Major histocompatibility complex (MHC) variation and susceptibility to the sea louse *Lepeophtheirus salmonis* in Atlantic salmon *Salmo salar*

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ABSTRACT: The relationship between genetic variation in major histocompatibility complex (MHC) Class I and II genes and susceptibility to sea lice *Lepeophtheirus salmonis* (Krøyer) in Atlantic salmon *Salmo salar* (L.) was studied in cage-reared post smolts. Polymorphic repeat markers located in the 3' untranslated regions (3UTR) of the genes Sasa-UBA (MHC Class I) and Sasa-DAA (MHC Class II) were screened in 1004 fish sampled from 11 full-sibling families. This gave rise to a total of 7 and 5 alleles, and 17 and 13 genotypes respectively. Significant relationships between both Sasa-UBA-3UTR and Sasa-DAA-3UTR genotypes and abundance of lice were observed within the pooled material, within individual families, and within the pooled material with both markers combined. However, most of these associations were either weak, linked with variation in fish size among genotypes, or influenced by family background genome. Nevertheless, within one family, the Sasa-DAA-3UTR 248/278 genotype displayed a significantly higher (33%) abundance of lice compared with the Sasa-DAA-3UTR 208/258 genotype, and this difference was not influenced by fish size. Consequently, the results of this study indicate a link between MHC Class II and susceptibility to lice.

KEY WORDS: *Lepeophtheirus salmonis* · Salmon · MHC · Susceptibility · Resistance

INTRODUCTION

The sea lice *Lepeophtheirus salmonis* (Krøyer) and *Caligus elongatus* (Nordmann) continue to represent a major challenge for the salmon farming industry. Atlantic salmon *Salmo salar* (L.) smolts placed into sea-cages often become infected with lice, and the farmer needs to control this infection level. Although a range of techniques are employed to control sea lice infections, including the use of cleaner wrasse (Deady et al. 1995, Tully et al. 1996, Treasurer 2002), the primary method involves the use of oral or bath-administered chemotherapeutics (Wootten et al. 1982, Roth et al. 1993, Mordue & Pike 2002).

Examples of sea lice displaying reduced sensitivity to chemotherapeutic agents have been documented (Jones et al. 1992, Treasurer et al. 2000, Tully & McFadden 2000, Sevatdal & Horsberg 2003, Fallang et al. 2004), and it is likely that this will become more extensive with continued widespread use of these agents. It is apparent that alternative sea lice control strategies are required for the future of salmon aquaculture. Vaccines (Roper et al. 1995, Raynard et al. 2002, Frost et al. 2006) in addition to selective breeding aimed to reduce the sensitivity of farmed salmon to sea lice infection (Mustafa & MacKinnon 1999, Glover et al. 2004a, 2005, Kolstad et al. 2005, Glover & Skaala 2006) are being considered as alternative control measures.

In an opportunistic study of genetic variation in the susceptibility of Atlantic salmon to *Caligus elongatus*, Mustafa & MacKinnon (1999) estimated heritability in susceptibility to this trait to be 0.22. In the first published estimate of susceptibility to *Lepeophtheirus salmonis*, Glover et al. (2005) estimated broad-sense heritability to be 0.07 (±0.02) among 30 full-sibling Atlantic salmon families reared in a single cage. The third estimate of susceptibility to sea lice in Atlantic agriculture.
salmon was published by Kolstad et al. (2005). These authors produced multiple estimates of heritability for total abundance of _L. salmonis_ in several year classes of Atlantic salmon, giving heritability values from 0.06 (±0.04) to 0.19 (±0.03), with a mean from 3 year classes of 0.14 (±0.02). Taken together, data from these studies indicate that there is sufficient genetic variation to decrease the susceptibility of farmed Atlantic salmon to sea lice through selective breeding. However, the estimated degree of heritability for this trait may be regarded as relatively low compared with many other traits currently selected for in salmon breeding (Gjedrem 2000).

