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Nucleotide variation and balancing selection at the *Ckma* gene in Atlantic cod: Analysis with multiple merger coalescent models

A high-fecundity organisms, such as Atlantic cod, can withstand substantial natural selection and can at any time simultaneously replace alleles at a number of loci due to their excess reproductive capacity. High-fecundity organisms may reproduce by sweepstakes leading to highly skewed heavy-tailed offspring distribution. Under such reproduction the Kingman coalescent of binary mergers breaks down and models of multiple merger coalescent are more appropriate. Here we study nucleotide variation at the *Ckma* (Creatine Kinase Muscle type A) gene in Atlantic cod. The gene shows extreme differentiation between the North (Canada, Greenland, Iceland, Norway, Barents Sea) and the South (Faroe Islands, North-, Baltic-, Celtic-, and Irish Seas) with a between regions $F_{ST} > 0.8$ whereas neutral loci show no differentiation. This is evidence for natural selection. The protein sequence is conserved by purifying selection whereas silent and non-coding sites show extreme differentiation. Relative to outgroup the site-frequency spectrum has three modes, a mode at singleton sites and two high frequency modes at opposite frequencies representing divergent branches of the gene genealogy that is evidence for balancing selection. Analysis with multiple-merger coalescent models can account for the high frequency of singleton sites and indicate reproductive sweepstakes. Coalescent time scales with population size and with the inverse of variance in offspring number. Parameter estimates using multiple-merger coalescent models show fast time-scales. Time-scales of mitochondrial DNA are about square root of the effective population size and time-scales of nuclear genes are much faster.

1 Nucleotide Variation and Balancing 2 Selection at the *Ckma* gene in Atlantic cod: 3 Analysis with multiple merger coalescent 4 models

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8 ABSTRACT

A high-fecundity organisms, such as Atlantic cod, can withstand substantial natural selection and can at any time simultaneously replace alleles at a number of loci due to their excess reproductive capacity. High-fecundity organisms may reproduce by sweepstakes leading to highly skewed heavy-tailed offspring distribution. Under such reproduction the Kingman coalescent of binary mergers breaks down and models of multiple merger coalescent are more appropriate. Here we study nucleotide variation at the *Ckma* (Creatine Kinase Muscle type A) gene in Atlantic cod. The gene shows extreme differentiation between the North (Canada, Greenland, Iceland, Norway, Barents Sea) and the South (Faroe Islands, North-, Baltic-, Celtic-, and Irish Seas) with a between regions $F_{ST} > 0.8$ whereas neutral loci show no differentiation. This is evidence for natural selection. The protein sequence is conserved by purifying selection whereas silent and non-coding sites show extreme differentiation. Relative to outgroup the site-frequency spectrum has three modes, a mode at singleton sites and two high frequency modes at opposite frequencies representing divergent branches of the gene genealogy that is evidence for balancing selection. Analysis with multiple-merger coalescent models can account for the high frequency of singleton sites and indicate reproductive sweepstakes. Coalescent time scales with population size and with the inverse of variance in offspring number. Parameter estimates using multiple-merger coalescent models show fast time-scales. Time-scales of mitochondrial DNA are about square root of the effective population size and time-scales of nuclear genes are much faster.

9
10 Keywords: Balancing Selection, *Ckma*, Atlantic cod, multiple-merger coalescent, time scales

11 INTRODUCTION

12 High fecundity translates into large excess reproductive capacity that would allow
13 organisms to withstand substantial natural selection enabling them to bear the entailing
14 high genetic load. Based on the concept of the cost of natural selection (Haldane,
15 1957) high-fecundity organisms relative to low-fecundity organisms should at any time
16 be able to adapt a larger proportion of their genome to meet various environmental
17 challenges. Trying to explain the paradox of sexual reproduction Williams (1975) in his

Sex and Evolution book argues that high-fecundity coupled with type III survivorship of heavy mortality of young may be able to pay the 50% fitness cost of meiosis. He developed several models, such as the Elm/Oyster and the Cod/Starfish models, which emphasize the importance of high-fecundity for selection. Williams also discussed the concept of reproductive sweepstakes. There is no heritability of fitness and sexual reproduction continuously assembles Sisyphean genotypes. The distribution of offspring numbers is highly skewed, heavy-tailed and with high variance (lognormal). That is Williams's fitness distribution. The environment factors are envisioned to act in a sequence of selective filters. With only a few factors (e.g. temperature, salinity, etc) there nevertheless can be an enormous number of different sequences of selective filters (environments) that do not recur. Hence a winning genotype is not permanent and must be continuously reassembled. Natural selection increases the variance in offspring number and thereby reduces effective population size genome-wide. Neutral variation will therefore drift faster under pervasive natural selection.

Coalescent theory (Kingman, 1982a,b) traces the genealogy of a sample and is very useful for making inference of molecular data. However, in an extreme case under a winner-take-all sweepstakes reproduction all samples would coalesce immediately in the previous generation (Árnason, 2004) and there would be no variation. The Kingman coalescent, which is derived from (Wright/Fisher) models of low fecundity non-skewed offspring distributions, assumes a bifurcating genealogy and is not appropriate for reproduction of this kind (Eldon and Wakeley, 2006; Schweinsberg, 2003; Wakeley, 2013). Some organisms may exhibit both high fecundity and highly skewed offspring distributions. For these organisms the Λ coalescent allowing multiple mergers of ancestral lineages (Pitman, 1999; Sagitov, 1999; Donnelly and Kurtz, 1999; Eldon and Wakeley, 2006; Schweinsberg, 2003; Sargsyan and Wakeley, 2008) or Ξ coalescent allowing simultaneous multiple mergers of ancestral lineages (Schweinsberg, 2000; Möhle and Sagitov, 2001) may be more appropriate. Wakeley (2013) gives an overview of the development of coalescent theory in new directions. There is also active development of statistical inference methods associated with multiple merger coalescents (e.g. Birkner et al., 2013b, 2014). Studies on the high fecundity organisms Pacific oyster *Crassostrea gigas* (Hedgecock and Pudovkin, 2011) and Atlantic cod *Gadus morhua* (Árnason and Pálsson, 1996; Árnason et al., 1998, 2000; Carr and Marshall, 1991a; Carr et al., 1995; Pepin and Carr, 1993; Árnason, 2004) have provided data for a number of tests of some of the new coalescent models (Eldon and Wakeley, 2006; Eldon, 2011; Eldon and Degnan, 2012; Steinrücken et al., 2013; Birkner et al., 2013b). Atlantic cod thus provides a model for studies of multiple merger coalescent. In this paper we apply some of these new methods for Λ coalescents in a study of balancing selection at a gene showing extreme spatial differentiation in Atlantic cod.

A dense genomic map of genetic variation in humans (and in model organisms) allows scanning the genome for signatures of natural selection (Voight et al., 2006; Sabeti et al., 2007; Storz, 2005). Asking what percentage of the human genome shows footprints of selection depends on the density of the maps and sensitivity of the various methods used (Voight et al., 2006; Sabeti et al., 2007; Storz, 2005). It is safe to say that only a small percentage of single nucleotide polymorphisms (SNPs) show footprints of selection in the low fecundity humans (Akey, 2009; Pickrell et al., 2009). For microsatellite loci 2% (13/624) were detected as outliers when African and non-African

human populations were compared (Storz et al., 2004). In contrast, comparable genome level studies in Atlantic cod find that 11% (26 out of 235) of independent SNPs (Moen et al., 2008) are F_{ST} outliers (by method of Beaumont and Nichols, 1996) and 4% SNPs (70 out of 1641 Bradbury et al., 2010) are Bayscan outliers (by method of Foll and Gaggiotti, 2008) likely undergoing selection. Similarly one fourth of microsatellite loci in Atlantic cod (Nielsen et al., 2006) are F_{ST} outliers. This supports our thesis that a considerable fraction of the Atlantic cod genome may be simultaneously under selection for different adaptations.

