

1 **RADseq and mate choice assays reveal unidirectional gene flow among three**
2 **lamprey ecotypes despite weak assortative mating: insights into the**
3 **formation and stability of multiple ecotypes in sympatry.**

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20 Running title: Gene flow among lamprey ecotypes.

21 ABSTRACT

22 Adaptive divergence with gene flow often results in complex patterns of variation within taxa
23 exhibiting substantial ecological differences among populations. One example where this may
24 have occurred is the parallel evolution of freshwater-resident nonparasitic lampreys from
25 anadromous parasitic ancestors. Previous studies have focused on transitions between these
26 two phenotypic extremes but here we considered more complex evolutionary scenarios where
27 an intermediate freshwater form that remains parasitic is found sympatrically with the other
28 two ecotypes. Using population genomic analysis (Restriction Associated DNA sequencing)
29 we found that a freshwater-parasitic ecotype was highly distinct from an anadromous-parasitic
30 form ($Q_{\text{lake-P}} = 96.8\%$, $F_{\text{st}} = 0.154$), but that a freshwater-nonparasitic form was almost
31 completely admixed in Loch Lomond, Scotland. Demographic reconstructions indicated that
32 both freshwater populations likely derived from a common freshwater ancestor. However,
33 while the nonparasitic ecotype has experienced high levels of introgression from the
34 anadromous-parasitic ecotype ($Q_{\text{anad-P}} = 37.7\%$), there is no evidence of introgression into the
35 freshwater-parasitic ecotype. Paradoxically, mate choice experiments predicted high potential
36 for gene flow: males from all ecotypes were stimulated to spawn with freshwater-parasitic
37 females, which released gametes in response to all ecotypes. Differentially fixed single
38 nucleotide polymorphisms identified genes associated with growth and development, which
39 could possibly influence the timing of metamorphosis, resulting in significant ecological
40 differences between forms. This suggests that multiple lamprey ecotypes can persist in
41 sympatry following shifts in adaptive peaks, due to environmental change during their repeated
42 colonization of post-glacial regions, followed by periods of extensive gene flow among such
43 diverging populations.

44

INTRODUCTION

45

46 Colonization of novel environments is predicted to promote intraspecific phenotypic variation
47 as a consequence of intense competition for limiting resources (van Valen 1965). Viewed
48 across longer periods of evolutionary time, this process manifests as a continuum of increasing
49 differentiation; ranging from resource polymorphisms within panmictic populations, to
50 complete and irreversible reproductive isolation between species (Hendry *et al.* 2009). Thus
51 the separation of one lineage of organisms from another (speciation) can be thought of most
52 simply as a consequence of adaptation to a different environment; which is made easier when
53 coupled with barriers to movement between them. Adaptive genetic divergence, where
54 different alleles prove advantageous for groups living in different environments (ecological
55 speciation), is predicted to reduce gene flow in those areas of the genome associated with well-
56 adapted phenotypes (Schluter 2009). Reproductive isolation can, therefore, stem from
57 ecological speciation in any geographic context where two populations are diverging
58 (allopatric, parapatric, sympatric), even in the face of gene flow (Smadja & Butlin 2011).
59 However, ecological speciation requires that: *i*) populations persist after shifts in adaptive
60 peaks (e.g. due to environmental change or colonization of new environments; Pavey *et al.*
61 2010; Rogers *et al.* 2012), and *ii*) genetic divergence is associated with pre- or post-zygotic
62 reproductive isolation (Rundle & Nosil 2005; Nosil 2012).

63 Northern freshwater fishes make particularly good models for understanding this
64 continuum from ecotype formation through speciation. The advances and retreats of glaciers
65 during the Quaternary period (15,000 – 10,000 years ago) had a profound impact on
66 phylogeographic distributions (Petit *et al.* 2003; Rowe *et al.* 2004), with post-glacial range
67 expansion varying greatly across the northern hemisphere (Hays *et al.* 1976; Schluter 1996).
68 This would have created infrequent opportunity for secondary contact between populations

69 inhabiting discrete and partly isolated post-glacial lakes during this period (Taylor & McPhail
70 2000; Lyons 2003). Parallel colonization of newly available lakes and streams led to the
71 evolution of distinct gene pools within species for multiple fish groups (Rogers *et al.* 2013).
72 As a consequence, several post-glacial fishes are characterized by substantial levels of
73 polymorphism driven by ecological opportunity. Such polymorphisms exhibit both adaptive
74 divergence (e.g. physiological differences, changes in maturation timing) and population
75 persistence in the face of exposure to environmental change (e.g. dietary shifts) (Taylor 1999;
76 Schluter 2000). For example, whitefish (*Coregonus* spp.) are highly polymorphic, with
77 phenotypes readily identifiable by differences in head shape, gill raker number and body shape
78 (Svårdson 1979; Kahilainen & Østbye 2006). In particular, gill raker number is a heritable trait
79 influencing trophic ecology and plays a central role in whitefish diversification within lakes
80 (Rogers & Bernatchez 2007). Furthermore, Arctic charr (*Salvelinus alpinus*) is arguably the
81 most variable of vertebrates (Klemetsen 2013), with a large number of descriptions of discrete
82 trophic specialists characterized by morphological differences functionally associated with
83 alternative feeding strategies (e.g. Hooker *et al.* 2016), but constrained by ecological
84 opportunity (Recknagel *et al.* 2017). Populations at different stages of the speciation process
85 are useful for exploring mechanisms and constraints associated with phenotypic diversity and
86 the evolution and maintenance of polymorphisms. Thus, the adaptation of fishes to newly
87 available environments in recently glaciated freshwater systems, the rapid evolution of
88 reproductive isolation between ecotypes, and their persistence in the face of gene flow make
89 them excellent systems with which to investigate ecological speciation (Schluter 1996).

90 One conspicuous, yet understudied example of differentiation in response to ecological
91 opportunity has occurred among lampreys (Petromyzontiformes), where several genera contain
92 species pairs or sister species complexes, where a marine-migrating ancestral species has given

93 rise to a single, or multiple, freshwater-resident species (Zanandrea 1959; Vladykov & Kott
94 1979). These are often phenotypically and genetically similar to their anadromous counterparts
95 (Hardisty & Potter 1971), and may therefore represent diverging ecotypes of single species
96 (Yamazaki & Goto 2000; Salewski 2003; Docker 2009). Following a protracted larval stage,
97 all lampreys undergo metamorphosis and either: *i*) migrate to feed on other fish as ectoparasites
98 in freshwater/marine environments prior to maturation or, *ii*) bypass the juvenile trophic phase
99 and instead undertake sexual maturation soon after metamorphosis (Docker 2009). Lamprey
100 ecotypes thus exhibit a disparity in body size at maturity as a consequence of post-metamorphic
101 feeding (or the lack thereof) but most return to the upper reaches of streams to spawn (Moser
102 *et al.* 2015; Johnson *et al.* 2015). It is widely assumed that size-assortative mating has resulted
103 in pre-zygotic reproductive isolation between lamprey ecotypes spawning in syntopic habitats
104 (Hardisty 1963; Beamish & Neville 1992) but this has rarely been tested experimentally (but
105 see Beamish & Neville 1992).

106 European river (*Lampetra fluviatilis*) and brook lampreys (*L. planeri*) are common to
107 coastal watersheds in Europe and are representative of a typical lamprey species pair; with the
108 former being an anadromous parasite as an adult and the latter a freshwater-resident
109 nonparasitic species. Genetic differentiation between them is only apparent in former glacial
110 refugia (e.g. Iberia, Mateus *et al.* 2013), with no evidence for species-specific fixed genetic
111 differences elsewhere, possibly as a result of ongoing gene flow (Schreiber & Engelhorn 1998;
112 Espanhol *et al.* 2007; Pereira *et al.* 2011; Bracken *et al.* 2015; Rougemont *et al.* 2015, 2016,
113 2017; Mateus *et al.* 2016). This has led to the persistence of two alternative hypotheses: *i*) these
114 species are the result of very recent divergence (either a single divergence event or multiple
115 parallel events) or *ii*) that these lampreys comprise just a single species consisting of multiple
116 ecotypes (Espanhol *et al.* 2007). However, the situation could be even more complicated than

117 this because some catchments include more than just the two phenotypic extremes (parasitic
118 anadromous and nonparasitic freshwater forms): in some locations an intermediate form
119 (usually considered an ecotype of *L. fluviatilis*) that is freshwater resident but continues to feed
120 parasitically has been reported (Collett 1905; Tuunainen *et al.* 1980; Morris 1989; Inger *et al.*
121 2010; Tsimbalov *et al.* 2015).

122 Since all previous studies have focused on the two extremes, the relative importance of
123 two key life-history elements fundamental to lamprey speciation remains unresolved: the
124 transition from a migratory to a freshwater resident life cycle, and the switch from parasitic to
125 nonparasitic feeding strategies. Whether or not this two-part transition has resulted from a
126 single evolutionary step repeatedly among lamprey species pairs remains an unanswered but
127 important question. Although restricted to only a few large postglacial lakes in Europe, the
128 freshwater parasitic ecotype has frequently been hypothesized to represent a necessary
129 transitional stage during lamprey speciation in post-glacial regions (Zanandrea 1959, 1961;
130 Beamish 1985; Morris 1989; Salewski 2003; Hardisty 2006). Alternatively, such populations
131 could represent an independent evolutionary step that only persists under certain environmental
132 contexts (e.g. allopatry in deep post-glacial lakes). However, no studies to date have compared
133 genetic and behavioural isolation between all three ecotypes where they occur sympatrically to
134 distinguish between these hypotheses.