If a selective breeding program for reducing the susceptibility of farmed Atlantic salmon to lice is to be successful, there is a need to be able to accurately select families and individuals that display decreased sensitivity to lice. In addition to genetic background at the population (Glover et al. 2001, 2003, 2004b, Glover & Skaala 2006) and family level (Mustafa & MacKinnon 1999, Glover et al. 2005, Kolstad et al. 2005), a range of known and unknown factors influence sea lice infections in fish, including fish size (Jaworski & Holm 1992, Todd et al. 2000, Glover et al. 2001, 2003, 2004a,b, Tucker et al. 2002, Genna et al. 2005, Glover & Skaala 2006), stress (Johnson & Albright 1992a) and a combination of light, salinity and host velocity (Genna et al. 2005). Consequently, it has been suggested that simply selecting individual salmon and families that display lower than average sea lice infections compared with the population in which selection is to be practised may not give an optimal response to selection (Glover et al. 2005, Glover & Skaala 2006). Furthermore, accurately and consistently counting lice infections on thousands of salmon in a commercial breeding program in which other traits are also selected for may present considerable practical challenges.

Infections with a parasite (from virus and bacteria to complex eukaryotic parasites) will typically induce a host response. This response varies depending on the host-parasite interaction (i.e. whether it is an intracellular, internal or external parasite) and commonly involves both the innate and acquired immune system. For ectoparasites, the immune response must be effecuated through the skin and other parts of the host (e.g. blood) that the parasites encounter. The major histocompatibility complex (MHC) system is an important part of the vertebrate immune system, and it has been shown that MHC allele diversity is important for resistance against parasites (e.g. Wegner et al. 2006).

In salmonids, significant associations between MHC genotype and susceptibility to bacterial (Langefors et al. 2001, Grimholt et al. 2003) and viral (Ozaki et al. 2001, Palti et al. 2001, Grimholt et al. 2003, Miller et al. 2004, Kjoglum et al. 2006) pathogens have been documented. Data from other fish species have indicated that MHC variation may be linked with susceptibility to parasites (e.g. Kurtz et al. 2004, Simkova et al. 2006, Wegner et al. 2006). Thus far, the potential link between genetic variation in the MHC in Atlantic salmon and susceptibility to sea lice has not been investigated. Consequently, the aim of the present study was to investigate the relationship between genetic variation in the MHC in Atlantic salmon and susceptibility to sea lice infection.

**MATERIALS AND METHODS**

**Fish and rearing.** The salmon used in this study were 7th generation domesticated fish originating from a Norwegian salmon breeding program. The breeding population was first established in the early 1970s and is at present controlled by Aqua Gen AS. This has been and continues to be one of the major farmed strains in Norway. Further details of the genetic origin of fish comprising this stock can be obtained from Gjedrem et al. (1991). Fifteen full-sibling families were established from the breeding stock in the autumn of 2002. Fertilised eggs were incubated, hatched and reared in single-family tanks. In June 2003, approximately 200 ind. from each family were tagged with passive integrated transponders (PIT) and transferred to a single freshwater tank (2 m³) for continued rearing. On 4 May 2004, these fish were transferred to a single 1200 m³ marine net-pen at a breeding station located at Hemne, central Norway. In the cage the fish were fed a commercial diet (Biomar) by hand according to standard feeding tables utilised by Aqua Gen AS.

**Monitoring of lice infection and sampling.** Starting 2 wk post transfer to salt-water, a sample of 10 salmon were sampled by a hand-net from the cage on a weekly basis to inspect fish for lice infection. These individuals were removed from the experiment.

The experiment was terminated between 27 September and 2 October, when there was close to 100% prevalence of lice on sampled fish and mean abundance of adult and pre-adult lice was >5. Termination involved removing small groups of fish (5 to 10) from the cage by wet net and placing them into a large white bucket containing 30 l water and an overdose of the anaesthetic benzocaine. Sedated fish were killed by a sharp blow to the head and placed into individual white buckets. This operation was performed in a quick and efficient manner such that potential for loss of lice in the anaesthetic bath was minimised. Using a similar sampling strategy, Glover et al. (2003) observed sampling-induced lice losses of 2.8 and 3.4%. Fish placed in white buckets were immediately taken into the laboratory and examined. Fish length, weight and
PIT-tag identification number were noted, and numbers of lice recorded. The white bucket in which the fish was transported was inspected for loss of lice, and these were removed and added to the individual’s lice count. Initial inspection of the fish revealed that the mobile adult and pre-adult stages represented over 95% of the population of lice on these fish at the time of sampling. Consequently, only numbers of Lepeophtheirus salmonis and Caligus elongatus were recorded; louse sex and developmental stage were not recorded. In total, lice were counted on 1342 salmon excluding a small number of individuals that were not included in the data set (15 fish), which had either fallen onto the floor when sampling or contained unreadable PIT tags.