More than half of the 70 outliers in Bradbury et al. (2010) study of Atlantic cod show adaptive parallel clines related to temperature on both the western and eastern side of the Atlantic Ocean. They show that multiple genes, located in three independent linkage groups, are involved. There are single genes as well as blocks of genes in “genomic islands” (Bradbury et al., 2013; Hemmer-Hansen et al., 2013). Some of the genes or blocks of genes show clear spatial patterns while other genes show complex spatio-temporal patterns in contrast to no differentiation of non-outlier (neutral) loci (Poulsen et al., 2011; Therkildsen et al., 2013). For example a locality in West Greenland shows great similarity to coastal areas in Iceland, implying either parallel adaptation on a fine scale or patterns of gene flow that are hard to reconcile with geographic distance. Another study (Hemmer-Hansen et al., 2014) adds even more complexity of population structure at outlier loci with little or no difference at non-outlier neutral loci.

The Moen et al. (2008) study of differentiation among four Atlantic cod populations along the coast of Norway showed no differentiation among presumably neutral non-outliers loci with an average $\bar{F}_{ST} = 0.0012$. In contrast, the outlier loci, presumably under selection, the average $\bar{F}_{ST} = 0.27$ ranging from 0.08 to an extreme differentiation of 0.83, representing almost fixation of alternative alleles. We analyze nucleotide variation at a large fragment of the gene showing extreme spatial differentiation to understand the nature of selection. It is the *Ckma* gene encoding a muscle isoform A of creatine kinase.

Creatine kinases (CK) are crucially important in bioenergetic processes in cells and tissues (Wallimann et al., 1992, 2011). The creatine kinase/phosphocreatine system (CK/PCr) is an intracellular energy shuttle. CK generates Phosphocreatine (PCr) at the sites of ATP production in glycolysis and oxidative phosphorylation in mitochondria and regenerates ATP from PCr at subcellular sites of ATP use by ATPases. The physiological advantage is to provide a spatial and temporal energy buffer storing and releasing energy in and from PCr. Importantly the rate of intracellular diffusion of both Creatine (Cr) and PCr is one and three orders of magnitude faster than diffusion of ATP and ADP respectively.

Here we thus have a gene with a well defined and well understood function. The gene shows extreme spatial differentiation most likely due to selection considering the behavior of neutral non-outliers. We ask what a detailed analysis of nucleotide variation using methods of multiple merger Λ coalescents at the scale of the gene itself can tell us about the nature of selection.

105 MATERIALS AND METHODS

106 Population sampling

107 We randomly sampled 122 individual cod from various localities from the distributional
108 range of Atlantic cod (Figure S1). The samples come from our large sample database.
109 The localities are the waters around Newfoundland (New), Greenland (Gre), Iceland
110 (Ice), Faroe Islands (Far), Norway (Nor), and the Barents Sea, North Sea (Nse), Celtic
111 Sea (Cel), Irish Sea (Iri), Baltic Sea (Bal), and the White Sea (Whi).

112 For outgroup comparison we included samples of the sister taxa Arctic cod *Bore-*
113 *ogadus saida* (Bsa) and Greenland cod *G. ogac* (Gog) both sampled in Greenland
114 waters as well as Pacific cod *G. macrocephalus* (Gma) and Walleye pollock *Theragra*
115 *chalcogramma* (Gch) sampled from the Pacific ocean. Carr et al. (1999) and Pogson
116 and Mesa (2004) discuss the relationship and biogeography of these taxa. Coulson
117 et al. (2006) provide the most comprehensive account based on mitochondrial genomics.
118 They consider Arctic cod to be an outgroup for all these taxa. Atlantic cod and Walley
119 pollock are sister taxa and Pacific cod slightly more distant. Pacific cod and Walleye
120 pollock represent two separate but nearly simultaneous invasions of the Pacific with
121 the Atlantic cod vs. Pacific cod split dated at 4 mya and the Atlantic cod vs. Walleye
122 pollock split at 3.8 mya using conventional rates of mtDNA evolution (see time scales
123 below). They suggest a nomenclature revision from *Theragra chalcogramma* to *Gadus*
124 *chalcogrammus* for Walleye pollock. Greenland cod is a recent reinvasion of Pacific cod
125 into the Arctic and Coulson et al. (2006) consider it to be a subspecies of Pacific cod.

126 The Icelandic Committee for Welfare of Experimental Animals, Chief Veterinary
127 Office at the Ministry of Agriculture, Reykjavik, Iceland has determined that the research
128 conducted here is not subject to the laws concerning the Welfare of Experimental
129 Animals (The Icelandic Law on Animal Protection, Law 15/1994, last updated with
130 Law 157/2012). DNA was isolated from tissue taken from dead fish on board research
131 vessels. Fish were collected during the yearly surveys of the Icelandic Marine Research
132 Institute. All research plans and sampling of fish, including the ones for the current
133 project, have been evaluated and approved by the Marine Research Institute Board of
134 Directors. The Board comprises the Director General, Deputy Directors for Science and
135 Finance and heads of the Marine Environment Section, the Marine Resources Section,
136 and the Fisheries Advisory Section. Samples were also obtained from dead fish from
137 marine research institutes in Norway, the Netherlands, Canada and the US that were
138 similarly approved by the respective ethics boards. The samples from the US used in this
139 study have been described in Cunningham et al. (2009) and the samples from Norway
140 in Árnason and Pálsson (1996). The samples from Canada consisted of DNA isolated
141 from the samples described in Pogson (2001). The samples from the Netherlands were
142 obtained from the Beam-Trawl-Survey

143 ([http://www.wageningenur.nl/en/Expertise-Services/](http://www.wageningenur.nl/en/Expertise-Services/Research-Institutes/imares/Weblogs/Beam-Trawl-Survey.htm)
144 [Research-Institutes/imares/Weblogs/Beam-Trawl-Survey.htm](http://www.wageningenur.nl/en/Expertise-Services/Research-Institutes/imares/Weblogs/Beam-Trawl-Survey.htm))

145 of the Institute for Marine Resources & Ecosystem Studies (IMARES), Wageningen
146 University, the Netherlands, which is approved by the IMARES Animal Care Committee
147 and IMARES Board of Directors.

148 Molecular analysis

149 We used sequences associated with the Moen et al. (2008) high F_{ST} SNP's (Gm366-
150 0514 with an $F_{ST} = 0.83$, Gm366-1022 with an $F_{ST} = 0.82$, and Gm366-1073 with an
151 $F_{ST} = 0.82$) to make probes to search an Atlantic cod BAC library. We had positive
152 clones 454 sequenced (Microsynth) and obtained a 34223 bp scaffold containing the gene
153 of interest. From this sequence we generated primers (Table S1) for PCR amplifying a
154 4000 bp fragment for population studies. Our scaffold largely but not entirely aligned
155 to GeneScaffold 4232 of the Atlantic cod genome sequence (Star et al., 2011) (www.ensembl.org).
156

157 We Topo-TA cloned fragments into pCR XL-TOPO vector (Invitrogen). We se-
158 quenced clones with M13 primers and sequencing primers (Table S1) using BigDye
159 Terminator kit (Applied Biosystems) and performed sequencing on ABI 3100 and
160 ABI3500XL (Applied Biosystems) automated sequencers.

