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Evaluating Adaptive Divergence Between Migratory and Nonmigratory Ecotypes of a Salmonid Fish, <i>Oncorhynchus mykiss</i>

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ABSTRACT Next-generation sequencing and the application of population genomic and association approaches have made it possible to detect selection and unravel the genetic basis to variable phenotypic traits. The use of these two approaches in parallel is especially attractive in nonmodel organisms that lack a sequenced and annotated genome, but only works well when population structure is not confounded with the phenotype of interest. Herein, we use population genomics in a nonmodel fish species, rainbow trout (<i>Oncorhynchus mykiss</i>), to better understand adaptive divergence between migratory and nonmigratory ecotypes and to further our understanding about the genetic basis of migration. Restriction site-associated DNA (RAD) tag sequencing was used to identify single-nucleotide polymorphisms (SNPs) in migrant and resident <i>O. mykiss</i> from two systems, one in Alaska and the other in Oregon. A total of 7920 and 6755 SNPs met filtering criteria in the Alaska and Oregon data sets, respectively. Population genetic tests determined that 1423 SNPs were candidates for selection when loci were compared between resident and migrant samples. Previous linkage mapping studies that used RAD DNA tag SNPs were available to determine the position of 1990 markers. Several significant SNPs are located in genome regions that contain quantitative trait loci for migratory-related traits, reinforcing the importance of these regions in the genetic basis of migration/residency. Annotation of genome regions linked to significant SNPs revealed genes involved in processes known to be important in migration (such as osmoregulatory function). This study adds to our growing knowledge on adaptive divergence between migratory and nonmigratory ecotypes of this species; across studies, this complex trait appears to be controlled by many loci of small effect, with some in common, but many loci not shared between populations studied.

KEYWORDS smoltification life history variation genomics SNP salmonids

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Migration, the long-distance movement of organisms to take advantage of seasonal resources, has long interested both scientists and lay people alike. The cues that initiate the cascade of physiological, behavioral, and morphological changes that prepare an individual for migration are varied and complex (Dingle 2006). These cues interact with signals from the environment, such as temperature and food availability, that in concert influence the decision of an individual to migrate (Dingle 2006; Wilcove and Wikelski 2009). Heritability studies across taxonomic groups suggest that both propensity and timing of migration have a genetic component (see Pulido and Berthold 2003 for review), but we have limited insight into the genes and molecular mechanisms involved (Dingle 1991; Liedvogel et al. 2011). This
paucity of information limits our understanding of the evolution of migration and prevents us from asking such questions as: are the same genetic mechanisms involved in migration conserved across different taxonomic groups? Do the genes that predispose an individual to migrate get inherited as a migratory gene package? And what patterns do we see in genes associated with migration in closely related species that differ in their migratory strategies (Pulido and Berthold 2003; reviewed in Liedvogel et al. 2011)?

The salmonid fishes (salmon, trout and char) are an exemplary taxonomic group to study the genetic basis of migration. Multiple species have both migratory, ocean-going forms and resident individuals that stay in fresh water throughout their life cycle. *Onchorhyncus mykiss* is an especially attractive taxon as it exhibits both life history types, often within the same river. Previous studies in controlled breeding designs have identified quantitative trait loci (QTL) for migration-related traits in this species (Nichols et al. 2008; Hecht et al. 2012; Le Bras et al. 2011) and other salmonids (Norman et al. 2011) confirming a genetic basis to migratory related traits. In addition, genome-wide association approaches (GWAS) have been used and, like QTL studies, have pointed to many different regions of the genome being associated with migration (Hecht et al. 2013). However, both approaches have their limitations, especially when applied in natural systems. For example, QTL studies are limited to surveying genotype-phenotype variation in progeny derived from a very small subset of individuals sampled within or between population(s). Such sampling captures a limited amount of the total genetic variation that underlies the trait(s) of interest. Meanwhile, association analysis approaches suffer from high false-positive rates due to underlying population structure and/or kin relatedness (e.g., Price et al. 2006; Ingvarsson and Street 2011). These false-positive results can be corrected for accounting for population structure; however, such methods can both induce false-negative results (where true small effect associations are removed) where population structure is highly correlated with phenotype. The latter problem may be of special concern in salmonids because resident populations can be land locked by physical barriers that restrict gene flow with neighboring migratory populations (Limborg et al. 2012). Population genomic methods offer an alternative approach in the identification of genes that have adaptive significance. The basic premise of such methods is that allele frequencies at selected sites will differ from neutral sites. Recently such methods have been used to highlight regions of the genome under selection for specific traits (Dalziel et al. 2009; Nielsen et al. 2009; Hohenlohe et al. 2010). For example, genomic scans in marine and freshwater three-spined sticklebacks (*Gasterosteus aculeatus*) both confirmed, and discovered new loci showing signatures of selection between the two phenotypes (Hohenlohe et al. 2010). Popular approaches to finding loci under selection can be broadly split into “empirical” and “theoretical” approaches. Examples of empirical approaches include *F*_ST outlier methods that aim to locate outlier loci among many neutral loci (Beaumont and Nichols 1996; Luikart et al. 2003), whereas examples of theoretical tests include statistics such as Tajima’s D (e.g., Nielsen 2005; Nielsen et al. 2007) that test for departures from neutrality.