**MHC genotyping.** A total of 1004 individual fish selected randomly from 11 of the 15 families were selected for genotyping. Genomic DNA from these individuals was isolated in 96 well-plate format using the standard protocol for the Qiagen DNA isolation kit. The polymorphic repeats located in the 3’ untranslated regions (3UTRs) of Sasa-UBA (MHC Class I) and Sasa-DAA (MHC Class II) genes (Grimholt et al. 2002, Stet et al. 2002) were amplified using the fluorescently labelled sense primers 5’-GGAGAGCTGCCACGATGACCT-3’ and 5’-GATGGCAAAGAGGAAAGTGAG-3’ and the reverse primers 5’-CAATTACCACAAGGCCGCTC-3’ and 5’-TTGTTATGCTCTACTCTGAA-3’. The PCR conditions were 50 ng genomic DNA per 10 µl total reaction volume for 25 cycles at 56°C annealing temperature. Markers were analysed using automated ABI 377 machines (Applied Biosystems).

**Statistics.** All statistical analysis was performed in the program STATISTICA version 7.0 (StatSoft). Families were compared for infection levels and mean weight using ANOVA and ANCOVA where continuous variables were included in the statistical design. Significance tests were followed by Tukey’s post hoc test for unequal N. Correlation was used to describe potential relationships between fish size and infection level within each family, within the pooled material including all families, and using mean family values.

Abundance of lice for the Sasa-UBA-3UTR and Sasa-DAA-3UTR genotypes was analysed for each marker separately within families and within pooled families. These data were analysed by ANCOVA, where individual fish weight was implemented as the continuous predictor. Significant tests were investigated further by Tukey’s post hoc test for unequal N. In order to investigate the potential effect of variation within the 2 markers simultaneously on louse infection, composite genotypes (Sasa-UBA-3UTR–Sasa-DAA-3UTR) were established for all individuals. Composite genotypes represented by 20 or more fish within the pooled material were compared with each other (in terms of abundance of lice) by ANCOVA. This analysis was restricted to pooled family data. In families displaying significant relationships between genotype and abundance of lice, G-tests were used to test for random sampling of lice.

## RESULTS

### General infection data

A total of 1342 fish originating from 15 full-sibling families were sampled for lice. At the time of sampling, Caligus elongatus displayed a very low abundance on the experimental fish and only 45 parasites were recorded in the entire material. Consequently, data from *C. elongatus* were excluded from statistical analysis. The summary statistics of the infection data for Lepeophtheirus salmonis are presented in Table 1. Significant differences in mean weight (ANOVA, $F_{14,1327} = 33.5$, $p < 0.0001$) and abundance of *L. salmonis* (ANOVA, $F_{14,1327} = 9.3$, $p < 0.0001$) were observed among the families. Mean weight varied by a factor of 2.47 among the families, whilst mean *L. salmonis* abundance varied by a factor of 1.36. Out of 105 pairwise post hoc comparisons for mean family abundance of *L. salmonis*, 25 were significant, indicating that a number of families were responsible for the observed trend. Within families, abundance of *L. salmonis* among individual salmon varied greatly. The range in abundance of *L. salmonis* varied from between 11 and

