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Patterns of variability at the major histocompatibility class II alpha locus in Atlantic salmon contrast with those at the class I locus

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Abstract In order to investigate the mechanisms creating and maintaining variability at the major histocompatibility (MH) class II alpha (*DAA*) locus we examined patterns of polymorphism in two isolated Atlantic salmon populations which share a common post-glacial origin. As expected from their common origin, but contrary to the observation at the MH class I locus, these populations shared the majority of *DAA* alleles: out of 17 sequences observed, 11 were common to both populations. Recombination seems to play a more important role in the origin of new alleles at the class II alpha locus than at the class I locus. A greater than expected proportion of sites inferred to be positively selected (potentially peptide binding residues, PBRs) were found to be involved in recombination events, suggesting a mechanism for increasing MH variability through an interaction between recombination and natural selection. Thus it appears that although selection and recombination are important mechanisms for the evolution of both class II alpha and class I loci in the Atlantic salmon, the pattern of variability differs markedly between these classes of MH loci.

Keywords MHC · Class II alpha · Recombination · Natural selection · Atlantic salmon

Introduction

The genes of the major histocompatibility complex (MHC) are among the best studied loci in vertebrates as they play a critical role in triggering the immune response (Klein 1986) and because they are among the most variable genes known (Parham and Ohta 1996). Numerous studies suggest that these high levels of variability are maintained by natural and sexual selection in a wide range of organisms (reviewed by Apanius et al. 1997; Jordan and Bruford 1998; Bernatchez and Landry 2003). MHC variation is generated and new alleles created by point mutation and by recombination and gene conversion events between existing alleles (Belich et al. 1992; Watkins et al. 1992; Parham and Ohta 1996; Martinsohn et al. 1999). Most of the amino acid variation is concentrated at peptide binding residues (PBRs): those amino acids responsible for binding peptides derived from pathogens. The high frequency of non-synonymous substitutions per site in PBR codons (Hughes and Nei 1988, 1989), long persistence times of alleles (Takahata 1990) and patterns of intra-population allelic variation indicate that MHC genes are under balancing selection (Apanius et al. 1997; Hedrick 1999), probably in relation to pathogen exposure (Edwards and Hedrick 1998; Hedrick 2002; Penn et al. 2002).

Most knowledge of the structure and function of MHC genes come from studies in mammals, particularly in humans (Hughes and Nei 1988, 1989; Hughes et al. 1994; Parham and Ohta 1996) and many studies in other vertebrates assume that the mechanisms that create and maintain MHC variability are similar across taxa. PBRs, for example, are commonly deduced by comparing MHC sequences from a species of interest with human counterparts (e.g. Grimholt et al. 1993; Kim et al. 1999; Hoelzel et al. 1999; Hambuch and Lacey 2002), where the crystal structure has been analysed (Brown et al. 1993). Often, patterns of selection on MHC genes are then studied with respect to PBRs/non-PBRs identified in

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this manner. However, methods for detecting positive selection that do not require a priori identification of candidate sites have recently been developed (Yang and Bielawski 2000).

Further insight into the molecular evolution of MHC genes can be obtained from the study of other vertebrate species, especially those with a relatively simple MHC structure, such as amphibians (Nonaka et al. 1997), chickens (Kaufman et al. 1999) or salmonids (Shum et al. 2001). Salmonids in general, and Atlantic salmon in particular, have proved to be a good model to analyse the mechanisms of MHC evolution because (1) as in the rest of the teleosts, class I and class II loci are not physically linked, allowing independent evolution of both classes of genes (Grimholt et al. 2002) [as the class I and class II genes do not form a complex they are known as MH genes in teleosts (Stet et al. 2003)], (2) single class I and class II MH loci are expressed (Shum et al. 2001; Stet et al. 2002; Grimholt et al. 2002), making analyses simpler, and (3) most of the present distribution of the species was colonized from refugia after the last glaciation (Hewitt 1999), making possible the study of adaptation of similar lineages colonising different environments and the analysis of molecular evolution on a relatively short temporal scale.