The advent of genotype-by-sequencing methods has made it possible to identify a large number of polymorphisms throughout a genome at a relatively low cost, and in doing so, has made genome-wide population genomic and association tests possible in nonmodel organisms (Baird et al. 2008; Miller et al. 2007; references in Seeb et al. 2011). The use of restriction site–associated DNA (RAD) tag sequencing is especially attractive because tens of thousands of SNPs can be found relatively quickly and cheaply (reviewed in Rowe et al. 2011 and Davey et al. 2011). For example, RAD tag sequencing has made it possible to construct linkage maps (Amores et al. 2011; Baxter et al. 2011), conduct population genomic studies (Hohenlohe et al. 2010), perform QTL analyses (Chuttimanitsakun et al. 2011; Miller et al. 2012; Pfender et al. 2011; Hecht et al. 2012), and perform GWAS analysis (Hecht et al. 2013) in organisms that previously did not have extensive SNP resources.

In this study, we used Illumina RAD tag methods in two populations of *O. mykiss* to identify genetic loci associated with migration and residency using population genomic approaches to question whether the same markers are divergent between migratory and resident ecotypes in each geographic location (i.e., common mechanisms). In the absence of a fully sequenced and annotated genome, this study illustrates the ability to conduct whole-genome studies on an ecologically and evolutionarily important trait with the aid of high-throughput genomic sequencing.

**MATERIALS AND METHODS**

**Sampling and DNA extraction**

Resident and migratory individuals were sampled from each of two populations: Sashin Creek, Alaska and Little Sheep Creek, Oregon. Hereafter, the migrant steelhead trout form of *O. mykiss* will be referred to as migrant trout and the resident rainbow trout form as resident trout. Upon returning from the ocean to their natal streams to spawn, all migrants were sampled at sexual maturity. Resident trout from the Sashin Creek, Alaska population also were sampled at sexual maturity (tested for the expression of gametes). The Sashin Creek residents are separated from returning migrants by two barrier waterfalls. All migrants sampled were downstream of these waterfalls, whereas all residents sampled were upstream (see Thrower et al. 2004 for more details). The migrant and resident individuals in the Little Sheep Creek population are not separated by barrier waterfalls (see Bernson et al. 2011 for more details) and so samples of both phenotypes were taken from the same area. Resident trout from the Little Sheep Creek, Oregon, population were determined to be resident as determined by size threshold (minimum fork length of 160 mm). Both river catchments have active hatchery programs; to limit effects of selection due to hatchery programs, only wild fish were sampled (upon release, all hatchery fish have the adipose fin removed). At sampling, the fork length (tip of snout to the fork of the caudal fin) and sex of the fish was recorded, and a portion of the caudal fin was removed and stored in ethanol for DNA extraction. The fish were immediately released after processing. DNA was extracted for a total of 386 fish (105 Sashin Creek migrants, 90 Sashin Creek residents, 100 Little Sheep Creek migrants, and 91 Little Sheep Creek residents) using either standard phenol:chloroform procedures (Wasko et al. 2003) or QIAGEN DNeasy Tissue Extraction kits (QIAGEN Inc., Valencia, CA). All Sashin Creek residents were sampled in 2008 and 2010, and Sashin Creek migrants were sampled in all years from 2006 to 2010. Little Sheep Creek samples for both migrants and residents were collected in 2011.

**RAD library construction**

A total of 500 ng of DNA was used from each sample (*n* = 386 in total) to construct RAD tag libraries for sequencing as described in Miller et al. (2012). To summarize, DNA from each individual was digested with *SfiI* and then barcoded with a unique 6-base sequence 3’ to the *SfiI* cut site. Barcoded DNA from 15 samples was pooled and fragmented using the Sonic Ruptor 400 (Omni International, Kennesaw, GA). Fragments were size selected between 300 and
Bioinformatics

Sequenced libraries were processed as described by Miller et al. (2012) using Perl scripts and Novocraft software (scripts are available as supplemental material in Miller et al. (2012)). To summarize, reads were trimmed to 89 bp to remove barcodes and the SbfI overhang and then quality filtered. Reads were assembled de novo to construct a database for identifying SNPs. The SNP database was constructed from 20 individuals: 10 from Sashin Creek (5 migrants and 5 residents) and 10 from Little Sheep Creek (5 migrants and 5 residents). To find loci variable both within and between populations, SNPs were found by identifying polymorphisms in three separate comparisons of these individuals: one constructed from all 20 fish, one from the 10 fish from Little Sheep Creek, and one from the 10 fish from Sashin Creek. Unique alignments of identical sequences with fewer than five reads and more than 200 reads were discarded. Retained alignments were compared for SNPs, which were tentatively separated into two alleles at the same locus. Candidate SNP loci required a minimum depth of coverage of five reads for both alleles. This procedure was repeated for all three databases, and the resulting SNP files for each database were combined to create a reference SNP database, removing redundant loci.