<table>
<thead>
<tr>
<th>Family</th>
<th>N</th>
<th>Weight g (SE)</th>
<th>Mean louse abundance (SE)</th>
<th>Median louse abundance</th>
<th>Range</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>87</td>
<td>461.8 (18.4)</td>
<td>19.9 (0.65)</td>
<td>20.0</td>
<td>7–42</td>
</tr>
<tr>
<td>2</td>
<td>106</td>
<td>766.9 (16.7)</td>
<td>23.5 (0.59)</td>
<td>24.0</td>
<td>7–45</td>
</tr>
<tr>
<td>3</td>
<td>82</td>
<td>518.6 (19.0)</td>
<td>19.6 (0.67)</td>
<td>19.0</td>
<td>4–31</td>
</tr>
<tr>
<td>4</td>
<td>113</td>
<td>446.9 (16.2)</td>
<td>19.2 (0.57)</td>
<td>19.0</td>
<td>7–38</td>
</tr>
<tr>
<td>5</td>
<td>76</td>
<td>371.7 (19.7)</td>
<td>19.9 (0.70)</td>
<td>18.0</td>
<td>6–56</td>
</tr>
<tr>
<td>6</td>
<td>122</td>
<td>451.3 (15.6)</td>
<td>18.5 (0.55)</td>
<td>18.5</td>
<td>7–44</td>
</tr>
<tr>
<td>7</td>
<td>64</td>
<td>309.9 (21.5)</td>
<td>19.1 (0.75)</td>
<td>18.0</td>
<td>9–32</td>
</tr>
<tr>
<td>8</td>
<td>92</td>
<td>441.4 (18.0)</td>
<td>17.5 (0.63)</td>
<td>17.0</td>
<td>6–53</td>
</tr>
<tr>
<td>9</td>
<td>94</td>
<td>557.7 (17.7)</td>
<td>23.3 (0.62)</td>
<td>23.0</td>
<td>9–47</td>
</tr>
<tr>
<td>10</td>
<td>51</td>
<td>423.9 (24.1)</td>
<td>21.4 (0.84)</td>
<td>21.0</td>
<td>9–35</td>
</tr>
<tr>
<td>11</td>
<td>97</td>
<td>502.1 (17.5)</td>
<td>22.1 (0.61)</td>
<td>22.0</td>
<td>9–41</td>
</tr>
<tr>
<td>12</td>
<td>108</td>
<td>411.9 (16.5)</td>
<td>20.4 (0.58)</td>
<td>21.0</td>
<td>9–39</td>
</tr>
<tr>
<td>13</td>
<td>88</td>
<td>461.6 (18.3)</td>
<td>20.6 (0.64)</td>
<td>20.0</td>
<td>11–35</td>
</tr>
<tr>
<td>14</td>
<td>104</td>
<td>401.6 (16.9)</td>
<td>17.3 (0.59)</td>
<td>17.0</td>
<td>4–52</td>
</tr>
<tr>
<td>15</td>
<td>58</td>
<td>467.6 (22.6)</td>
<td>21.3 (0.79)</td>
<td>21.0</td>
<td>9–33</td>
</tr>
</tbody>
</table>
35 within the family displaying the least variation to between 4 and 52 in the family displaying the greatest variation.

Fish size is an important factor influencing infection level with sea lice (Jaworski & Holm 1992, Todd et al. 2000, Glover et al. 2001, 2003, 2004a,b, Tucker et al. 2002, Glover & Skaala 2006). Within the pooled material, including data from all 15 families, a weak but nevertheless significant positive relationship between individual fish weight and abundance of Lepeophtheirus salmonis was observed ($R^2 = 0.07, N = 1342, p < 0.0001$). In addition, a significant positive relationship between mean family weight and mean family abundance of $L. salmonis$ was observed ($R^2 = 0.43, N = 15, p < 0.0001$) (Fig. 1). The relationship between individual fish size and abundance of $L. salmonis$ was investigated within each of the 15 families separately. Correlations ranged from $R^2 = 0.0002 (p = 0.89)$ to $R^2 = 0.19 (p < 0.0001)$, with 7 of the 15 tests giving significant p-values with 95% confidence. When the significance level was adjusted for multiple independent tests (15 tests, new significance level $p = 0.003$), 4 of the 15 tests were still significant, indicating that some families displayed significant relationships between fish weight and abundance of $L. salmonis$. All significant relationships were positive, i.e. larger fish displayed a higher abundance of $L. salmonis$.

