

2018-11

# Neutral variation does not predict immunogenetic variation in the European grayling (*Thymallus thymallus*) - implications for management

Huml, JV

<http://hdl.handle.net/10026.1/12250>

---

10.1111/mec.14864

Molecular Ecology

Wiley

---

*All content in PEARL is protected by copyright law. Author manuscripts are made available in accordance with publisher policies. Please cite only the published version using the details provided on the item record or document. In the absence of an open licence (e.g. Creative Commons), permissions for further reuse of content should be sought from the publisher or author.*

1 **Title:**

2 **Neutral variation does not predict immunogenetic variation in the European**  
3 **grayling (*Thymallus thymallus*) – implications for management**

4

5 Huml J.V.<sup>1,3</sup>, Taylor M.I.<sup>2</sup>, Harris W. Edwin<sup>1</sup>, Sen R.<sup>1</sup>, Ellis J.S<sup>3</sup>.

6

7 Accepted 29-8-2018

8

9 Addresses:

10 1 School of Science & Environment, Manchester Metropolitan University, Oxford  
11 Road, Manchester, M15 6BH, UK

12 2 School of Biological Sciences, University of East Anglia, Norwich Research Park,  
13 Norwich, NR4 7TJ, UK

14 3 School of Biological and Marine Sciences, University of Plymouth, Drake Circus,  
15 Plymouth, PL4 8AA, UK

16

17 Keywords: major histocompatibility complex; conservation genetics; population  
18 augmentation; European grayling (*Thymallus thymallus*); amplicon sequencing

19

20 Corresponding author: Vanessa Huml, School of Biological and Marine Sciences,  
21 Plymouth University, Drake Circus, Plymouth, PL4 8AA, UK. Email:

22 [vanessa.huml@plymouth.ac.uk](mailto:vanessa.huml@plymouth.ac.uk), Fax: none

23

24

25 Running title: Neutral and functional variation in grayling

26

27

## 28 Abstract

29 Preservation of genetic diversity is critical to successful conservation and there is  
30 increasing demand for the inclusion of ecologically meaningful genetic information in  
31 management decisions. Supportive breeding programmes are increasingly  
32 implemented to combat declines in many species, yet their effect on adaptive genetic  
33 variation is understudied. This is despite the fact that supportive breeding may  
34 interfere with natural evolutionary processes. Here, we assessed the performance of  
35 neutral and adaptive markers (Major Histocompatibility Complex; MHC) to inform  
36 management of European grayling (*Thymallus thymallus*), which routinely involves  
37 supplementation of natural populations with hatchery-reared fish (stocking). This  
38 study is the first to characterize MH II DAA and DAB loci in grayling and to  
39 investigate immune genetic variation in relation to management practice in this  
40 species. High-throughput Illumina sequencing of 'introduced', 'stocked native' and  
41 'non-stocked native' populations revealed significantly higher levels of allelic richness  
42 and heterozygosity for MH markers than microsatellites exclusively in non-stocked  
43 native populations. Likewise, significantly lower differentiation at the MH II than for  
44 microsatellites was apparent when considering non-stocked native populations, but  
45 not stocked populations. We developed a simulation model to test the effects of  
46 relaxation of selection during the early life stage within captivity. Dependent on the  
47 census population size and stocking intensity, there may be long-term effects of  
48 stocking on MH II, but not neutral genetic diversity. This is consistent with our  
49 empirical results. This study highlights the necessity for considering adaptive genetic

50 variation in conservation decisions and raises concerns about the efficiency of  
51 stocking as a management practice.

52

## 53 Introduction

54 In order to manage biodiversity in the light of elevated rates of species extinctions  
55 (Ceballos et al. 2015), it is acknowledged that the consideration of genetic variation  
56 is crucial (Pertoldi et al., 2007; Sgrò et al., 2011). In the short-term, the loss of  
57 genetic variation directly impacts population viability due to negative effects  
58 associated with inbreeding depression (Spielman et al., 2004). In the long-term,  
59 populations are expected to persist in a changing environment only if they harbour  
60 sufficient adaptive potential (Duploux et al., 2013). The management of adaptive  
61 genetic variation is therefore at the core of conservation genetics (Allendorf et al.,  
62 2010). Indeed, there is great promise in measuring adaptive genetic variation  
63 because it makes the consideration of evolutionary dynamics possible, which may  
64 greatly improve the effectiveness of conservation planning (Brodersen and  
65 Seehausen, 2014). Assessing adaptive genetic variation directly is important  
66 because neutral variation may be affected differently by demographic processes  
67 (e.g. through bottlenecks (Ejsmond and Radwan, 2011; Sutton et al., 2011)), thus  
68 conservation decisions based solely on assessment of neutral variation may be  
69 poorly informed. Whilst neutral marker surveys continue to be in wide use in  
70 conservation genetics due to their convenience, repeatability and low cost,  
71 increasingly, there are calls to study the dynamics of functional genetic variation  
72 underlying ecologically meaningful traits in conservation genetic studies (Piertney  
73 and Webster, 2010). Despite the great promise in monitoring and managing adaptive

74 genetic variation, it has only become feasible to do so at a large scale in recent  
75 years due to the increasing accessibility of whole genomic screening techniques  
76 (Koboldt et al., 2013).

77

78 The assessment of adaptive genetic variation is particularly important where species  
79 management includes captive breeding. Although captive breeding is an important  
80 management tool to reduce biodiversity loss (Frankham, 2008; Griffiths and  
81 Pavajeau, 2008), it can interfere with adaptive processes (Ayllon et al., 2006). A  
82 good example is supportive breeding where adults are maintained temporarily in  
83 captivity to produce offspring that are released into the wild population. Neff et al.  
84 (2011) found that evidence for successful restoration of stable populations through  
85 supportive breeding is rare. Failing to preserve adaptive genetic variation is  
86 potentially one of the main causes of the ineffectiveness of current supportive  
87 breeding programmes, but more evidence is required to assess this (Neff et al.,  
88 2011). Supportive breeding is predicted to affect both neutral and adaptive genetic  
89 diversity in some contexts (like the reduction of the effective population size through  
90 unequal reproductive contributions of hatchery fish (Ryman and Laikre, 1991)), but in  
91 others may only affect adaptive and not neutral variation. For example, both the lack  
92 of natural selection acting on early life stages (Lynch and O'Hely, 2001) (which might  
93 be particularly important in species with high rates of juvenile mortality (de Eyto et  
94 al., 2011)), and the lack of natural mate choice in supportive breeding programmes  
95 can interfere with the preservation of adaptive genetic variation (Quader, 2005). It is  
96 therefore crucial to enhance our understanding of the effects of supportive breeding  
97 on adaptive genetic variation for this management technique to become a more  
98 fruitful conservation tool.

100 An important adaptive marker in the context of optimizing fitness of offspring from  
101 artificial breeding programmes is the immune related Major Histocompatibility  
102 Complex (MHC)(Hedrick, 2003; Pitcher & Neff, 2007; Ujvari & Belov, 2011). Protein  
103 products of the MHC have a central role in the recognition and elimination of foreign  
104 peptides and pathogens (Zinkernagel and Doherty, 1974). A large body of evidence  
105 demonstrates an association between MHC variation or specific MHC variants with  
106 overall or pathogen specific resistance (e.g. Evans and Neff, 2009; Meyer-Lucht and  
107 Sommer, 2005; Miller et al., 2004; Savage and Zamudio, 2011). Pathogen-mediated  
108 selection through negative frequency dependent selection and heterozygote  
109 advantage are thought to be the main mechanisms maintaining high diversity in the  
110 MHC and can lead to habitat specific MHC gene diversity (Eizaguirre, Lenz, Kalbe, &  
111 Milinski, 2012). Sexual selection has also been implicated in maintaining  
112 polymorphism through MHC-mediated mate choice in a range of taxa (Consuegra  
113 and Leaniz, 2008; Setchell et al., 2010; Strandh et al., 2012). Whilst the MHC does  
114 not represent overall adaptive variation, the loss of variation at this marker can have  
115 a strong negative effect on fitness, e.g. in inbred populations (Arkush et al., 2002),  
116 and standing genetic variation at the MHC is particularly important in the context of  
117 developing resistance to emerging disease (Dionne et al., 2009). The MHC is  
118 therefore widely recognized as a key marker for monitoring adaptive genetic  
119 variation in a conservation context (Eyto et al., 2007; Sommer, 2005; Ujvari and  
120 Belov, 2011).

122 Supportive breeding is becoming a widespread tool to re-invigorate species of  
123 conservation concern (Manlick et al., 2017; Moorkens, 2018; Tapley et al., 2015). It  
124 is a particularly common management strategy in salmonids (Fraser, 2008), so they  
125 are a good model to investigate its effectiveness in meeting conservation goals.  
126 European grayling (*Thymallus thymallus*) is a non-anadromous salmonid fish species  
127 with a wide distribution, ranging from France and Great Britain in the West to the  
128 Ural mountains in the East and from Montenegro in the South to Scandinavia in the  
129 North (Gum et al., 2009). The species is listed as protected in Appendix II of the Bern  
130 convention (Swatdipong et al., 2010) and UK populations are considered  
131 endangered (Dawnay et al., 2011). A number of pathogens and parasites are known  
132 to infect grayling (Dorovskikh and Stepanov, 2009; Pylkkö et al., 2006), likely  
133 imposing selection pressures on natural populations. There are also emerging  
134 threats to grayling such as proliferative kidney disease (Wahli et al., 2002) and the  
135 spread of disease from fish farms (Algöet et al., 2009). European grayling exhibit a  
136 high degree of spatial genetic structure across their natural range in the UK and  
137 continental Europe (Dawnay et al., 2011; Koskinen et al., 2002). In the UK, limited  
138 gene flow was detected between populations and four demographic clusters have  
139 been identified (Dawnay et al. 2011). To compensate for declines, supportive  
140 breeding (stocking) is a common practice to manage natural populations (Dawnay et  
141 al., 2011; Persat et al., 2016) and stocking policy in the UK has been altered in  
142 response to neutral genetic data in order to avoid homogenisation of genetically  
143 differentiated populations (Environmental Agency, 2011; Dawnay et al., 2011).  
144 However, so far only neutral genetic markers have been assessed and there is no  
145 information available on adaptive genetic variation.

147 Here, we combine empirical and modelling approaches to examine functional MHC  
148 genetic diversity of the class II  $\alpha$ -chain (DAA) and  $\beta$ -chain (DAB) in European  
149 grayling and its relationship with neutral genetic diversity. Specifically, we test (i) the  
150 degree to which neutral genetic variation reflects MHC genetic variation; (ii) the  
151 degree to which risk assessment of population viability and the definition of  
152 management units based on neutral genetic variation are consistent with results from  
153 MHC genetic variation; (iii) if there is an effect of management on neutral and  
154 adaptive genetic variation; (iv) using forward simulations we explore whether these  
155 effects can be explained by the lack of natural selection acting on hatchery produced  
156 offspring during captivity.

157

## 158 Materials and Methods

### 159 *Tissue Samples*

160 Thirty-seven to forty individuals from each of twelve populations were used for this  
161 study. These samples are a subset of those previously genotyped at ten  
162 microsatellite loci by Dawnay et al. (2011, see Figure 1). Five of the sampled  
163 populations are classified as 'non-stocked native' (Dee, Severn, Ure, Wye and  
164 Wylfe), four as 'stocked native' (Aire, Derbyshire Derwent, Dove and the Hampshire  
165 Avon) and three as introduced populations (Clyde, Eden and Itchen) (Dawnay et al.,  
166 2011). These populations represent all four demographic units (DUs) that were  
167 identified by Dawnay et al. (2011). The introduced populations are thought to be  
168 sourced from the Dove and Derbyshire Derwent, with one or more introductions  
169 taking place over the past 200 years (Ibbotson et al., 2001; Wilson, 1963). For the  
170 stocked native populations detailed information available on exact timing and



171 numbers of stocking events is limited. In the River Aire stocking was performed  
172 every year between 2006-2009, releasing between 1000 and 2000 individuals  
173 (Environmental Agency UK, personal communication). In the River Dove 1500  
174 individuals were stocked in 2007 (Environmental Agency UK, personal  
175 communication). The age of stocked fish was between ~6-18 months (0+ or 1+). The  
176 likely provenance of the stocked fish is the river Test (Dawnay et al., 2011).

177

### 178 *MH II target loci*

179 In teleost fish, class I and class II major histocompatibility genes are not within one  
180 complex like in other vertebrates and hence are designated as MH (Stet et al.,  
181 2003). Our methods target variation at the MH class II  $\alpha$ -chain (DAA) and  $\beta$ -chain  
182 (DAB), covering most of the class II peptide binding region (PBR). Primer sequences  
183 for the DAA exon were based on published primers developed for brown trout  
184 (*Salmo trutta*; Stet et al., 2002; amplicon length: ~213 bp). Previously described  
185 primers for the  $\beta$ 1 domain encoded by exon 2 of the DBB gene, involved in peptide  
186 binding, were modified from those described by Pavey et al. (2011) (forward: 5'-  
187 ATGTTTTTCCTTTTAGATGGATATTTT -3', reverse: 5'- GTCTTATCCAGTACGACAC  
188 -3'; amplicon length: ~286 bp).

189

### 190 *NGS library preparation*

191 Tagged sequencing was used in a nested PCR, with the outer primer containing the  
192 Illumina adapter sequences and tags and the inner primer the target-specific  
193 sequence (after Lange et al., 2014). This allows different inner primers to be used

194 with the same set of tagged outer primers and is therefore flexible and cost-efficient.  
195 The assay was designed as a one-step PCR on a Fluidigm Access Array microfluidic  
196 chip (Lange et al., 2014), but was modified here to a two-step PCR for conventional  
197 thermocyclers.