161 For neutral locus comparisons we applied the same methods and sequenced 711 bp
162 of the Hemoglobin α 2 (*HbA2*) locus (Halldórsdóttir and Árnason, 2009a,b; Borza et al.,
163 2009) and 1021 bp of the myoglobin (*Myg*) locus (Lurman et al., 2007). The *HbA2* data
164 were of 114 Atlantic cod individuals and 13 individuals of various sister taxa. The *Myg*
165 data were from 45 Atlantic cod individuals and two individuals of Pacific cod. Other
166 sister taxa did not amplify for *Myg*. The *HbA2* and *Myg* individuals covered much the
167 same geographic localities as *Ckma*.

168 All sequences have been deposited in Genbank with *Ckma* accession numbers
169 KM624178 – KM624309, *HbA2* accession numbers KM624310 – KM624436, and *Myg*
170 accession numbers KM624437 – KM624483.

171 Statistical analysis

172 We base called, assembled and edited sequence reads using phred, phrap and
173 consed (Ewing et al., 1998; Ewing and Green, 1998; Gordon et al., 1998). We
174 aligned sequences using muscle (Edgar, 2004), inspected alignments using seaview
175 (version 4) (Gouy et al., 2009) and generated maximum likelihood trees with phym1
176 (Guindon and Gascuel, 2003) under seaview. We used R (R Core Team, 2013) and
177 the ape, pegas, seqinr, ade4, adegenet, and LDheatmap packages (Paradis
178 et al., 2004; Paradis, 2010; Charif and Lobry, 2007; Dray and Dufour, 2007; Jombart and
179 Ahmed, 2011; Shin et al., 2006) and various function written by us for managing, ana-
180 lyzing, and plotting the data. We used the MLHKA program (Wright and Charlesworth,
181 2004) for a maximum likelihood HKA test (Hudson et al., 1987) based on the Kingman
182 coalescent.

183 By PCR amplifying and cloning of fragments polymerase copy errors in the PCR
184 reaction inevitably will be found in clones. The coalescent methods are especially
185 sensitive to singleton variants and errors that would enter into the data as singleton
186 variants should be removed. To remove PCR errors and ensure authenticity of natural
187 variation among individuals we sequenced three clones from each individual. We
188 claim that taking three clones is sufficient to eliminate PCR errors among clones of an
189 individual and yield a consensus sequence of one allele from that individual. Two of
190 the three clones will be of the same allele (the same chromosome). The third clone is
191 expected to be of that same allele in one half cases and of the alternative allele from the
192 other chromosome in one half cases. In the first case a consensus sequence will be a

193 true consensus of that allele. In the second case a consensus sequence will be a true
194 consensus except at sites where the third clone (alternative allele) matches one of the
195 other clones. That is when a naturally occurring site variant or a PCR error in the third
196 clone matches a PCR error in one of the other two clones. This scenario is expected to
197 be a rare event. The effect of such a rare event would be to generate variation that would
198 look like recombination thus, if anything, reducing measures of linkage disequilibrium.

199 We thus got consensus sequences for a number of individuals. We visually inspected
200 all variant sites using the above mentioned tools. To maximize the number of individ-
201 uals and the size of the sequenced fragment we struck a balance between number of
202 individuals and quality of sequence. We removed individuals with a short sequences and
203 eliminated regions with a phred quality less than 30. We thus ended up with consensus
204 sequences of three clones from each of 122 Atlantic cod and 10 individuals of sister taxa
205 covering three fragments of the gene (Figure S2) concatenated to give a total sequence
206 of 2500 bp.

207 We analyzed sequence variation for statistics of neutrality and selection using DNAsp
208 (Rozas et al., 2003) and R functions. Site frequency spectra are a most important
209 summary statistics for coalescent analysis of nucleotide data (Wakeley, 2009). We
210 analyzed site frequency spectra using the Kingman coalescent (Kingman, 1982a) and
211 statistical methods developed for multiple merger Λ coalescents (Birkner et al., 2013b).

212 RESULTS

213 Gene and protein

214 The gene is *Ckma* encoding creatin kinase muscle isoform a (CKMA). The locus
215 is 3604 base pairs (bp) in GeneScaffold 4232 (coordinates 332764 to 336367, gene
216 name ENSGMOG00000008778 in the cod genome, www.ensembl.org Star et al.
217 (2011)). The gene has seven exons (Figure S2). Ensemble reports 382 amino acids
218 (aa). However, both genescan (<http://genes.mit.edu/GENSCAN.html>)
219 and fgenesh (www.softberry.com) predicted 381 aa and our analysis of our own
220 data confirmed that. The www.ensembl.org sequence adds a Glycine (G) residue
221 in position 323 apparently due to incorrect splicing at the junction of the last two exons.

222 For mapping the gene the SNP locus cgpgmo-S497 at position 19.5 in linkage
223 group CGP16 is found in a partial cDNA mRNA sequence (Genbank accession number
224 EX184243) (Hubert et al., 2010; Borza et al., 2010) matching the *Ckma* gene. We take
225 that as the location of the gene.

226 There are seven paralogous genes found in the Atlantic cod genome (www.ensembl.org).
227 encoding mitochondrial, brain and muscle isoforms. The protein sequence of the
228 two alleles *A* and *B* in Atlantic cod and of all the sister taxa studied were of the CKMA
229 isoform (Figure S3). The variation reported is thus from orthologous genes.

230 Nucleotide variation and divergence

231 The variants of *Ckma* in Atlantic cod fell into two distinct and divergent groups which
232 we refer to as *A* and *B* alleles (Figures 1 and S4). They were fixed for a C vs T at site
233 1732 in the concatenated sequence (Table S2). There also were nearly fixed differences
234 between the alleles at 19 additional sites (Figure 2 and Table S2).

235 The divergence of the *A* and *B* alleles has arisen after the speciation between *Gadus*

236 *morhua* and its Pacific sister species *G. macrocephalus* or *G. chalcogrammus*. The gross
237 and net nucleotide divergence between the *A* and *B* alleles was about one half that of the
238 divergence between the sister taxa (Table S3). The *Ckma* and *HbA2* divergences between
239 the sister taxa are very similar but the *Myg* divergence is about twice that (Figure S5 and
240 Table S3). Contra Coulson et al. (2006) the maximum likelihood tree for *Ckma* (Figure 1)
241 and divergence estimates (Table S3) imply that separation of *G. chalcogrammus* predates
242 the separation of *G. macrocephalus* and *G. morhua*. Similarly, the *HbA2* locus showed
243 the same pattern that *G. chalcogrammus* is outside of *G. macrocephalus* and *G. morhua*
244 (Figure S6). Unfortunately the *Myg* locus did not yield sequences for *G. chalcogrammus*.

245 All summary statistics showed high variation for *Ckma* (Table 1). In particular
246 nucleotide diversity $\hat{\pi}$ was high relative to the scaled population size $\hat{\theta}_S$ resulting in a
247 non-significant Tajima's \hat{D} . This was due to the great number of high heterozygosity
248 sites nearly fixed between the two alleles (Figure 2 and Table S2). Considering the North
249 and South population and the *A* and *B* alleles separately there was much less variation.
250 Although there were several polymorphic sites within both *A* and *B* alleles (Figure 2
251 and Table S2) nucleotide diversity was lower than for the entire sample and the relative
252 difference of $\hat{\pi}$ and $\hat{\theta}_S$ for each allele was greater resulting in negative and significant
253 Tajima's \hat{D} . The *HbA2* gene had a very low haplotype and nucleotide diversity but
254 disparity with $\hat{\theta}_S$ gave overall a negative and significant Tajima's \hat{D} . In congruence with
255 divergence measures the *Myg* locus had high haplotype and nucleotide diversity, albeit
256 lower than *Cmka*, but overall a negative and significant Tajima's \hat{D} .