Despite variable relationships between individual fish size and abundance of Lepeophtheirus salmonis at the individual, family, and pooled levels, the observed variation in mean $L. salmonis$ abundance among families was still significant when the analyses were performed with fish weight as a continuous predictor (ANCOVA, family effect: $F_{14,1326} = 5.9, p < 0.0001$; however, weight had a more significant influence on the infection level (ANCOVA, individual weight: $F_{1,1326} = 45.4, p < 0.0001$).

Genotyping results

A random selection of individuals ($N = 1004$) from 11 of the 15 families were chosen for genotyping. Not all individuals produced readable genotypes for both Sasa-UBA-3UTR and Sasa-DAA-3UTR. Consequently, the numbers of individuals analysed for either of these 2 markers are lower than the total number of individuals selected for genotyping. These individuals include samples that failed to produce readable genotypes after one set of re-runs and were thus excluded from the analyses. Numbers of individual fish analysed per family for each marker ranged from 72 to 107.

For Sasa-UBA-3UTR, a total of 7 alleles ranging from 314 to 336 bp were detected within the 11 families, giving 17 genotypes across families. For Sasa-DAA-3UTR, 5 alleles were detected in the range from 208 to 278 bp, giving a total of 13 genotypes across families. Fish were sorted into genotypes both within families and among families in order to investigate potential effects of MHC genotype on abundance of Lepeophtheirus salmonis.

Genotype vs. abundance of Lepeophtheirus salmonis: families pooled

The relationship between genotype vs. fish weight and genotype vs. abundance of Lepeophtheirus salmonis for both markers are presented in Fig. 2. Significant differences in abundance of lice among genotypes for Sasa-UBA-3UTR were observed (ANCOVA, $F_{16} = 2.2$, $p = 0.0046$); however, weight explained a large proportion of the variation ($F_{1} = 63.8, p < 0.0001$). Pair-wise post hoc tests for abundance of $L. salmonis$ revealed that all significant pair-wise tests involved the homozygous genotype 318/318. This genotype displayed a significantly higher abundance of $L. salmonis$ in 9 of the 16 pair-wise tests against other genotypes. No other genotype pairs were significantly different from each other. The 318/318 genotype was only present in Family 2. Coincidently, this family (Table 1) and genotype displayed the highest average weight of all families and genotypes (Fig. 2).

Significant differences in abundance of lice for the Sasa-DAA-3UTR genotype was observed in the pooled material (ANCOVA, $F_{12} = 2.1$, $p = 0.016$); however, similar to the results for Sasa-UBA-3UTR, weight was of greater significance ($F_{1} = 90.0, p < 0.0001$) (Fig. 2). In
this instance, none of the post hoc pair-wise tests for abundance of lice were significant. For both Sasa-UBA-3UTR and Sasa-DAA-3UTR, a pattern suggesting a link between average weight for fish displaying a given genotype and abundance of *Lepeophtheirus salmonis* for that genotype is evident (Fig. 2).

**Genotype vs. abundance of *Lepeophtheirus salmonis*: within families**

Significant relationships between Sasa-UBA-3UTR or Sasa-DAA-3UTR genotype and abundance of lice were observed in 3 of the 11 families studied (Table 2). In Family 3, a significant relationship between abundance of *Lepeophtheirus salmonis* and genotype was observed for Sasa-UBA-3UTR (ANCOVA, $F_{\text{Genotype}} = 6.9$, $p = 0.01$, $F_{\text{Weight}} = 11.6$, $p = 0.001$). A significant relationship between the abundance of *L. salmonis* and the Sasa-DAA-3UTR genotype was also observed in Family 3 (ANCOVA, $F_{\text{Genotype}} = 3.2$, $p = 0.027$, $F_{\text{Weight}} = 11.4$, $p = 0.001$). Pair-wise post hoc tests indicated significant differences in abundance of lice between genotypes 208/228 and 208/248 ($p = 0.045$) and between genotypes 208/248 and 228/258 ($p = 0.003$). In Family 4, a significant relationship between the Sasa-UBA-3UTR genotype and abundance of *L. salmonis* was observed (ANCOVA, $F_{\text{Genotype}} = 3.1$, $p = 0.029$, $F_{\text{Weight}} = 2.0$, $p = 0.16$). Pair-wise post hoc tests indicated significant differences in abundance of lice between genotypes 322/336 and 314/336 ($p = 0.04$) and between genotypes 316/322 and 322/336 ($p = 0.03$). In Family 6, a significant relationship between the Sasa-DAA-3UTR genotype and abundance of *L. salmonis* was observed (ANCOVA, $F_{\text{Genotype}} = 4.5$, $p = 0.005$, $F_{\text{Weight}} = 16.7$, $p < 0.0001$). Pair-wise post hoc tests indicated that the genotypes 208/258 and 248/278 were significantly different ($p = 0.004$). In
all 4 instances where significant trends within family were observed, a G-test was applied to test for random sampling of alleles within each family. Results of all G-tests demonstrated random sampling (all p-values > 0.1).