198 Library preparation using PCR was performed as follows. Inner target-specific PCR  
199 was performed with a total volume of 6  $\mu$ l, containing 3.75 mM MgCl<sub>2</sub>, 0.2 mM each  
200 dNTP, 4% DMSO, 0.2  $\mu$ M of each target-specific primer, FastStart High Fidelity  
201 Reaction Buffer and 0.15 U of FastStart High Fidelity Blend Enzyme (Roche/Sigma  
202 Aldrich) on Prime (Bibby) PCR cyclers or in a ABI 1 PCR cycler. Amplification used a  
203 thermal profile of: 95°C for 10 min, followed by 15 cycles at 95°C for 25 s, target-  
204 specific temperature and annealing time and 72°C for 90 s, and a final extension at  
205 72°C for 5 min. Target-specific temperatures and annealing times were 59°C for 60 s  
206 (DAA locus) or 60°C for 45 s (DAB). PCR products were diluted (1:20) in H<sub>2</sub>O and 3  
207  $\mu$ l used as template in the second PCR which was carried out in a total volume of 7  
208  $\mu$ l containing 3.75 mM MgCl<sub>2</sub>, 0.2 mM each dNTP, 4% DMSO, 0.1  $\mu$ M of each outer  
209 primer, FastStart High Fidelity Reaction Buffer and 0.25 U of FastStart High Fidelity  
210 Blend Enzyme (Roche/Sigma Aldrich). The thermal profile of the second PCR was  
211 95°C for 10 min, followed by 27 cycles at 95°C for 25 s, 60°C for 60 s and 72°C for  
212 90 s, and a finishing step at 72°C for 5 min. Amplification success was verified on  
213 20% of samples using 1% agarose gels. All samples were prepared in independent  
214 replicates along with ten randomly distributed negative controls for each locus, which  
215 represent samples without DNA input. Subsequently, PCR products were pooled per  
216 locus for each population prior to purification using AmpureXP (Beckmann and  
217 Coulter) and quantification using a Qubit 3.0 Fluorometer (Thermo Fisher Scientific).  
218 All populations were then pooled for each locus (equimolar concentrations) and run

219 on an Agilent 2100 Bioanalyzer to check product size and successful removal of  
220 unincorporated adapters and primers. Samples were then pooled in equal  
221 concentrations across loci and sequenced using an Illumina Miseq Nano (250bp  
222 paired end).

223

#### 224 *Data analysis*

225 For quality control, all reads with a quality score below 20 in more than 90 percent of  
226 the sequence were filtered using the Filter by Quality tool on Galaxy Server (Goecks  
227 et al., 2010). Only sequences with both paired-end reads of sufficient quality were  
228 retained and aligned to each other using Mothur (Schloss et al., 2009). Primer  
229 mismatches (>1bp) and frame-shifts were filtered and examined for repeated  
230 sequences that could be derived from co-amplifying pseudo-genes. Read counts  
231 were adjusted if a variant (unique sequence) was present in a negative control. In  
232 this case, the highest read count of the variant observed in a control was subtracted  
233 from all amplicons where this variant was detected. Genotyping was performed if  
234 amplicons had a minimum of twenty reads. Because of the challenges associated  
235 with genotyping highly variable multi-gene families such as the MHC (Lighten et al.,  
236 2014a), like the distinction between natural recombinants and artificial chimeras, our  
237 genotyping approach builds upon the combination of two previously described  
238 pipelines to allow for high genotyping confidence. Briefly, genotyping was primarily  
239 done following the method described by Sommer et al. (2013), which is based on the  
240 comparison between replicate samples and also accounts for differences in allele  
241 amplification efficiency. Where the most frequent variant within one amplicon was  
242 not present within the technical replicate, an assignment error was assumed and the

243 individual excluded from the analysis. Where no replicate sample was available the  
244 genotyping methods described by Lighten et al. (2014b) were used as an additional  
245 criterion to assure genotyping confidence. Non-replicated genotyping estimates were  
246 only considered if they were consistent between the ‘Sommer’ and ‘Lighten’  
247 estimate. The methods described by Lighten et al. (2014b) are based on the  
248 calculation of the degree of change (DOC-method) between variants and the  
249 comparison of read numbers to expectations under a number of alternative copy  
250 number scenarios (CNV method). The CNV-method was also applied to compare the  
251 effect of control read subtraction on the overall fit of the data to specific copy number  
252 scenarios. In this case up to five loci were considered. Here, the F-ratio test was  
253 used to decide whether control read subtraction resulted in significantly lower  
254 variance and better fit.

255

### 256 *Genetic diversity analysis*

257 Summary statistics of genetic diversity were calculated for all populations.  
258 Conformity to Hardy-Weinberg equilibrium and allele frequency difference amongst  
259 populations were investigated using the Fisher’s exact test implemented in Genepop  
260 (Rousset, 2008) using 10,000 dememorizations, 100 batches and 10,000 iterations  
261 per batch. Observed and expected heterozygosity were calculated in GenAlex 6.5  
262 (Peakall and Smouse, 2012). F-statistics (Weir and Cockerham, 1984) and allelic  
263 richness were calculated using Fstat (Goudet, 2001). Whilst Weir and Cockerham’s  
264  $F_{ST}$  can be biased by differences in mutation rates (Hedrick 1999), which can be  
265 elevated for microsatellite markers, here variation at microsatellites was not higher  
266 than for MH markers, so that  $F_{ST}$  was considered sufficient to reflect differentiation

267 (Whitlock, 2011). Tests of significant differences of  $F_{IS}$  estimates and of  $F_{ST}$   
268 estimates were based on 24000 randomisations and 66000 permutations  
269 respectively. The peptide binding region (PBR) was inferred by alignments of  
270 grayling MH II sequences to human HLA sequences (Brown et al., 1993). Amino acid  
271 (AA) diversity was calculated for the PBR as p-distance within and across  
272 populations in MEGA 7.021 and also as average pairwise p-distance across  
273 individuals for the whole sequence and only for the PBR (Kumar et al., 2016). A  
274 Mann-Whitney-Wilcoxon test was used to compare the pairwise p-distance across  
275 individuals for the whole sequence and only for the PBR.

276 MH locus data were compared to neutral microsatellite diversity for the same  
277 populations to evaluate how well neutral genetic variation reflects ecologically  
278 meaningful genetic variation. We conducted a simulation analysis by sequentially  
279 removing two microsatellite loci and measuring their correlation with the remaining  
280 eight loci over 1000 bootstrap cycles for all standard measures of genetic diversity to  
281 assess our ability to detect significance.

282 To detect differences in functional relative to neutral genetic variation across  
283 management classes, differences in observed and expected heterozygosity,  
284 inbreeding coefficient  $F_{IS}$  and allelic richness between MH II and microsatellites were  
285 tested for non-stocked native, stocked native and introduced populations. This was  
286 done using a clustered Mann-Whitney-Wilcoxon test implemented in the R-package  
287 'clusrank', using the Datta-Satten method and 1000 bootstrap cycles (Jiang et al.,  
288 2017), to account for the dependency of measurements derived from the linked DAA  
289 and DAB genes respectively. The Kruskal-Wallis test was used to identify differences  
290 in measurements of genetic diversity across management classes for each marker  
291 type. Pairwise  $F_{ST}$  estimates were compared using a Mann-Whitney-Wilcoxon test

292 between MH II and microsatellite loci, using (i) all populations, (ii) non-stocked native  
293 and stocked native or (iii) only non-stocked native populations. For all tests involving  
294 multiple comparisons, the Benjamini-Hochberg method was used to correct for  
295 multiple testing (Hochberg and Benjamini, 1990). To assess whether population  
296 structure reflected by neutral markers is supported by adaptive genetic  
297 differentiation, a neighbour-joining phylogenetic tree was built based on Nei's genetic  
298 distance (Nei, 1972) in PHYLIP using a consensus of 2000 bootstrapped replications  
299 for all genes studied (Felsenstein, J, 1989). An analysis of molecular variance was  
300 done for both microsatellite and MH II data in GenAlex 6.5 (Peakall and Smouse,  
301 2012).

302

### 303 *Inference of selection*

304 Recent effects of selection on each gene and population were evaluated in  
305 ARLEQUIN 3.5 (Excoffier and Lischer, 2010) using a Ewens-Watterson  
306 homozygosity test (Ewens, 1972; Watterson, 1978). The Ewens-Watterson test  
307 compares allele frequencies observed within each population to those expected  
308 under neutrality for populations at mutation-drift equilibrium. The test assumes  
309 population equilibrium and is sensitive to demographic changes. During population  
310 bottlenecks low frequency alleles are lost at a higher rate, producing allele  
311 frequencies that are more even than expected under neutrality (Ewens, 1972;  
312 Watterson, 1978). Similarly population expansion leads to an increase in low  
313 frequency alleles and lower heterozygosity than expected under neutral-equilibrium  
314 (Meyer et al., 2006). In order to distinguish demographic and selective forces and  
315 their effect on allele frequency changes a Ewens-Watterson test was also performed

316 on the microsatellite data from Dawnay et al. (2011) for all populations studied.  
317 Where recent demographic events are the reason for deviations from neutrality both  
318 neutral and adaptive markers are expected to be affected, whilst selection is  
319 expected to only affect MH II markers. Following Larson et al. (2014) alpha margins  
320 of 10% ( $p < 0.1$ ,  $p > 0.9$ ) were considered as evidence of selection, because of the  
321 limited statistical power of the Ewens-Watterson test in detecting weak or moderate  
322 selection (Ewens, 1972) .

323

### 324 *Simulations*

325 We implemented a simulation model using simuPop, version 1.1.8.3 (Peng and  
326 Kimmel, 2005), in order to investigate whether the lack of natural selection during  
327 early life-stages of hatchery reared juveniles could result in changes in observed and  
328 expected heterozygosity in supplemented populations where population census size  
329 differs (script available on request). We assumed a natural population with constant  
330 size and with an age class structure as described for grayling populations in  
331 Woolland and Jones (1975). We assumed age-dependent female fecundity (Charles  
332 et al., 2006). We assumed allele frequencies for DAA and DAB MH loci were the  
333 same as our estimates of the native Dee population (this study). We used a  
334 heterozygote advantage model for offspring survival with a selection coefficient of  
335 0.1, which is within the range reported for loci under balancing selection in natural  
336 populations (0.05-0.15, Aguilar et al., 2004) and a model without selection to  
337 represent a comparable neutral marker reference. Thus, the probability of survival  
338 was given by the average fitness value across the two MH loci divided by the sum of  
339 probabilities across all individuals of a certain age class. We did not evaluate the

340 scenario of using foreign stocks and introducing potentially maladaptive alleles as  
341 this does not represent the generally recommended practice in a conservation  
342 context and is not the current practice of the Environmental Agency for grayling in  
343 the UK (Dawnay et al., 2011). After simulating the evolution of the population for ten  
344 years, ten adult males and females were selected randomly to produce the simulated  
345 hatchery offspring, before the adults were returned to the population. Selection-  
346 dependent survival on hatchery produced offspring was removed in the first year,  
347 before 1000 individuals at the age of 1 year were stocked into the source population.  
348 Of these individuals 50% were randomly removed from this cohort to simulate non-  
349 genetic effects of high initial mortality in stocked fish (Pedersen et al., 2003). This  
350 stocking procedure was simulated in three consecutive years and the allele  
351 frequencies in the population monitored for another ten years. Stocking intensity and  
352 frequency generally followed those actually practiced in the native stocked  
353 populations described above (Environmental Agency UK, personal communication).  
354 Stocking intensity was kept constant for different source population sizes of 500,  
355 750, 1500 and 2000, so that the ratio of naturally produced offspring surviving to an  
356 age of one year to those stocked that initially survived (50%) were roughly 0.5:1,  
357 0.8:1, 1.6:1 and 2:1 respectively. The population was replicated with or without  
358 stocking 100 times respectively. Observed and expected heterozygosity was  
359 compared between them, across the following ten years after stocking, using a  
360 Mann-Whitney-Wilcoxon test. Differences between stocked and non-stocked  
361 replicates were also tested for significance in each year after stocking for neutral and  
362 MH markers using a Mann-Whitney-Wilcoxon test.

363



## 364 Results

### 365 *Quality control*

366 The Illumina Nano run resulted in  $n=1,227,780$  reads. A small number of reads were  
367 observed in negative controls: the mean reads for these across 10 control samples  
368 respectively were  $10 \pm 26$  (total 103) and  $1.8 \pm 6$  (total 19), representing 0.02% and  
369 0.005% of the total reads for the DAA and DAB locus respectively. The fit of the  
370 overall dataset to specific copy number scenarios was significantly better after  
371 control read subtraction (F-test:  $F=0.67$ ,  $p<.0005$ ). Genotypes were obtained for a  
372 total of 389 individuals for the DAA and 359 individuals for the DAB locus. Of these  
373 82% were derived from replicated samples for the DAA and 52% for the DAB locus.  
374 Several samples were excluded from the analysis due to a potential assignment  
375 error (DAA  $n=7$ ; DAB  $n=1$ ), where the most frequent variant within one amplicon was  
376 not present in the replicate. The genotypes of most individuals were consistent with  
377 the single classical class II locus system found within other salmonids (Stet et al.,  
378 2002). However, individuals for the DAA ( $n=4$ ; 1%) and DAB ( $n=1$ ; 0.3%) loci  
379 exhibited three alleles and were excluded from subsequent analysis. The mean per  
380 amplicon coverage was 143 for the DAA locus and 89 for the DAB locus. For the  
381 DAA and DAB locus 15 and 10 alleles were identified, of which 14 and 10 encoded  
382 different protein sequences respectively.