A comparison of the allelic composition at the MH class I locus between two geographically separated Atlantic salmon populations isolated since the last glaciation (15,000 years ago) showed that despite their common origin, the populations possessed almost non-overlapping sets of alleles although they share major ancient allelic lineages (Consuegra et al., submitted). As the substitution rate of single base mutations is relatively slow, even at MHC loci (Klein and O'Huigin 1994), the most plausible mechanism for the rapid divergence of alleles in these populations is recombination between alleles. Strong evidence for recombination was found, particularly in the regions close to and including sites under positive selection and therefore potential PBRs. It is not known, however, whether the class II loci (*Sasa-DAA* and *Sasa-DAB*), that are tightly linked to each other but not linked to class I (Grimholt et al. 2000; Stet et al. 2002), are evolving the same way in the short term. In particular, it is not known whether the role of recombination in generating new alleles is as important in class II as it appears to be in class I for salmonid fish (Shum et al. 2001; Consuegra et al., submitted).

Class I and class II MHC genes are known to evolve differently not only in primates (Boyson et al. 1996) but also in other vertebrates such as cyprinids (Kruiswijk 2002) and sparrows (Bonneaud et al. 2004), although in different ways. In primates, class II loci show trans-species sharing of allelic lineages that is not seen at class I loci (Boyson et al. 1996; Seddon and Ellegren 2002). In contrast, in cyprinids and salmonids class I alleles represent highly divergent and ancient allelic lineages while class II alleles are more recent (Shum et al. 2001; Stet et al. 2002; Kruiswijk 2002).

Here we analyse variation at the class II alpha locus (*Sasa-DAA*) in two recently diverged Atlantic salmon populations, using maximum likelihood methods for detecting molecular

adaptation and recombination to examine if (1) both populations differ in allelic composition at the class II alpha locus (as for class I), (2) there is evidence of positive selection on any sites that may indicate that they are putative PBRs, and (3) recombination at the PBRs is involved in creating new alleles.

Materials and methods

Samples

Juvenile Atlantic salmon were sampled from four west coast Irish rivers with natural populations of Atlantic salmon: Owenmore, Owenduff, Burrishole and Carrowiskey. Samples of white muscle or adipose fins were stored in 95% ethanol while anterior kidney tissue was stored in RNAlater buffer (Qiagen) for subsequent extraction of RNA.

DNA isolation, cDNA synthesis, amplification and sequencing

Genomic DNA was isolated from muscle samples using the GeneClean DNA Purification kit (Qiagen), resuspended in 100 μ l of elution buffer and stored at 4°C until use in PCR amplifications.

Total RNA was extracted from anterior kidney tissue of 17 individuals using the Purescript RNA Isolation kit from GENTRA (Gentra Systems, Minneapolis, Minn., USA) and 11 μ l of purified RNA digested with DNase I was used to synthesise first strand cDNA using the First-Strand cDNA Synthesis kit (Amersham Pharmacia Biotech UK). First strand cDNAs were used as templates for PCR amplification of the Atlantic salmon β -actin locus to check for possible genomic DNA contamination (the presence of an intron between the primers results in products of different size in genomic and cDNA) with the primers (Act fwd 5'-ATGGAAGATGAAATCGCCGC-3' and Act_rev 5'-TGC CAGATCTTCTCCATGTCG-3'). Samples that gave a band of the correct size (~200 bp), with no evidence of genomic DNA contamination (a product of ~450 bp), were then used for amplification of the MHC class II alpha locus.

A region of ~214 bp of the cDNA was amplified with a 50:50 mix of the following primers: DAAexon2_fwd: 5'-GGTTTCTTTTCTCAGTTCTGC-3', and DAAexon2_rev: 5'-CTTCTCTCTTACCTATTTTCTTCTTG-3'. This region spans most of exon 2 (α_1 domain) of the class II *Sasa-DAA* locus (Grimholt et al. 2002). The final amplification volume was 25 μ l, distributed as follows: 16.6 μ l sterilized distilled water, 2.5 μ l 10 \times amplification buffer, 3.5 μ l 2 mM dNTPs, 1.5 μ l 50 mM MgCl₂, 0.075 μ l of each primer (100 pM), 0.5 μ l DMSO and 1.25 U of *Taq* polymerase (Invitrogen). PCR conditions were 95°C for 5 min, then 5 cycles of 94°C for 30 s, 60°C for 30 s and 72°C for 1 min, then another 5 cycles of 94°C for 30 s, 55°C for 30 s and 72°C for 1 min and 25 cycles of 94°C for 30 s, 50°C for 30 s and 72°C for 1 min followed by an extension of 72°C for 10 min.

