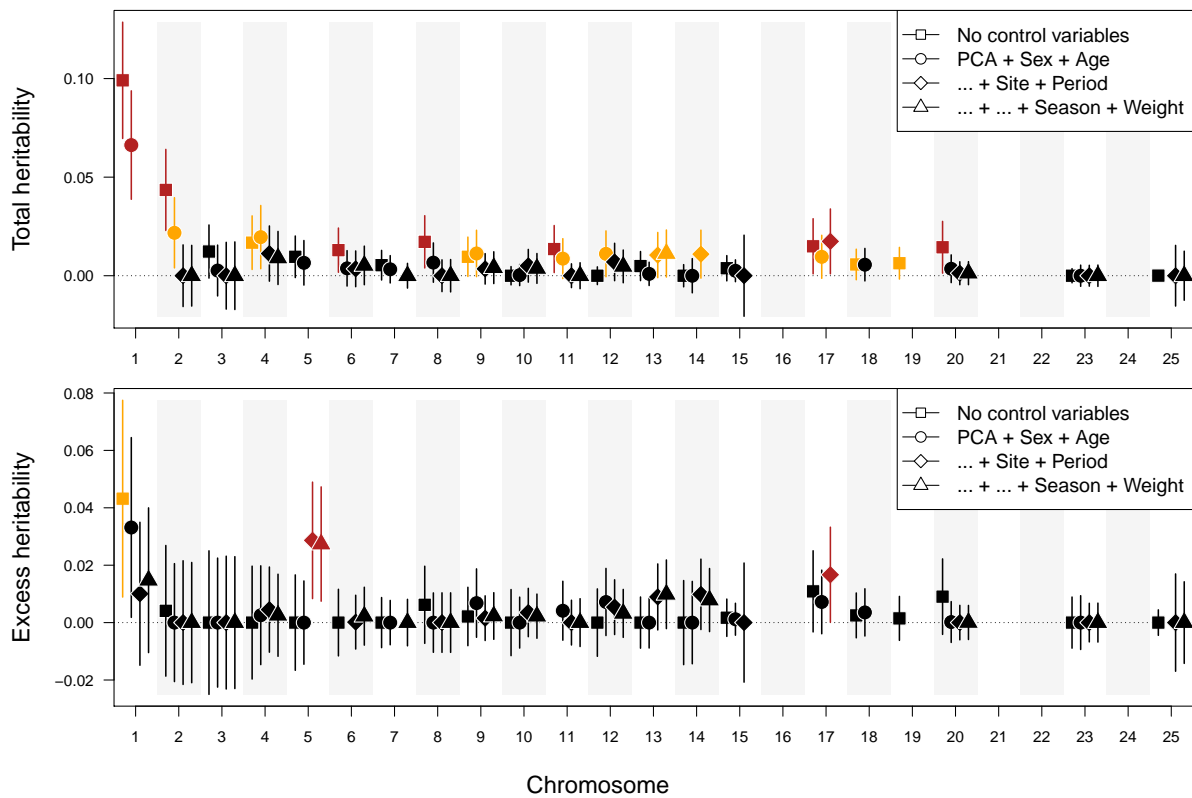


Figure 4: Chromosome-specific total heritability (top panel) and excess heritability of nematode burden unaccounted for by gene content (bottom panel). Heritability was partitioned across chromosomes using linear mixed models incorporating chromosome-specific genetic variance and three nested sets of control variables. Both panels display heritability estimates (\pm SE) with colour-coded statistical significance (black: $p > 0.1$; orange: $p \leq 0.1$; red: $p \leq 0.05$).

Table 2: Characterisation of SNPs significantly associated with nematode burden using genome-wide association. Statistical significance is indicated by asterisks for genotypes and alleles across five negative binomial models. Each model incorporates sampling site, period and season alongside eight principal components of the genetic relationship matrix as control variables. Additional control variables are indicated (MC: principal component following multiple correspondence analysis on age, sex and condition index). A liberal heritability estimate is indicated as the coefficient of determination (r^2) of a simple OLS model with the SNP as the sole predictor. Minor allele frequency (MAF), observed (H_O) and expected (H_E) heterozygosities are presented with the P -value for Hardy-Weinberg equilibrium (P_{HWE}). Chicken chromosome number and functional information on proximal ENSEMBL genes are given below.