383

### 384 *Genetic diversity*

385 The total number of alleles per population ranged from 2-7 for the DAA locus and 2-6  
386 for the DAB locus. Two populations, the stocked native Aire (AIR) and non-stocked

387 native Severn (SEV), showed significant heterozygote deficits (Table 1) and  
388 departure from HWE for the DAB gene.  $F_{IS}$  estimates ranged from -0.22 to 0.33 for  
389 the DAA locus and from -0.2 to 0.6 for the DAB locus. Average AA diversity within  
390 the PBR was 0.11 for DAA and 0.41 for DAB across all populations. There was  
391 greater within-population than between-population AA diversity for all but the Eden  
392 population for the DAB locus (Table 1). Average pairwise AA distance across  
393 individuals was significantly higher for the PBR than for the whole sequence for both  
394 the DAA and DAB locus (Mann-Whitney-Wilcoxon test:  $p < 0.001$ ).

395 No significant correlations were observed between microsatellite and MH expected  
396 heterozygosity (Spearman:  $\rho = 0.26$ ,  $p = 0.22$ ), observed heterozygosity  
397 (Spearman:  $\rho = 0.39$ ,  $p = 0.06$ ), allelic richness (Spearman:  $\rho = 0.36$ ,  $p = 0.09$ )  
398 or  $F_{IS}$  (Spearman:  $\rho = 0.21$ ,  $p = 0.32$ ). However, the results of the simulation  
399 analysis indicated that correlations between random subsets of two and ten of the  
400 microsatellite markers resulted in lower values of  $\rho$  in 17.6% for expected  
401 heterozygosity, 54% for observed heterozygosity, 40% for allelic richness and 7% for  
402  $F_{IS}$ . Thus, there is insufficient power to detect correlations between MH and  
403 microsatellite loci.

404

405 Expected heterozygosity and allelic richness differed significantly for the MH II  
406 among management classes (Kruskall Wallis test:  $p = 0.003$ ,  $p = 0.007$ , Figure 2).  
407 Introduced populations showed the lowest diversity and native non-stocked  
408 populations the highest. This pattern was not evident for microsatellite loci. Expected  
409 heterozygosity and allelic richness were significantly higher for MH II genes than for  
410 microsatellites in non-stocked native populations (clustered Mann-Whitney-Wilcoxon

411 test:  $p = 0.002$ ,  $p = 0.008$ ; Figure 2). This was not the case for the other  
412 management classes. No significant differences were observed between  
413 management classes for observed heterozygosity,  $F_{IS}$  values or effective population  
414 size which was inferred from microsatellites (Dawnay et al., 2011). Percentages of  
415 molecular variance were 68% within populations, 30% among populations and 2%  
416 among individuals for microsatellites and were 51% within populations, 23% among  
417 populations and 26% among individuals for the MH II genes.

418

#### 419 *Population differentiation*

420 We found a significant correlation between MH II and microsatellite pairwise  $F_{ST}$   
421 estimates (Mantel test DAA:  $P = 0.001$ ,  $r = 0.55$ ; DAB:  $p = 0.02$ ,  $r = 0.38$ ; Figure 3).  
422 Pairwise  $F_{ST}$  estimates were not significantly different between MH II and  
423 microsatellite estimates across all populations and when comparing non-stocked  
424 native and stocked native populations (Figure 4A, B). However, considering only  
425 non-stocked native populations pairwise  $F_{ST}$  estimates were significantly lower for  
426 MH II genes than for microsatellites (Pairwise Mann-Whitney-Wilcoxon; DAA:  $p =$   
427  $0.009$ , DAB:  $p = 0.02$ ) (Figure 4C).

428

429 Pairwise  $F_{ST}$  estimates significantly greater than zero were found for most population  
430 pairs for all genes (Table 2). Dawnay et al. (2011) identified four demographic units  
431 based on microsatellites (A-D). Un-rooted neighbour-joining phylogenetic trees  
432 suggest a similar pattern of population sub-groups for MH II genes as for neutral  
433 markers (Figure 5). However, the Dee population groups with cluster C rather than A

434 and the Derbyshire Derwent with A rather than D, where they were grouped for  
435 neutral markers (Figure 5).

436

### 437 *Selection*

438 For the stocked native Aire (AIR) and Dove (DOV) and the introduced Clyde (CLD)  
439 populations no evidence for selection was identified by the Ewens-Watterson test for  
440 any MH gene and microsatellite results suggested a recent population decline  
441 (Supporting Information 1). For the non-stocked native Dee (DEE), Severn (SEV)  
442 and Wylfe (WLA/B), as well as for the introduced Eden (EDN) and stocked native  
443 Hampshire Avon (HAV) populations allele frequencies deviated significantly from  
444 expectations under neutrality for both MH II genes and microsatellites, but in each  
445 case the difference between observed and expected allele frequencies was greater  
446 for microsatellites, indicating a dominant effect of a recent population decline  
447 (Supporting Information 1). Populations that did not show larger significant  
448 differences between observed and expected allele frequencies for microsatellites  
449 than for MH genes were the non-stocked native Ure (URE) and Wye (WYE)  
450 populations. The DAA locus showed evidence for balancing selection for the Ure  
451 population and the DAB locus for the Wye population.

452

### 453 *Simulations*

454 Simulating the effect of stocking between neutral and MH II markers for different  
455 population census sizes, showed that at very low population sizes (census size 500)  
456 neutral markers are affected more strongly than MH II markers, as measured by

457 stronger reductions in observed and expected heterozygosity (Figure 6). There was  
458 no significant difference in MH observed heterozygosity between stocked and non-  
459 stocked replicates at this population size. In all other cases, there was a significant  
460 reduction of MH heterozygosity (observed and expected, Figure 6) after stocking.  
461 Comparing the marker types, for higher population sizes (census size 750, 1500,  
462 2000) the effect of stocking was significantly stronger on MH expected  
463 heterozygosity than on microsatellites in all cases and for observed heterozygosity  
464 for population census sizes of 750 and 1500 (Figure 6). For population sizes of 1500  
465 and 2000 there was no significant effect of stocking on expected heterozygosity on  
466 neutral markers and for observed heterozygosity a significant effect of stocking was  
467 found only for a population size of 2000 (Figure 6). Looking at the effect of stocking  
468 separately for each year, shows that a persisting significant effect on MH  
469 heterozygosity is observed at a population size of 750 (Figure 7B).

470

## 471 Discussion

472 In order to maintain adaptive genetic variation in threatened populations, it is  
473 important to understand how management impacts on functional genetic diversity  
474 and evolutionary processes. In this study we compared the performance of neutral  
475 and functional markers in informing conservation and management decisions, using  
476 salmonids as a model to evaluate the effect of supportive breeding on these different  
477 types of genetic markers. Measurements of genetic diversity at functional MH loci  
478 could not be predicted by neutral markers. Across different grayling population  
479 management classes only non-stocked native populations showed evidence for  
480 selection maintaining higher levels of variation at the MH II than at neutral loci. We

481 implemented a simulation model to test if the removal of natural selection on early  
482 life-stages within the hatchery could explain our empirical results. A significant  
483 reduction in MH diversity but not neutral diversity was predicted by our model at  
484 intermediate population sizes. This is consistent with our empirical results. Further, a  
485 significant reduction in the response to selection resulting from supportive breeding  
486 was predicted by all simulated scenarios. Overall, our results show clear differences  
487 between functional versus neutral genetic loci, confirming the imperative to use  
488 adaptive genetic markers to inform conservation decisions (Piertney and Webster,  
489 2008; Sutton et al., 2011). Our results have clear implications for population  
490 management involving augmentation, calling into question its efficiency in supporting  
491 long term viable populations with high adaptive potential.

492

493 We found significant differences in allelic richness and expected heterozygosity for  
494 MH II genes, but not microsatellites, across management classes. Although the  
495 lowest diversity was found in introduced populations, which might be expected as the  
496 consequence of a bottleneck this result is not supported by neutral markers. The loss  
497 of diversity in introduced populations was specific to the MH II. Explanations other  
498 than founder effects must explain the loss of variation. In a similar study looking at  
499 population genetic variation in translocated rainbow trout (*Onchorhynchus mykiss*),  
500 Monzón-Argüello et al. (2013) found low MH II diversity relative to neutral markers.  
501 These authors attribute this to selection pressures against MH alleles that perhaps  
502 did not provide a selective advantage in the novel environment into which they were  
503 introduced. Such habitat specific adaptations, where there is a fitness advantage of  
504 local genotypes, have been found at the MH II in river and lake populations of three-  
505 spined stickleback (Eizaguirre et al., 2012b). Our study further underlines that, using

506 neutral markers as a surrogate of adaptive genetic variation is unreliable. This  
507 observation has been demonstrated in a range of taxa, e.g. mammals (Aguilar et al.,  
508 2004), other salmonids (Dionne et al., 2007) and birds (Hartmann et al., 2014).  
509 Specific consideration of adaptive markers and likely impacts of demographic history  
510 and management on them needs to be a routine part of conservation genomic  
511 research.

512

513 Few studies (e.g. Schenekar and Weiss, 2017) have focussed on assessing  
514 adaptive versus neutral variation specifically in supportive breeding programmes, a  
515 practice becoming increasingly adopted as means to effectively manage population  
516 declines. Here, significantly higher genetic diversity (allelic richness and expected  
517 heterozygosity) of MH genes than microsatellites was observed in non-stocked  
518 native populations. Non-stocked native populations also showed significantly lower  
519 differentiation for MH II markers than neutral markers. This suggests that balancing  
520 selection is acting to retain variation at the MH in natural populations, but that this is  
521 not the case for introduced or stocked native populations. On the one hand, stocked  
522 native populations with reduced MH II diversity could be undergoing drift and this has  
523 removed variation more rapidly at MH II loci due to the combined effect of drift and  
524 selection (Ejsmond and Radwan, 2011; Sutton et al., 2011). This seems unlikely  
525 because Dawnay et al. (2011) found evidence for bottlenecks in ten populations and  
526 these included populations of all three categories (introduced, stocked native and  
527 non-stocked native). Thus, there is no evidence that most grayling populations  
528 selected for stocking suffered recent and severe population decline, or exhibit lower  
529 effective population sizes (Dawnay et al., 2011). Therefore, considering a direct

530 effect of the stocking process on the efficiency of selection to act within the  
531 supplemented population is consistent with our results.

532

533 We found no correlation between MH II and neutral markers using any measure of  
534 genetic diversity, however we show that for the number of microsatellite markers  
535 genotyped (twelve) by Dawnay et al. (2011) and two MH loci, the statistical power for  
536 detection is insufficient to identify differences between functional versus neutral loci  
537 we analysed. This highlights the importance of caution when making inferences of  
538 overall genetic diversity from only a low number of markers (DeWoody and  
539 DeWoody, 2005) and the importance of considering power explicitly when designing  
540 a programme of sampling. However, we found evidence for recent selection on MH  
541 loci as outlined above. Additionally, we found higher within-population amino acid  
542 (AA) diversity than between populations, and significantly more even allele  
543 frequencies than expected, while we did not observe this for microsatellites.

544

545 The levels of diversity reported for the MH II here, compare to those of other  
546 salmonids, where generally both the alpha and beta chain show similar levels of  
547 diversity (Gómez et al., 2010). This contrasts with other vertebrates ( e.g. chicken,  
548 Salomonsen et al., 2003; humans, Reche & Reinherz, 2003), where the alpha chain  
549 shows much lower levels of diversity. Two populations, Aire and Severn, significantly  
550 departed from HWE due to heterozygote deficiency at the DAB locus (Table 1).  
551 Whilst a technical cause, such as allelic drop-out at the DAB locus, cannot be ruled  
552 out, elevated (but not significant)  $F_{IS}$  vales have also been found for the same  
553 populations at the DAA locus. This is not consistent with uneven reproductive



554 success between families or a Wahlund effect (Wahlund, 1928) because the pattern  
555 is not also shown by neutral loci. To observe higher levels of inbreeding for genes  
556 under balancing selection than for neutral markers seems counterintuitive. However,  
557 a loss of diversity under the simultaneous effects of selection and drift has been  
558 shown both empirically and theoretically (Ejsmond and Radwan, 2011; Sutton et al.,  
559 2011). Additionally, MH II mediated mate choice is not necessarily disassortative,  
560 seeking highest offspring dissimilarity, but assortative, where particular alleles confer  
561 highest resistance, e.g. under frequency dependent selection (Eizaguirre et al.,  
562 2009). We report more than 20 times higher proportion of molecular variance found  
563 among individuals for the MH II than for microsatellites, which supports an important  
564 role of frequency dependent or heterogeneous selection in space and time opposed  
565 to overdominant selection. Thus, unbalanced reproductive success for particular MH  
566 II genotypes, resulting in elevated inbreeding, might be expected, particularly where  
567 competition for mating opportunities is high (Milinski, 2006). This can be for example  
568 the case when spawning grounds are scarce (Castric et al., 2002), which has been  
569 documented for the Severn population (Lewis, 2006).

570

571 Our ability to directly compare native populations before and after stocking is limited  
572 by lack of pre- and post-stocking samples. Our simulation model addresses this  
573 focussing on plausible genetic effects caused by unequal reproductive contributions  
574 of hatchery reared young in relation to naturally produced offspring and the different  
575 selective environments experienced by each respectively.