Three genotype datasets were created: (1) all fish from both populations (n = 386 individuals); (2) only fish from Sashin Creek, Alaska (n = 195); and (3) only fish from Little Sheep Creek, Oregon (n = 191). All individuals were genotyped against the database using Bowtie and custom Perl scripts (Miller et al., 2012), which report the number of reads for alleles A and B at each SNP. Genotypes were scored only if a minimum total read count of 8 was met. The log_{10} ratio of the number of reads aligning to allele A/ allele B was used to separate homozygotes and heterozygotes. Heterozygotes were determined if the log_{10} ratio was between −0.61 and 0.61 (one SD around the mean) with AA homozygotes scored if the ratio was greater than 0.9 and BB homozygotes if the ratio was less than −0.9 (as in Hecht et al., 2012). Reads that fell outside these thresholds were removed.

SNP filtering

Salmonids have experienced a relatively recent whole-genome duplication (between 25 and 100 million years ago; Allendorf and Thorgaard 1984), which can lead to the erroneous identification of candidate SNPs from the alignment of paralogs that are very nearly identical (Everett et al., 2011; Miller et al., 2012). Paralogous sequence variants (PSVs) are false SNPs that should be removed from the data set. As also described by Miller et al. (2012), we performed RAD tag sequencing on two doubled haploid *O. mykiss* that should have no heterozygous positions within any one individual genome; any sequence producing a heterozygous genotype in doubled haploid *O. mykiss* is identified as a PSV and were removed. Candidate PSVs have also been shown to produce an excess of heterozygous genotypes (M. Everett, unpublished data; Hohenlohe et al., 2011). Therefore, any SNP that produced an observed number of heterozygous genotypes greater than 90% was removed. SNPs were further filtered by minimum minor allele frequency <2%, that were missing >50% genotypes and were out of Hardy Weinberg Equilibrium (−log_{10} P = > 5.0) as measured in each dataset. The combined dataset also was filtered to only include markers that were polymorphic in both the Sashin Creek and Little Sheep Creek populations.

Genomic ordering and annotation of SNPs

To provide information on the genomic location of SNPs in this study, polymorphic RAD tags were aligned to two recent *O. mykiss* linkage maps produced from RAD tag SNPs (Miller et al., 2012; Hecht et al., 2012) and a draft version of the *O. mykiss* genome, using Bowtie. In both cases, RAD tags were considered matches if they only produced one significant hit (allowing 2 mismatches). The draft version of the *O. mykiss* genome consists of sequenced scaffolds some of which have been linked to the physical contigs sequenced at the Clemson University Genomics Institute center (Palti et al., 2011). To annotate the genomic regions linked to the top significant SNPs in this study, each scaffold associated with a filtered RAD tag was annotated using BLASTn (minimum e-value = 1 × 10^{−20}, minimum of 85 bp of continuous alignment (tags are 89 bp after quality-filter). Identity was confirmed if the tag only produced a significant hit (e-value < 1.0 × 10^{−20}, at least 85 bp of continuous alignment) to one scaffold. Scaffolds with significant hits were then annotated using BLASTn against the nr database and BLASTx against the zebrafish genome (http://useast.ensembl.org/info/data/ftp/index.html) and a compiled annotated salmonid EST database [downloaded from cGRASP (web.uvic.ca/grasp/) and NAGRP (www.csrees.usda.gov/nea/animals/in_focus/an_breeding_if.nagrp.html); date of download March 27, 2012]. Minimum requirements for annotation was a contiguous alignment of 300 bp, with 85% sequence similarity and a minimum e-value of 1 × 10^{−20}. As only approximately one quarter of the SNPs were mapped we will report both the number of mapped SNPs and the total number of SNPs that were significant for any statistical test.

Population genomic analyses

Individual global and pairwise (between residents and migrants) F_{ST} was estimated for each locus in GenAlEx 6.4 (Peakall and Smouse 2006), as was mean expected (H_{E}) and observed (H_{O}) heterozygosity and the number of private alleles. To identify outlier markers that may be candidates for selection, LOSITAN (Beaumont and Nichols 1996; Antao et al., 2008) was used with filtered loci from Sashin Creek and Little Sheep Creek with the following parameters: 55,000 simulations, confidence interval of 0.995, and a false discovery rate of 0.05. Simulated F_{ST} was 0.032 for Sashin Creek and 0.001 for Little Sheep Creek, and estimated F_{ST} was 0.069 and 0.003 for the two populations, respectively. We used loci with a probability greater than 0.995 to infer candidates for positive selection and neutral loci were defined as having a probability between 0.9 and 0.1. Tajima’s D was calculated to estimate the allele frequency spectrum within each of the filtered RAD tags. PoPoolations (version 2.1; Kofler et al., 2011) was used to estimate within population Tajima’s D by pooling 50 samples from each of the four subpopulations (Sashin Creek and Little Sheep Creek, migrants and residents) and using a window size of 100 base pairs, a minimum rare allele count of 2, a minimum depth of coverage of 4 and a population size estimate of 50, the same parameters were used to estimate nucleotide diversity (Watterson’s theta; Watterson 1975). To generate a smooth, genome-wide distribution of these statistics, a kernel smoothing average using a Gaussian function as described in Hohenlohe et al. (2010). This approach was used only on markers
with mapping information. Linkage disequilibrium (LD) was calculated as $D'$ (Lewontin 1964) between all pairs of SNPs that were matched to loci in Miller et al. (2012). The chromosome and distance between markers was inferred from Miller et al. (2012). Tests were run separately for each population, and average $D'$ per chromosome was calculated. A two-tailed $t$-test was used to test the hypothesis that average chromosome-wide $D'$ was different between the two populations. $D'$ calculations were performed in JMP Genomics (SAS Inc., Cary, NC).