		SNPs (genotype/allele)				
Model	n	X1407	X5104	X1375	X2298	X2277
1: +sex + age	695	** / **	** / **	** / **	** / n.s.	* / n.s.
2: +sex + age + mass	693	** / **	** / **	** / **	** / n.s.	* / n.s.
3: +sex + age + condition	649	* / *	n.s. / *	n.s. / n.s.	* / n.s.	n.s. / n.s.
4: +MC1 + MC2 + MC3	649	* / *	n.s. / *	n.s. / n.s.	* / n.s.	n.s. / n.s.
5: +MC1 + MC2 + MC3 (imputed)	695	** / **	** / **	** / *	** / n.s.	* / n.s.
Heritability (OLS r^2)		0.028	0.006	0.010	0.004	0.001
MAF		0.155	0.126	0.283	0.313	0.484
H_O		0.258	0.217	0.406	0.408	0.534
H_E		0.262	0.220	0.406	0.430	0.500
P_{HWE}		0.665	0.728	1	0.184	0.081
Chromosome	5	13	1	1	20	10
Type of mutation	intronic	non-coding	intronic	intronic	non-coding	non-coding
Proximal genes (distance)	MAPKBP1	RAPGEF6 (6 kbp), FNIP1 (23 kbp)	KLHL34	KLHL34	CEBPB (30 kbp); PTPN1 (127 kbp)	NR2F2 (540 kbp); MCTP2 (215 kbp)
Ensembl ENSGALG gene ID	ENSGALG000000008692	ENSGALG00000006569; ENSGALG00000017462	ENSGALG00000016388	ENSGALG00000008014; ENSGALG00000008010	ENSGALG00000007000; ENSGALG00000006981	ENSGALG00000007000; ENSGALG00000006981
Key GENEONTOLOGY term	GO:0005515 protein binding	GO:0007264 small GTPase mediated signal transduction; GO:0002327 immature B cell differentiation	GO:0005515 protein binding	GO:0006954 inflammatory response; GO:0008286 insulin receptor signaling pathway	GO:0006629 lipid metabolic process; GO:0019722 calcium-mediated signaling	GO:0006629 lipid metabolic process; GO:0019722 calcium-mediated signaling
Physiological process	immune system	immune system	protein ubiquitination	immune system; energy metabolism	immune system; energy metabolism; immune system	energy metabolism; immune system

** $q \leq 0.05$; * $q \leq 0.1$; n.s.: $q > 0.1$

Table 3: Heritability estimates of nematode burden based on additive genetic variance derived from identity-by-state genetic relationship matrices. Genome-wide heritability estimates (h^2) are presented with standard errors (SE), and statistical significance is indicated by the likelihood-ratio test statistic LRT (compared to a model without the genetic variance component) and the associated P -value. Alongside, the sum of chromosome-specific heritability estimates is given with results of three independent linear regressions on chromosome-specific numbers of genes, numbers of genotyped SNPs, or the interaction among both predictors. All results are presented for four nested models with increasing numbers of control variables.

Model	Genome-wide h^2				Chromosome-specific h^2				Genes-SNPs interaction
	h^2	SE	LRT	P	$\sum h^2$	Number of genes	Number of SNPs		
No control variables	0.294	0.041	108.643	$\ll 0.001$	0.290	$F_{1,18} = 55.39; P \ll 0.001$	$F_{1,18} = 60.62; P \ll 0.001$	$F_{3,16} = 51.21; P < 0.001$	
+PCA + sex + age	0.182	0.044	25.347	$\ll 0.001$	0.184	$F_{1,17} = 37.58; P \ll 0.001$	$F_{1,17} = 38.84; P \ll 0.001$	$F_{3,15} = 35.93; P < 0.001$	
+PCA + sex + age + site + period	0.075	0.040	4.558	0.016	0.070	$F_{1,14} = 0.16; P = 0.697$	$F_{1,14} = 0.23; P = 0.639$	$F_{3,12} = 0.45; P = 0.722$	
+PCA + sex + age + site + period + season + mass	0.070	0.040	3.970	0.023	0.039	$F_{1,12} = 0.01; P = 0.999$	$F_{1,12} = 0.01; P = 0.951$	$F_{3,10} = 0.901; P = 0.474$	

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Supplementary materials

- S1: Characterisation of 384 SNPs based on orthology with the chicken genome
- S2: Distribution of sample sizes and parasite load across sampling-related and phenotypic categories
- S3: SNP linkage disequilibrium and distributions of sample inbreeding coefficients (F_{IS}) and relatedness.
- S4: GWA model statistics and tests for SNP Hardy-Weinberg equilibrium
- S5: Manhattan plots of all five models used for GWA analysis
- S6: Full GENEONTOLOGY terms in *biological process* ontology of genes linked to SNPs identified through GWA
- S7: Sensitivity analysis of genome-wide heritability estimation to SNP density

Data accessibility

- Genotype data (DataDryad: doi 10.5061/dryad.02pr5)
- Metadata (sampling/phenotype information) (DataDryad: doi 10.5061/dryad.02pr5)

Author contributions

SBP conceived and designed the study. MAW and MCJ performed field and lab work. MCJ and AD developed SNP markers. MAW analysed the data. MAW and SBP wrote the manuscript.

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