576

577 The key outcome of our simulation is that the consideration of stocking intensity in  
578 relation to naturally produced offspring is crucial to reduce negative long-term effects  
579 on adaptive genetic diversity. Across taxa, empirical and simulated data need to be  
580 obtained to establish the effects of supplementary breeding on adaptive genetic  
581 variation that underlies fitness. For example, our results show that stocking can have  
582 a strong effect on genetic variation at lower population census sizes (500 and 750)  
583 where the stocking intensity exceeded numbers of naturally produced offspring. The  
584 largest effect of drift was observed at a low census size (500), where the differential  
585 effect of stocking was larger at neutral loci than at MH II loci (Figure 6). With  
586 increasing population census size the effect of drift decreased and in all other cases  
587 variation at the MH II was lost at a higher rate than at neutral markers as a result of  
588 stocking. At a population census size of 750, heterozygosity remained significantly  
589 lower for the MH II in year wise comparisons, but not neutral markers even ten years  
590 after stocking (Figure 7B). Marie et al. (2010) also reported that the loss of genetic  
591 integrity correlates with stocking intensities in brook charr (*Salvelinus fontinalis*).  
592 Even for larger census sizes (1500 and 2000) where the ratio of naturally produced  
593 offspring to stocked offspring was high there was a large effect on MH loci but a  
594 negligible effect on neutral markers. However, the effect was weaker than for smaller  
595 population sizes as would be theoretically predicted. As populations with low census  
596 size would be most likely to be considered for stocking it is important to notice their  
597 vulnerability to genetic deterioration. Furthermore, our results are likely to  
598 underestimate the role of selection, because MH related mate choice was not  
599 considered, though it has been shown to maintain MH diversity in teleost (Consuegra  
600 and Leaniz, 2008; Eizaguirre et al., 2009). Also, we focus specifically on the effect of  
601 the removal of natural selection within the artificial rearing environment and do not

602 consider adaptation to the hatchery environment, which would be likely to further  
603 exacerbate differences in adaptive genetic diversity.

604

605 It is interesting from a theoretical point of view that a lower differential effect of  
606 stocking on the MH compared to neutral markers was observed at a census size of  
607 500. This could be explained by the expectation that in the population prior to  
608 stocking, MH diversity is lost at a greater rate than neutral genetic diversity through  
609 the combined forces of drift and selection (Ejmond and Radwan, 2011; Sutton et al.,  
610 2011). Considering that selection becomes less efficient through stocking fish that  
611 have not experienced natural selection at the early life stage, the supplemented  
612 population is partly alleviated from this additional force, so that the difference  
613 between stocked and non-stocked MH diversity is smaller than that observed for  
614 neutral markers.

615

616 In the simulation model, we assume that a limited number of individuals is selected  
617 as brood stock (20 individuals), which reflects current practice. However, selecting a  
618 larger number of individuals would likely retain more genetic diversity within the  
619 hatchery brood stock and reduce the effect of drift. Also, we assume an initial  
620 mortality of stocked fish of 50%, as described in *Salmo trutta* (Pederson, 2003). The  
621 survival of stocked grayling has shown to be highly variable, e.g. in some places  
622 natural populations do not show any signs of introgression with hatchery stocks,  
623 whilst in other places the original population was completely replaced (Persat et al.,  
624 2016). Evidence that stocked grayling within UK rivers do survive and contribute to  
625 the population is provided by the observation that genetic relationships of stocked

626 populations agreed with stocking records and through the recapture of stocked  
627 individuals (Dawnay et al., 2011). Given the uncertainty around exact rates of initial  
628 survival of stocked fish, the most relevant parameters here are the ratios of  
629 supplemented individuals relative to the number of offspring naturally produced  
630 within the recipient population. This also allows for high transferability of the model  
631 predictions to other systems and specific cases.

632

633 Whilst the MH represents only a specific locus of adaptive importance, our findings  
634 may have implications for loci under selection in a broader sense. Both our empirical  
635 results and simulations suggest a dilution effect through the supplementation of a  
636 natural population with individuals reared within an artificial environment, which  
637 adversely affects the efficiency of selection. Though we evaluated a case of  
638 balancing selection, a reduction in the efficiency of selection to act upon a population  
639 might be expected to extend to other types of selection. As dynamic adaptive  
640 responses are crucial under the pressure of current rates of environmental change  
641 conservation management should carefully evaluate the possible inference with  
642 natural evolutionary processes. In this context it is important to assess the rate of  
643 natural production of a population, which in the case of grayling is frequently  
644 restricted by habitat deterioration, which reduces the availability of suitable spawning  
645 grounds (Nykänen and Huusko, 2002). This will much better inform the number of  
646 individuals to supplement into a natural population and offers the possibility to  
647 implement measures of habitat restoration as a first resort where natural  
648 reproductive capacities are not fully exploited. This reflects the general need for a  
649 more comprehensive evaluation of potential risks and benefits from *ex situ* versus *in*

650 *situ* management practices before these are implemented in a conservation context  
651 (Dolman et al., 2015).

652

## 653 Conclusions

654 MH II genes in non-stocked native populations of European grayling showed higher  
655 variation than was predicted by microsatellites. We also found significant differences  
656 at MH loci between different population management regimes (introduced, stocked  
657 native and non-stocked native populations) which were not detected by neutral  
658 markers. Our findings highlight the importance of using functional genetic markers to  
659 inform the conservation management of genetic diversity (Kirk and Freeland, 2011;  
660 Piertney and Webster, 2008). We present evidence consistent with selection  
661 maintaining genetic variation in functional loci for non-stocked native populations,  
662 which is aligned with results from our simulation model. Simulation results suggest  
663 selection is less efficient to maintain genetic variation at functional loci in stocked  
664 populations, while the effect is negligible in neutral loci.

665 Our findings have implications for population conservation management where  
666 translocation, reintroduction or population augmentation is practised. Our results  
667 highlight the need for a clear understanding of the interaction of selective processes  
668 with management actions. Conservation programmes need to more explicitly  
669 incorporate and consider possible interference with natural evolutionary and adaptive  
670 processes during supplementation, especially considering the current rate of  
671 environmental change.

672

## Acknowledgements

This project was funded by a MMU internal studentship as part of VH's PhD research. We would also like to thank the Grayling Research Trust for financial support for this project. Thanks are also given to Prof. Steven Weiss and Dr. Henri Persat for their input and guidance throughout the study and Dr. Lewis Spurgin for provision of an R script to perform the power analysis. Further, we also want to thank Dr. Mairi Knight and five anonymous reviewers for their insightful comments that greatly improved the manuscript. The project would not have been possible without the support of the anglers that provided samples for this study.

## References

- Aguilar, A., Roemer, G., Debenham, S., Binns, M., Garcelon, D., Wayne, R.K., 2004. High MHC diversity maintained by balancing selection in an otherwise genetically monomorphic mammal. *Proc. Natl. Acad. Sci.* 101, 3490–3494.
- Algöet, M., Bayley, A.E., Roberts, E.G., Feist, S.W., Wheeler, R.W., Verner-Jeffreys, D.W., 2009. Susceptibility of selected freshwater fish species to a UK *Lactococcus garvieae* isolate. *J. Fish Dis.* 32, 825–834.
- Allendorf, F.W., Hohenlohe, P.A., Luikart, G., 2010. Genomics and the future of conservation genetics. *Nat. Rev. Genet.* 11, 697–709.
- Arkush, K.D., Giese, A.R., Mendonca, H.L., McBride, A.M., Marty, G.D., Hedrick, P.W., 2002. Resistance to three pathogens in the endangered winter-run chinook salmon (*Oncorhynchus tshawytscha*): effects of inbreeding and major histocompatibility complex genotypes. *Can. J. Fish. Aquat. Sci.* 59, 966–975.
- Ayllon, F., Martinez, J.L., Garcia-Vazquez, E., 2006. Loss of regional population structure in Atlantic salmon, *Salmo salar* L., following stocking. *ICES J. Mar. Sci. J. Cons.* 63, 1269–1273.
- Brodersen, J., Seehausen, O., 2014. Why evolutionary biologists should get seriously involved in ecological monitoring and applied biodiversity assessment programs. *Evol. Appl.* 7, 968–983.
- Brown, J.H., Jardetzky, T.S., Gorga, J.C., Stern, L.J., Urban, R.G., Strominger, J.L., Wiley, D.C., 1993. Three-dimensional structure of the human class II histocompatibility antigen HLA-DR1. *Nature* 364, 33–39.
- Castric, V., Bernatchez, L., Belkhir, K., Bonhomme, F., 2002. Heterozygote deficiencies in small lacustrine populations of brook charr *Salvelinus fontinalis* Mitchill (Pisces, Salmonidae): a test of alternative hypotheses. *Heredity* 89, 27–35.
- Ceballos, G., Ehrlich, P.R., Barnosky, A.D., García, A., Pringle, R.M., Palmer, T.M., 2015. Accelerated modern human-induced species losses: Entering the sixth mass extinction. *Sci. Adv.* 1.
- Charles, S., Mallet, J.-P., Persat, H., 2006. Population Dynamics of Grayling: Modelling Temperature and Discharge Effects. *Math. Model. Nat. Phenom.* 1, 31–48.
- Consuegra, S., Leaniz, C.G. de, 2008. MHC-mediated mate choice increases parasite resistance in salmon. *Proc. R. Soc. Lond. B Biol. Sci.* 275, 1397–1403.

- Dawnay, N., Dawnay, L., Hughes, R.N., Cove, R., Taylor, M.I., 2011. Substantial genetic structure among stocked and native populations of the European grayling (*Thymallus thymallus*, Salmonidae) in the United Kingdom. *Conserv. Genet.* 12, 731–744.
- de Eyto, E., McGinnity, P., Huisman, J., Coughlan, J., Consuegra, S., Farrell, K., O'Toole, C., Tufto, J., Megens, H.-J., Jordan, W., Cross, T., Stet, R.J.M., 2011. Varying disease-mediated selection at different life-history stages of Atlantic salmon in fresh water. *Evol. Appl.* 4, 749–762.
- DeWoody, Y.D., DeWoody, J.A., 2005. On the Estimation of Genome-wide Heterozygosity Using Molecular Markers. *J. Hered.* 96, 85–88.
- Dionne, M., Miller, K.M., Dodson, J.J., Bernatchez, L., 2009. MHC Standing Genetic Variation and Pathogen Resistance in Wild Atlantic Salmon. *Philos. Trans. Biol. Sci.* 364, 1555–1565.
- Dionne, M., Miller, K.M., Dodson, J.J., Caron, F., Bernatchez, L., 2007. Clinal Variation in Mhc Diversity with Temperature: Evidence for the Role of Host–Pathogen Interaction on Local Adaptation in Atlantic Salmon. *Evolution* 61, 2154–2164.
- Dolman, P.M., Collar, N.J., Scotland, K.M., Burnside, R.J., 2015. Ark or park: the need to predict relative effectiveness of ex situ and in situ conservation before attempting captive breeding. *J. Appl. Ecol.* 52, 841–850.
- Dorovskikh, G.N., Stepanov, V.G., 2009. Structure of component parasite communities in the grayling, *Thymallus thymallus* L. (Salmoniformes, Thymallidae), and minnow, *Phoxinus phoxinus* L. (Cypriniformes, Cyprinidae), from the upper reaches of the Pechora River. *Biol. Bull.* 36, 298–306.
- Duplouy, A., Ikonen, S., Hanski, I., 2013. Life history of the Glanville fritillary butterfly in fragmented versus continuous landscapes. *Ecol. Evol.* 3, 5141–5156.
- Eizaguirre, C., Lenz, T.L., Kalbe, M., Milinski, M., 2012a. Rapid and adaptive evolution of MHC genes under parasite selection in experimental vertebrate populations. *Nat. Commun.* 3, 621.
- Eizaguirre, C., Lenz, T.L., Kalbe, M., Milinski, M., 2012b. Divergent selection on locally adapted major histocompatibility complex immune genes experimentally proven in the field. *Ecol. Lett.* 15, 723–731.
- Eizaguirre, C., Yeates, S.E., Lenz, T.L., Kalbe, M., Milinski, M., 2009. MHC-based mate choice combines good genes and maintenance of MHC polymorphism. *Mol. Ecol.* 18, 3316–3329.
- Ejmond, M.J., Radwan, J., 2011. MHC diversity in bottlenecked populations: a simulation model. *Conserv. Genet.* 12, 129–137.
- Evans, M.L., Neff, B.D., 2009. Major histocompatibility complex heterozygote advantage and widespread bacterial infections in populations of Chinook salmon (*Oncorhynchus tshawytscha*). *Mol. Ecol.* 18, 4716–4729.
- Ewens, W.J., 1972. The sampling theory of selectively neutral alleles. *Theor. Popul. Biol.* 3, 87–112.
- Excoffier, L., Lischer, H.E.L., 2010. Arlequin suite ver 3.5: a new series of programs to perform population genetics analyses under Linux and Windows. *Mol. Ecol. Resour.* 10, 564–567.
- Eyto, E. de, McGinnity, P., Consuegra, S., Coughlan, J., Tufto, J., Farrell, K., Megens, H.-J., Jordan, W., Cross, T., Stet, R.J.M., 2007. Natural selection acts on Atlantic salmon major histocompatibility (MH) variability in the wild. *Proc. R. Soc. B Biol. Sci.* 274, 861–869.
- Felsenstein, J., 1989. PHYLIP - Phylogeny Inference Package (Version 3.2). *Cladistics* 5, 163–166.
- Frankham, R., 2008. Genetic adaptation to captivity in species conservation programs. *Mol. Ecol.* 17, 325–333.
- Fraser, D.J., 2008. How well can captive breeding programs conserve biodiversity? A review of salmonids. *Evol. Appl.* 1, 535–586.