**RESULTS**

**SNP filtering**

Twenty-six lanes of sequencing were conducted using various Illumina instruments (see Supporting Information, file S1). The average number of quality-filtered reads per individual was 3.17 million ($\pm$1.26 million (SEM)). Illumina reads for each individual (quality filtered and trimmed reads) are deposited at Data Dryad (doi:10.5061/dryad.c6fb2). A total of 30,642 unique candidate biallelic SNPs were identified from 26,459 RAD tags (22,590 tags had a single SNP, 2796 tags had 2 SNPs, and 820 tags had 3 SNPs). Removing SNPs that were heterozygous in either of the sequenced doubled haploids or that showed an observed heterozygosity $\geq 90\%$ reduced the total number of SNPs to 28,863. Removing SNPs with $>50\%$ missing genotypes, minor allele frequency $<0.02$, and markers out of Hardy-Weinberg equilibrium (and only including markers that were polymorphic in both populations for the combined dataset) further reduced the number of SNPs to 7920, 6755, and 5185 for the Sashin Creek, Little Sheep Creek, and the combined datasets, respectively. RAD tags were compared with Miller et al. (2012) and Hecht et al. (2012a), and positive matches were named the same with new SNPs following the same nomenclature (see File S2).

**Genomic ordering and annotation of SNPs**

A total of 1990 (25% of total) filtered SNPs were matched to two RAD tag–based genetic linkage maps (Miller et al. 2012; Hecht et al. 2012). This number does not include SNPs that matched multiple tags in Miller et al. (2012). In this way, synteny between this study and previous linkage and QTL mapping studies in *O. mykiss* could be inferred from shared microsatellite and SNP markers. In *O. mykiss*, linkage groups have further been matched to physical chromosomes by fluorescence *in situ* hybridization (Phillips et al. 2006).

**Population genomics between phenotypes**

$F_{ST}$ between migrants and residents revealed different patterns of variation in the two populations. In Sashin Creek, mean $F_{ST}$ between migrants and residents was 0.062 with a range from 0 to 0.459. Mean $F_{ST}$ between migrants and residents was much lower in the Little Sheep Creek population (0.003) and ranged from 0 to 0.218. The global comparison between migrants and residents (using both populations) produced a mean $F_{ST}$ of 0.075, with values ranging from 0 to 0.510. In $F_{ST}$ outlier analyses from LOSITAN, candidates for balancing selection could not be distinguished from loci with an $F_{ST}$ of zero and so were not considered further. LOSITAN identified 772, 61, and 55 loci ($P > 0.995$) as candidates for positive selection between residents and migrants in the Sashin Creek, Little Sheep Creek, and global datasets, respectively. Fourteen loci were significant in both the Sashin Creek and Little Sheep Creek samples, and four loci were in common between the global and Sashin Creek datasets. LOSITAN also was used to identify neutral loci, which were defined as those with probabilities between 0.9 and 0.1 (as in Hess et al. 2013). We found 5752, 6047, and 3801 loci that fell within this range in the Sashin Creek, Little Sheep Creek, and global populations, respectively. Mapping information was obtained for 178 outlier loci in the Sashin Creek dataset, with at least one significant outlier locus on every chromosome (see Figure 1A). Kernel smoothing showed an increase in the pairwise $F_{ST}$ on Omy2, Omy7, and Omy8. A total of 11 loci mapped to eight chromosomes in the Little Sheep Creek analysis of which Omy4 (4) and Omy23 (2) contained multiple outliers. However, kernel smoothing failed to find any chromosomes that seemed to show a significant enrichment for outlier loci (Figure 1B).