135 In the Loch Lomond catchment, Scotland, U.K., which is a large post-glacial lake basin,
136 three lamprey ecotypes occur: anadromous parasitic (which we refer to as anad-P), freshwater-
137 resident parasitic (lake-P) and freshwater-resident non-parasitic (non-P) (Morris 1989).
138 Analysis of mtDNA variation (2077 bp concatenated across the *ND3*, *cyt b* and *ATPase 6/8*
139 genes) from across this catchment area identified 7 haplotypes among 10 individuals; with a
140 single haplotype that was shared among all three ecotypes, one haplotype that was shared

141 between lake-P and non-P individuals and the rest unique to either anad-P or non-P individuals
142 (Hume 2013; Table 7.2). While this could suggest that the two freshwater-resident forms
143 shared a common ancestor more recently than the anad-P ecotype, the sampling was too
144 restricted to draw definitive conclusions. Moreover, analysis of mtDNA precludes
145 interpretation of hybridization and introgression, which has been found to have played an
146 important role in the differentiation of other lamprey populations (Rougemont *et al.* 2015,
147 2016, 2017). In this study, we thus focused on nuclear variation using restricted-associated
148 DNA sequencing, combined with behavioural observations of mating preferences. We use
149 samples collected from a single stream in the Loch Lomond catchment, supporting spawning
150 populations of all three ecotypes, to investigate the evolutionary forces resulting in the
151 formation of ecotypes in sympatry. Our aims were to: *i*) test the extent of differentiation among
152 the three lamprey ecotypes using a combination of population genomics and behavioural assays
153 to test patterns of reproductive isolation; *ii*) reconstruct the evolutionary history of these
154 ecotypes; and *iii*) identify regions of the genome associated with adaptive divergence in
155 relation to migration and feeding transitions.

156

157

METHODS

158 **Collection & maintenance of experimental animals**

159 Between October and April three ecotypes of lampreys were captured in static, double-funnel
160 traps (Morris & Maitland 1987) as they migrated upstream to spawning grounds in the Endrick
161 Water, Loch Lomond, Scotland, U.K. (56°3'17.3" N, 4°27'16.2" W) (Fig. 1). Twelve
162 individuals (six male, six female) each of: *i*) anadromous-parasitic *Lampetra fluviatilis*
163 (hereafter 'anad-P'); *ii*) freshwater resident-parasitic *L. fluviatilis* (hereafter 'lake-P'); and *iii*)

164 freshwater resident-nonparasitic *L. planeri* (hereafter ‘non-P’) were separated based on
165 published phenotypic descriptions (Morris 1989; Renaud 2011). Specifically, anad-P and non-
166 P can be easily distinguished by their discrete body sizes as adults (non-P rarely exceeds 170
167 mm whereas anad-P can exceed 490 mm total length (TL) (Renaud 2011). Lake-P is readily
168 separated from the others due its black coloration compared with olive-brown colors of the
169 other two (Morris 1989). Each ecotype was held separately in 175 L tanks using water drawn
170 from Loch Lomond at ambient temperature in a flow-through system, and exposed to artificial
171 light that tracked natural photoperiod. The same individuals ($n = 36$) were used in all
172 methodological approaches outlined below.

173

174 **1. Differentiation of Ecotypes**

175 *Genetic comparison of ecotypes*

176 *DNA sequencing and bioinformatic processing*

177 A small piece of tissue was removed from the second dorsal fin of each individual and stored
178 in 95% ethanol at -20°C until required. DNA was extracted using Qiagen DNeasy Blood &
179 Tissue Kits (QIAGEN Sample & Assay Technologies, Copenhagen, Denmark) following the
180 manufacturer’s instructions.

181 Extracted samples ($n = 36$; at least 5 µg per sample) were sent to the GenePool (now
182 Edinburgh Genomics, University of Edinburgh) where a paired-end restriction-site associated
183 DNA (RAD) library was prepared using *SbfI* enzyme digestion (Baird *et al.* 2008). The library
184 was sequenced on an Illumina HiSeq v3 on a single sequencing lane, with a target read length
185 of 100 bp. A total of *c.* 305 million reads were generated and analysed using STACKS (v.
186 0.99994, Catchen *et al.* 2011). Reads were first assigned to each of the sequenced individuals,

187 identified by a specific barcode at the beginning of each read and trimmed to a final length of
188 93 bp. All individual reads were then merged into stacks of loci, by allowing identical reads
189 and those with up to two variant positions to be merged into a single locus. Since there is not
190 a closely related reference genome (closest relative sequenced is *Petromyzon marinus*, ~16
191 million years divergence [Kuraku *et al.* 2006]), loci were assembled *de novo*. A catalogue of
192 loci was built based on all individuals, and each individual's loci were then matched to this
193 catalogue. Each locus was assigned to a specific ID and contained information on which, and
194 how many, individuals contained that locus, what the respective coverage was, and whether it
195 contained single nucleotide polymorphisms (SNPs). Average coverage per locus and individual
196 was 53.4x (median: 48.4x; standard deviation: 23.0x). Loci with coverage higher than 8x,
197 present in all three ecotypes, and in at least 50% of the individuals of each ecotype were
198 extracted for further genomic analyses. A single individual ('B09 Anad') of the anad-P ecotype
199 was excluded from all further genetic analyses because of limited data (median coverage 5x).

200

201 *Differentiation of ecotypes*

202 A total of 7678 loci were analysed using STRUCTURE (v. 2.3.4, Pritchard *et al.* 2000). For
203 polymorphic loci containing more than one SNP, only the first was included to minimize the
204 number of linked markers. Polymorphic loci that contained a SNP differentially fixed between
205 ecotypes and outlier loci (see *Detecting fixed SNPs and outlier loci*) were excluded from further
206 analysis, since these loci might be under selection and would not represent neutral genetic
207 differentiation between ecotypes. Population differentiation was assessed using a model of
208 admixture and correlated allele frequencies. A total of 500,000 Markov-chain Monte Carlo
209 (MCMC) repetitions were performed after discarding a burn-in of 100,000. We tested for up
210 to five different genetic clusters ($K = 1-5$) in four consecutive independent runs. To compare

211 likelihoods between different numbers of genetic clustering, ΔK was estimated (Evanno *et al.*
212 2005). Using the most likely number of clusters identified, admixture of each individual was
213 assessed using the Q -value in STRUCTURE, which indicates the proportion of each
214 individual's genome assigned to each of the pre-defined genetic clusters. In addition, a co-
215 ancestry matrix and population structure were estimated using FineRADStruture (Malinsky *et*
216 *al.* 2018), using default parameters. Degrees of admixture between individuals and genetic
217 clusters were compared to assess introgression and hybridization at the individual level and
218 ecotype/genetic clusters.

219 Additionally, a Principal Coordinate Analysis (PCoA) was performed on the same set
220 of polymorphic loci with the programme GenAlEx (v. 6.5., Peakall & Smouse 2012), with
221 interpolation of missing genotypes. We estimated the genetic variance explained by
222 differentiation among ecotypes using the AMOVA F_{st} implemented in GenAlEx and using
223 1000 permutations. The proportion of heterozygous loci (H_{obs}) was calculated for each
224 individual from the three ecotypes separately, and then compared between ecotypes. On
225 average, 16,541 loci (including monomorphic sites) were extracted per individual (9,177 -
226 19,440 loci) to derive these estimates.

227

228 ***Behavioural comparison of ecotypes***

229 *Preparation of experimental animals*

230 Following fin-clipping for genetic analysis all lamprey were returned to holding tanks, where
231 they were examined periodically to assess the progress of sexual maturation. Sexually mature
232 female lampreys were identified when they became swollen with eggs that were visible through
233 a patch of translucent skin near the cloacal opening and had developed a post-cloacal fin-fold.

234 Sexually mature male lampreys were identified by an obvious genital papilla that extended
235 several millimeters from the cloacal opening. All individuals were fully ripe (ovulating and
236 spermiating) prior to undertaking experiments.

237 Each of the 36 individuals from the three ecotypes (six males, six females per ecotype)
238 used for genetic analyses were included in mate choice tests. Lampreys were anaesthetized
239 using a benzocaine solution, measured to the nearest 1 mm L_T and had a Visible Implant Alpha
240 tag inserted beneath the skin on the dorsal surface of the branchial chamber to allow for easy
241 identification during/after the trial period. Prior to inclusion in tests, lampreys were held in 10
242 L tanks as same-sex, same-ecotype pairs to prevent spawning. These tanks were maintained
243 under the same conditions as the holding tanks.

244

245 *Experimental design*

246 Behavioural differences between ecotypes were examined by employing a multi-male mate
247 choice design, commonly used for testing for the presence of pre-zygotic barriers to gene flow
248 in closely related taxa (e.g. Kozak *et al.* 2013). An artificial stream measuring 5.72 m in length
249 was used to simulate natural conditions as closely as possible during each trial. The base of the
250 stream was covered to a depth of approximately 4 cm by gravel (0.5 – 2 cm diameter) collected
251 locally. The stream was partitioned by fine mesh (1 mm) screens, creating six discrete
252 experimental arenas measuring 91 cm long by 58 cm wide. Water from Loch Lomond was
253 pumped through the stream at velocities of 5 – 20 cm s^{-1} , and temperatures ranged 8.5 – 11.5°C,
254 which are typical of the conditions experienced by these lamprey in the wild (Maitland 1980).
255 Artificial, low-light levels on a natural photoperiod were maintained throughout to aid
256 observations.

257 Mate choice trials comprised a single focal female and three males (one representing
258 each ecotype). The members of each of these male triads remained constant throughout. Thus,
259 each of the 18 females was exposed to the same six male triads, resulting in a total of 108 trials.
260 For each trial, groups of the four experimental animals were placed in separate stream sections
261 and allowed to acclimate for five minutes before observations began. Each trial lasted six hours,
262 during which time all reproductive activity was recorded (direct observations). To prevent
263 exhaustion of the female's egg stock, they were removed after one trial and allowed to rest for
264 a minimum of six hours. Males were rested after two consecutive trials. Only females from the
265 same ecotype were tested in successive trials, to reduce any possible residual effects from
266 pheromones, or other chemical stimuli. Following each trial, the stream section was examined
267 for eggs, which were collected using a siphon. The gravel was then scoured to remove any
268 traces of nests, and that section allowed to dry for at least 12 hours. After all six females from
269 an ecotype were tested with the six male triads, the stream water was drained for a period of
270 24 hours, and the gravel scoured before refilling with fresh drawn water. The experimental
271 period lasted 15 days, during which no experimental animal died.