- Gómez, D., Conejeros, P., Marshall, S.H., Consuegra, S., 2010. MHC evolution in three salmonid species: a comparison between class II alpha and beta genes. *Immunogenetics* 62, 531–542.
- Goudet, J., 2001. FSTAT, a program to estimate and test gene diversities and fixation indices (version 2.9.3). <http://www2.unil.ch/popgen/softwares/fstat.htm>.
- Griffiths, R.A., Pavajeau, L., 2008. Captive Breeding, Reintroduction, and the Conservation of Amphibians. *Conserv. Biol.* 22, 852–861.
- Gum, B., Gross, R., Geist, J., 2009. Conservation genetics and management implications for European grayling, *Thymallus thymallus*: synthesis of phylogeography and population genetics. *Fish. Manag. Ecol.* 16, 37–51.
- Hartmann, S.A., Schaefer, H.M., Segelbacher, G., 2014. Genetic depletion at adaptive but not neutral loci in an endangered bird species. *Mol. Ecol.* 23, 5712–5725.
- Hedrick, P., 2003. The major histocompatibility complex (MHC) in declining populations: an example of adaptive variation. *Conserv. Biol. Ser. Camb.* 97–113.
- Hochberg, Y., Benjamini, Y., 1990. More powerful procedures for multiple significance testing. *Stat. Med.* 9, 811–818.
- Ibbotson, A.T., Cove, R.J., Ingraham, A., Gallagher, M., Hornby, D.D., Furse, M., Williams, C., 2001. A review of grayling ecology, status and management practice: recommendations for future management in England and Wales. Environment Agency.
- Kirk, H., Freeland, J.R., 2011. Applications and implications of neutral versus non-neutral markers in molecular ecology. *Int. J. Mol. Sci.* 12, 3966–3988.
- Koboldt, D.C., Steinberg, K.M., Larson, D.E., Wilson, R.K., Mardis, E.R., 2013. The Next-Generation Sequencing Revolution and Its Impact on Genomics. *Cell* 155, 27–38.
- Koskinen, M.T., Nilsson, J., Veselov, A.J., Potutkin, A.G., Ranta, E., Primmer, C.R., 2002. Microsatellite data resolve phylogeographic patterns in European grayling, *Thymallus thymallus*, Salmonidae. *Heredity* 88, 391–401.
- Kumar, S., Stecher, G., Tamura, K., 2016. MEGA7: Molecular Evolutionary Genetics Analysis Version 7.0 for Bigger Datasets. *Mol. Biol. Evol.* 33, 1870–1874.
- Lange, V., Böhme, I., Hofmann, J., Lang, K., Sauter, J., Schöne, B., Paul, P., Albrecht, V., Andreas, J.M., Baier, D.M., Nething, J., Ehninger, U., Schwarzelt, C., Pingel, J., Ehninger, G., Schmidt, A.H., 2014. Cost-efficient high-throughput HLA typing by MiSeq amplicon sequencing. *BMC Genomics* 15, 63.
- Larson, W.A., Seeb, J.E., Dann, T.H., Schindler, D.E., Seeb, L.W., 2014. Signals of heterogeneous selection at an MHC locus in geographically proximate ecotypes of sockeye salmon. *Mol. Ecol.* 23, 5448–5461.
- Lighten, J., van Oosterhout, C., Bentzen, P., 2014a. Critical review of NGS analyses for de novo genotyping multigene families. *Mol. Ecol.* 23, 3957–3972.
- Lighten, J., van Oosterhout, C., Paterson, I.G., McMullan, M., Bentzen, P., 2014b. Ultra-deep Illumina sequencing accurately identifies MHC class IIb alleles and provides evidence for copy number variation in the guppy (*Poecilia reticulata*). *Mol. Ecol. Resour.* 14, 753–767.
- Lynch, M., O’Hely, M., 2001. Captive breeding and the genetic fitness of natural populations. *Conserv. Genet.* 2, 363–378.
- Manlick, P.J., Woodford, J.E., Gilbert, J.H., Eklund, D., Pauli, J.N., 2017. Augmentation provides nominal genetic and demographic rescue for an endangered carnivore. *Conserv. Lett.* 10, 178–185.
- Meyer, D., Single, R.M., Mack, S.J., Erlich, H.A., Thomson, G., 2006. Signatures of Demographic History and Natural Selection in the Human Major Histocompatibility Complex Loci. *Genetics* 173, 2121–2142.
- Meyer-Lucht, Y., Sommer, S., 2005. MHC diversity and the association to nematode parasitism in the yellow-necked mouse (*Apodemus flavicollis*). *Mol. Ecol.* 14, 2233–2243.
- Milinski, M., 2006. The major histocompatibility complex, sexual selection, and mate choice. *Annu Rev Ecol Evol Syst* 37, 159–186.



- Miller, K.M., Winton, J.R., Schulze, A.D., Purcell, M.K., Ming, T.J., 2004. Major Histocompatibility Complex Loci are Associated with Susceptibility of Atlantic Salmon to Infectious Hematopoietic Necrosis Virus. *Environ. Biol. Fishes* 69, 307–316.
- Moorkens, E.A., 2018. Short-term breeding: releasing post-parasitic juvenile *Margaritifera* into ideal small-scale receptor sites: a new technique for the augmentation of declining populations. *Hydrobiologia* 810, 145–155.
- Neff, B.D., Garner, S.R., Pitcher, T.E., 2011. Conservation and enhancement of wild fish populations: preserving genetic quality versus genetic diversity *Canadian Journal of Fisheries and Aquatic Sciences*, 68(6), pp.1139-1154.
- Nei, M., 1972. Genetic Distance between Populations. *Am. Nat.* 106, 283–292.
- Nykänen, M., Huusko, A., 2002. Suitability criteria for spawning habitat of riverine European grayling. *J. Fish Biol.* 60, 1351–1354.
- Peakall, R., Smouse, P.E., 2012. GenAEx 6.5: genetic analysis in Excel. Population genetic software for teaching and research—an update. *Bioinformatics* 28, 2537–2539.
- Pedersen, S.S., Dieperink, C., Geertz-Hansen, P., 2003. Fate of stocked trout *Salmo trutta* L. in Danish streams: Survival and exploitation of stocked and wild trout by anglers. *Int. J. Ecohydrol. Hydrobiol.* 1, 39–50.
- Peng, B., Kimmel, M., 2005. simuPOP: a forward-time population genetics simulation environment. *Bioinformatics* 21, 3686–3687.
- Persat, H., Mattersdorfer, K., Charlat, S., Schenekar, T., Weiss, S., 2016. Genetic integrity of the European grayling (*Thymallus thymallus*) populations within the Vienne River drainage basin after five decades of stockings. *Cybium* 40, 7–20.
- Pertoldi, C., Bijlsma, R., Loeschcke, V., 2007. Conservation genetics in a globally changing environment: present problems, paradoxes and future challenges. *Biodivers. Conserv.* 16, 4147–4163.
- Piertney, S.B., Webster, L.M.I., 2010. Characterising functionally important and ecologically meaningful genetic diversity using a candidate gene approach. *Genetica* 138, 419–432.
- Pitcher, T.E., Neff, B.D., 2007. Genetic quality and offspring performance in Chinook salmon: implications for supportive breeding. *Conserv. Genet.* 8, 607–616.
- Pylkkö, P., Suomalainen, L.-R., Tirola, M., Valtonen, E.T., 2006. Evidence of enhanced bacterial invasion during *Diplostomum spathaceum* infection in European grayling, *Thymallus thymallus* (L.). *J. Fish Dis.* 29, 79–86.
- Quader, S., 2005. Mate choice and its implications for conservation and management. *Curr. Sci.* 89, 1220–1229.
- Rousset, F., 2008. genepop'007: a complete re-implementation of the genepop software for Windows and Linux. *Mol. Ecol. Resour.* 8, 103–106.
- Ryman, N., Laikre, L., 1991. Effects of Supportive Breeding on the Genetically Effective Population Size. *Conserv. Biol.* 5, 325–329.
- Salomonsen, J., Marston, D., Avila, D., Bumstead, N., Johansson, B., Juul-Madsen, H., Olesen, G.D., Riegert, P., Skjødt, K., Vainio, O., Wiles, M.V., Kaufman, J., 2003. The properties of the single chicken MHC classical class II  $\alpha$  chain (*B-LA*) gene indicate an ancient origin for the DR/E-like isotype of class II molecules. *Immunogenetics* 55, 605–614.
- Savage, A.E., Zamudio, K.R., 2011. MHC genotypes associate with resistance to a frog-killing fungus. *Proc. Natl. Acad. Sci.* 108, 16705–16710.
- Schenekar, T., Weiss, S., 2017. Selection and genetic drift in captive versus wild populations: an assessment of neutral and adaptive (MHC-linked) genetic variation in wild and hatchery brown trout (*Salmo trutta*) populations. *Conserv. Genet.* 18, 1011–1022.
- Setchell, J.M., Charpentier, M.J.E., Abbott, K.M., Wickings, E.J., Knapp, L.A., 2010. Opposites attract: MHC-associated mate choice in a polygynous primate. *J. Evol. Biol.* 23, 136–148.
- Sgrò, C.M., Lowe, A.J., Hoffmann, A.A., 2011. Building evolutionary resilience for conserving biodiversity under climate change. *Evol. Appl.* 4, 326–337.

- Sommer, S., 2005. The importance of immune gene variability (MHC) in evolutionary ecology and conservation. *Front Zool* 2, 16.
- Sommer, S., Courtiol, A., Mazzoni, C.J., 2013. MHC genotyping of non-model organisms using next-generation sequencing: a new methodology to deal with artefacts and allelic dropout. *BMC Genomics* 14, 542.
- Spielman, D., Brook, B.W., Frankham, R., 2004. Most species are not driven to extinction before genetic factors impact them. *Proc. Natl. Acad. Sci.* 101, 15261–15264.
- Stet, R.J.M., Kruiswijk, C.P., Dixon, B., 2003. Major Histocompatibility Lineages and Immune Gene Function in Teleost Fishes: The Road Not Taken. *Crit. Rev. Immunol.* 23, 473–488.
- Strandh, M., Westerdahl, H., Pontarp, M., Canbäck, B., Dubois, M.-P., Miquel, C., Taberlet, P., Bonadonna, F., 2012. Major histocompatibility complex class II compatibility, but not class I, predicts mate choice in a bird with highly developed olfaction. *Proc. R. Soc. Lond. B Biol. Sci.* 279, 4457–4463.
- Sutton, J.T., Nakagawa, S., Robertson, B.C., Jamieson, I.G., 2011. Disentangling the roles of natural selection and genetic drift in shaping variation at MHC immunity genes. *Mol. Ecol.* 20, 4408–4420.
- Swatdipong, A., Primmer, C.R., Vasemägi, A., 2010. Historical and recent genetic bottlenecks in European grayling, *Thymallus thymallus*. *Conserv. Genet.* 11, 279–292.
- Tapley, B., Bradfield, K.S., Michaels, C., Bungard, M., 2015. Amphibians and conservation breeding programmes: do all threatened amphibians belong on the ark? *Biodivers. Conserv.* 24, 2625–2646.
- Ujvari, B., Belov, K., 2011. Major Histocompatibility Complex (MHC) Markers in Conservation Biology. *Int. J. Mol. Sci.* 12, 5168–5186.
- Wahlund, S., 1928. Zusammensetzung von Populationen und Korrelationserscheinungen vom Standpunkt der Vererbungslehre aus betrachtet. *Hereditas* 11, 65–106.
- Watterson, G.A., 1978. The Homozygosity Test of Neutrality. *Genetics* 88, 405–417.
- Weir, B.S., Cockerham, C.C., 1984. Estimating F-Statistics for the Analysis of Population Structure. *Evolution* 38, 1358–1370.
- Whitlock, M.C., 2011. G'ST and D do not replace FST. *Mol. Ecol.* 20, 1083–1091.
- Wilson, T.K., 1963. How our rivers got their grayling. *Fishing*, 9–10.
- Woolland, J.V., Jones, J.W., 1975. Studies on grayling, *Thymallus thymallus* L., in Llyn Tegid and the upper River Dee, North Wales. *J. Fish Biol.* 7, 749–773.
- Zinkernagel, R.M., Doherty, P.C., 1974. Restriction of in vitro T cell-mediated cytotoxicity in lymphocytic choriomeningitis within a syngeneic or semiallogeneic system. *Nature* 248, 701–702.

### *Data Accessibility*

Genotype data is available from DRYAD under doi:10.5061/dryad.dj625ng; raw reads obtained from the Illumina Nanorun are deposited in the NCBI SRA archive under accession number SRP155806;

### *Author contributions*

V.H. performed the MHC genotyping laboratory work, conducted the data analysis and simulations and wrote the manuscript. All authors assisted in the research design and editing of the manuscript. J.S.E. conceived of the project.