The average Tajima’s $D$ statistics for the Sashin Creek migrant population was $-0.417$ (ranging from $-4.003$ to $3.902$) with 742 loci that differed significantly from neutral expectations. Of these, 109 showed a positive departure from neutrality, or loci that showed an excess of common polymorphisms compared with neutral expectations, whereas 633 loci showed a negative departure from neutrality. These results strongly suggest an excess of low-frequency polymorphisms suggesting population expansion and/or strong effects of purifying selection. Mapping information was obtained for 169 loci that different from neutrality. Of these, 28 had a significant positive value, and 141 had a significantly negative value. Loci with positive Tajima’s $D$ values were distributed on 16 chromosomes, whereas all chromosomes had at least one locus with a significant negative $D$ value. Kernel smoothing approaches in Sashin Creek migrants suggest Omy2, Omy7, Omy22, and the sex chromosome contain peaks of negative $D$ values, whereas Omy20 and Omy22 contain peaks of positive $D$ values; note, however, that the magnitude of the positive peaks is smaller than the negative peaks (see Figure 2A). Also note that the majority of the distribution is toward a negative Tajima’s $D$. The Sashin Creek resident population produced an average Tajima’s $D$ of $-0.003$ and ranged from $-3.294$ to 0.798. Only five loci departed from neutrality, all of which produced a negative $D$ value and all of which mapped. The kernel smoothing approach showed the distribution of Tajima’s $D$ values was roughly around 0, suggesting neutrality (data not shown) in the resident Sashin Creek population. We combined data from both the migrant and resident individuals within the Little Sheep Creek population as Tajima’s $D$ is most powerful at the population level (Tajima 1989) and principle component analysis suggested strongly that both phenotypes form one population (see below). Average Tajima’s $D$ was $-0.145$ with values ranging from $-3.36$ to 4.13 with 296 markers departing from neutrality (269 negative Tajima’s $D$ and 27 positive Tajima’s $D$). Mapping information was obtained for 54 loci with a significant Tajima’s $D$ value. Of these, 52 mapped loci had a significant negative value and two a significant positive value. The negative markers were distributed on every chromosome except Omy13, Omy21, Omy10, Omy24, and Omy26. The two positive loci were on Omy2 and Omy16. Kernel smoothing suggested that Omy12 contained a peak of negative loci with Omy8 and Omy23 smaller peaks. Only Omy2 seemed to show a positive peak (Figure 2B). For details of all significant loci from population genomic analyses see File S3.

Patterns of observed heterozygosity varied across the genome. The Sashin Creek migrants showed an average observed heterozygosity of 0.229, with peaks of heterozygosity on Omy6, Omy8 and Omy24 and lows on Omy3 and Omy5 (Figure 3A). The Sashin Creek residents had an average observed heterozygosity of 0.246, and appeared to show more variation between chromosomes with peaks of heterozygosity on Omy2, Omy8, Omy12, Omy20, and
Figure 1 $F_{ST}$ for markers that were mapped to two linkage maps of the *O. mykiss* genome (Miller et al. 2012; Hecht et al. 2012). Markers in red represent significant outliers as determined by LOSITAN (for details, see the section Materials and Methods), black line represents the kernel smoothed distribution of all markers (both outliers and “neutral” loci). Chromosomes are marked by alternate shading with some of the chromosomes labeled. (A) $F_{ST}$ between Sashin Creek residents and Sashin Creek migrants and (B) $F_{ST}$ between Little Sheep Creek residents and Little Sheep Creek migrants.
Omy22 (Figure 3B). The Little Sheep Creek samples has an average observed heterozygosity of 0.252 and showed little variation between chromosomes with only Omy27 showing an increase in the observed heterozygosity and no chromosomes showing a large reduction in heterozygosity (see Figure 3C). The Sashin Creek migrants produced an average nucleotide diversity (Watterson’s theta) of 0.008, the Sashin Creek residents 0.0001 and the Little Sheep Creek samples 0.003. Kernel smoothing methods showed that the Sashin Creek migrants have greater-than-average nucleotide diversity on Omy4, Omy6, and Omy10 and reduced levels on Omy2, Omy5, Omy7, Omy12, and Omy24 (Figure 4A). The Sashin Creek residents showed a near 0 nucleotide diversity that did not show much variation between chromosomes (data not shown). The Little Sheep Creek samples showed a generally higher diversity on Omy8, Omy13, and Omy23 and a reduction on Omy10, Omy5 and the sex chromosome (Figure 4B).

Figure 2 Tajima’s D results for markers that mapped to two linkage maps of the O. mykiss genome (Miller et al. 2012; Hecht et al. 2012). Markers in blue produced a significant positive Tajima’s D and markers in red a significantly negative Tajima’s D. The black line represents the kernel-smoothed average Tajima’s D for all mapped markers. Chromosomes are alternately shaded with a subset labeled. (A) Tajima’s D calculated within the Sashin migrants, (B) within the Little Sheep Creek samples (combined migrants and residents).
Figure 3 The kernel-smoothed average observed heterozygosity in markers, which mapped to two linkage maps of the *O. mykiss* genome (Miller et al. 2012; Hecht et al. 2012). The kernel-smoothed average for all markers that mapped is presented for the (A) Sashin migrants, (B) Sashin residents, and (C) Little Sheep Creek samples. Chromosomes are alternately shaded.
Figure 4  Nucleotide diversity as calculated by Watterson’s theta in markers that mapped to two linkage maps of the O. mykiss genome (Miller et al. 2012; Hecht et al. 2012). The kernel-smoothed average for all markers that mapped is presented for (A) Sashin Creek migrants and (B) Little Sheep Creek samples. Chromosomes are alternately shaded with some chromosomes labeled.
We compared the private allele density (alleles that are unique to one population) between the resident and migrant populations in Sashin Creek. A total of 988 loci had private alleles of which 981 had alleles that were unique to the migrant population and 7 loci had alleles that were unique to the resident population. For Tajima’s D estimates in the Little Sheep Creek population the migrant and resident phenotypes were combined as they are one population. A total of 52 loci had private alleles of which 23 were unique to the migrant phenotype and 19 were unique to the resident phenotype.