With the exception of the observation in Family 6, all genotypes displaying significantly lower lice abundances as demonstrated though pair-wise post hoc tests were also smaller in average weight compared with the genotypes from which they differed (Table 2). In Family 6, no differences in mean weight were observed between the pair of genotypes 208/258 and 248/278 (Table 2).

Table 2. *Salmo salar* and *Lepeophtheirus salmonis*. Families in which significant associations between sea lice abundance and genotype were detected for Sasa-UBA-3UTR or Sasa-DAA-3UTR

<table>
<thead>
<tr>
<th>Family</th>
<th>Marker</th>
<th>Genotype</th>
<th>Abundance of sea lice N (SD)</th>
<th>Mean weight g (SD)</th>
<th>Observed no. of fish</th>
</tr>
</thead>
<tbody>
<tr>
<td>3</td>
<td>Sasa-UBA-3UTR</td>
<td>322/334</td>
<td>21.4 (0.8) 548.8 (25.7)</td>
<td>39</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>334/334</td>
<td>17.5 (0.9) 474.2 (26.7)</td>
<td>36</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>Sasa-DAA-3UTR</td>
<td>208/228</td>
<td>19.9 (1.2) 590.9 (35.9)</td>
<td>19</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>208/248</td>
<td>15.2 (1.3) 419.3 (41.9)</td>
<td>14</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>228/258</td>
<td>21.7 (1.0) 534.6 (32.7)</td>
<td>23</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>248/258</td>
<td>18.8 (1.3) 481.6 (39.2)</td>
<td>16</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>Sasa-UBA-3UTR</td>
<td>314/316</td>
<td>19.8 (1.2) 402.4 (30.5)</td>
<td>24</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>314/336</td>
<td>20.5 (1.1) 484.8 (27.8)</td>
<td>29</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>316/322</td>
<td>20.9 (1.3) 488.6 (33.5)</td>
<td>20</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>322/336</td>
<td>16.0 (1.2) 416.7 (31.2)</td>
<td>23</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>Sasa-DAA-3UTR</td>
<td>208/248</td>
<td>19.0 (1.0) 463.3 (18.3)</td>
<td>31</td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td>208/258</td>
<td>16.3 (1.1) 450.3 (20.8)</td>
<td>24</td>
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<td>21.7 (1.2) 455.0 (22.8)</td>
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</tr>
<tr>
<td></td>
<td></td>
<td>258/278</td>
<td>17.9 (0.9) 446.6 (17.5)</td>
<td>34</td>
<td></td>
</tr>
</tbody>
</table>

In the pooled material, a total of 12 composite genotypes (Sasa-UBA-3UTR–Sasa-DAA-3UTR) were exhibited in over 20 individuals. Only these composite genotypes were analysed in the pooled material for the potential relationship between composite genotype and susceptibility to *Lepeophtheirus salmonis*. Significant overall differences in abundances of *L. salmonis* were observed among the 12 composite genotypes (ANCOVA, F = 2.1, p = 0.022); however, fish weight determined a large proportion of the observed variation (F = 34.3, p < 0.0001) (Fig. 3). Post hoc tests revealed that the composite Genotype 6 (322/334–258/278) displayed a significantly lower abundance of *L. salmonis* than did composite Genotype 8 (322/334–208/258), but no other pair-wise comparisons were significant. Investigating further, the composite Genotype 322/324–258/278 consisted of fish from Families 2 (n = 7, mean abundance = 21.3), 8 (n = 9, mean abundance = 13.3) and 14 (n = 8, mean abundance 18.0), whilst composite Genotype 322/334–208/258 consisted of fish from Families 8 (n = 2, mean abundance = 17.0), 9 (n = 7, mean abundance = 23.4) and 11 (n = 17, mean abundance = 23.8). Clearly, the low abundance of *L. salmonis* observed on the 9 fish from Family 8 was responsible for creating the difference in abundance of lice between the composite genotypes.