### *Supporting Information*

**1.** *Results of the Ewens-Watterson test for MH and microsatellite markers*

Tables:

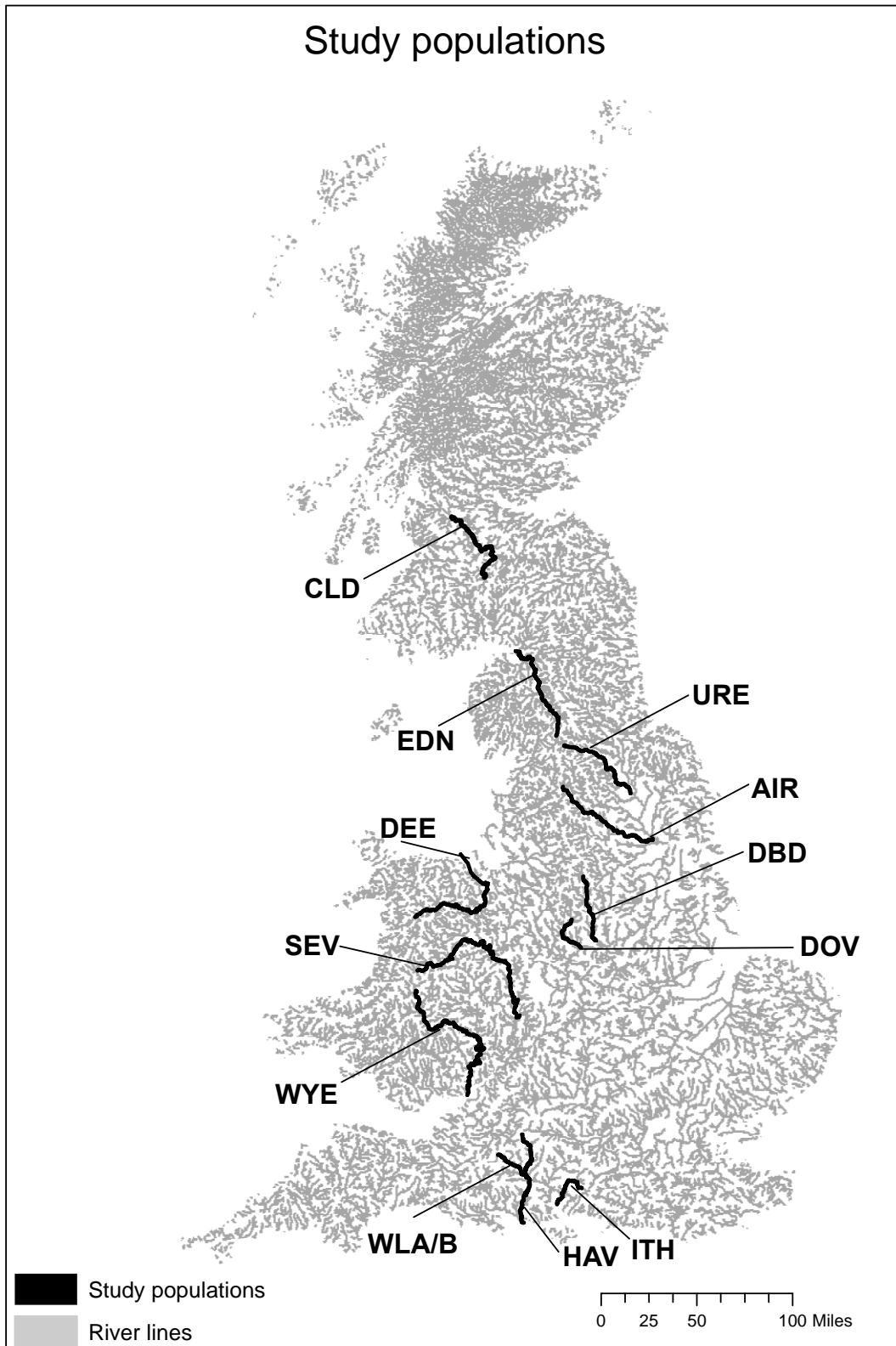
**Table 1:** Sample locations with population classification and summary of genetic diversity for microsatellite markers (from Dawnay et al.(2011)), MH class II  $\alpha$  (DAA) and  $\beta$  chain (DAB). Population classifications are given for non-stocked native (N), stocked native (NS) and introduced (I) populations; number of genotyped samples (N) and allelic richness (Na), expected heterozygosity (He), observed heterozygosity (Ho), inbreeding coefficient ( $F_{IS}$ ), with values showing significant deviation from Hardy-Weinberg equilibrium after Hochberg-Bonferroni correction in bold and number of private alleles (NP) are given. For microsatellite markers estimated effective population size ( $N_e$ ), where (\*) indicates the detection of a bottleneck, is given; for MH loci average amino acid (AA) distance of alleles (PBR) within populations are given and mean pairwise AA distance across individuals for the whole sequence and only for the PBR; population abbreviations are followed as in Dawnay et al (2011)

Microsatellites (from Dawnay et al. (2011))								DAA							DAB								
pop	class	N	$N_e$	Na	He	Ho	$F_{IS}$	N	Na	He	Ho	$F_{IS}$	NP	Mean AA distance alleles PBR	Mean pairwise AA distance all/PBR	N	Na	He	Ho	$F_{IS}$	NP	Mean AA distance alleles PBR	Mean pairwise AA distance all/PBR
CLD	I	64	68.6*	2.3	0.39	0.37	0.05	40	2	0.31	0.38	-0.22	0	0.44	0.06/0.18	37	2	0.29	0.35	-0.2	0	0.5	0.07/0.18
EDN	I	45	48.7*	2.5	0.4	0.38	0.04	33	3	0.36	0.24	0.33	0	0.29	0.03/0.08	36	3.5	0.34	0.25	0.26	0	0.4	0.04/0.11
ITH	I	50	86.6	2.5	0.39	0.38	0.02	34	2	0.42	0.29	0.31	0	0.25	0.04/0.08	20	2	0.41	0.45	-0.1	0	0.65	0.1/0.29
DEE	N	52	43.2*	3.5	0.54	0.51	0.04	27	6.7	0.8	0.74	0.07	0	0.3	0.08/0.2	26	5.9	0.77	0.54	0.3	0	0.42	0.08/0.21
SEV	N	39	40.8	2.8	0.42	0.41	0.03	31	3	0.53	0.39	0.27	0	0.29	0.04/0.12	30	3.9	0.57	0.23	<b>0.6*</b>	0	0.42	0.03/0.1
URE	N	58	62.5	2.9	0.35	0.32	0.09	31	6	0.8	0.58	0.28	4	0.27	0.08/0.18	30	5	0.59	0.4	0.32	2	0.47	0.08/0.2
WYE	N	55	121	3	0.4	0.4	0	30	3.7	0.64	0.63	0.02	0	0.28	0.06/0.18	22	3	0.65	0.55	0.16	0	0.47	0.08/0.23
WLA/B	N	48/51	33.5*	2.2	0.34	0.32	0.05	34	4	0.73	0.68	0.07	0	0.29	0.08/0.22	25	4	0.75	0.56	0.26	0	0.43	0.09/0.25
AIR	NS	39	63.9	3	0.45	0.44	0.02	31	4	0.59	0.42	0.3	1	0.23	0.05/0.1	29	4	0.54	0.28	<b>0.5*</b>	1	0.42	0.05/0.13
DBD	NS	39	36.9	2.7	0.42	0.45	0.07	35	3.4	0.54	0.57	-0.06	2	0.32	0.1/0.26	36	2	0.5	0.58	-0.18	0	0.5	0.11/0.29
DOV	NS	50	64.4	2.6	0.35	0.32	0.04	26	2.8	0.3	0.23	0.23	0	0.33	0.04/0.1	35	2.9	0.3	0.26	0.14	0	0.51	0.05/0.13
HAV	NS	58	32.5*	2.5	0.42	0.45	0.07	37	4.9	0.73	0.7	0.04	0	0.34	0.09/0.25	33	4.9	0.75	0.73	0.03	0	0.46	0.12/0.33

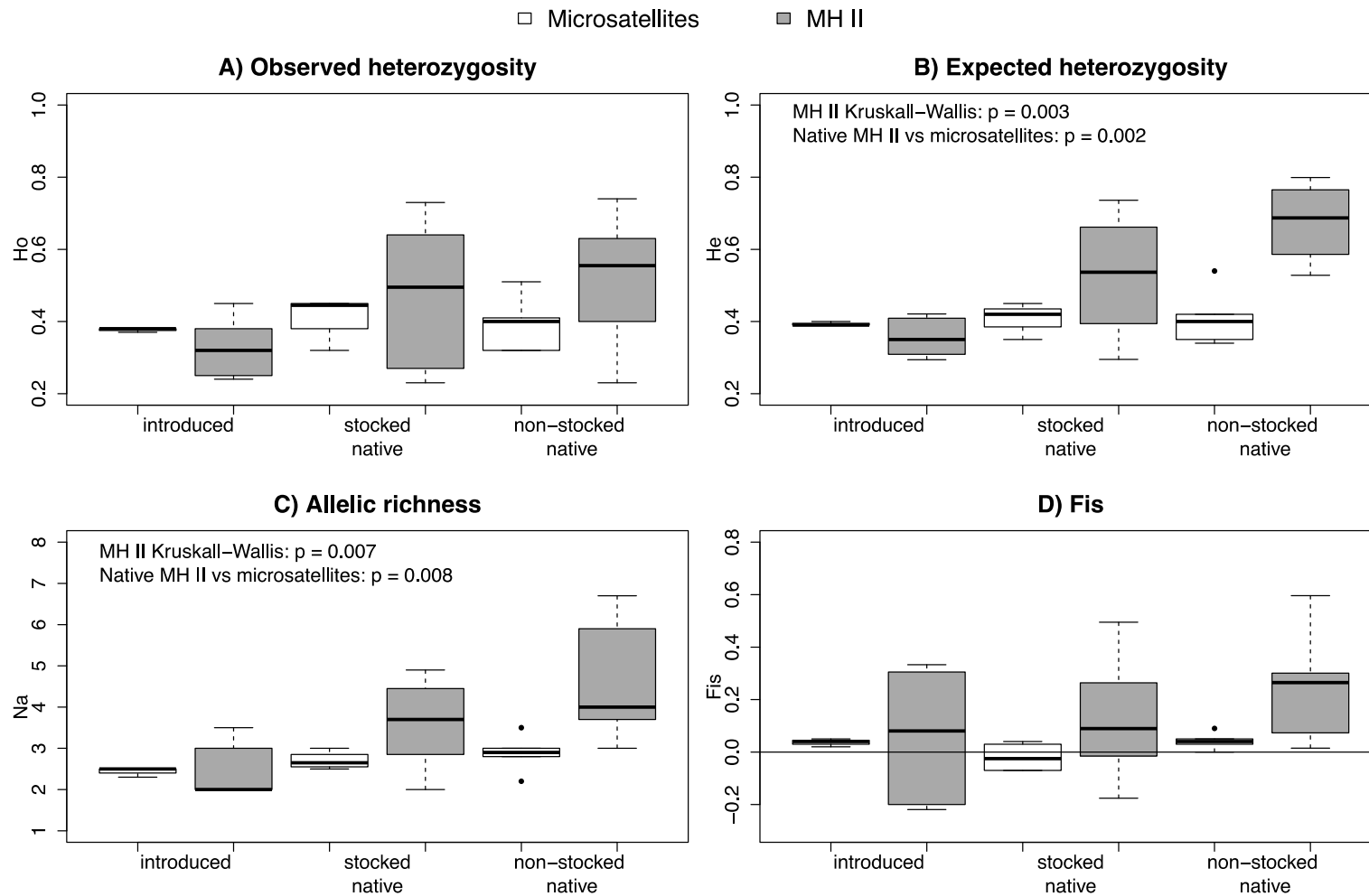
**Table 2:** Pairwise linear  $F_{st}$  ( $F_{st}/(1-F_{st})$ ) between all populations for each gene individually

DAA	AIR	CLD	DEE	DBD	DOV	EDN	HAV	ITH	SEV	URE	WYE
CLD	<b>0.81</b>										
DEE	<b>0.28</b>	<b>0.29</b>									
DBD	<b>0.53</b>	<b>0.14</b>	<b>0.16</b>								
DOV	<b>0.74</b>	-0.01	<b>0.26</b>	0.15							
EDN	<b>0.69</b>	0.01	<b>0.22</b>	<b>0.15</b>	-0.01						
HAV	<b>0.33</b>	<b>0.31</b>	<b>0.05</b>	<b>0.21</b>	<b>0.28</b>	<b>0.25</b>					
ITH	<b>0.65</b>	<b>1.10</b>	<b>0.26</b>	<b>0.74</b>	<b>1.07</b>	<b>0.95</b>	<b>0.33</b>				
SEV	<b>0.73</b>	<b>0.95</b>	<b>0.32</b>	<b>0.28</b>	<b>0.96</b>	<b>0.85</b>	<b>0.48</b>	<b>1.04</b>			
URE	<b>0.28</b>	<b>0.50</b>	<b>0.15</b>	<b>0.31</b>	<b>0.45</b>	<b>0.42</b>	<b>0.19</b>	<b>0.53</b>	<b>0.47</b>		
WYE	<b>0.49</b>	<b>0.52</b>	<b>0.13</b>	<b>0.27</b>	<b>0.49</b>	<b>0.36</b>	<b>0.26</b>	<b>0.71</b>	<b>0.23</b>	<b>0.28</b>	
WLA/B	<b>0.39</b>	<b>0.78</b>	<b>0.10</b>	<b>0.48</b>	<b>0.72</b>	<b>0.66</b>	<b>0.09</b>	<b>0.21</b>	<b>0.57</b>	<b>0.27</b>	<b>0.40</b>
DAB	AIR	CLD	DEE	DBD	DOV	EDN	HAV	ITH	SEV	URE	WYE
CLD	<b>0.99</b>										
DEE	<b>0.45</b>	<b>0.26</b>									
DBD	<b>0.68</b>	0.10	<b>0.14</b>								
DOV	<b>0.66</b>	0.00	<b>0.17</b>	0.07							
EDN	<b>1.03</b>	-0.01	<b>0.27</b>	<b>0.15</b>	0.01						
HAV	<b>0.50</b>	<b>0.37</b>	<b>0.09</b>	<b>0.25</b>	<b>0.26</b>	<b>0.39</b>					
ITH	<b>0.69</b>	<b>0.87</b>	<b>0.27</b>	<b>0.53</b>	<b>0.59</b>	<b>0.91</b>	<b>0.29</b>				
SEV	<b>0.89</b>	<b>0.98</b>	<b>0.26</b>	<b>0.37</b>	<b>0.74</b>	<b>1.04</b>	<b>0.50</b>	<b>0.71</b>			
URE	<b>0.43</b>	<b>0.12</b>	<b>0.13</b>	<b>0.15</b>	<b>0.06</b>	<b>0.12</b>	<b>0.19</b>	<b>0.39</b>	<b>0.64</b>		
WYE	<b>0.72</b>	<b>0.76</b>	<b>0.13</b>	<b>0.37</b>	<b>0.55</b>	<b>0.76</b>	<b>0.37</b>	<b>0.54</b>	<b>0.09</b>	<b>0.43</b>	
WLA/B	<b>0.52</b>	<b>0.63</b>	0.07	<b>0.39</b>	<b>0.43</b>	<b>0.67</b>	0.01	<b>0.28</b>	<b>0.52</b>	<b>0.28</b>	<b>0.38</b>