The PCR products were run in a 1% agarose gel, bands of the expected size were excised and DNA was purified using the Qiaquick Gel Purification Kit (Qiagen) and resuspended in 30 μ l of elution buffer. The purified products were sequenced on both strands using the same forward and reverse exon 2 primers with the ABI Prism BigDye Terminator Cycle Sequencing Kit diluted with Better Buffer (Microzone) following the manufacturer's protocol and sequences were resolved on an ABI Prism 377 automated sequencer. The sequences were compared with previously described Atlantic salmon sequences (Stet et al. 2002; Megens et al., in preparation). In the case of previously undescribed sequences the purified products were cloned into the pCR2.1 plasmid vector (TA-cloning kit, Invitrogen) and transformed in INF'strain of *Escherichia coli*. Plasmid DNA from at least 5 colonies per individual was isolated using the Quiaprep Spin Miniprep Kit (Qiagen) and sequenced as described above.

Sequence analysis

Only sequences represented by at least two clones from independent PCRs were considered in subsequent analyses. Sequences from class II alleles were aligned with Sequencher (Genecodes) software and BioEdit v5.0.9 (using the Clustal W program included in the package). MEGA 2.1 (Kumar et al. 2001) was used to calculate the gamma distance from the amino acid sequences and to build a neighbour-joining phylogenetic tree with 1,000 bootstrap iterations to assess support for nodes in the phylogeny. *Onchorynchus mykiss* class II alpha (*Onmy-DAA*) sequences (GenBank accession nos. AJ251431–33) were used as an outgroup to root the tree (Grimholt et al. 2000). As in previous studies (Stet et al. 2002; Grimholt et al. 2002), alleles were defined on the basis of deduced amino acid sequences, not on nucleotide sequence.

Detecting positive selection

The ratio of non-synonymous/synonymous substitutions ($\omega=d_N/d_S$) is the most common measure used for detecting positive selection acting on protein coding genes. A ratio of $\omega>1$ is interpreted as evidence that non-synonymous mutations result in fitness advantages and are fixed at a higher rate than synonymous mutations (are positively selected). Neutral amino acid changes will result in $\omega=1$ while amino acid sites under purifying selection will produce ratios $\omega<1$. As a high number of the sites in the protein will be invariant due to structural constraints, classical analysis comparing rates of non-synonymous and synonymous substitutions in the complete coding sequence can be inadequate to detect positive selection. To overcome this problem, we used maximum-likelihood models of codon substitution to address the question of whether the rate of non-synonymous substitution (d_N) is greater than the rate of synonymous substitution (d_S) over the entire set of se-

quences, taking into account the phylogenetic structure of the sequences. To detect positive selection we used different codon-based models that allow for variable selection among sites as recommended by Yang et al. (2000) and implemented in the program CODEML of the PAML 3.14 package (Yang 1997). Five different models that allow for different intensity of selection among sites (and deduced from the data) were tested. We compared the scenario where non-synonymous mutations are either neutral or deleterious (models M1 and M7, respectively) with models that allow for positive selection including an additional category for advantageous substitutions (models M2, M3 and M8). Three of the models assume a discrete distribution of the ω statistic (dN/dS) among sites: M1 (neutral) assumes two categories of sites conserved ($\omega=0$) and neutral ($\omega=1$); M2 (selection) includes an additional category of sites with ω estimated from the data; M3 (discrete) assumes a discrete distribution of K different ω ratios. Two additional models assume a continuous distribution for heterogeneous ω ratios among sites: M7 (beta) that assumes a beta distribution and does not allow for positively selected sites and M8 (beta and ω) that accounts for positively selected sites ($\omega>1$). Nested models can be compared in pairs using the likelihood ratio test (LRT): twice the log-likelihood difference is compared with a χ^2 distribution with degrees of freedom equal to the difference in the number of parameters between both models. The null model has a fixed $\omega=1$ while the alternative models have an estimate of ω as a free parameter. In this way, the more general models M2 and M3 can be tested against M1 and M8 against M7. Maximum likelihood trees to provide the phylogenetic information were constructed using DNAML from PHYLIP (Felsenstein 1989). A Bayesian approach implemented in CODEML was used to identify residues under positive selection in the $\alpha 1$ domain and sites with a posterior probability $>95\%$ were considered as positively selected under the model that best fitted the data.