257 There were five non-synonymous changes segregating as singleton sites within
258 Atlantic cod (Tables S4 and S2). Two of these were also segregating as singletons
259 within *B. saida* and *G. macrocephalus* and one other singleton was also found in *G.*
260 *macrocephalus*. *B. saida* was fixed for a Glycine (GGT codon) for which the other
261 taxa have a Glutamine (CAG codon) with changes in all three sites of the respective
262 codon (aa number 242). Assuming independent mutations and depending on the path of
263 evolution of that particular codon all three changes may have been non-synonymous.

264 There was considerable linkage disequilibrium (LD) throughout the gene (Figure S7).
265 The high heterozygosity sites nearly fixed between the alleles were influential in gener-
266 ating LD between sites throughout the gene.

267 The results of a maximum likelihood HKA test of selection that is based on the
268 Kingman coalescent (Wright and Charlesworth, 2004) gave a selection parameter $k =$
269 2.12 in the direction of balancing selection (Table S5). However, the results were not
270 statistically significant possibly because of too high variation among the presumed
271 neutral loci (*HbA2* and *Myg*) used for comparison in the test.

272 Spatial differentiation

273 The variation was spatially patterned. The *A* allele was nearly fixed in an area that
274 we call South (Faroe Islands, North Sea, Baltic Sea, Celtic Sea and Irish Sea) at a
275 frequency of 97% (Table S6). Conversely the *B* allele was at a high frequency of
276 92% in an area that we call North ranging from the Northwest (Nova Scotia and
277 Newfoundland in Canada) through Greenland, Iceland, Norway, Barents Sea and the
278 White Sea. The differentiation was evident in interlocality F_{ST} values (Table S7).
279 There was no significant differentiation among localities within either the North or the
280 South but very high and significant differentiation between North and South localities.

281 Similarly, there was great differentiation between the *A* and *B* alleles (Table S8). This
282 was in stark contrast to the lack of differentiation at the *HbA2* and *Myg* loci (Table S9).

283 The high differentiation was mostly due to the great number of fixed or nearly fixed
284 sites between the two alleles (Figure 2 and Table S2). Three of the sites were the SNPs
285 already found by Moen et al. (2008) with an $F_{ST} = 0.82$. The high frequency sites
286 showed indications of recombination between the *A* and *B* alleles (see for example
287 patterns of segregating sites for individuals 105698, 124401, 105657, 200500, 118129,
288 119535, 118147, and 106620 in Table S2).

289 There were also several high heterozygosity polymorphic sites within both the *A*
290 and *B* alleles (Figure 2). This variation, however, did not show geographical patterns
291 (Table S2). For example sites 1050 and 1428 mutated relative to outgroup within the *A*
292 alleles were found among individuals from Iceland, White Sea, Celtic Sea, Faroe Islands
293 and the Baltic. Similarly within the *B* alleles high heterozygosity sites 656, 691, 1340,
294 and 1444, which were mutated relative to the outgroup, were all widespread among
295 North localities ranging from the Northwest to the Northeast Atlantic (Figure S1).

296 Site frequency spectra

297 The unfolded site frequency spectrum for the *Ckma* gene was trimodal (Figure 3), with
298 a mode at singleton sites, a mode at 43, and a mode at 79. The latter modes were at
299 opposite frequencies out of a total of 122 and represented the *A* and *B* lineages of the
300 genealogy. The Kingman coalescent did not fit the data well. Both the Beta($2 - \alpha, \alpha$)
301 and point-mass coalescent models gave a much better fit (Table S10) in particular by
302 capturing the singleton class. None of the coalescent models captured the modes at 43
303 and 79.

304 In contrast the site frequency spectra for the *HbA2* and *Myg* genes were L shaped
305 with a high peak at singleton sites (Figures S8 and S9). Again the Kingman coalescent
306 did not fit well but both multiple merger coalescent models captured the high frequency
307 of singleton sites.

308 The site frequency spectra of the *A* and *B* alleles alone were bimodal with a high
309 singleton class and peaks around 40 and 78 respectively (Figure S10). The 40 and
310 78 modes came about because most of the high frequency and high heterozygosity
311 sites that separate the two alleles were not fixed within each allele presumably due to
312 recombination (Table S2).

313 Coalescent parameter estimates

314 Following Birkner et al. (2013b) we used the ℓ^2 distance, the sum of the squared
315 differences between the observed and expected site frequency spectrum (scaled with the
316 number of segregating sites), for estimating parameters of two Λ coalescent models, $\hat{\alpha}$
317 for the Beta($2 - \alpha, \alpha$) and $\hat{\psi}$ for the point-mass coalescent (Table 2 and Figures S11
318 and S12). The Kingman coalescent, a null model for which $\alpha = 2.0$, had the highest ℓ^2
319 indicating worst fit among the models. The *HbA2* and *Myg* loci had an $\hat{\alpha} = 1.00$ and a
320 $\hat{\psi} = 0.23$. The *Ckma* locus had overall a considerably higher α and lower ψ . The two
321 alleles separately were in the direction of the presumed neutral loci *HbA2* and *Myg*.

322 For comparison we also estimated the parameters for the entire dataset of mtDNA
323 variation in the North Atlantic (Árnason, 2004) and the various subsamples making up
324 that total sample using the unfolded site frequency spectrum with *G. macrocephalus*

as outgroup (Table 2 and Figure S12). Previously these have been analysed using the folded site frequency spectrum (see for example Birkner et al., 2013b; Steinrücken et al., 2013). For the total sample, spanning a similar geographic range as the nuclear genes, the parameter estimates differed from the nuclear loci with $\hat{\alpha} = 1.53$ and $\hat{\psi} = 0.01$. The large samples from Newfoundland and Iceland and the sample from the Faroe Islands gave similar values. The values for Greenland, Norway, White Sea, and Baltic Sea were much closer to the results for the Kingman coalescent ($\alpha = 2.0$). For these localities homoplasies were relatively somewhat more frequent in the data than for the total and the large samples. Homoplasies will reduce the number of singletons and move such sites towards the right tail of the site frequency distribution. This explains the higher values for these localities.

DISCUSSION

Genes and proteins

The CKMA protein is highly conserved among the taxa. The single aa difference between *B. saida* and the other species presumably is adaptive with all sites of the codon having changed. The few aa variants were all singletons in the sample. In fact most of the variation is in non-coding regions and all the high heterozygosity sites in coding regions are synonymous changes. Given the high conservation of the protein and the high variation among silent and non-coding sites that are indicative of the mutational pressure the singleton non-synonymous changes are likely slightly deleterious and will be removed by purifying selection. Some or even all of the silent and non-coding differences between the *A* and *B* alleles may be functional control elements important in expression in different tissues or under different environments. The potential functional differences remain to be studied.

The *HbA2* and *Myg* genes have well defined functions. They are likely under purifying selection. They were taken as independent genes in separate linkage groups for comparison. A caveat is that genetic variation at unlinked sites may be correlated and not independent in high fecundity populations with skewed distribution of offspring (Eldon and Wakeley, 2008; Birkner et al., 2013a). The question remains, however, whether and to what extent such dependence impacts inference.

Allele divergence and spatial differentiation

Three possible scenarios and explanations for the great divergence of the *A* alleles and *B* alleles, their spatial differentiation, and the trimodal site-frequency spectrum will now be considered.