272 Lamprey spawning comprises several discrete behaviours (Johnson *et al.* 2015).
273 Typical spawning occurs within nests created in shallow gravel beds and begins when females
274 attach to an object immediately upstream of the nest. A male then glides along the female's
275 body, attaches to the female's head and wraps his tail around her trunk, forming a tight loop
276 that acts to express the ova from her body. Both partners vibrate rapidly as gametes are released
277 and fertilization takes place externally within the nest. The complete sequence of behaviours
278 leading to potential production of progeny thus begins with a male attaching to a female and
279 culminating in the release of gametes by both sexes. However, male attachment does not
280 always follow a gliding movement along the female's body, a female may not respond to a

281 male's attachment, or she may reject a male by detaching from the nest. Therefore, we
282 considered "successful spawning" as a combined measure of female and male motivation for
283 mating and "frequency of mating attempts" to quantify stimulation of male reproductive
284 behaviours in the presence of particular females.

285

286 *Statistical analysis*

287 A generalized linear mixed modeling (GLMM) approach was used to determine whether
288 successful spawning (female gamete release) or spawning attempts (stimulation of male
289 reproductive behaviours) varied in relation to female or male ecotype, or differences in size
290 between potential mating partners. Spawning success and whether or not a spawning attempt
291 was made by a given male when presented with a given female (classified as binary response
292 variables) were considered as separate response variables and analysed using the binomial
293 family using the *glmer* function, in the *lmer* package of the statistical programming
294 environment R (R Core Development Team, 2017). Given that ecotypes differed in size, we
295 specifically tested whether interactions between female ecotype*male ecotype, female
296 ecotype*size difference or male ecotype*size difference contributed significantly to the
297 observed variation. Female ecotype, male ecotype, size difference between potential partners
298 and the three interactions were treated as fixed effects. Since females were tested independently
299 and so could exhibit a choice but males were tested in trios and always had only one potential
300 female to mate with, female ID and male trio were considered as random effects, to account
301 for repeated measures. Model selection was performed using likelihood ratio tests, with fixed
302 variables not significantly altering the likelihood removed from the final model. All random
303 effects were included in the final model, to account for any variance they contributed.

304 The frequency of spawning attempts by males was also analysed as an integer trait and
305 analysed using the *glmer.nb* function in R, using a negative binomial as the underlying
306 distribution. We also tested whether the amount of admixture displayed by individuals
307 contributed to male mating attempts and successful spawning (both treated as binary
308 responses), by considering “own Q” values (i.e. the proportion of their genome attributed to
309 the dominant genotype found in their own ecotype) from the STRUCTURE analysis. We
310 included the random effects of female_ID and male_trio and considered the fixed effects of
311 female own_Q, male own_Q and their interactions. We also used a contingency chi-squared
312 analysis to determine whether there was a deviation from random mating when differences in
313 the relative frequency of activities in each of the ecotypes were accounted for.

314

315 **2. Estimating evolutionary history and admixture between ecotypes**

316 We assessed the likelihood of twelve alternative scenarios of lamprey demographic history by
317 testing all plausible evolutionary relationships between ecotypes and time points, direction and
318 extents of gene flow (Fig. 2). We used FASTSIMCOAL2 (Excoffier *et al.* 2013) to compare
319 all alternative scenarios of past history for ecotype evolution. The dataset included 7665
320 biallelic loci (i.e. excluding loci with more than two variants). We performed 25 runs of
321 100,000 simulations for each of the twelve evolutionary scenarios. Subsequently, we selected
322 the run with the highest likelihood score for each scenario and used the parameters estimated
323 as priors for an additional 1 million simulations to obtain final maximum likelihoods for each
324 scenario. We compared these likelihoods and calculated the AIC to select the best model. To
325 assess how robust the results obtained from FASTSIMCOAL2 analyses were, we performed
326 additional simulations using the Approximate Bayesian Computation software DIYABC
327 (Cornuet *et al.* 2014). Scenarios seven to twelve included gene flow not resulting in the

328 formation of a new lineage, an option currently not implemented in DIYABC and therefore not
329 included here. For the first six scenarios, Hudson's algorithm was applied to the simulated SNP
330 dataset (Fig. 2). We generated 5 million datasets that were compared to the empirical data to
331 assess how likely each of these scenarios was. This was done by computing posterior
332 probabilities (PP) for each model using the direct estimate implemented in the programme
333 (number of times simulated data of a model is closest to the observed data) and a logistic
334 regression estimate (regression of proportion of the model supported and distance of the
335 simulated data to the observed data) including 95% confidence intervals. Estimates from the
336 DIYABC simulations were compared to the FASTSIMCOAL2 likelihoods (Table S1). In
337 addition, we performed a TREEMIX search. First, migration events were set to 0 (no
338 admixture) and model residuals compared. Migration was then increased until model residuals
339 were equivalent between ecotypes (Pickrell & Pritchard 2012). Three-population f_3 statistics
340 (Reich *et al.* 2009) were estimated to assess whether lamprey ecotypes were significantly
341 admixed. Negative values in f_3 and in the Z-score indicate that a lineage has experienced
342 admixture events.

343

344 **3. Predicting candidate genes associated with adaptive divergence**

345 To identify genomic regions that may have facilitated adaptive differentiation, candidate SNPs
346 fixed between ecotypes were detected by extracting loci that were differentially fixed in at least
347 two ecotypes (for example 'A' in one, 'G' in another and 'A/T' in the third). Included loci had
348 to be present in at least two ecotypes and in at least four individuals per ecotype, resulting in a
349 final set of 7726 SNPs. In a second approach, a global outlier analysis was performed using
350 the programme BayeScan (v. 2.1, Foll & Gaggiotti 2008), with default conditions using loci
351 with a minor allele frequency (MAF) of 0.05. Additionally, outlier analyses between migration

352 strategy (anadromy *vs.* residency) and feeding type (parasitic *vs.* nonparasitic) were performed.
353 All loci that had a probability higher than $P = 0.5$ were treated as putative outliers.

354 Outlier loci and those that were differentially fixed between ecotypes were individually
355 checked against the Ensembl (v. 71, Flicek *et al.* 2014) sea lamprey (*P. marinus*) genomic
356 database to identify whether they were present in transcribed or non-transcribed genomic
357 locations. If they were part of a transcribed gene, the gene name was recorded and its function
358 and process was checked on the National Center for Biotechnology information (NCBI) gene
359 database (<http://www.ncbi.nlm.nih.gov/gene/>). Genomic location and gene ontology were
360 similarly checked for the SNPs differentially fixed between ecotypes.

361 Catalogued loci were identified from the first read and then assembled into contigs
362 using both the first and the second reads from the paired-end sequencing. A BLAST search
363 (<http://blast.ncbi.nlm.nih.gov/Blast.cgi>) was performed to match those loci to the genome of
364 the distantly related sea lamprey to identify their genomic location. At present, the sea lamprey
365 genome consists of more than 2500 unlinked scaffolds, making it difficult to gain a general
366 overview of the distribution of identified loci across the genome. However, we used the linkage
367 groups identified in a previous RAD analysis of the sea lamprey (Smith *et al.* 2014) to label
368 each scaffold that we found with a RAD marker and its position on the respective linkage
369 group. If several markers were matched to a single scaffold, that scaffold was tabulated with
370 the marker with the lowest position. To visualize if genomic regions showed evidence of
371 selection in the global analysis and the ecotype contrasts, RAD loci were indexed with
372 positional information and their F_{st} values were plotted along the genetic map.

373

374

RESULTS

375 **1. Differentiation Among Ecotypes**

376 *Genetic comparison of ecotypes*

377 Three genetic clusters were strongly supported by the STRUCTURE analysis ($\Delta K = 118$, vs. <
378 53 in $K = 2, 4$ and 5; Figs. 3, S1, S2). Although these clusters were broadly consistent with the
379 three ecotypes (anad-P, lake-P and non-P), levels of admixture varied significantly between
380 them (Fig. 3). The highest degree of genetic distinctiveness was exhibited by the lake-P ecotype
381 ($Q_{\text{lake-P}} = 96.8\%$), which showed low levels of admixture with the other two ecotypes ($Q_{\text{anad-P}}$
382 $= 1.7\%$, $Q_{\text{non-P}} = 1.5\%$). In comparison, the non-P ecotype was almost completely admixed.
383 Genotypes within the non-P ecotype were also assigned at relatively high frequency to both
384 anad-P ($Q_{\text{anad-P}} = 37.7\%$) and lake-P ($Q_{\text{lake-P}} = 15.2\%$) STRUCTURE clusters. However, for
385 anad-P individuals, one of the females and one male also showed high levels of admixture with
386 the non-P type ($Q_{\text{anad-P}} = 64\%, 51\%$; $Q_{\text{non-P}} = 35\%, 41\%$), while all other individuals showed a
387 minimum of 91% match to their own ecotype (Tables S2, S3).

388 Three individuals assigned to the non-P ecotype ('B43 Resi', 'B44 Resi' and 'B45 Resi')
389 were genetically assigned to the lake-P cluster (Fig. 4, Fig. S4). In addition, no individual
390 morphologically assigned to the non-P ecotype based on these prior assumptions was found to
391 have a non-admixed genome (highest individual value, $Q_{\text{non-P}} = 54.1\%$). Finally, all individuals
392 morphologically assigned to the anad-P ecotype were found to fall within the anad-P genetic
393 cluster, and levels of admixture with the other two ecotypes were low (average $Q_{\text{lake-P}} = 2.78\%$,
394 average $Q_{\text{non-P}} = 8.74\%$). Estimates of heterozygosity corroborated these observed patterns,
395 with the non-P ecotype exhibiting significantly higher values compared with the other two
396 ecotypes (Tukey's HSD, $P < 0.0001$, Fig. S2).