**Linkage disequilibrium**

From the 1990 filtered SNPs that were confidently matched to a previous linkage map (Miller et al. 2012), average chromosome-wide $D'$ ranged from 0.659 ± 0.013 in the Sashin Creek fish to 0.603 ± 0.009 in the Little Sheep Creek population. The greater LD observed in the Sashin Creek is significantly different than that observed in Little Sheep Creek ($P < 0.001$). Average chromosome-wide $D'$ varied between 0.54 for Omy28 and 0.82 for Omy26 in samples from Sashin Creek, and between 0.49 for Omy21 and 0.68 for Omy25 and Omy22 in Little Sheep Creek fish.

**BLAST analysis**

A total of 221 RAD tags that contained both SNPs that produced a significant statistic from population genetic analyses, and mapping information from previous linkage maps (Miller et al. 2012; Hecht et al. 2012), were BLASTed to scaffolds from a draft version of the *O. mykiss* genome. Of which, 101 RAD tags produced a unique match to one scaffold (e ≤ 1.0^-20; see File S4). Of these 64 scaffolds produced significant matches to genes with annotation (see Table 1 for results of BLASTn analysis and Table 2 for results of BLASTx analysis).

**DISCUSSION**

Migration is a central behavior in the ecology and evolution of many organisms. Both migration and dispersal have a heritable component in animals as diverse as crickets (Roff and Fairbairn 2007) and birds (Hansson et al. 2003; Pasinelli et al. 2004). In salmonids, the heritability of migratory-related traits appears to be high (e.g., proportion of brood smolting in *O. mykiss*; $h^2=0.726$; Thrower et al. 2004 and smolt timing in Atlantic Salmon, *Salmo salar*; Páez et al. 2011), strongly suggesting some level of underlying genetic control. To determine the genetic basis to migration, the authors of previous studies have either taken a “bottom-up” approach (i.e., investigating candidate genes for variation associated with migration) or a “top-down” approach such as by constructing genetic crosses that vary in a phenotype associated with migration. In this study, we used population genomics methods to identify loci showing unique patterns of divergence between migratory and nonmigratory ecotypes in natural populations of *O. mykiss*, supporting “top-down” approaches as a tractable way to detect loci under selection in natural populations.

| SNP ID      | Chr | Scaffold               | BLASTn Hit        | e-Value |
|-------------|-----|-----------------------|-------------------|---------|
| R22186      | 1   | MSSRT009C_scaffold_1734_1 | LDA gene for MHC class I antigen, allele: | 0       |
| R33240      | 1   | MSSRT018H_scaffold_1292_1 | GH2 growth hormone 2 gene | 0       |
| R10206      | 1   | MSSRT043A_scaffold_1632_1 | Cytochrome P450 family 1 subfamily B polypeptide 1 (cyt1b1) | 0       |
| R51388      | 1   | MSSRT138E_scaffold_1568_1 | Cyclin D1 (ccnd1) | 0       |
| R34273      | 2   | MSSRT001A_scaffold_1294_1 | Malate dehydrogenase | 1E-90   |
| R41089      | 3   | MSSRT092A_scaffold_2616_1 | CD83 (Omy-CD83) gene | 0       |
| R29125      | 5   | MSSRT037B_scaffold_1504_1 | Olfactory receptor family C subfamily 4 member 5 gene | 0       |
| R49596      | 5   | MSSRT081B_scaffold_1813_1 | Dax-1 (Dax1) gene, complete cds | 0       |
| R08400      | 7   | MSSRT085C_scaffold_1375_1 | Alpha-globin and beta-globin, clone 3 | 0       |
| R19618      | 7   | MSSRT095B_scaffold_2418_3 | (ADP-ribose) polymerase family | 0       |
| R00565      | 7   | MSSRT119G_scaffold_1110_1 | Adiponectin receptor protein 1 (adr1) | 0       |
| R00141      | 9   | MSSRT005H_scaffold_1052_1 | MHC class I a region | 0       |
| R13477      | 9   | MSSRT043B_scaffold_2167_1 | MHC class I antigen (Sasa-UDA) gene | 0       |
| R17494      | 9   | MSSRT043G_scaffold_1780_1 | Dnaj homolog subfamily C member 5 | 0       |
| R15095      | 11  | MSSRT043C_scaffold_1956_1 | Myosin regulatory light chain 2 | 3E-129  |
| R21238      | 12  | MSSRT087F_scaffold_1654_1 | S toll-like receptor 8a2 (TLR8a2) | 0       |
| R35677      | 12  | MSSRT107C_scaffold_1855_1 | ADP-riboisylated factor 1 | 0       |
| R20262      | 13  | MSSRT013E_scaffold_1784_1 | Immunoglobulin heavy chain (igd-A) gene | 0       |
| R19667      | 14  | MSSRT001A_scaffold_1328_1 | Delta-6 fatty acyl desaturase defad_a gene | 0       |
| R40426      | 14  | MSSRT056B_scaffold_1710_1 | MHC class I a region | 0       |
| R50712      | 14  | MSSRT084F_scaffold_1265_2 | Tapasin-B (TAPBP) | 0       |
| R07628      | 17  | MSSRT033D_scaffold_1722_1 | Inhibitor of differentiation 1C (ID1C) gene | 0       |
| R14298      | 17  | MSSRT108H_scaffold_1570_1 | GH1 interferon alpha 1-like gene | 0       |
| R50966      | 18  | MSSRT045C_scaffold_1546_1 | Myostatin 2b (MSTN2) | 0       |
| R32416      | 18  | MSSRT049G_scaffold_1774_1 | Na,K-transporter alpha subunit isoform 1b/i (ATP1A1B/i) | 0       |
| R28876      | 19  | MSSRT009E_scaffold_1603_1 | Steroidogenic acute regulatory protein (star) gene | 0       |
| R30814      | 20  | MSSRT031B_scaffold_1485_1 | GH1 interferon alpha 1-like gene | 0       |
| R30448      | 20  | MSSRT098B_scaffold_1661_1 | Gonadotropin subunit beta-2 (gthb2) | 0       |
| R27417      | 21  | MSSRT083G_scaffold_1866_1 | Chaperonin gene, complete cds | 0       |
| R40451      | 23  | MSSRT001B_scaffold_1711_1 | Homeobox protein hoxd13aa (hoxd13aa) | 0       |
| R00446      | 23  | MSSRT058G_scaffold_1786_1 | Growth hormone 2 gene, complete cds | 0       |
| R31852      | 25  | MSSRT097D_scaffold_1534_1 | Potassium-transporting atpase subunit beta-m (at1b4) | 0       |
| R04223      | sex | MSSRT014E_scaffold_2046_1 | Carbonyl reductase/20beta-hydroxysteroid dehydrogenase B | 0       |
| R37330      | sex | MSSRT111B_scaffold_1390_1 | Polyunsaturated fatty acid elongase (elvol5a) gene | 0       |
Table 2 BLASTx hits for scaffolds that contain SNPs that produced an outlier locus in $F_{ST}$ analysis or a significant departure from neutrality