First, there is the possibility of recent admixture of anciently separated and divergent gene pools that have come together in a hybrid zone of secondary contact (Bowcock et al., 1991; Bernardi et al., 1993; Guinand et al., 2004). The spatial patterns of genetic separation between the South (Faroe Islands, North Sea, Baltic Sea, Celtic Sea and Irish Sea) and the North (Nova Scotia and Newfoundland, Greenland, Iceland, Norway, Barents Sea, and White Sea) could be taken as evidence for this. The South is a shallow water environment whereas the the North has more diversity of depth ranging from shallow to deep waters. Differences in temperature, salinity and other environmental factors is correlated with the North South difference. The great nucleotide divergence

between the North and the South would imply either that this is an ancient divergence or even a not-so-ancient divergence driven by strong selection over a shorter time. If the time of separation of *G. morhua* and *G. macrocephalus* and *G. chalcogrammus* is taken at 3.8–4.0 Mya (Coulson et al., 2006) the time of separation of the *A* and *B* clades would then be 2 Mya based on the nucleotide divergence of the *A* and *B* clades which we show is one half that of the sister taxa. An even lower divergence time of 2.1 Mya has been suggested (Pogson and Mesa, 2004) that would still leave the divergence of the *A* and *B* clade at 1 Mya. These divergence times, however, are all based on the Kingman coalescent and time scales of the multiple merger coalescent are discussed below.

A counter argument is that isolation and admixture are part of the breeding structure of a population leaving genome-wide impacts (Wright, 1931). Under this scenario different genes should be concordant in their behavior (Bernardi et al., 1993). The *Hba2* and the *Myg* show no differentiation between the North and the South. Also the non-outlier SNPs in Moen et al. (2008) show no differentiation whereas three SNPs of the *Ckma* gene show high and extreme F_{ST} . Similarly, Bradbury et al. (2010) find that non-outlier SNPs show no differentiation although other SNPs show differentiation from parallel adaptation to temperature on the eastern and western side of the Atlantic Ocean. Nielsen et al. (2003) describe a pattern of microsatellite variation in a transition area between the Baltic and Danish Belt Sea which they interpret as a hybrid zone. There is no evidence for a hybrid zone at that location in the *Ckma* data. In fact, specific variants within the *A* allele are widely distributed among localities in the South including the Baltic Sea. This implies gene flow among localities in the South. Similar patterns within *B* alleles imply gene flow among localities in the North. If indeed there is a hybrid zone for the *Ckma* gene it would lie between the Faroe Islands on one hand and Iceland and north and middle Norway on the other hand. It is not a parsimonious explanation to consider there to be multiple hybrid zones of secondary contact within distribution of the species.

For comparison one can consider the *Pan I* locus (Fevolden and Pogson, 1995, 1997) that clearly is under selection (Pogson, 2001; Pogson and Mesa, 2004) related to depth and fisheries (Sarvas and Fevolden, 2005; Case et al., 2005; Árnason et al., 2009). At face value the locus shows similar differentiation between north and south (Sarvas and Fevolden, 2005) as the *Ckma* locus. However, the details differ. The *Pan I B* allele which is adapted to the deep (Pampoulie et al., 2007; Árnason et al., 2009) is largely absent from the South. However, there is no particular *Pan I A* allele that characterizes the South (Hernandez and Árnason, 2014). The *Pan I B* allele, which is found in the North and in deep water, is much less variable than the *Pan I A* alleles (Pogson, 2001; Hernandez and Árnason, 2014). This is opposite to what we find for the *Ckma A* alleles (the South allele) which has less variation than the *Ckma B* allele (Figure 1) although this is not seen in the summary statistics (Table 1) because of greater recombinational variation at the base of the *A* clade (Table S2). Also the *Pan I* locus variation is more related to depth than to geography (Árnason et al., 2009). Under the admixture scenario these two loci (and all loci showing genome wide effects) are expected to show the same pattern.

Overall, therefore, we find that the *Ckma* gene does not fit the scenario of ancient divergence of gene pools and admixture in secondary contact.

413 Site frequency spectra

414 The trimodal site frequency spectrum is not predicted by any of the coalescent mod-
415 els considered here, the Kingman coalescent and the two Λ coalescent models, the
416 Beta($2 - \alpha, \alpha$) (Schweinsberg, 2003) and the point-mass coalescent (Eldon and Wake-
417 ley, 2006). Under the Λ coalescent at most a single multiple merger event occurs at any
418 one time. The distribution of family size is of interest and the parameter α influences
419 the probability of getting large families. Under the Beta($2 - \alpha, \alpha$) coalescent model the
420 probability of a family size of k or more viable offspring decays like $k^{-\alpha}$ (Schweinsberg,
421 2003) in the limit of a large k . The pool of viable offspring is then resampled to form the
422 next generation under the same conditions. For the Kingman coalescent $\alpha \geq 2$ and there
423 is little chance of seeing large families. For the Beta($2 - \alpha, \alpha$) coalescent $1 \leq \alpha < 2$ and
424 the lower α the greater is the chance of seeing a large family (Schweinsberg, 2003). The
425 ψ parameter of the point-mass coalescent (Eldon and Wakeley, 2006) similarly measures
426 the proportion of the population that is the offspring of a single individual and is thus an
427 indicator of reproductive sweepstakes. Our estimates of ψ indicate reproductive sweep-
428 stakes at the neutral loci and within the *A* and *B* alleles of *Ckma*. Balancing selection
429 at *Ckma* lessens the effects of sweepstakes reproduction. Sweepstakes reproduction
430 has been detected in other high fecundity organisms (Hedgecock and Pudovkin, 2011;
431 Harrang et al., 2013).

432 Under the more general Ξ coalescent $0 < \alpha < 1$ (Schweinsberg, 2000) there can
433 be many large families independently in each generation. It would seem that this
434 process could generate multimodal site frequency spectra. Indeed in simulations of Ξ
435 coalescence site frequency spectra can display multiple modes (Bjarki Eldon personal
436 communication). This question needs further theoretical work. In terms of the concept
437 of sweepstakes reproduction multiple local sweepstakes could have this effect on the
438 site frequency spectrum. Under local sweepstakes genetic structure may be ephemeral
439 (Johnson and Wernham, 1999). Whether this affects the location of the modes and the
440 exact shape of the site frequency spectrum under Ξ coalescent is not known. However,
441 one would not expect build-up of sites around a specific mode of the site frequency
442 spectrum or of two modes at opposite frequencies as at *Ckma*. Also there should be
443 no particular or regular geographical pattern. We, therefore, think that bumps in the
444 site frequency spectrum under Ξ coalescent is not a good explanation for the *Ckma*
445 spectrum.

446 Balancing selection

447 Balancing selection generates long branches in the genealogy and neutral variation
448 accumulates on the branches. The balanced functional types (the *Ckma A* and *B* alleles
449 in this case) act as they were separate and isolated populations accumulating neutral
450 variation. Recombination can bring variation from one branch to another acting like
451 migration that brings alleles from one population to another (Charlesworth et al., 1997,
452 2003; Charlesworth, 2006). However, the molecular signatures of balancing selection
453 depend on many factors. Is it a long standing, even trans-species, polymorphism such as
454 *MHC* in human and chimpanzee (Fan et al., 1989; Nei and Hughes, 1991) or *Cathelicidin*
455 in gadids (Halldórsdóttir and Árnason, 2014)) or is it very recent? Examples of the
456 latter are human glucose 6 phosphate dehydrogenase (G6PD) (Verrelli et al., 2002), and
457 hemoglobin β *S* (Curat et al., 2002) and hemoglobin β *E* (Ohashi et al., 2004) and

spatially divergent selection of lactase persistence (Tishkoff et al., 2007; Ranciaro et al., 2014) in which a particular allele sweeps a chromosomal segment to an intermediate equilibrium frequency. In these instances recombination has not had time to break up LD which can extend over large regions. There is very little variation among the new alleles while the alternative chromosomes show much more variation in this region representing the standing variation in the population at the start of the partial sweep.