397 Consistent with the STRUCTURE analysis, the PCoA returned three separate clusters,
398 consistent with the three ecotypes (Fig. 4b). The anad-P and lake-P ecotypes were clearly
399 differentiated on the axis that explained the majority of genetic variation (PC1, 33.5%).
400 Although the non-P ecotype was not clearly differentiated on PC1, having an intermediate
401 position between the other two, the second axis clearly separated the non-P ecotype from the
402 others (PC2, 17.9%). However, in accordance with STRUCTURE analysis, the same three
403 individuals (all males) previously mentioned assigned to the non-P ecotype clustered with
404 individuals belonging to the lake-P ecotype. Notably, all three of these individuals spawned
405 with females of all three ecotypes but showed a much higher number of attempted matings
406 with lake-P females than non-P or anad-P (Table S2). Euclidean distances derived from the
407 PCoA were smallest between the non-P and lake-P ecotype, and greatest between the lake-P
408 and anad-P ecotypes.

409 Pairwise population differentiation estimates (F_{st}) indicated that the lake-P ecotype was
410 highly differentiated from the anad-P ecotype ($F_{st} = 0.154$), whereas the non-P ecotype was
411 less differentiated from both anad-P ($F_{st} = 0.062$) and lake-P ecotypes ($F_{st} = 0.049$). These data
412 are consistent with the population genetic analysis that revealed high levels of admixture in the
413 non-P ecotype.

414

415 ***Behavioural comparison of ecotypes***

416 Average body length of anad-P individuals was 296 mm (range = 258-327 mm), for lake-P,
417 221 mm (range = 193-283 mm), and for non-P 146 mm (range = 129-164 mm), with males
418 being smaller than females for the lake-P (214 mm vs 228 mm) and anad-P (294 mm vs 299
419 mm) but not non-P (both 145 mm on average) individuals tested, confirming that ecotype was

420 not independent of size difference. However, mean body size differences did not preclude
421 between-type mating attempts. Overall, spawning activity by males was recorded on 966
422 occasions from 72 trials ($n = 108$ trials total; Tables 1, S2, S3); all males tested attempted to
423 spawn with females of more than one ecotype and all attempted to spawn with both anad-P and
424 lake-P females (see Totals under Male Mating Attempts in Table 1). In contrast, few males
425 showed reproductive responses to nonparasitic females (5/6 non-P males and a single lake-P
426 male; Table S3). Three of the males phenotypically classified as non-P that spawned the most
427 frequently ('B43', 'B44' and 'B45') were genetically assigned as lake-P in both the PCA and
428 admixture analyses ($Q_{\text{lake-P}} = 94\%$, 88% , and 90% , respectively); they all showed a strong
429 preference for lake-P females but also attempted to mate with the other two ecotypes (Table
430 S4).

431 There was more substantial variation among females in their propensity to release
432 gametes following initiation of reproductive behaviour by males but there was little evidence
433 of assortative mating by ecotype (Table S4). Release of gametes by females occurred in 32
434 pairings: eight anad-P females (stimulated by four anad-P and four lake-P males); 23 lake-P
435 (eight non-P, 11 lake-P and four non-P males); and only one non-P (who was stimulated only
436 by one of her own conspecifics).

437 The qualitative patterns were supported by the GLMM analyses, which took into
438 account the non-independence of individual females and male trios and attempted to
439 disentangle the effects of size difference and ecotypes. When male mating attempts were
440 treated as a binary response variable, the best-fitting model based on likelihood ratio tests
441 included female ecotype, male ecotype and their interaction, but did not include an individual
442 effect of size difference (Table S5). The significant interaction was driven by an increase
443 between lake-P females and non-P males (estimate = 1.4373; $P = 0.0446$) compared to the

444 intercept. Similar conclusions were reached when considering spawning frequency under a
445 negative binomial distribution but there was no effect of male ecotype*size difference ($df = 2$,
446 $\chi^2 = 3.711$, $P = 0.1637$; AIC = 1130.3) and a much larger difference between the effects of
447 female ecotype*size difference ($df = 2$, $\chi^2 = 13.624$, $P = 0.0011$, AIC = 1120.4) and female
448 ecotype*male ecotype ($df = 4$; $\chi^2 = 20.386$, $P = 0.0004$, AIC = 1117.6).

449 For “successful” spawning in terms of female gamete release, since only a single non-
450 P female spawned, statistical comparisons were made only between anad-P and lake-P females.
451 No significant interactions were found but the final model included the individual effects of
452 female ecotype ($df = 1$, $\chi^2 = 5.3177$, $P = 0.021$, AIC = 146.82), male ecotype ($df = 2$, $\chi^2 =$
453 11.609 , $P = 0.003$, AIC = 150.63) and size difference ($df = 1$, $\chi^2 = 5.8003$, $P = 0.016$, AIC =
454 146.34). More lake-P females spawned than anad-P (estimate = 3.636; $P = 0.00355$); both lake-
455 P (estimate = -2.602; $P = 0.01359$) and non-P (estimate = -6.209; $P = 0.00144$) males induced
456 females to spawn less than anad-P males. Increasing size difference decreased female gamete
457 release (estimate = -0.073; $P = 0.0717$).

458 Considering admixture of interacting mating partners on male gamete release
459 (considered as a binomial variable), the GLMM indicated no significant interaction between
460 female_own Q and male_own Q or male_own Q on its own but males attempted to mate
461 significantly more with less admixed females (i.e., with a higher own_Q; LR compared to
462 intercept model: $df = 1$, $\chi^2 = 23.963$, $P = 9.8 \times 10^{-7}$; estimate = 5.847; $P = 4.15 \times 10^{-9}$). This
463 effect was not apparent when excluding nonparasitic females, who were the most admixed.
464 For female gamete release, there was a significant interaction between male and female
465 admixture ($df = 1$, $\chi^2 = 8.3915$, $P = 0.00377$). Considering only parasitic females, there was no
466 significant interaction but both male (LR: $df = 1$, $\chi^2 = 7.0274$, $P = 0.008$; estimate = 1.9884, P

467 = 0.01564) and female (LR: $df = 1$, $\chi^2 = 7.3425$, $P = 0.0067$; estimate = 51.3658, $P = 0.01374$)
468 admixture decreased successful mating in terms of female gamete release. Only a single anad-
469 P female (B03) showed extensive admixture ($Q_{\text{anad-P}} = 64\%$; $Q_{\text{non-P}} = 35\%$) and this individual
470 failed to spawn with any male. None of the other anad-P females showed q values lower than
471 0.91 and the lake-P females were highly “pure” ($q = 0.95-1$). Similarly, only a single anad-P
472 male (B05) was extensively admixed ($Q_{\text{anad-P}} = 51\%$; $Q_{\text{non-P}} = 41\%$); he only induced a single
473 female (lake-P) to release gametes, although he attempted to mate with females from all
474 ecotypes.

475

476 **2. Estimating evolutionary history and admixture between ecotypes**

477 Although there was some inconsistency among the best-fitting models found using different
478 approaches to predict the past history of ecotype evolution, all analyses supported the
479 conclusion that a strictly bifurcating scenario was unlikely. There was less consensus on
480 whether both parasitic forms contributed to introgression into non-P, or only one form.

481 Coalescent simulations using FASTSIMCOAL2 showed that all strictly bifurcating
482 tree-like scenarios, which do not include introgression between ecotypes (i.e. scenarios *i – iii*),
483 were unlikely. Of the simulations allowing admixture, scenario *v* was most likely, suggesting
484 that the anad-P split first from the ancestor of the two freshwater resident ecotypes, with a
485 subsequent split between the lake-P and non-P ecotypes and introgression from anad-P into
486 non-P. The second best model ($\Delta\text{AIC} = 8.4$ compared to scenario *v*, Table S2) was scenario
487 *vii*, which is similar but also includes secondary contact and gene flow from lake-P into non-P
488 after divergence. All other tested models were considerably less likely ($\Delta\text{AIC} > 10$, Table S2).
489 For the DIYABC analyses, posterior probability estimates supported scenario *iv* as the most

490 likely demographic history supported by the data (direct approach: PP = 0.703, CI = 0.576-
491 0.829; logistic approach: PP = 0.682, CI = 0.465-0.885; see Fig. 2; Table S1). This suggests
492 that the non-P ecotype arose after splitting from the anad-P ecotype, followed by substantial
493 introgression with the lake-P ecotype, sometime after it split from the common ancestor with
494 anad-P. In agreement with the coalescent analysis, all strict isolation scenarios were unlikely.
495 However, scenario *i*, implying a common ancestor for the two freshwater resident ecotypes,
496 showed the second highest PP in the logistic approach (PP = 0.318, CI = 0.114-0.535). In the
497 direct approach, the second best model was scenario *vi* suggesting that non-P originated *via*
498 hybridization between anad-P and lake-P (PP = 0.180, CI = 0.074-0.287). Substantial
499 admixture was further corroborated by the TREEMIX analysis, which suggested unidirectional
500 gene flow from the anad-P to the non-P ecotype, also supporting scenario *v*. Three-population
501 f_3 statistics and the *Z*-score also suggested significant levels of introgression from anad-P and
502 lake-P into non-P ($f_3 = -0.0032$; *Z*-score = -5.39), but no significant signals of admixture in the
503 other two ecotypes ($f_3 > 0.0191$; *Z*-score > 19.82).