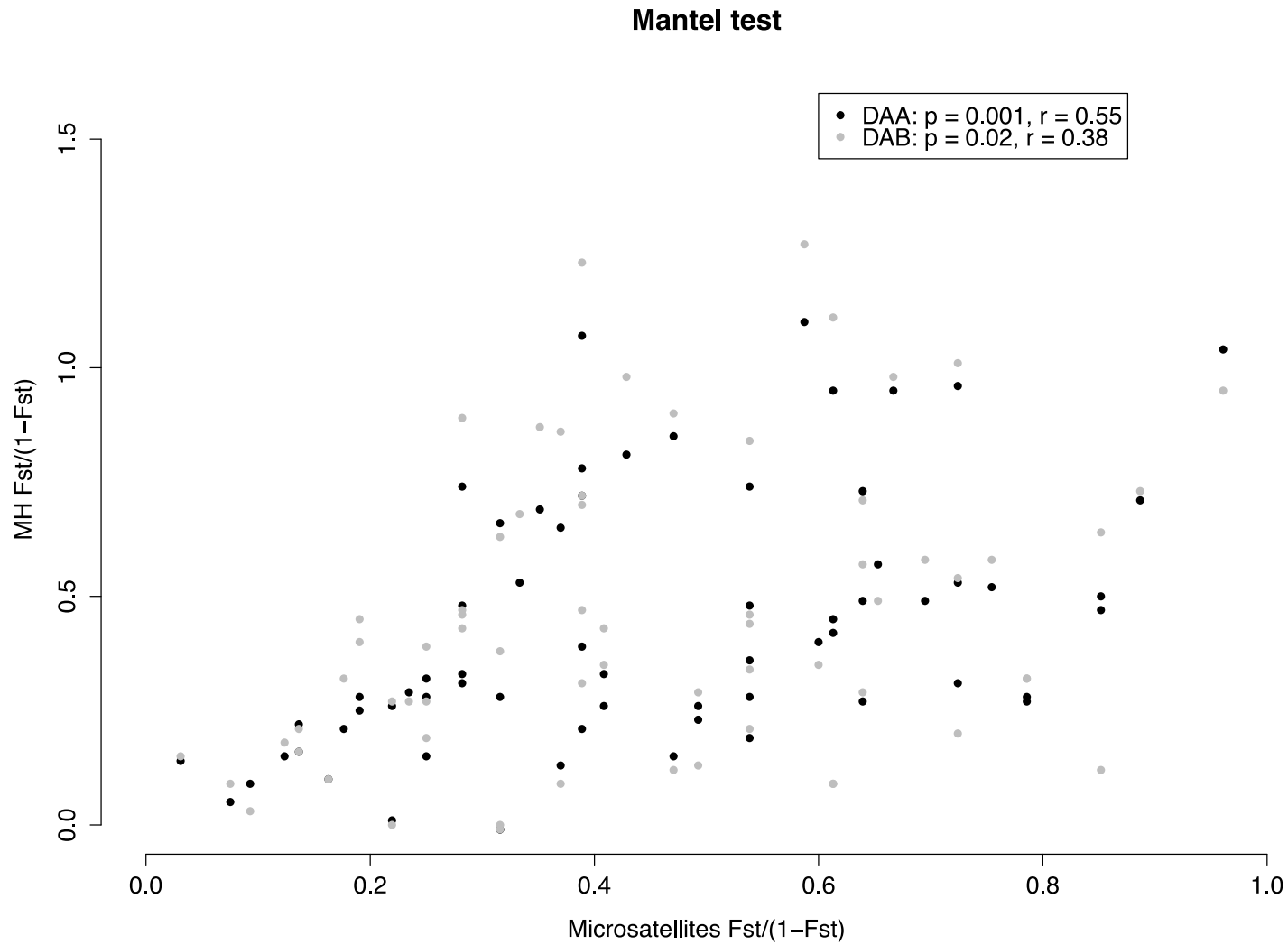
Figures



**Figure 1:** Populations genotyped at MH II DAA and DAB markers in this study in black: Clyde (CLD), Dee (DEE), Derbyshire Derwent (DBD), Dove (DOV), Eden (EDN), Hampshire Avon (HAV), Itchen (ITH), Severn (SEV), Ure (URE), Wye (WYE), Wylfe (WLA/B); River lines across the UK are shown in grey



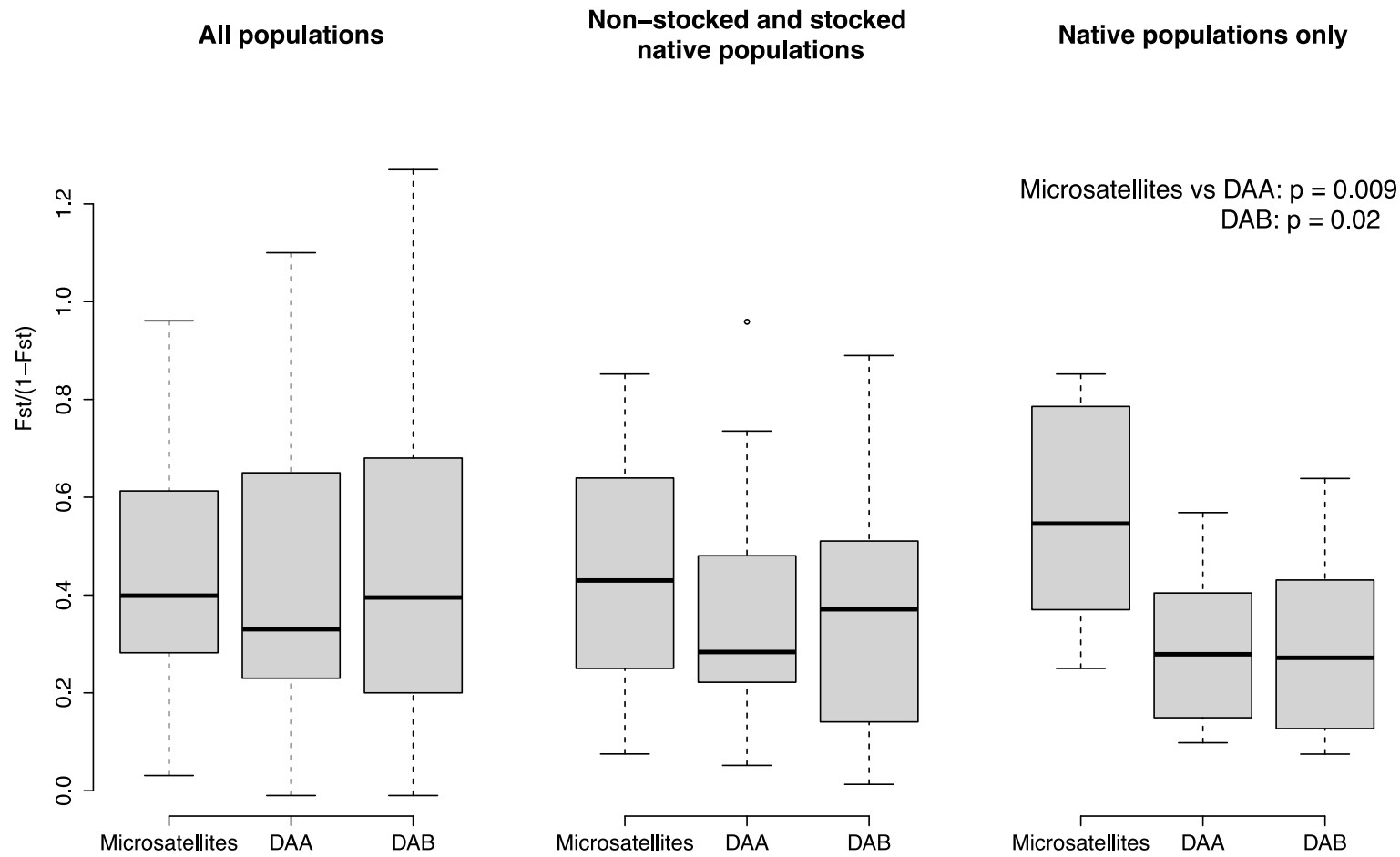
**Figure 2:** Comparison between microsatellite and MH II measurements of genetic diversity (A: observed heterozygosity, B: expected heterozygosity, C: allelic richness ( $N_a$ ) and D: inbreeding coefficient ( $F_{IS}$ )) across management classes consisting of introduced, stocked native and non-stocked native populations; Significant differences after correction for multiple testing are shown for Kruskal-Wallis tests across management classes and clustered Mann-Whitney-Wilcoxon test between markers within each management class;



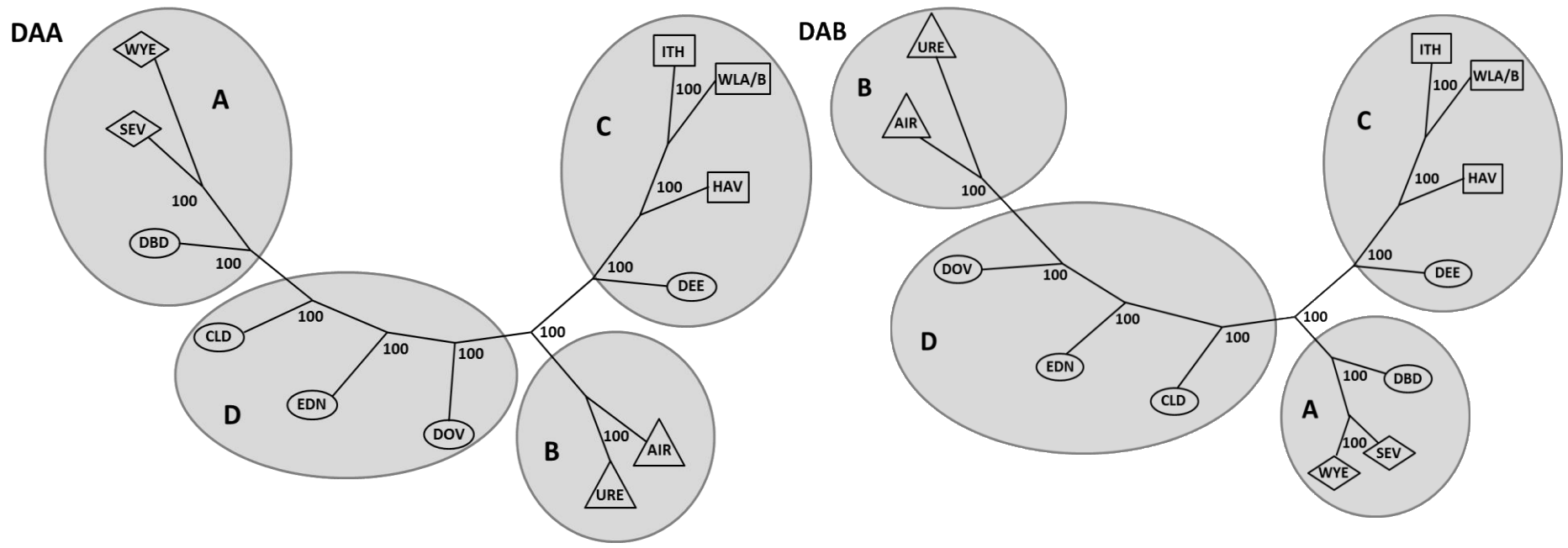
**Figure 3:** Relationship of pairwise  $F_{ST}/(1-F_{ST})$  of microsatellites and MH II DAA and DAB genes: P values are given with Spearman correlation coefficients in significant cases;