The effects of a long standing single locus balancing selection will extend only short distances with free recombination and will be difficult to detect (Wiuf and Hein, 1999). If, however, there are obvious signs of a long standing balanced polymorphism it is likely due to a build-up of co-adapted complexes of epistatic interactions among multiple sites and/or suppression of recombination (Wiuf and Hein, 1999). The concept of a supergene of multiple co-adapted sites possibly locked together by structural variation (Thompson and Jiggins, 2014) such as found in butterfly mimicry (Joron et al., 2011) is relevant. There also can be both partial and complete selective sweeps of new types within each allele of a supergene. Such intra-allelic selective sweeps would reduce variation within and increase variation between alleles. Such reduction of variation could look similar to that for a recent balanced polymorphism except that it would not be limited to one functional type. Thus Pogson (2001) argues that he has detected on-going partial sweeps within each of the two *Pan I* alleles of Atlantic cod.

Pogson and Mesa (2004) further argue that the *Pan I* polymorphism is older than speciation of Atlantic cod and Walleye pollock, the closest relatives. The *Pan I* locus is in a “genomic island” (Bradbury et al., 2013; Hemmer-Hansen et al., 2013) a potential supergene of co-adapted complexes possibly locked together by structural variation. Looking in detail at variation at 12.5 kb region Hernandez and Árnason (2014) find large number of differences between the two functional *Pan I* types that are too extensive to be a partial sweep of a new allele. Such variation is likely to be built up over some time by selection (see time scales below). This is in face of considerable gene flow implied by lack of differentiation of neutral loci (Moen et al., 2009; Bradbury et al., 2010; Eiríksson and Árnason, 2013; Hemmer-Hansen et al., 2014). Similarly, the wide distribution of variants within both the *A* and *B* alleles of *Ckma* implies gene flow among localities within South and within North areas. The recombinant haplotypes between the *A* and *B* alleles of *Ckma* imply gene flow between the South and the North localities.

The coalescent used here are models of neutrality. One could argue that it is not appropriate to apply such neutral models to the *Ckma* locus that is already suspected to be under selection. However, understanding how the locus deviates from neutrality is important for understanding the pattern of selection. Under the neutral theory (Kimura, 1983) polymorphism within species is the transient phase of molecular evolution that leads to divergence between species. This is the rationale for the HKA test of selection or neutrality (Hudson et al., 1987) that neutrally evolving genomic regions should have the same proportion of polymorphism to divergence. Balancing selection would tend to increase the level of polymorphism within species relative to divergence between them. The results of HKA test are in the direction of balancing selection. The HKA test shows a relative slowing down of divergence to rate of polymorphism at the *Ckma* locus.

Similarly we consider the peaks in the site frequency spectrum of the *Ckma* gene to be evidence for balancing selection. The trimodal site frequency spectrum with two high frequency peaks at opposite frequencies that fold into one peak in a folded

site frequency spectrum points to the build-up of variation over time. Under a recent balanced polymorphism scenario, such as *G6PD* and β globins in humans, there would be one peak at a particular frequency in the site frequency spectrum representing all sites at which the new allele differs from the ancient alleles. There could be multiple peaks representing high frequency polymorphisms among the ancient alleles. However, they are not expected to be at opposite frequencies to the frequency of the new allele. We, therefore, argue that the pattern at *Ckma* represents a balanced polymorphism that has been built up over time.

Coalescent parameter estimates and time scales

The question of coalescent time scale, however, must be considered. Under the Kingman coalescent time is measured in terms of N/σ^2 , population size scaled by the variance of family size (Sagitov, 1999; Árnason, 2004; Tavaré, 2004). With a Poisson distribution of family size $\sigma^2 = 1$ for a constant size haploid population so times scales with N . In an extreme winner-take-all sweepstakes $\sigma^2 = N$ and a sample would coalesce in the previous generation and there would be no variation (Árnason, 2004). In more realistic multiple merger coalescent models the time scale is the quantity $c_N = \frac{E(v_1-1)^2}{N-1}$ where c_N is the probability of two lineages coalescing in the previous generation in a haploid population of fixed size N and v_1 is the random number of offspring of individual 1 (Sagitov, 1999). In general the time scale of multiple merger coalescent models can be much shorter than for Kingman coalescent. Under the Beta($2 - \alpha, \alpha$) coalescent model time scales with $N^{\alpha-1}$ (Schweinsberg, 2003; Birkner et al., 2014). For this model our estimates of α for the nuclear genes are quite low which implies very short time scales. The neutral genes would seem to coalesce in the very recent past. The *A* and *B* alleles of *Ckma* run on very similar time scales to the neutral genes and the locus itself at a slower rate due to the balancing selection with a time scale approximately the cube root of the effective population size N_e . The mitochondrial DNA runs at yet another and slower time scale. For mtDNA time scales with approximately the square root of N . Predicted turnover of alleles is faster and ages of alleles shorter under multiple merger coalescent (Eldon, 2012). Different populations and species may run on different time scales (Eldon and Degnan, 2012) complicating divergence time estimates. Estimates based on Kingman coalescent of divergence times of Atlantic cod populations (Bigg et al., 2008) or divergence of gadid taxa (Coulson et al., 2006) may therefore be too high and may need revision.

Conclusion

The *Ckma* protein coding sequence is conserved between all but the most distantly related Arctic cod. The amino acid variants are all singletons in the sample. Based on these facts we conclude that the protein coding sequence is under purifying selection. At the same time silent and non-coding variation at the locus shows extreme spatial differentiation with an F_{ST} greater than 0.8 between the North and the South regions. The regulatory function of this variation is unclear. We argue that the high and locus-specific F_{ST} , the highest seen so far for any locus and any spatial comparison in Atlantic cod, indicates that selection and not admixture of anciently divergent gene pools is responsible. Selection is likely to be very strong. It follows that *Ckma* (or an extremely tightly linked locus) is the focus of selection because the highest F_{ST} indicates the site

548 of action of selection (Nielsen, 2005). Some of the variation may be neutral having
549 risen in frequency within the balanced functional allele where it arose (Charlesworth,
550 2006). Alternatively some of the variation may be due to selection building co-adapted
551 complexes (Thompson and Jiggins, 2014). In addition to a peak at singleton sites,
552 characteristic of multiple-merger coalescent, the site frequency spectrum has two high-
553 frequency modes at opposite but matching frequencies representing the two branches of
554 the genealogy. This pattern is further support for balancing selection. Finally time scales
555 faster under multiple-merger than the Kingman coalescent. Our estimates of parameters
556 of multiple-merger Λ coalescent show that time-scales are fast.