504

505 **3. Predicting candidate genes associated with adaptive divergence**

506 Consistent with the STRUCTURE analyses, the majority of differentially fixed SNPs ($n = 10$,
507 from 16 SNPs) were identified between anad-P and lake-P ecotypes (Table S6). None of the
508 substitutions in these differentially fixed SNPs were non-synonymous. A global outlier
509 analysis detected 60 out of 5286 loci that had a probability > 50% of being outliers based on
510 comparison between the three ecotypes (Fig. 5). Out of those 60 outliers, four mapped to the
511 lamprey genome and were located in a gene (Table S7). Two of these were involved in
512 cytoskeleton binding and transport, while the other two were involved in RNA binding or
513 apoptosis and viral infection. An outlier analysis contrasting anadromous (anad-P) and resident

514 individuals (lake-P and non-P) detected a total of 13 loci (total $n = 2509$), including eleven of
515 the outlier loci detected in the global analysis plus an additional two outliers. Four of these
516 outliers mapped to three different genes, with two outliers mapping to a different position in
517 the same gene (Table S7). There was one synonymous and one non-synonymous substitution
518 in this gene (CD 109). The three genes are reported to be involved in protein maturation and
519 lipidation, apoptosis and central nervous system development. A third analysis, comparing
520 parasitic (anad-P and lake-P) and non-parasitic (non-P) individuals, identified 22 outliers from
521 a total of 3263 loci, of which four were shared with the outliers detected in the global analysis
522 (Fig. 5). Four of these mapped to a gene, including one that is part of the dystrophin protein
523 family (RAD-ID: 89575). Dystrophin is part of a protein complex connecting the cytoskeleton
524 to the extracellular matrix (Pasternak *et al.* 1995). The other three genes are involved in
525 transmembrane transport, embryo implantation, extracellular matrix organization and DNA
526 replication.

527 When contrasting F_{st} values between anadromous (anad-P) and resident individuals
528 (lake-P and non-P), there was no apparent pattern along the linkage groups and outliers were
529 randomly distributed (Fig. 5). In contrast, there were several loci with higher F_{st} values on
530 linkage group six (LG6) when contrasting parasitic (anad-P and lake-P) and non-parasitic
531 individuals (non-P). A t-test revealed significantly higher F_{st} values in LG6 compared with the
532 rest of the genome ($t = 5.47$, $P < 0.001$).

533

534

DISCUSSION

535 Our study has shown that three highly differentiated ecotypes of lampreys occurring
536 sympatrically in the Loch Lomond catchment exhibit substantial variability in genetic and

537 behavioural differentiation that does not conform to current taxonomic schemes. Similar
538 conclusions have been drawn previously, for the species considered in this study and for others
539 (Yamazaki & Goto 1997, 2000; Kucheryavyy *et al.* 2007, 2016; Artamonova *et al.* 2011;
540 Yamazaki *et al.* 2011; Spice *et al.* 2014; Makhrov & Popov 2015; Vatandoust *et al.* 2015;
541 Rougemont *et al.* 2015, 2016; Mateus *et al.* 2016). However, the inclusion of a freshwater-
542 feeding parasitic ecotype (lake-P) from a post-glacial lake basin in this current study enables
543 new insights into the evolutionary history, possible process of divergence, and taxonomic
544 distinctiveness of alternative life history strategies in lampreys that has not been possible
545 previously. Specifically, we were able to consider genomic differentiation associated with two
546 distinct evolutionary transitions: from anadromy to freshwater-residency and from parasitic to
547 nonparasitic life history strategies.

548 Because freshwater-parasitic lamprey ecotypes are putatively intermediate in life
549 history strategy between marine-parasitic and freshwater nonparasitic, it has been frequently
550 postulated that they represent a necessary transitional stage during the colonization of post-
551 glacial regions (Zanandrea 1959, 1961; Beamish 1985; Morris 1989; Salewski 2003; Hardisty
552 2006). Our data are somewhat consistent with this hypothesis, as demographic reconstructions
553 suggests a common ancestry for both freshwater ecotypes. However, the phenotype of that
554 ancestor – and whether or not it was parasitic – remains uncertain. Consistent with studies
555 conducted in other regions (Rougemont *et al.* 2015, 2017), we found that the nonparasitic
556 freshwater ecotype was genetically introgressed with the other two forms. However,
557 intriguingly, the freshwater parasitic ecotype in our study showed the highest degree of
558 reproductive isolation, despite appearing highly attractive to other ecotypes during spawning
559 and being intermediate in life-history and morphology. In nature, these freshwater parasitic
560 ecotypes of typically anadromous parasitic species occur only rarely (e.g. Maitland 1980;

561 Tsimbalov *et al.* 2015), which could suggest a fitness cost relative to the more commonly found
562 non-parasitic freshwater forms. Alternatively, populations such as the lake-P ecotype of Loch
563 Lomond could represent a life history transition that only evolves under certain geographic
564 contexts (e.g. allopatry in large lake basins containing abundant foraging opportunity during
565 glacial advance-retreat, Collett 1905; Tuunainen *et al.* 1980; Morris 1989; Inger *et al.* 2010;
566 Tsimbalov *et al.* 2015). We suggest that the formation of lamprey ecotypes in Loch Lomond,
567 and the European river and brook lamprey “species pair” more generally, are dependent on an
568 interplay between ecological opportunity and introgression.

569

570 ***Differentiation and introgression of lamprey ecotypes***

571 Genetic variability, based on RAD sequencing, is not consistent with currently recognized
572 taxonomic units. The lake-P ecotype, currently considered *L. fluviatilis*, was found to be more
573 highly genetically distinct from the anad-P ecotype (also considered *L. fluviatilis*) than the non-
574 P ecotype (considered *L. planeri*), which was highly admixed with the other two. Demographic
575 analyses ruled out that the two parasitic ecotypes (anad-P and lake-P) share a more recent
576 common ancestor than with the non-P ecotype (Fig. 3, Table S1). Intriguingly, three males that
577 would have been classified as non-P based on their morphology showed a Q value of less than
578 11% for their own ecotype but greater than 89% for the lake-P ecotype. This could suggest that
579 the nonparasitic phenotype is controlled by a relatively small portion of the genome, which
580 was supported by our finding of a single genomic region strongly associated with the transition
581 to nonfeeding. There was no evidence for admixture in the lake-P individuals but a single male
582 and a single female that were morphologically anad-P showed extensive levels of admixture
583 with the non-P ecotype. Additionally, the lake-P form appears to contain a subset of mtDNA
584 haplotypes found in the non-P population from Loch Lomond (Table S8; data from Hume

585 2013). Only two mtDNA haplotypes were found in the lake-P individuals sampled: one that
586 was found in all Scottish populations sampled and one (which showed a single bp mutation
587 from the common haplotype) that was shared with non-P but was not found among the anad-P
588 individuals sampled. Although based on only a small sample size, both anad-P and non-P from
589 the same catchment showed more mtDNA variation than the lake-P ecotype, suggesting that
590 the latter could have experienced a more substantial population bottleneck in the relatively
591 recent past or less historical introgression relative to the other two ecotypes.

592 Across Europe, anad-P (*L. fluviatilis*) and non-P (*L. planeri*) lampreys appear to have
593 resulted from multiple and parallel speciation events (Mateus *et al.* 2016; Rougemont *et al.*
594 2016). Based on mtDNA and nuclear gene markers, no species-specific fixed differences have
595 been found (Schreiber & Engelhorn 1998; Espanhol *et al.* 2007; Pereira *et al.* 2011) and neither
596 do they exhibit consistent microsatellite allele frequency differences across their range
597 (Bracken *et al.* 2015; Rougemont *et al.* 2015; Mateus *et al.* 2016). A detailed comparison
598 among sympatric and parapatric populations of the two “species” in France using both
599 microsatellite genotypes (Rougemont *et al.* 2015) and genome-wide, single nucleotide
600 polymorphisms based on RAD sequencing (Rougemont *et al.* 2016) provides little evidence
601 for a restriction of gene flow between them. Secondary contact and introgression between
602 lamprey ecotypes and demographic models with asymmetric gene flow from the anad-P into
603 the non-P were also evident from these locations (Rougemont *et al.* 2017), consistent with our
604 data from Loch Lomond. Hybridization and introgression are increasingly recognized as
605 evolutionary forces creating diversity and possibly promoting the evolution of differentiated
606 ecotypes (e.g. charr, Garduño-Paz *et al.* 2012; Jacobs *et al.* 2017; crater lake cichlids, Martin
607 *et al.* 2015; whitefish, Rougeux *et al.* 2017). For example, secondary contact and introgression
608 between evolutionary lineages of post-glacial fishes seems to frequently result in ecological

609 divergence and the formation of new ecotypes (Jacobs *et al.* 2017; Rougeux *et al.* 2017).
610 Overall, phylogeography indicates that gene flow between anad-P (*L. fluviatilis*) and non-P
611 ecotypes (*L. planeri*) is opportunistic, and introgression likely varies in accordance with
612 contemporary and historic barriers (physical or environmental) separating these migratory and
613 resident ecotypes (Bracken *et al.* 2015; Mateus *et al.* 2016).

614 Given this high potential for gene flow and sensitivity to ecological disturbance, most
615 ecotypes formed in sympatry are thus unlikely to attain the status of “good biological species”
616 (Mallet 2008; Hendry *et al.* 2009; Weissing *et al.* 2011; Lowry 2012; Østman *et al.* 2014;
617 Rudman & Schluter 2016). However, revealing those circumstances that do lead to
618 reproductive isolation is critical to understanding speciation. In our study, despite clear genetic
619 differentiation of the lake-P ecotype from the other two, there was little evidence of pre-zygotic
620 isolation among them, due particularly to the high attractiveness of lake-P females to all males
621 and their willingness to mate with any of the three ecotypes tested. Thus, there does indeed
622 appear to be the potential for gene flow among ecotypes if they were to spawn in syntopic
623 habitats. The inclusion of an intermediate sized lake-P ecotype enables more fine-scale testing
624 of the prediction that size-assortative mating is the primary driver of reproductive isolation
625 between diverging lamprey ecotypes (Beamish & Neville 1992). We found evidence for an
626 interaction between body size difference and female ecotype, which explained nearly as much
627 variation as the interaction between male and female ecotypes for male mating behaviours –
628 but, there was no separate effect of body size when the latter interaction was accounted for. In
629 our assays, the interaction was driven largely by reduced mating between small non-P and large
630 anad-P individuals; anad-P females did not prefer their own conspecifics over lake-P males
631 and anad-P males attempted to mate with females from all ecotypes. Thus, while size is

632 certainly a factor in lamprey mate choice, the relative size differences between ecotypes does
633 not preclude the potential for gene flow.