| SNP ID | Chr | Scaff            | GB             | Hit                          | e-Value |
|--------|-----|------------------|----------------|------------------------------|---------|
| R24286 | 1   | MMSRT043A_scaff_1632_1 | NP_001167620.1 | Cytochrome P450, family 1, subfamily B, polypeptide 1 | 0       |
| R39129 | 2   | MMSRT081F_scaff_1690_1 | ACQ58433.1     | C-C chemokine receptor type 9 | 1E-123  |
| R44099 | 5   | MMSRT068D_scaff_2079_1 | NP_001167113.1 | Sulfate transporter          | 0       |
| R27510 | 5   | MMSRT074G_scaff_1637_1 | NP_001104690.1 | Growth arrest-specific 1a precursor | 9E-79   |
| R41103 | 5   | MMSRT133E_scaff_1870_1 | NP_001177233.1 | Hemicentin-1                 | 4E-68   |
| R10656 | 8   | MMSRT111H_scaff_1511_1 | NP_001100745.1 | Fucosyltransferase 9         | 3E-146  |
| R00710 | 9   | MMSRT030E_scaff_2197_1 | ACN10547.1     | Leucine-rich repeat-containing protein 8D | 0       |
| R35771 | 10  | MMSRT055B_scaff_1860_1 | XP_003455229.1 | Alpha-2 adrenergic receptor  | 0       |
| R18859 | 10  | MMSRT122H_scaff_1570_1 | ATP-sensitive inward rectifier potassium channel 15   | 1E-163  |
| R08544 | 11  | MMSRT057C_scaff_1604_1 | XP_003451939.1 | Olfactory receptor 51B2      | 4E-84   |
| R03392 | 12  | MMSRT097H_scaff_1700_1 | ABC43375.1     | Odorant receptor              | 1E-73   |
| R19667 | 14  | MMSRT001A_scaff_1328_1 | XP_001192036.2 | Netrin-1-like                 | 1E-82   |
| R28018 | 15  | MMSRT079E_scaff_1499_1 | XP_003448843.1 | Potassium voltage-gated channel subfamily D | 0       |
| R15666 | 16  | MMSRT106A_scaff_1950_1 | XP_003977854.1 | Olfactomedin-like protein 3B  | 4E-94   |
| R05928 | 17  | MMSRT061F_scaff_1802_1 | XP_004070492.1 | Neurogenic differentiation factor 4 | 3E-63   |
| R24186 | 17  | MMSRT125C_scaff_1898_1 | CBN80811.1     | Sodium voltage-gated channel  | 0       |
| R03565 | 18  | MMSRT049G_scaff_1774_1 | ADD60471.1     | Na+/K+ atpase alpha           | 6E-135  |
| R41273 | 19  | MMSRT141G_scaff_1929_1 | NP_001018411.1 | Fidgetin                     | 5E-146  |
| R01482 | 19  | MMSRT050F_scaff_1759_1 | ACS35075.1     | Thrombospondin-1a            | 1E-117  |
| R12769 | 20  | MMSRT062C_scaff_2490_1 | XP_004084735.1 | Metabotropic glutamate receptor 1 | 5E-109  |
| R03286 | 20  | MMSRT124H_scaff_1869_1 | XP_003404464.1 | Dystonin-like isoform 2       | 2E-148  |
| R30464 | 23  | MMSRT038D_scaff_2090_1 | NP_00157024.1  | Forkhead box protein A2       | 1E-153  |
| R00544 | 25  | MMSRT062C_scaff_2440_1 | XP_003448092.1 | Prolactin-18                  | 3E-71   |
| R14689 | 25  | MMSRT110B_scaff_3626_2 | ADYS118.1      | Ryanodine receptor 3 [Opsanichthyis bidens] | 1E-134  |
| R50870 | 25  | MMSRT038D_scaff_1820_1 | XP_003222987.1 | Cadherin-8                    | 2E-71   |
| R20892 | 26  | MMSRT029B_scaff_2020_1 | XP_003763770.1 | Transcription factor SOX-14    | 4E-79   |
| R05730 | sex| MMSRT014A_scaff_2072_2 | AA63316.1      | Adrenergic, alpha-2D, receptor b | 8E-173  |
| R34426 | sex| MMSRT022B_scaff_1619_1 | NP_003452446.1 | Sodium/calcium exchanger 1    | 9E-173  |
| R03937 | sex| MMSRT041G_scaff_1450_1 | NP_001121839.1 | Prostaglandin E2 receptor EP4 subtype | 1E-100 |
| R46658 | sex| MMSRT072E_scaff_1452_1 | EKIC18115.1    | Sperm-associated antigen 16    | 0       |