We performed an analysis of sequence variability using a variability metric (V) (Reche and Reinherz 2003) that is formally similar to the Shannon entropy index (Shannon 1949) and that allows identification of variable amino acid residues. For a multiple protein sequence alignment the modified Shannon entropy (V) for every site follows the equation:

$$V = - \sum_{i=1}^M P_i \log_2 P_i$$

Where P_i is the fraction of residues of amino acid type i , and M is the number of amino acid types (20). Values of V range from 0 (only one residue in present at that position) to 4.322 (all 20 residues equally represented in that position). Amino acid sites with $V>1.0$ are considered variable, whereas those with $V<1$ are considered conserved. Variable amino acid residues estimated in this way may be functionally relevant in immune recognition through involvement in peptide contact (i.e. potential PBRs) (Stewart et al. 1997).

acid residues were polymorphic (21.1%). Only one nucleotide change was synonymous and every sequence therefore corresponded to a different allele.

Sequences from both populations were pooled in the rest of the analyses in order to determine the number of shared sequences and assess similarity between unique and common sequences.

Phylogenetic analysis

We performed phylogenetic analyses by building a neighbour-joining tree based on gamma distance (Fig. 2). Branch lengths were generally short, but three possible allelic lineages may be defined although they were supported by low bootstrap values. The newly identified Irish sequences cluster in one of the lineages (I) consisting of two sub-lineages (*Sasa-DAA*0301*, *Sasa-DAA*0302*, *Sasa-DAA*0303*, *Sasa-DAA*0304*, *Sasa-DAA*0305* and *Sasa-DAA*0601*, *Sasa-DAA*1201*, *Sasa-DAA*1202*) differing from the Norwegian sequences only by a few (2–5) base changes. The sequences absent in the Irish population (*Sasa-DAA*0801*, *Sasa-DAA*0901*, *Sasa-DAA*1101*), on the other hand, are distributed across the other three clusters.

Patterns of positive selection

Maximum likelihood models that allow positive selection fitted the data significantly better than those that assume only neutral or conserved mutations (Table 1). LRT tests suggested that model M2, which allows for positive selection, fitted the data better than model M1 (which considers only conserved and neutral sites) ($P < 0.001$). Estimates using the M2 model suggest that 20% of sites in the $\alpha 1$ domain were under strong positive selection ($\omega = 32.7$) and the rest of the sites were under purifying selection ($\omega = 0$). All variable sites were identified as positively selected by this model. Model M3, which assumes three site classes, fitted the data significantly better than M1 ($P < 0.001$) and M2 ($P < 0.01$). The results of model M3 suggested that 4.2% of the sites in the $\alpha 1$ domain were under strong positive selection ($\omega = 46.2$).

The LRT test comparing the two models that assume a beta distribution of ω over sites (M7 and M8) indicated that M8 (which allows for selection) fitted the data better than M7 (which does not allow selection) ($P < 0.001$). Estimates from M8 indicate that 16% of the sites are under strong positive selection in the sequences ($\omega = 44.8$).

Results from the different models indicate that there is variable selective pressure across sites of the MHC class II alpha sequences and the presence of a number of posi-

Fig. 2 Phylogenetic tree of $\alpha 1$ domain amino acid sequences of the MHC class II gene of Atlantic salmon for Irish and Norwegian populations based on gamma distance with $a = 1.53$. The reliability of the cluster analysis was tested by 1,000 bootstrap iterations and the results are shown in the nodes. Sequences in *bold* are unique to Irish populations and sequences in *italics* are unique to Norwegian populations