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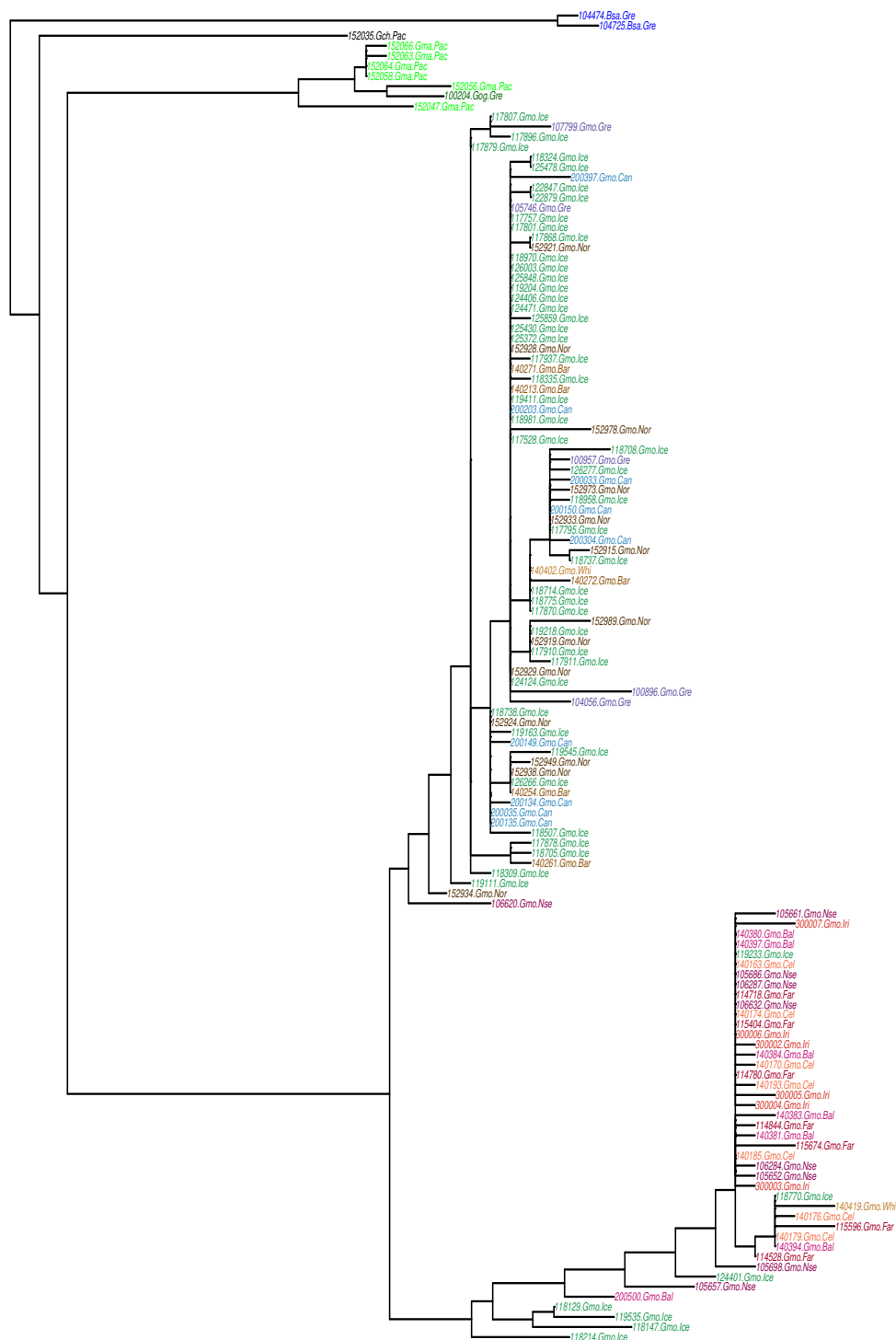


Figure 1. Maximum likelihood tree of *Ckma* variation among 122 individual Atlantic cod and 10 individuals of sister taxa. Localities and color codes as in Figure S1.

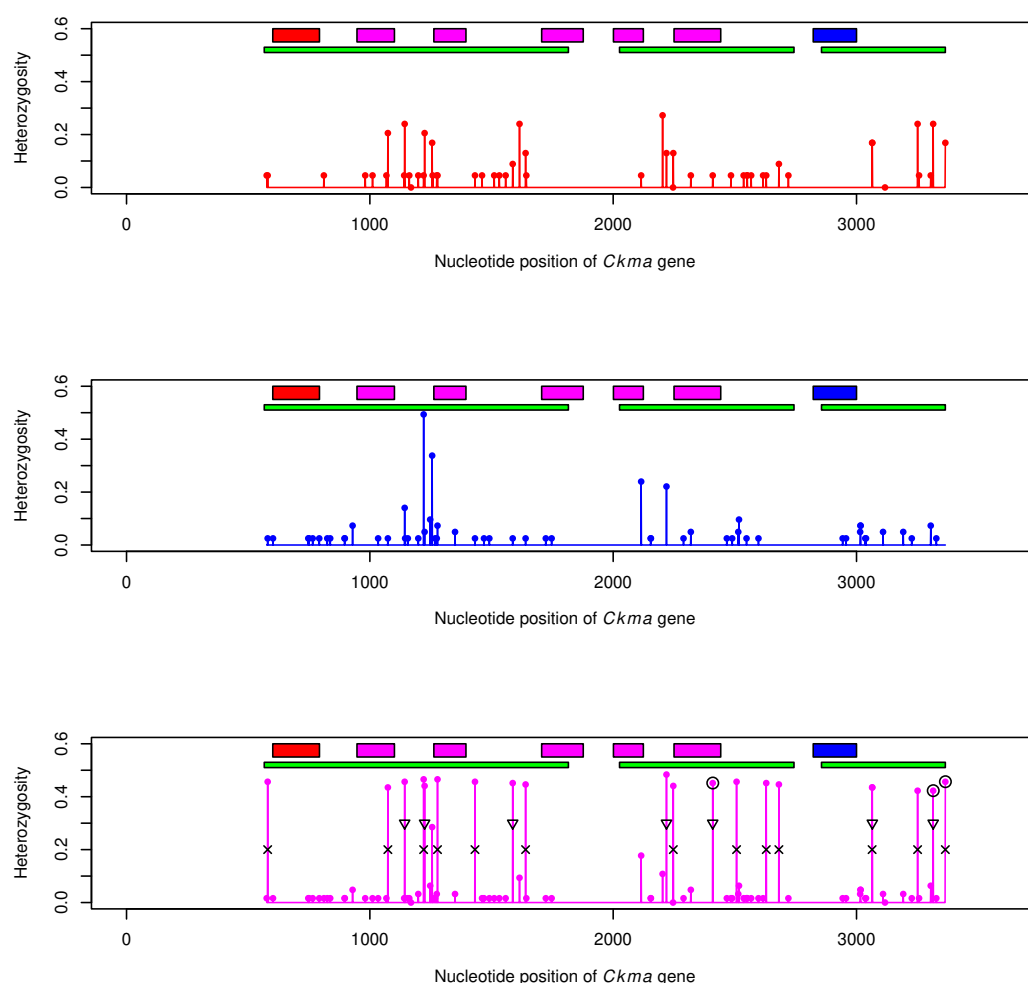


Figure 2. Heterozygosity per nucleotide site of *Ckma* locus among *A* alleles (red top panel, $n = 43$), *B* alleles (blue middle panel, $n = 79$), and all individuals combined (magenta bottom panel, $n = 122$). Boxes represent exons, start (red), internal (magenta) and terminal (blue). Green boxes represent sequenced fragments trimmed to Phred score of at least 30. The black circles mark the three SNPs of Moen et al. (2008), *Gm366-0514* locus with an $F_{ST} = 0.83$, *Gm366-1022* locus with an $F_{ST} = 0.82$, and *Gm366-1073* with an $F_{ST} = 0.82$ from left to right respectively. Crosses mark mutant sites relative to outgroup that were fixed or nearly fixed among *A* alleles. Triangles mark mutant sites relative to outgroup that have were fixed or nearly fixed among *B* alleles. *Gadus macrocephalus* individual 152047 was outgroup.