634

635 ***Factors influencing ecological speciation***

636 The patterns of introgression among lamprey ecotypes in this study invokes three non-mutually
637 exclusive hypotheses generally explaining ecological speciation in sympatry: *i*) temporal or
638 spatial isolation restricts gene flow between ecotypes under current natural conditions; *ii*)
639 frequency-dependent encounter rates influence mate choice between ecotypes; and *iii*) hybrids
640 between ecotypes suffer high fitness costs as a consequence of intrinsic or extrinsic post-
641 zygotic barriers. Our results are consistent with a combination of these factors influencing the
642 parallel differentiation of ecotypes in lampreys, which could be more broadly representative of
643 post-glacial expansion of freshwater fishes.

644 Firstly, the importance of geographical isolation to the acquisition of adaptive
645 differences between populations via divergent selection has been recognized since Darwin's
646 time (Mallet 2008). The role of physical separation of lamprey populations in facilitating
647 genetic divergence has been highlighted previously, where migratory ecotypes become
648 separated from stream-resident forms as they pursue foraging opportunities in large river
649 systems, lakes, or marine environments (e.g. due to geological or climate shifts, construction
650 of in-stream barriers, Zanandrea 1959; Hardisty & Potter 1971; Kan & Bond 1981; Yamazaki
651 *et al.* 2011; Bracken *et al.* 2015). While some intriguing patterns of parallel speciation and
652 adaptation have been revealed previously by genetic analysis, suggesting some degree of
653 isolation between lamprey ecotypes (Mateus *et al.* 2013, 2016; Bracken *et al.* 2015; Rougemont
654 *et al.* 2015; 2016), there is strong evidence that lampreys can continue to exhibit multiple

655 ecotypes yet exist in true sympatry (Beamish 1987; Yamazaki *et al.* 1998; Kucheryavyy *et al.*
656 2007; this study). Post-glacial fishes, particularly Arctic charr, exemplify this “problem”
657 (Klemetson 2010), with evidence of ongoing gene flow between charr ecotypes apparent in
658 some regions (e.g. Salisbury *et al.* 2017). However, site fidelity and associated assortative
659 mating among ecotypes of charr is required to maintain discrete forms in sympatry (Jonsson &
660 Jonsson 2001). Observations in nature have provided evidence against spatial or temporal
661 segregation of many lamprey ecotypes on spawning grounds (Morman 1979; Manion &
662 Hanson 1980; Kucheryavyy *et al.* 2007; Cochran *et al.* 2008; Lasne *et al.* 2010). Furthermore,
663 sneak male mating tactics among ecotypes and between species have also been documented
664 (Cochran 2008; Hume *et al.* 2013a). Thus, strict reproductive isolation is an unsatisfactory
665 explanation for the maintenance of discrete lamprey ecotypes across entire geographic ranges.
666 However, recent work suggests that in fact more variable and larger ecosystems can support
667 more variable populations or multiple ecotypes as a consequence of greater ecological
668 opportunity (Mahler *et al.* 2013; Præbel *et al.* 2013; Wellborn & Langerhans 2014; Recknagel
669 *et al.* 2017). This suggests that the Loch Lomond system, a large and deep lake itself, is an
670 important but uncommon ecological opportunity driving the formation of the two freshwater
671 parasitic lamprey ecotypes in Europe.

672 Secondly, sexually reproducing organisms can benefit from selecting between multiple
673 potential mates; but being too choosy can incur high fitness costs in terms of potential loss of
674 reproductive opportunity, energetic costs and high mortality (Andersson 1994). One important
675 factor influencing mate choice is encounter rate, where low rates increase the risks of not
676 mating and often results in reduced choosiness (Jennions & Petrie 1997). Based on sampling
677 the Loch Lomond catchment during the lamprey spawning migration, the anad-P ecotype is
678 rare, the non-P ecotype the commonest, and the lake-P ecotype somewhat intermediate (Morris

679 & Maitland 1987; Morris 1989; Adams *et al.* 2008; Hume 2011, 2013). These relative
680 frequencies are consistent with the relative rarity of freshwater parasitic populations of
681 lampreys compared to non-feeding forms in northern lakes in general (e.g. Maitland 1980;
682 Tsimbalov *et al.* 2015). This suggests that continuing to feed in freshwater represents a strategy
683 that only evolves in response to particular ecological opportunity (Schluter 2000; Siwertsson
684 *et al.* 2010; Lowry 2012; Recknagel *et al.* 2017); principally, in river basins with large lakes
685 and abundant foraging opportunity (Collett 1905; Tuunainen *et al.* 1980; Morris 1989; Inger
686 *et al.* 2010; Tsimbalov *et al.* 2015). There could also be a fitness disadvantage relative to
687 nonparasitic forms in having to exploit new food resources (i.e. freshwater fishes) in a finite
688 environment. If we assume that weak assortative mating occurs naturally in this system (i.e.
689 not an artifact of behavioural assays in this study) then the likelihood of anad-P and lake-P
690 ecotypes mating with the more common non-P ecotype seems high. Thus, if mate choice
691 operates by lampreys only choosing the "correct" ecotype when encounter rates with them are
692 frequent, then logically spawning with the other two should occur when encounter rates with
693 their own kind are infrequent. This is consistent with Rougemont *et al.* (2017) who found
694 higher effective population sizes for anad-P compared to non-P forms, as well as asymmetric
695 introgression between ecotypes. Hybridization has been associated with a scarcity of
696 conspecifics (Mayr 1963, 1970; Avise & Saunders 1984; Leaniz & Verspoor 1989; McGowan
697 & Davidson 1992; Wirtz 1999; Randler 2002; Quilodr an *et al.* 2014). However, the influence
698 of encounter rate between ecotypes on mate choice has rarely been experimentally tested
699 (Willis *et al.* 2011). Our results are consistent with relative frequency of the three ecotypes
700 during the spawning season explaining the patterns of introgression observed.

701 Thirdly, intrinsic post-zygotic barriers such as hybrid inviability, or extrinsic barriers
702 such as maladaptation of hybrids to parental environments, may also contribute to reproductive

703 isolation between ecotypes (Mallet 2008; Hendry *et al.* 2009; Lowry 2012). Loss of genetic
704 compatibility during adaptive divergence is a principal cause of reproductive isolation (Coyne
705 & Orr 2004). When alleles with beneficial effects in one population are recombined in a hybrid
706 genome with alleles originating at different loci in a second population, hybrid inviability can
707 occur (Burke & Arnold 2001; Presgraves *et al.* 2003). The lack of introgression in the lake-P
708 lamprey ecotype and low admixture in the anad-P individuals in this study could be the result
709 of high mortality of hybrids that adopt a migratory-parasitic life history strategy.
710 Metamorphosis in lampreys is a highly complex, protracted event requiring drastic changes in
711 physiology and anatomy (Manzon *et al.* 2015), and conceivably is a source of substantial
712 mortality of parasitic lampreys. Although there is evidence for high survivorship of hybrid
713 embryos between lamprey ecotypes in the laboratory (Enequist 1937; Piavis *et al.* 1970;
714 Beamish & Neville 1992; Hume *et al.* 2013b; Rougemont *et al.* 2015), the viability of adults,
715 as well as F1 or F2 hybrids remains untested. The low frequency of hybrids between lampreys
716 in the wild has been suggested to be a consequence of selection against those hybrids
717 (Rougemont *et al.* 2017). It is predicted that where incipient speciation is occurring selection
718 against hybrids should also have an ecological component (Schluter 1996); where hybrids
719 experience reduced foraging efficiency (e.g. Hatfield & Schluter 1999) and/or suffer higher
720 rates of mortality (e.g. Nilsson *et al.*, 2017). Therefore, offspring resulting from inter-ecotype
721 matings between lampreys in Loch Lomond may suffer higher fitness costs when they adopt
722 migratory-parasitic strategies, compared to the relatively sedentary and cryptic life cycle of
723 nonparasitic lampreys.

724

725 *Adaptive divergence of lamprey ecotypes*

726 Despite evidence for gene flow between parasitic and nonparasitic ecotypes, there is still
727 substantial neutral and adaptive genetic differentiation between them. Neutral differentiation
728 may reflect the limited dispersal capability of lampreys confined to freshwater systems (e.g.
729 Taylor *et al.* 2012), as anadromous lampreys consistently exhibit high levels of gene flow
730 among populations across a variety of spatial scales (Spice *et al.* 2012; Hess *et al.* 2013;
731 Yamazaki *et al.* 2014; Bracken *et al.* 2015). The identities of transcribed genes that are
732 differentially fixed, or are genomic outliers, in anadromous-parasitic vs resident-parasitic and
733 resident-nonparasitic ecotypes are potentially revealing in regards to adaptive differences in
734 life history strategy between them.

735 In our study, we found candidate genes associated with different important
736 physiological functions that could differ in relation to a transition between feeding
737 environments. For example, the gene *cd109* may influence growth rate, as it interacts
738 selectively with transforming growth factor- β , a multifunctional peptide that controls
739 proliferation, differentiation and other functions in many cell types (e.g. Hockla *et al.* 2010).
740 Another important factor in the transition between anadromous and resident life history
741 strategies is the immune response (Mateus *et al.* 2013). Communities of pathogens in marine
742 and freshwater environments differ substantially, therefore resident and anadromous lampreys
743 will experience different selection pressures for pathogen resistance. We identified the gene
744 *nckap-1* as a genomic outlier between anadromous and both resident ecotypes. This gene has
745 been previously identified as crucial in resistance against a bacterial pathogen in channel
746 catfish *Ictalurus punctatus* (Zhou *et al.* 2017). This emphasizes the power of outlier analyses,
747 even using genome sampling methods like RAD sequencing in revealing potential genomic
748 regions involved in functionally important traits for adapting to new environments. We do not
749 imply that any of the SNPs identified here are causal mutations for adaptive phenotypes, but

750 encourage that future studies investigate genes and genomic regions associated with adaptive
751 divergence identified here using more fine-scale genomic data.