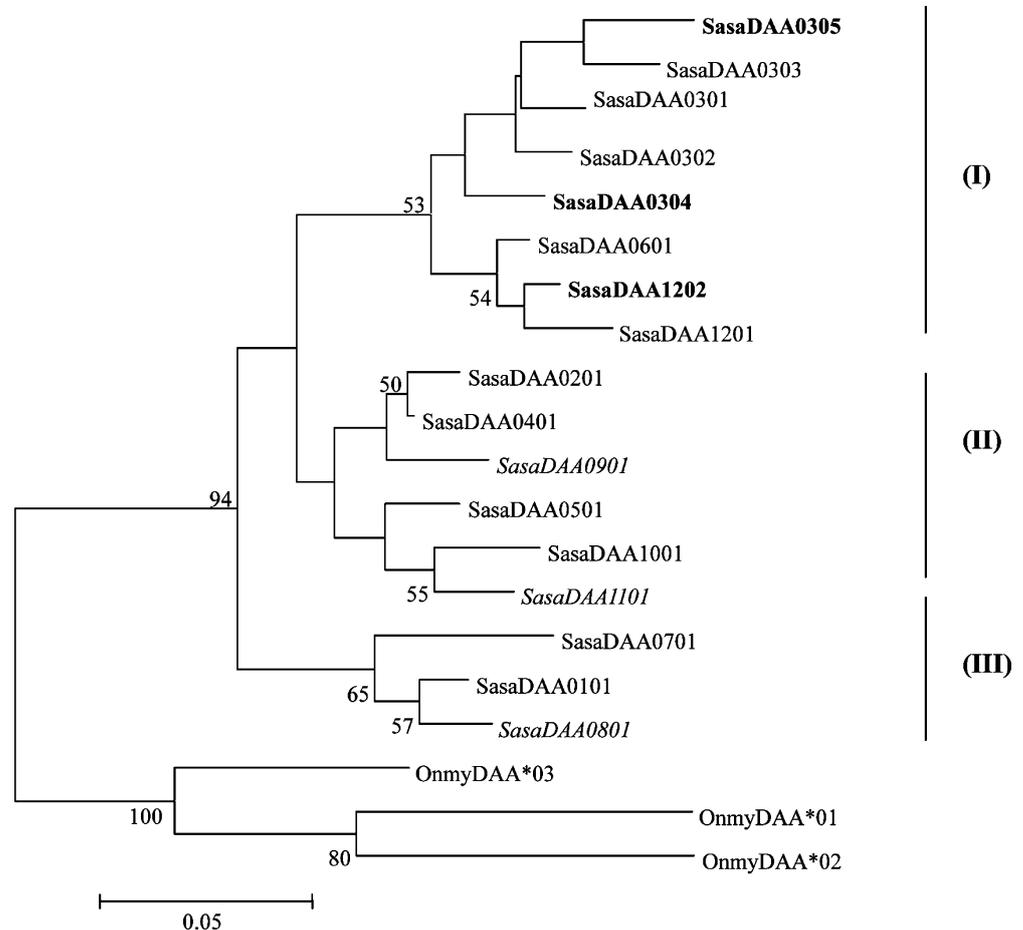


Table 1 Log-likelihood values (L) and parameter estimates (ω =non-synonymous/synonymous rates ratio estimates; p_i =fraction of the codons having as discrete ω or following a ω distribution, p & q =statistical parameters of the beta distribution) under random-sites

Model	L	Estimates of Parameters	Positively selected sites
M1	-566.74	$p_0=0.751$ ($\omega_0=0$) $p_1=0.249$ ($\omega_1=1$)	Not allowed
M2	-530.01	$p_0=0.793$ $p_1=0.000$ $p_2=0.207$ ($\omega_2=32.7$)	8L**, 9D**, 19W**, 26Q**, 30V**, 31A**, 41F**, 42P**, 44F**, 47Q**, 54V**, 57G**, 61K**, 62C**
M3	-525.05	$p_0=0.006$ ($\omega_0=0.341$) $p_1=0.831$ ($\omega_1=0.342$) $p_2=0.042$ ($\omega_2=46.2$)	8L**, 9D**, 26Q*, 30V**, 41F**, 42P**, 44F**, 47Q**, 54V**, 61K**, 62C**
M7	-567.08	$p=0.007$ $\omega=0.202$ $q=0.025$	Not allowed
M8	-525.98	$p_0=0.836$ $p_1=0.163$ ($\omega=44.8$) $p=49.697$ $q=99.000$	8L**, 9D**, 26Q*, 41F**, 42P**, 44F**, 47Q**, 54V**, 61K**, 62C**

models for the Irish and Norwegian class II MH alleles. Positively selected sites (* represents 95%, ** represents 99%) were identified by a Bayesian method implemented in CODEML. Amino acid positions refer to those in *Sasa-DAA*0101*

tively selected sites. The patterns of distribution of positively selected sites were consistent among models (M2, M3 and M8), although a larger number of sites was inferred under model M2 than the alternative models. The results from the M8 model, which were the most conservative, are presented in Fig. 3.

Following the criteria of Reche and Reinherz (2003) for HLA, four of the sites identified as positively selected were also identified as highly polymorphic ($V>1$) by the entropy analysis, and on this basis could be considered as potential PBRs. All other positively selected sites were classified as polymorphic ($V>0.5$) (Fig. 3).