DISCUSSION

To our knowledge, this study represents the first investigation of MHC variation and susceptibility to the sea louse *Lepeophtheirus salmonis*. Differences in abundance of *L. salmonis* were observed among genotypes for the polymorphic markers Sasa-UBA-3UTR (MHC Class I) and Sasa-DAA-3UTR (MHC Class II). Differences were observed within individual families, within the pooled material, and among composite genotypes within the pooled material. However, the majority of the observed trends were linked with variation in mean fish size between genotypes. Nevertheless, a relationship between the Sasa-DAA-3UTR genotype and abundance of lice within Family 6 was observed, and this was not influenced by fish size or family background genetic variation. Consequently, these data provide evidence suggesting that MHC is linked with susceptibility to *L. salmonis*. However, the extent and significance of this involvement, and whether or not this represents direct or indirect involvement, is difficult to conclude from this study.

The link between the polymorphic repeat marker and coding region, in particular that for Sasa-UBA, is unpredictable and varies among strains and populations. A recombination signal exists in the intron between the α1 and α2 domains so that α1 domains are shifted between different α2 domains and downstream sequences, explaining part of this instability (Shum et al. 2001). However, recombination within the marker itself has been observed, with one Sasa-UBA allele having different markers in 2 different populations (authors’ unpubl. data). Previous studies also showed that 1 marker may also be linked to multiple alleles, as for instance the SasaUBA-3UTR 322 bp marker, which represented the alleles UBA*0201, UBA*0301, UBA*0401, UBA*0801 and UBA*1201 (Grimholt et al. 2003). There is no knowledge on linkage between marker and coding region for the group of fish in the present study. The SasaDAA-3UTR alleles 208, 228, 248, 258, 278, 288 bp have previously been linked with the α-β haplotypes DAA*0201/DAB*0201, DAA*0101/DAB*0801, DAA*0301/DAB*0401, DAA*0401/DAB*0701, DAA*0501/DAB*0301 and DAA*0601/DAB*0601, respectively. Whether or not this stability exists within the present population is unknown; however, if it did, then the genotype displaying lowest abundance of lice in Family 6 would be DAA*0201/DAB*0201–DAA*0401/DAB*0701. In addition, it is not possible to exclude the possibility that the families challenged displayed different but ‘functionally similar’ alleles within these markers by chance. Consequently, the possibility cannot be excluded that a stronger link between MHC and abundance of lice would be observed by performing the analysis based on coding region, or by performing this study on a group of fish displaying different or greater diversity of alleles (such as a wild population).

Associations between allelic variation in the MHC in Atlantic salmon and susceptibility to bacterial (Langefors et al. 2001, Grimholt et al. 2003) and viral (Ozaki et al. 2001, Palti et al. 2001, Grimholt et al. 2003, Miller et al. 2004, Kjoglum et al. 2006) pathogens have been reported. In addition, a statistical relationship between severity of amoebic Gill disease (AGD) and MHC variation has been reported in Atlantic salmon (Wynne et al. 2007). The present experiment was similar in design to the study elucidating the link between MHC variation and susceptibility to both furunculosis and infectious salmon anemia (ISA) (Grimholt et al. 2003), and the link between MHC variation and susceptibility to AGD (Wynne et al. 2007). However, within the present study, a fewer number of families and larger number of fish representing each family were prioritised for screening. This design was chosen owing to the fact that the within-family component of variation in abundance of *Lepeophtheirus salmonis* is much greater than the between-family abundance of *L. salmonis* (Glover et al. 2005, present study), in addition to the fact that the within-family analysis is less influenced by background family genetic variation.