752 Outlier analyses also revealed potential differences related to the transition to a
753 nonparasitic life cycle. A single region of the genome (linkage group 6) revealed significant
754 differences between parasitic ecotypes and the nonparasitic form and these may be consistent
755 with directional selection in lamprey feeding mode despite high levels of gene flow (e.g. Nosil
756 & Crespi 2004). The linkage of genes in so-called genomic islands may facilitate the
757 establishment of reproductive isolation between ecotypes and eventually lead to speciation
758 (Feder & Nosil 2010). The *reck* gene is associated with linkage group 6 and is implicated in
759 the organization of extracellular matrix and cell migration. Knockdown experiments of this
760 gene in zebrafish result in impaired vascular integrity and lack of dorsal root ganglia
761 formations, suggesting that *reck* is important during embryonic development (Prendergast *et*
762 *al.* 2012). This gene is also suggested to have a crucial role during embryonic development in
763 other vertebrates (de Oliveira *et al.* 2010; Yamamoto *et al.* 2012). Two additional genes,
764 *scn4aa* and an orthologue of *rev3l*, were detected as genomic outliers. The former gene codes
765 for a sodium-channel protein and is involved in development of the heart, while the latter is a
766 DNA polymerase also known to function in embryonic development. In mice, gene mutations
767 of *rev3l* resulted in smaller neonates with retarded development (Wittschieben *et al.* 2000).
768 Although speculative at present, genes such as these may be vital in lamprey embryological
769 development, and alterations of their expression could impact metamorphosis and potentially
770 be involved in the transition from a parasitic to nonparasitic life history strategy.

771

772 ***Conclusions***

773 Our data suggest that freshwater-resident feeding forms of the typically anadromous *Lampetra*
774 *fluviatilis* evolve and persist only in large post-glacial lake basins throughout Europe. In Loch
775 Lomond, this ecotype (lake-P) is genetically distinct from the anadromous form (anad-P),
776 whereas a nonparasitic ecotype (non-P here; often referred to as *L. planeri*) was introgressed
777 with the other two. Although we were unable to fully resolve the question of whether lake-P
778 ecotypes are a necessary transitional step in the evolution of nonparasitism among lampreys
779 more generally, we found both lake-P and non-P ecotypes in Loch Lomond do share a common
780 freshwater ancestor. This suggests ecological opportunity, followed by variation in the
781 probability of introgression, strongly influences the extent of divergence between sympatric
782 lamprey ecotypes.

783

784

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789

790

DATA ACCESSIBILITY

791 Data can be accessed using BioProject ID PRJNA488599.

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1123

1124 Figure 1. Photograph of three lamprey ecotypes collected in the Loch Lomond basin and used
1125 in genetic and behavioural comparisons. (A) is nonparasitic (non-P), (B) is freshwater parasitic
1126 (lake-P), and (C) is anadromous parasitic (anad-P).

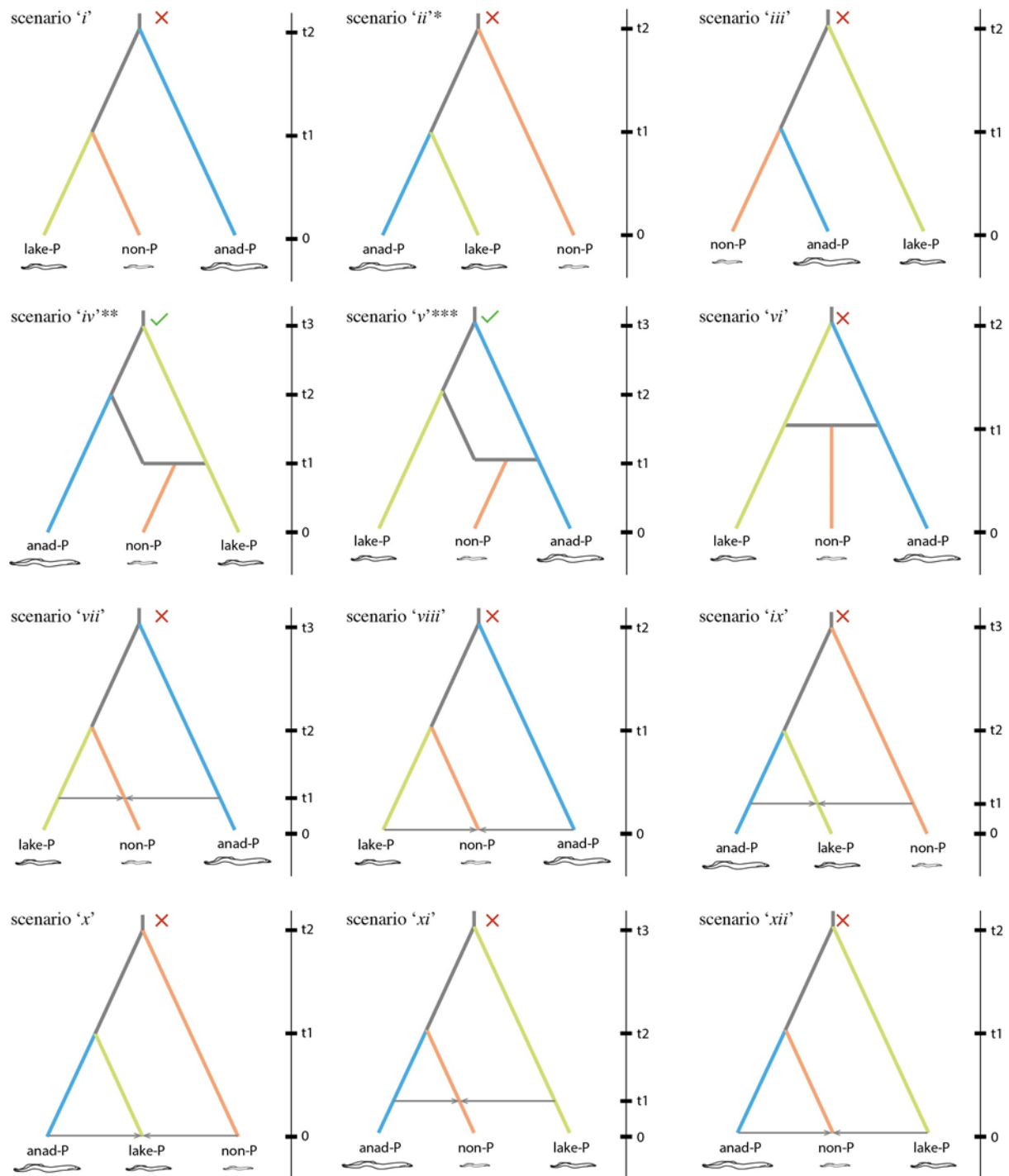
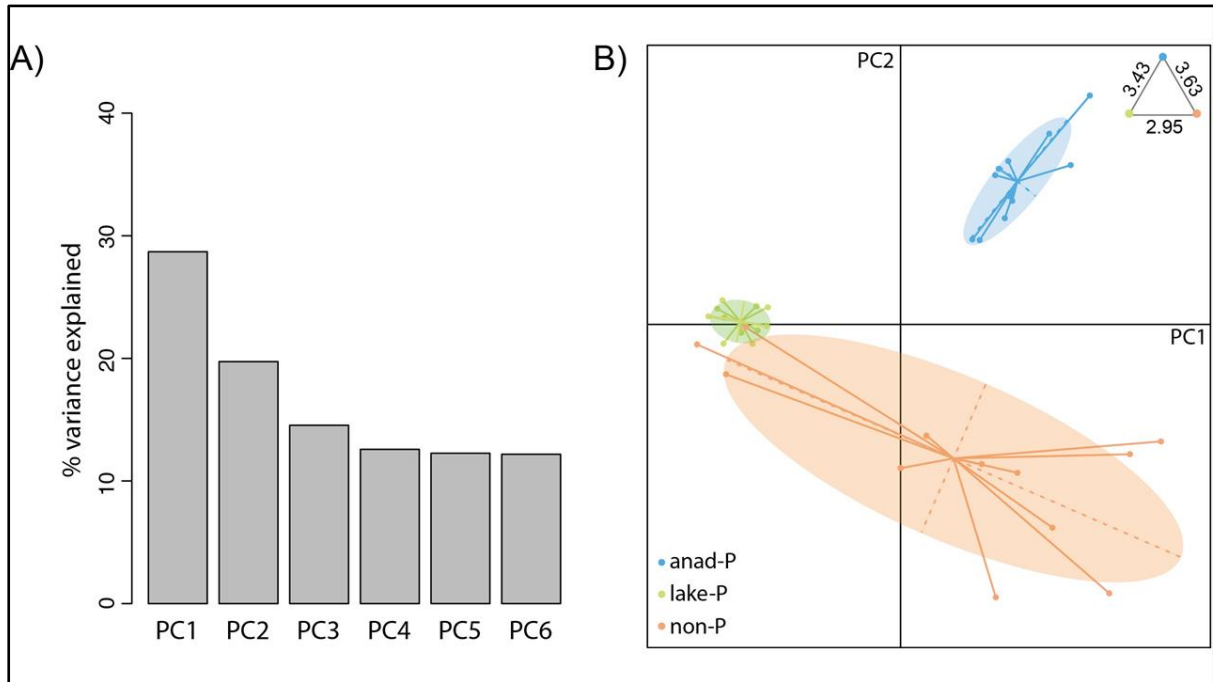