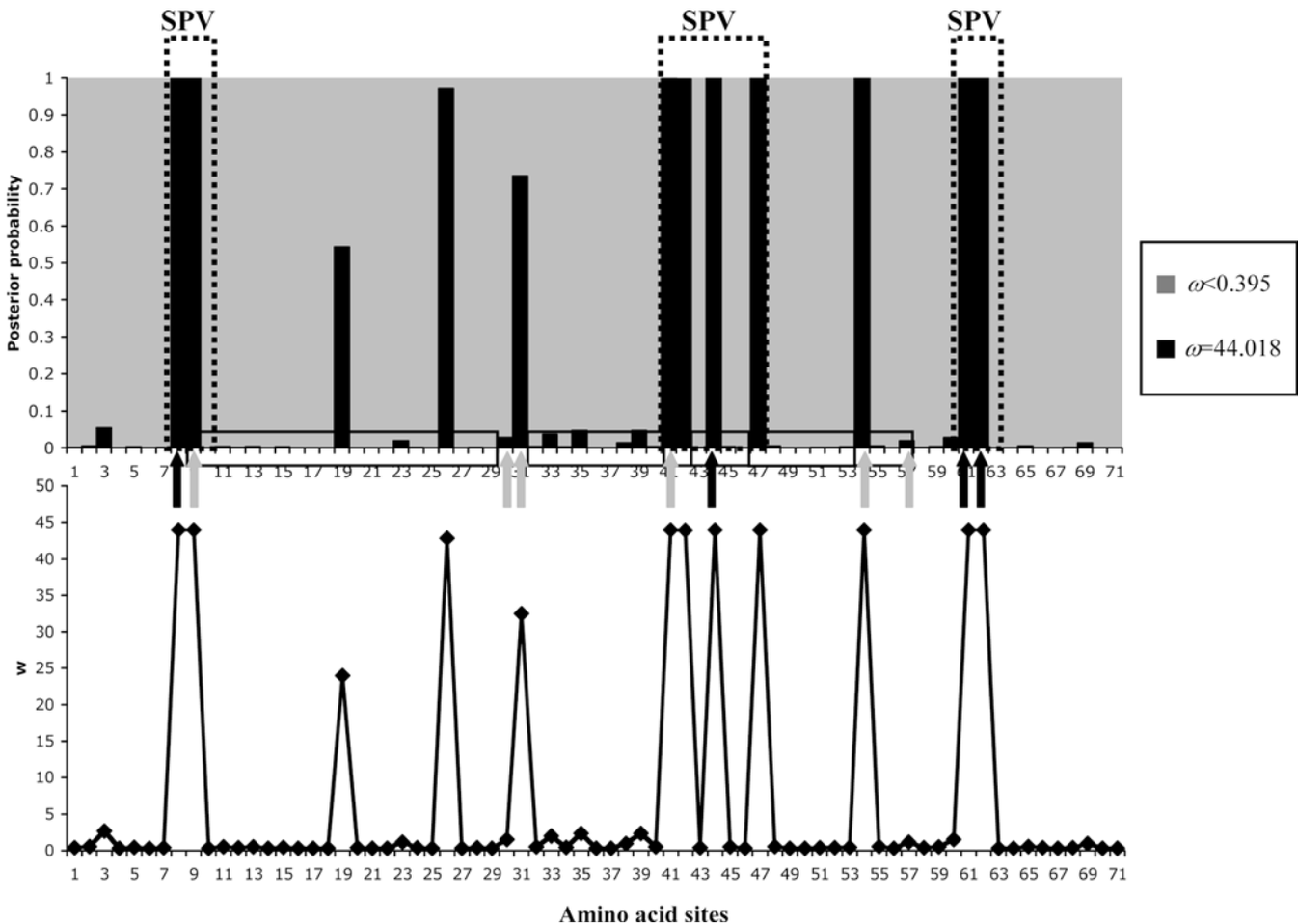


Fig. 3 Posterior probabilities of site classes for sites along the MH class II alpha 1 domain under the random-sites model M8 (beta & ω). Ten equal-probability categories were used to approximate the beta distribution (Yang et al. 2000), so that the model has 11 categories.