In order to verify the influence of the Sasa-DAA-3UTR genotype on the abundance of *Lepeophtheirus salmonis*, further experimentation is required. Verification of MHC involvement could be achieved by a strategy similar to that used to verify specific MHC Class I and II allele combinations affecting resistance to ISA in Atlantic salmon (Kjoglum et al. 2006) or, alternatively, by testing within multiple families that display the Sasa-DAA-3UTR 208/258 and 248/278 heterozygotes.

For furunculosis, the heritability of susceptibility (see Gjedrem 2000) is considerably higher than that reported for sea lice (Mustafa & MacKinnon 1999, Glover et al. 2005, Kolstad et al. 2005) where non-genetic effects dominate. Clearly, finding a link between potential genetic markers and susceptibility to *Lepeophtheirus salmonis* presents a more challenging task than it would in the case of a pathogen that displays a higher heritability of susceptibility and where the phenotype is more closely linked with the genotype. However, fundamental differences in host-pathogen relationships exist between external parasites such as sea lice, which are largely protected from the host’s immune system, and bacterial and viral pathogens that enter the organism or its cells. A recent study of AGD in Atlantic salmon demonstrated a link between MHC variation and severity of infection (Wynne et al. 2007). However, similar to the results of the present study, the associations observed by these
authors were weak. Nevertheless, the fact that a link between the MHC and parasite susceptibility has been observed in several fish species (e.g. Kurtz et al. 2004, Simkova et al. 2006) supports the idea that the MHC has the potential to be linked with susceptibility to *L. salmonis*.

Despite the fact that specific antibodies to *Lepeophtheirus salmonis* have been observed in Atlantic salmon (Grayson et al. 1991, 1995), the acquired immune response to sea lice is apparently weak. This is illustrated by the fact that Atlantic salmon are easily reinfected with high levels of sea lice (e.g. Glover et al. 2004a). In an experimental study of *L. salmonis* infection in coho salmon, Johnson & Albright (1992a) observed a significantly higher infection level in immune-suppressed fish compared with control fish, and that this difference was primarily mediated by the apparent suppression of inflammatory response and the development of epithelial hyperplasia. Likewise, in a comparison of susceptibility to *L. salmonis* among 3 salmonid species, Johnson & Albright (1992b) demonstrated that Atlantic salmon displayed a significantly greater infection level of *L. salmonis* compared with both coho and Chinook *Onchorhynchus tshawytscha* salmon, and that these differences were primarily related to the lack of host-tissue responses in Atlantic salmon compared with the other species.

In repeated challenge experiments (Glover et al. 2004a, Glover & Skaala 2006), individual salmon displaying lower than average infection levels in a first challenge (i.e. individuals identified as ‘resistant’) did not necessarily display a lower than average infection level in a second or third challenge. Consequently, unidentified factors in addition to fish size appear to be dominating in determining individual lice abundance levels in challenge tests in both cages and tanks. Clearly, it is not possible to exclude the possibility that the results of the present study may be stock specific. However, a combination of the low observed level of MHC involvement in *Lepeophtheirus salmonis* abundance, the fact that heritability of lice abundance is low or at best moderate (Mustafa & MacKinnon 1999, Glover et al. 2005, Kolstad et al. 2005), and that Atlantic salmon are regarded as highly susceptible to *L. salmonis*, it can be argued that it is unlikely that the MHC has any major effect on susceptibility to sea lice. This requires further study.

In summary, differences in abundance of *Lepeophtheirus salmonis* were observed among genotypes for the polymorphic repeat markers located in the 3UTR of the Sasa-UBA and Sasa-DAA genes. When associations caused by the link with fish size were excluded, the only clear evidence of interaction between MHC and abundance of *L. salmonis* was observed in a single family for Sasa-DAA-3UTR. It is clearly difficult to indicate the significance of this result, and further study is required to categorise the extent and nature of involvement of the MHC in susceptibility to sea lice.

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Glover et al.: MHC variation and susceptibility to L. salmonis

65


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