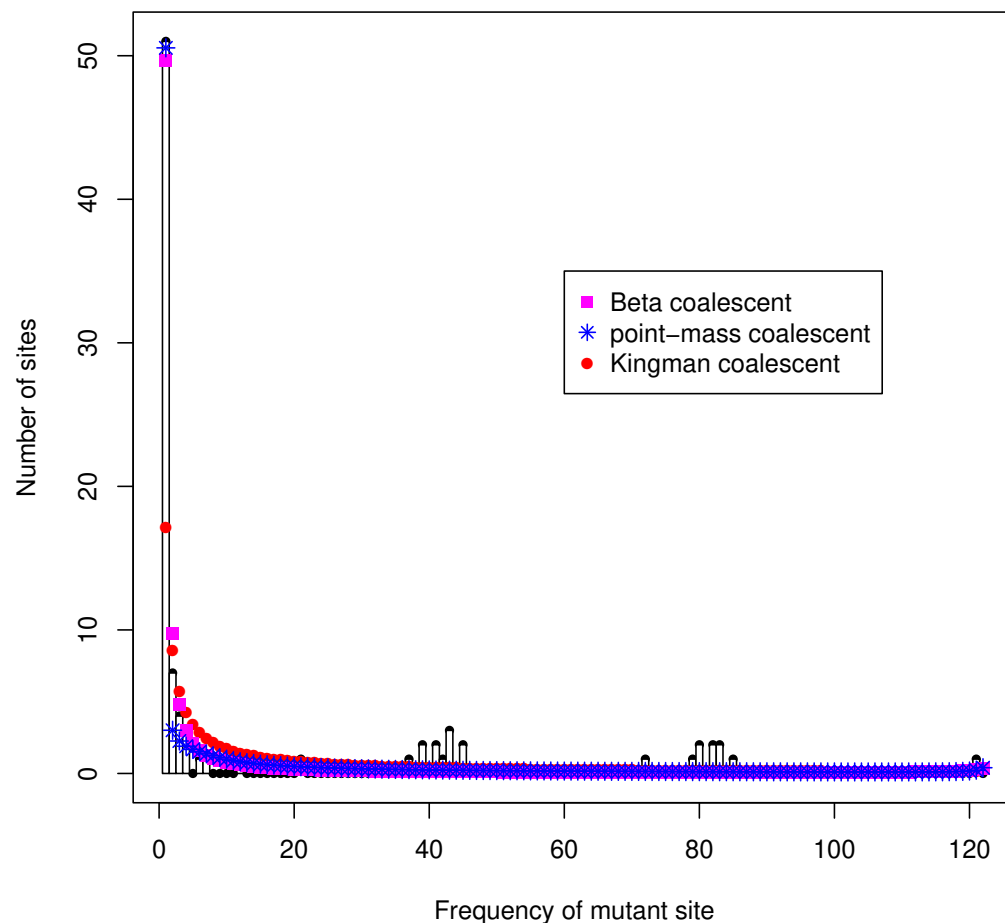


Figure 3. Unfolded site frequency spectrum of Atlantic cod *Ckma* gene. *Gadus macrocephalus* is outgroup. Number of individuals $n = 122$. Theroretical expectation under Kingman coalescent (red dots), Beta($2 - \alpha, \alpha$) coalescent (magenta squares), and point-mass coalescent (blue stars).

Table 1. Summary statistics of polymorphism of 2500 bp fragment of the *Ckma* gene, 711 bp fragment of the *HbA2* gene and 1021 bp fragment of the *Myg* gene in Atlantic cod.

Group	n	S	H	\hat{h}	\hat{K}	$\hat{\theta}_S$	$\hat{\pi}$	\hat{D}
<i>Ckma</i> all	122	87	72	0.959	10.62	0.0067	0.0043	−1.13 ^{ns}
<i>Ckma</i> North	86	65	51	0.941	5.12	0.0054	0.0015	−1.97 ^{ns}
<i>Ckma</i> South	36	45	23	0.891	3.61	0.0045	0.0015	−2.43 ^{**}
<i>Ckma</i> A allele	43	49	28	0.907	4.37	0.0047	0.0018	−2.20 ^{**}
<i>Ckma</i> B allele	79	53	44	0.930	3.10	0.0044	0.0013	−2.33 ^{**}
<i>HbA2</i> all	114	11	11	0.338	0.37	0.0030	0.0005	−2.09 [*]
<i>HbA2</i> North	95	9	9	0.347	0.39	0.0025	0.0005	−1.95 [*]
<i>HbA2</i> South	19	3	4	0.298	0.32	0.0016	0.0005	−0.95 ^{ns}
<i>Myg</i> all	45	30	24	0.901	2.74	0.0071	0.0028	−2.03 [*]
<i>Myg</i> North	36	28	20	0.894	2.65	0.0069	0.0027	−2.12 [*]
<i>Myg</i> South	9	10	7	0.944	3.22	0.0037	0.0033	−0.58 ^{ns}

Sample size n , number of segregating sites S , number of haplotypes H , haplotype diversity \hat{h} , average number of pairwise differences \hat{K} , scaled population size from S $\hat{\theta}_S$, nucleotide diversity $\hat{\pi}$, and Tajima's \hat{D} . ns is not significant, * represents $P < 0.05$, and ** represents $P < 0.01$.

Table 2. Parameter values minimizing the ℓ^2 distance (sum of squares) between observed and expected unfolded site frequency spectra for nuclear genes and for mtDNA variation of various localities.

Source	$\hat{\alpha}$	$\hat{\psi}$	$\ell^2(\hat{\alpha})$	$\ell^2(\hat{\psi})$	$\ell^2(0)$	n	Reference
Nuclear locus							
<i>Hba2</i>	1.000	0.230	0.035	0.016	0.431	113	This study
<i>Myg</i>	1.000	0.225	0.010	0.018	0.230	45	This study
<i>Ckma</i>	1.280	0.070	0.006	0.007	0.141	122	This study
<i>Ckma</i> ^A	1.100	0.170	0.017	0.012	0.161	43	This study
<i>Ckma</i> ^B	1.140	0.120	0.006	0.015	0.189	79	This study
Locality for mtDNA							
Newfoundland	1.550	0.015	0.014	0.028	0.084	378	Carr <i>et al.</i>
Greenland	1.945	0.005	0.072	0.071	0.072	78	Árnason <i>et al.</i> (2000)
Iceland	1.550	0.010	0.006	0.050	0.078	519	Árnason <i>et al.</i> (2000)
Norway	1.895	0.015	0.093	0.089	0.095	100	AP 1996
White Sea	2.000	0.005	0.551	0.554	0.551	109	Árnason <i>et al.</i> (1998)
Faroe Islands	1.555	0.050	0.059	0.055	0.093	74	SA 2003
Baltic Sea	2.000	0.005	0.105	0.109	0.105	109	Árnason <i>et al.</i> (1998)
Atlantic	1.530	0.010	0.006	0.055	0.249	1278	Árnason (2004)

Based on method of Birkner *et al.* (2013b). Parameters α of the Beta($2 - \alpha, \alpha$), and ψ of the point-mass coalescent and their respective ℓ^2 . The $\ell^2(0)$ is based on the Kingman coalescent for which $\alpha = 2$. For the mtDNA Carr *et al.* refers to Carr and Marshall (1991a,b); Carr *et al.* (1995); Pepin and Carr (1993), AP 1996 refers to Árnason and Pálsson (1996), and SA 2003 refers to Sigurgíslason and Árnason (2003).