Posterior means of ω , calculated as the average of ω over the 11 site classes, weighted by the posterior probabilities. (◻) Sites involved in recombination, (◻◻◻◻) areas of SPV, polymorphic sites determined by their entropy values (◻) ($V>0.5$), (◻◻) ($V>1.0$)

Analysis of recombination

A test of alternative models of sequence evolution (Modeltest v3.06, Posada and Crandall 1998) indicated that the sequences followed a F81 model of nucleotide substitution with a gamma distribution shape parameter $\alpha=1.536$ and those parameters were used to run the analysis of spatial phylogenetic variation in PLATO that was carried out for the whole set of sequences described in both Norwegian and Irish. The result of the test indicated that three regions of SPV (23–28, 123–141 and 182–187; Fig. 3).

The results from the maximum likelihood recombination test suggested significant evidence of recombination ($P<0.001$) in the class II alpha sequences. Estimated recombination rate in the population ($\rho=4$ $N_e=96$) was higher than the mutation rate (Watterson estimate for population rate of mutation, $\theta=6.5$). Moreover, the results of the recombination analysis from DnaSP carried out including all sequences pooled revealed that at least six recombination events have occurred and involved regions between 3 and 27 nucleotides long (Fig. 3).

Most of the sites identified as positively selected fall into a region of spatial variable region coinciding in many cases with sites involved in recombination (Fig. 3). However, significant evidence of recombination was observed after repeating the maximum likelihood recombination test without sites identified as positively selected ($P<0.001$). After removing sites identified as positively selected, the estimated recombination rate in the population ($\rho=23$) remained higher than the mutation rate ($\theta=2.7$). Only two of the six recombination events initially detected by DnaSP remained after excluding selected sites (regions from positions 23–89 and 94–123).

Discussion

Population variability

Irish and Norwegian Atlantic salmon populations had substantially overlapping sets of alleles, differing in only a few class II alpha alleles (14 alleles described in each population, with only 3 of them private to each). Only a single locus was expressed in all individuals, as reported previously in Norwegian farmed Atlantic salmon (Stet et al. 2002). The 19 alleles described here, including three new sequences found only in Irish populations to date, were polymorphic and differed from one another by ~9 substitutions on average, similar to values found previously in class II *DAA* genes in farmed Atlantic salmon (Stet et al. 2002). The fragment of the exon 2 sequenced corresponds to the region where PBRs are potentially located and hence is likely to constitute the most variable region of the gene (Stern et al. 1994).

The phylogeny of the exon 2 sequences revealed three possible allelic lineages of the *Sasa-DAA* locus in Atlantic salmon, although these lineages were supported by low bootstrap values. Branch lengths in each lineage were short, with low genetic distances suggesting a recent origin of the

alleles at this locus (Shum et al. 2001; Stet et al. 2002). The class II alleles described in this study in both Norwegian and Irish populations have low divergence and all three new Irish alleles are located in one of the pre-existing clusters of shared alleles.

MH class I versus class II patterns of variability

Contrary to the observation in class I, where Irish and Norwegian populations had almost non-overlapping sets of alleles (Consuegra et al., submitted), they share the majority of class II *DAA* alleles (11 of 17 alleles). Although class I and class II molecules have similar tertiary structures, they differ in the class of peptides that they bind and the proteolytic routes for processing them (Kaufman et al. 1994; Castellino et al. 1997; Gromme and Neefjes 2002). They also differ in the class of T cells that they react with (Housset and Malissen 2003). Different modes of binding the peptide by class I and class II molecules, relatively strict in class I and more permissive in class II, may be responsible for functional differences between the two classes of MHC molecule, e.g differences in positive and negative selection of T cells (Huseby et al. 2003). The different nature of peptide binding may be one reason for the different evolutionary rates between class I and class II genes, as has been proposed to explain the higher turnover of the class I genes with respect to the class II genes in primates (Go et al. 2003). Under this model, stricter peptide binding by class I molecules would promote a higher rate of change in order to adapt to new infections (Go et al. 2003). However, the nature of class I peptide binding specificities has not been determined in fish (Grimholt et al. 2002) and our present results for Atlantic salmon contrast with the situation in the Lake Tana barbs species flock where closely related species have been shown to share class I alleles but differ in their class II alleles. The class II alleles are completely partitioned among the 10 different species studied without any sharing between species (Kruiswijk 2002).