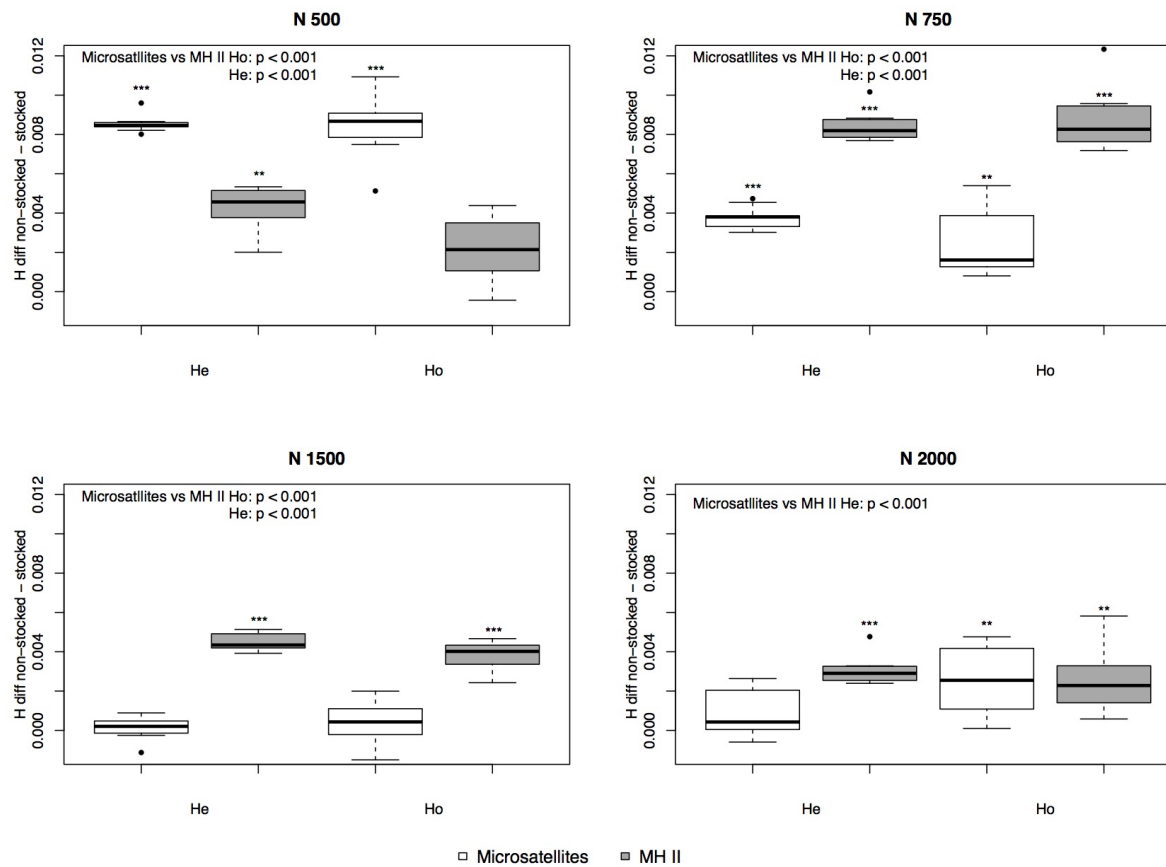
## Population divergence for different markers



**Figure 4:** Population divergence measured as pairwise  $F_{ST}/(1-F_{ST})$  for the different markers studied: P values of Mann-Whitney-Wilcoxon tests with significant differences between markers are given; A: between all populations; B: considering only native and native stocked populations; C: considering only purely native populations;



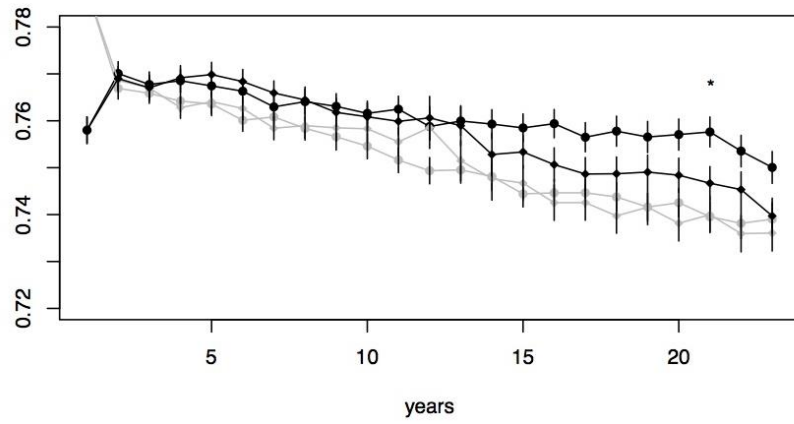
**Figure 5:** Unrooted phylogenetic trees based on Nei's genetic distance for DAA and DAB of the MH class II: Bootstrap support based on 2000 replicates is given; shapes around population abbreviations reflect the assignment to clusters based on neutral markers in Dawnay et al. (2011), with diamonds for cluster A, triangles for cluster B, boxes for cluster C and circles for cluster D;



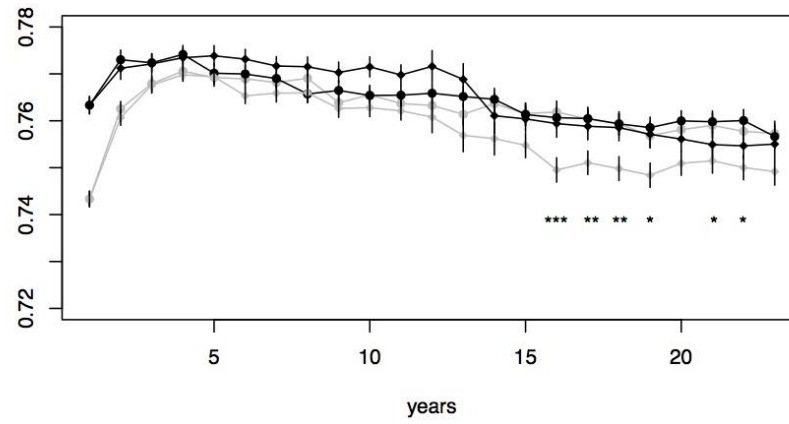
**Figure 6:** Difference in expected ( $H_e$ ) and observed heterozygosity ( $H_o$ ) between non-stocked and stocked simulated replicates (averaged across the ten following years after stocking events) are shown for neutral and MH markers at population census sizes of 500, 750, 1500 and 2000 (the ratio of naturally produced offspring to those stocked were roughly 0.5:1, 0.8:1, 1.6:1 and 2:1 respectively); Significant differences after correction for multiple testing are shown above plots with \*\* indicating  $p$ -values below 0.01 and \*\*\*  $p$ -values below 0.001; significant comparison between neutral and MH marker differences are printed;

**A) observed heterozygosity over time in stocked and non-stocked replicates**

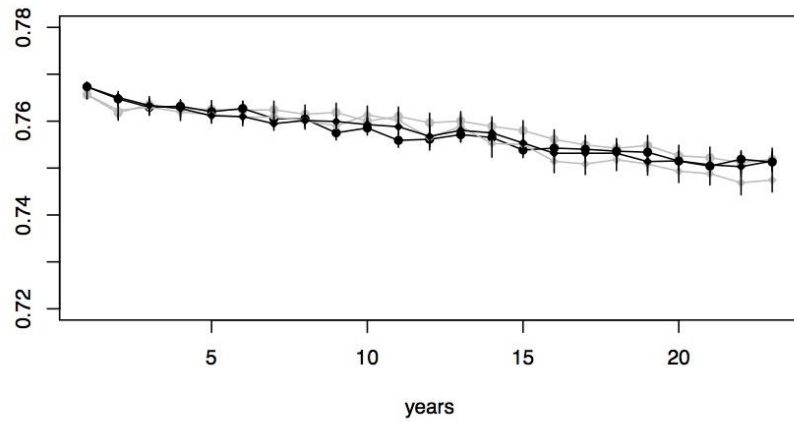
**N 500**



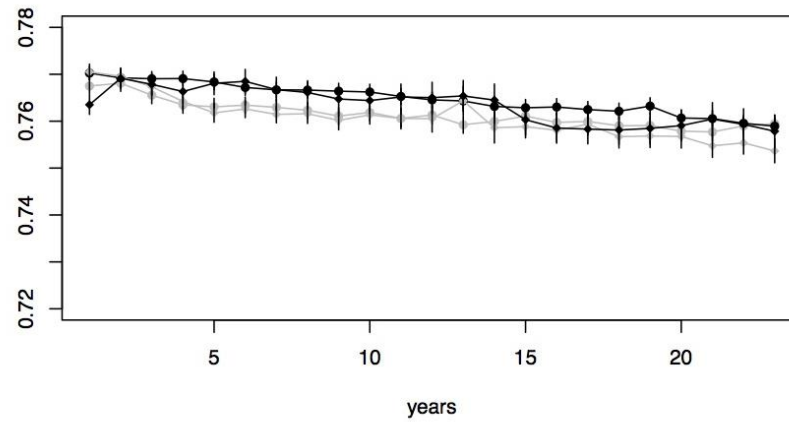
**N 750**



**N 1500**



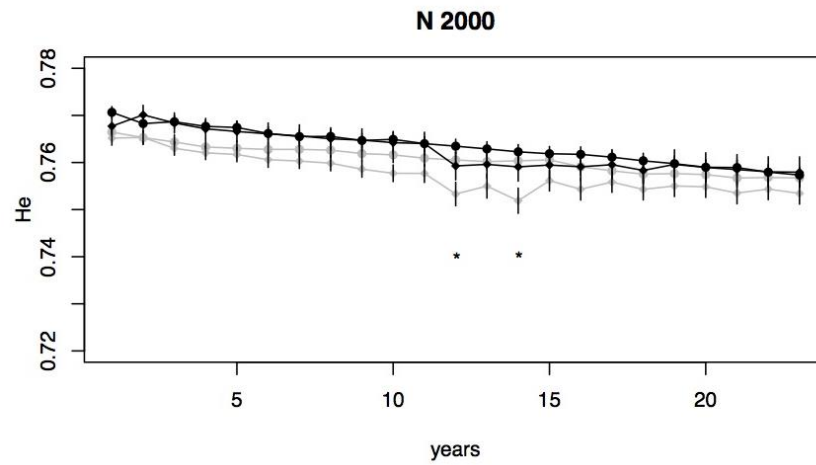
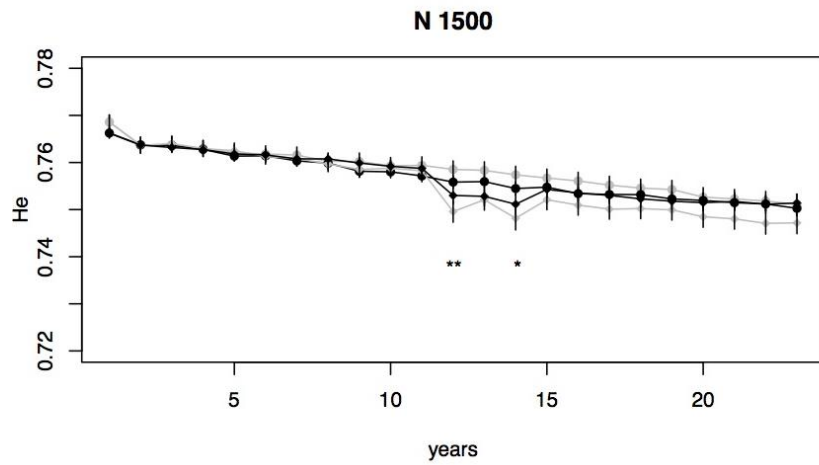
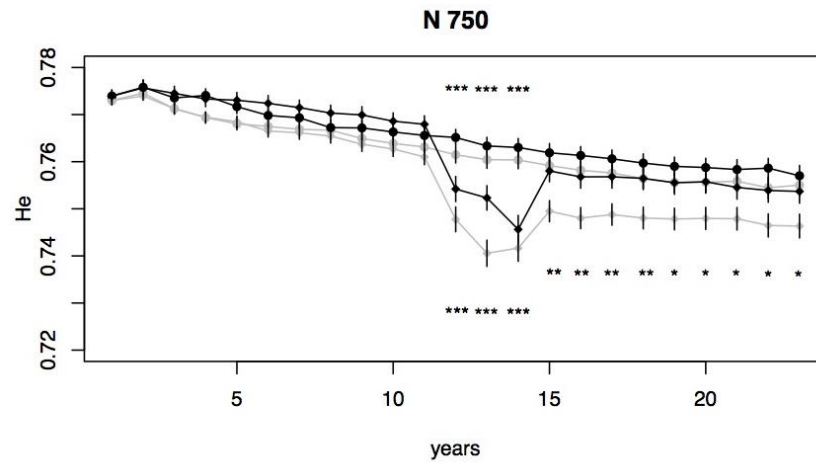
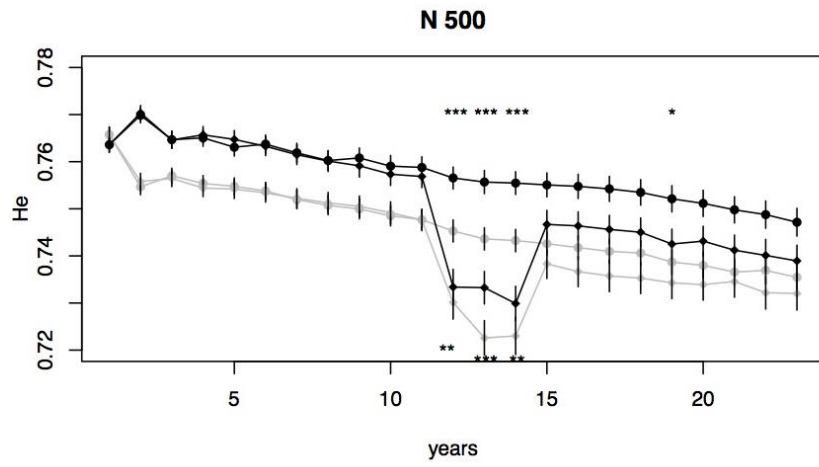
**N 2000**



● Microsatellites non-stocked  
◆ MH II stocked

◆ Microsatellites stocked  
● MH II non-stocked

**B) expected heterozygosity over time in stocked and non-stocked replicates**



- Microsatellites non-stocked
- ◆ Microsatellites stocked
- ◆ MH II stocked
- MH II non-stocked

Figure 7: Measurements of observed (A) and expected heterozygosity (B) are averaged across two MH II and two neutral loci and 100 replicates respectively; results are shown when stocking was implemented at year 11, 12 and 13 or when no stocking was implemented. Significant differences after correcting for multiple testing (Hochberg and Benjamini, 1990) between stocked and non-stocked replicates are indicated (Mann-Whitney Wilcoxon tests): \*\*\*  $p < 0.001$ , \*\*  $p < 0.01$ , \*  $p < 0.05$