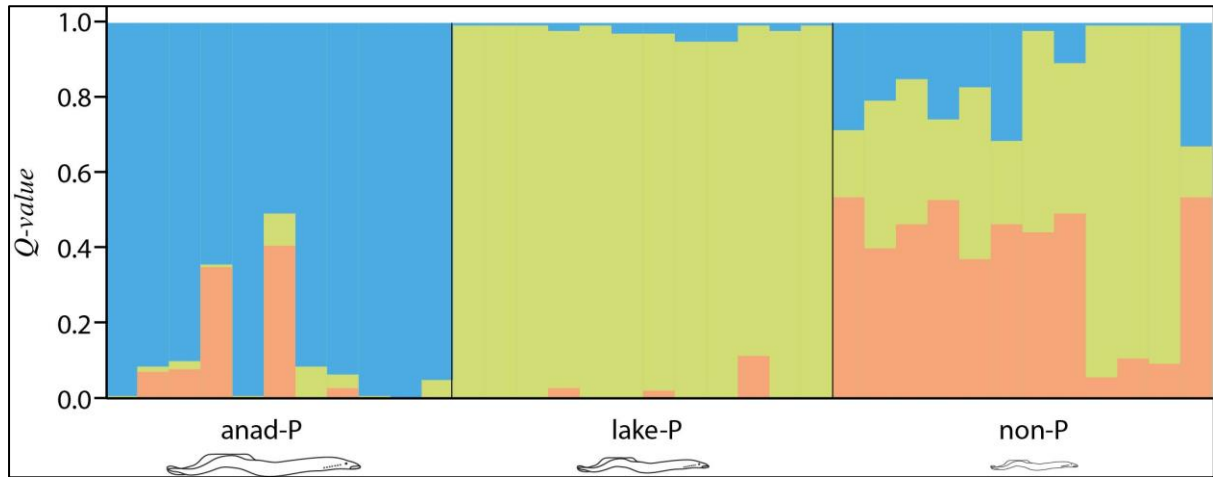
Figure 2. Hypothetical evolutionary scenarios for lamprey ecotypes tested using demographic reconstruction analyses. Scenarios *i - iii* are strictly bifurcating trees with no gene flow between ecotypes. Scenarios *iv* and *v* require an initial splitting event followed by a second split of the non-P ecotype and subsequent introgression into non-P from either parasitic form. Scenario *vi* suggests hybrid speciation between lake-P and anad-P, giving rise to the non-P ecotype. Scenarios *vii - xii* differ in the evolutionary relationships between ecotypes and include gene flow at different time points from both parasitic ecotypes into non-P. Scenarios with uneven number (*vii, ix, xi*) are evolutionary models with recent ongoing gene flow and those with even numbers (*viii, x, xii*) past gene flow from the parasitic into the non-parasitic ecotype. Scenarios with check mark were supported by analyses, and scenarios with an x-mark were not supported

in any analysis. * consistent with current taxonomy; ** supported by DIYABC analysis; *** supported by FASTSICMOAL2 and TREEMIX.



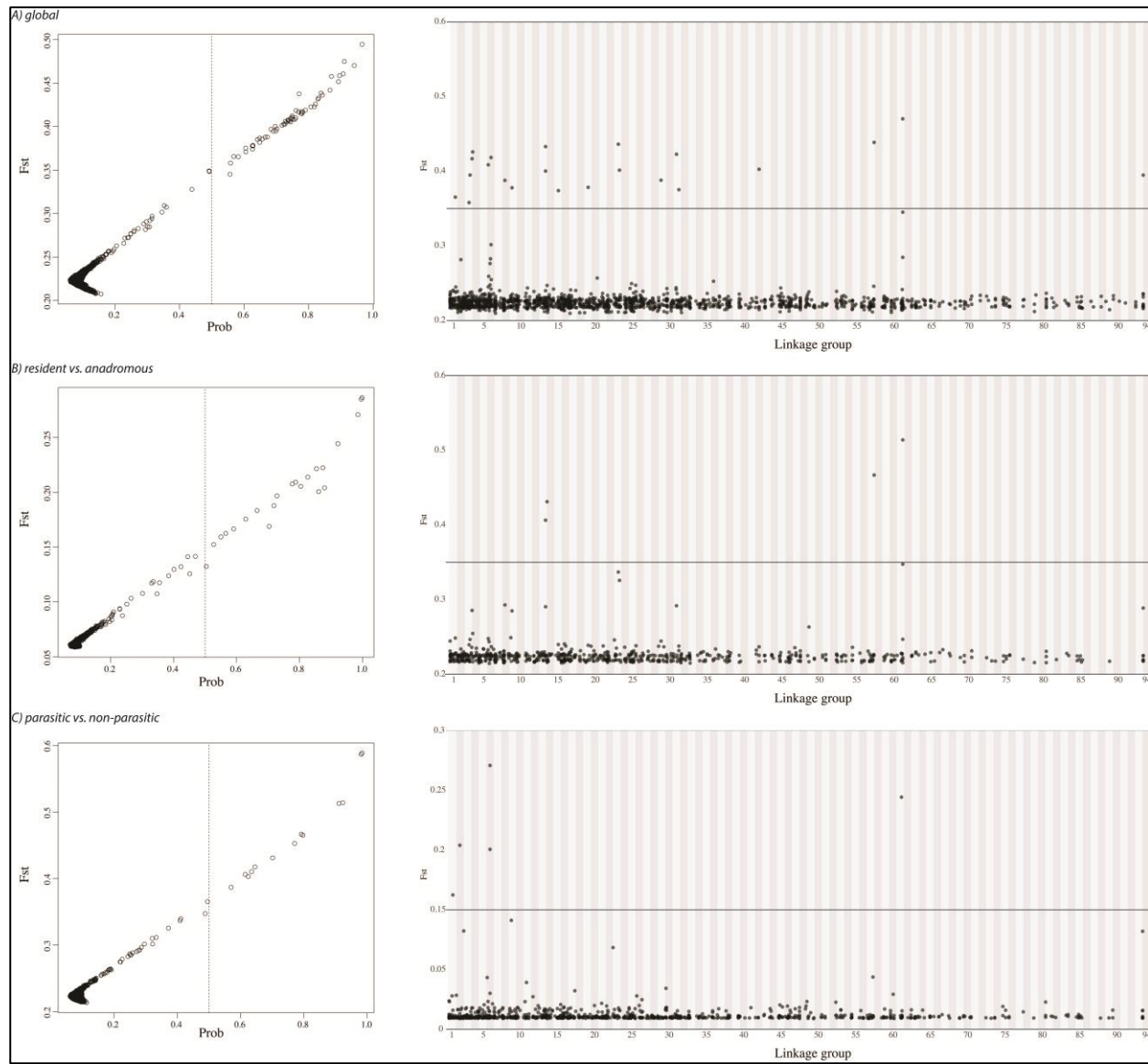
1127

1128 Figure 3. Multivariate analyses illustrating genetic differentiation between lamprey ecotypes.
 1129 Analyses were performed with the same 35 individuals. Panel A shows relative percentages of
 1130 variance explained by the first six principal components. The first two principal components
 1131 accounting for 48.44% of the total variance are shown in Panel B (anad-P = blue, lake-P =
 1132 green, non-P = orange), with the scores for each individual of the different ecotypes. Ellipses
 1133 drawn are confidence intervals including $\frac{2}{3}$ of the samples within each ecotype. Average
 1134 Euclidean distance between ecotypes are shown in the upper right corner of PCA plots.



1135

1136 Figure 4. Output from a STRUCTURE analysis using $K = 3$. See Figure S3 for comparisons
 1137 with $K = 4$ and 5. Each bar represents an individual and the relative proportion of its genotype
 1138 belonging to each of the three genetic clusters. The analysis was performed excluding
 1139 differentially fixed SNPs and outlier loci and includes a total of 7, 678 loci.



1141 Figure 5. Plot of F_{st} outliers and genome scan of all identified loci that matched to the indexed
1142 sea lamprey (*Petromyzon marinus*) genome. On the left, F_{st} values obtained from BAYESCAN
1143 outlier analyses from (A) global across all ecotypes; (B) anadromous *vs.* resident individuals;
1144 and (C) parasitic *vs.* nonparasitic individuals are displayed. Loci that fall right of the vertical
1145 lines have a probability greater than 50% of being an outlier. On the right, the position of each
1146 locus along the sea lamprey linkage groups (x-axis; values normalized to a fraction of 1 for
1147 each linkage group) and the respective F_{st} value (y-axis) are shown. Horizontal lines represent
1148 the probability threshold of 50% of being an outlier.

1149 Table 1. Proportion of mating events occurring within and between ecotypes. Male mating attempts occurred in 142/324 potential pairings,
 1150 whereas females only released gametes in 32. Shown are the proportion of pairings where males attempted to mate at least once (male mating
 1151 attempts), the frequency of male reproductive behaviours (male spawning frequency), and the pairings where females were stimulated to release
 1152 gametes (female gamete release). Anad-P = anadromous, parasitic; Lake-P = lake-resident parasitic; Non-P = lake-resident nonparasitic. The
 1153 “Pro. Males” columns show the proportion of males of each ecotype that attempted to mate, spawned, or stimulated female gamete release.

Female Ecotype	Male Mating Attempts				Male Spawning Frequency				Female Gamete Release			
	Anad-P	Lake-P	Non-P	Pro. Males	Anad-P	Lake-P	Non-P	Pro. Males	Anad-P	Lake-P	Non-P	Pro. Males
Anad-P	0.51	0.53	0.30	0.45	0.41	0.42	0.09	0.31	0.27	0.33	0.00	0.25
Lake-P	0.49	0.45	0.55	0.49	0.59	0.56	0.85	0.67	0.73	0.67	0.80	0.72
Non-P	0.00	0.02	0.16	0.06	0.00	0.02	0.06	0.03	0.00	0.00	0.20	0.03

1154