Patterns of positive selection and recombination

Our results suggested that the observed diversity in the class II alpha sequences may be to a large extent generated by positive selection on PBRs. Maximum likelihood models allowing selection fitted the data significantly better than models that considered only neutral or conserved sites. The selected sites identified by Bayesian analysis coincided largely with sites that aligned with PBRs in human sequences (Stet et al. 2002) and which represent most of the variability among the sequences. Ten sites were identified with a strong signal of positive selection by model M8 (Table 1; Fig. 3) from which five correspond to potential PBRs (41F, 44F, 54V, 61K, 62C) and one to a conserved residue (42P) according to the comparison of Stet et al. (2002) with *HLA-DRA* sequences. Models M2 and M3 included all the sites identified by M8; M3 added one more site and model M2 identified three additional sites, one of

them potentially a PBR site (57G) according to Stet et al. 2002. The fact that sites identified as PBRs by crystallography in humans have been independently identified as positively selected in Atlantic salmon strongly suggests that they are implicated in peptide binding in this species.

The observed sequence diversity is probably not only the result of point mutations but is also generated by recombination events. The higher estimated recombination rate in relation to the mutation rate in the population suggests that recombination has been an important force in creating the present allelic diversity. Even after removing positively selected sites, results from the maximum likelihood tests showed evidence for recombination in the sequences.

Three of the four regions of SPV identified by PLATO overlapped with sites involved in recombination (as identified by DnaSP). Moreover, the regions identified as areas of SPV included in all cases sites identified as positively selected. Although the method implemented by PLATO cannot discriminate between recombination and selection as the cause for SPV (Grassly and Holmes 1997), the fact that after removing all positively selected sites evidence for recombination remained, indicate that at least part of the SPV regions detected correspond to recombination events.

Positively selected sites coincided in half the cases with the regions involved in recombination deduced by DnaSP. The fact that there is a non-random association between sites showing recombination and selection ($G\text{-test}=11.02$, $P=0.001$) suggests that alleles with recombinant PBRs have been positively selected, thus increasing variability, as suggested by Otha (1996). More positively selected sites (10) were detected than recombination events (6), although the latter cover most of the sequence (66%).

However, the coincidence of sites identified as both involved in recombination and under diversifying selection could also be due to the effect of recombination on the maximum-likelihood methods. Recombination can introduce stochasticity in the phylogenies (Schierup and Hein 2000) making it difficult to infer the number of ancestral allelic lineages. Current phylogeny-based models of codon substitution that include heterogeneous selective pressures across sites do not take the effects of recombination into account, although Bayesian methods for inferring positively selected sites seem to be little affected by recombination (Anisimova et al. 2003). Although the number of sequences analysed is not particularly large ($n=17$), and they are not very divergent (tree length $S=1.0$), the strength of selection estimated for positively selected sites, their coincidence with human PBRs and their independent identification as polymorphic sites ($I>1$) suggest that our results are robust to the effects of recombination. In theory, it is possible that some of the sites identified by the M2 model and not by the M8 model are the result of the confounding effects of the recombination in the phylogeny. However, we have taken that into account by using the M8 model results (apparently less affected by recombination; Anisimova et al. 2003) to compare the results of recombination and selection tests.

In common with the situation for class I sequences, recombination involving the putative PBR sites seems to play an important role in the origin of variability in class II

sequences. The existence of a large intron between $\alpha 1$ and $\alpha 2$ domains of the salmonid MH class I gene, where potential PBRs are located, may provide an additional mechanism for increasing variability through allele shuffling (Shum et al. 2001; Grimholt et al. 2002) compared to class II, where potential PBRs are only encoded in exon 2. In fact, the number of recombination events at the class II alpha locus was lower than that observed for the same populations for the class I locus, even when considering both class I domains independently. This may be the result of a more recent origin of class II alleles compared to those at the class I locus; class I alleles may simply have maintained the signature of old recombination events and accumulated more variation (Takahata and Satta 1998). Also, point mutation rates at the class II locus were lower than recombination rates, contrary to that the situation found at the class I locus.

In summary, recombination seems to play an important role in the origin of new alleles at the class II alpha locus of Atlantic salmon, both in Irish and Norwegian natural populations. A relatively large proportion of positively selected sites (potential PBRs) were found to be involved in recombination events. Although similar processes (selection and recombination) appear to be shaping variability at the class I and class II alpha loci in Atlantic salmon, the resulting distribution of variability across populations is very different for the two classes of MH loci.

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