By using an unassembled scaffold version of the O. mykiss genome, we used kernel smoothing average methods to identify regions enriched for markers that deviated from the rest of the genome. The Sashin Creek population produced a large number of markers that suggest differentiation between migrants and residents in both population genetic and association approaches. The Sashin Creek $F_{ST}$ comparisons suggest loci on Omy2 and Omy7 show the greatest amount of differentiation between residents and migrants. A negative peak on Omy2 is also found in the Tajima’s $D$ analysis within the Sashin Creek migrants, suggesting either the effects of purifying selection or rapid population expansion after a bottleneck. The historical population size of the Sashin Creek migrants is unknown and of late the number of returning fish is decreasing (F. P. Thrower, unpublished data), suggesting negative departures from neutrality in the Sashin Creek migrants are caused by purifying selection. A reduction in the nucleotide diversity is also found in the Sashin Creek migrants on Omy2, suggesting reduced variation in this region of the genome, possibly due to purifying selection for alleles involved in anadromy.

By contrast the Little Sheep Creek samples did not show strong differentiation between the migrants and the residents on Omy2. There was a departure from neutrality in the positive direction on Omy2 (the opposite from the general trend) in the Tajima’s $D$ results, suggesting positive selection or population contraction. However, it is likely that selection on this locus is not connected to migration because, first, there was no such peak in the $F_{ST}$ analysis, and second, both resident and migrant samples were combined for the Little Sheep Creek Tajima’s $D$ analysis. The kernel smoothing $F_{ST}$ outlier analysis also suggested that Omy7 contains markers divergent between the resident and migrant populations in Sashin Creek, and like Omy2 these associations are coupled with a decrease in the Tajima’s $D$ value within the Sashin Creek migrants and a reduction in nucleotide diversity. The Little Sheep Creek samples, by contrast, did not show any regions of the genome suggestive of the effects of selection (or associations between genotype and phenotype; see File S5). Several other peaks of differentiation were noted from the Sashin Creek $F_{ST}$ outlier analysis, namely Omy4, Omy8, Omy12, Omy14, and Omy22. Although these peaks are lower than those associated with Omy2 and Omy7, they also exhibit a negative departure from neutrality in the Sashin Creek migrants and a neutral distribution in the Sashin Creek residents, suggesting there are loci on these chromosomes that also contribute to the genetic differentiation between migrants and residents. However, we were only able to map 1990 (25%) SNPs. It is therefore possible that we have missed regions of the genome that show signatures of selection.

Comparing our results with other studies is difficult, especially because there appear to be population-specific genetic effects associated with migration/residency. For example neither Nichols et al. (2008) nor Hecht et al. (2012a) found QTL for migratory-related traits (during the juvenile phase) on Omy2. However, Hecht et al. (2012a) did find QTL for shape differences on Omy7. Comparisons between our results and Hecht et al. (2012a) are more applicable because they used a QTL scan between markers segregating between an F2 cross in Sashin Creek. Of the other regions of the genome with elevated differentiation in Sashin Creek, Hecht et al. (2012a) found QTL on Omy4, Omy8, Omy12, Omy14, and Omy22. Although these peaks are lower than those associated with Omy2 and Omy7, they also exhibit a negative departure from neutrality in the Sashin Creek migrants and a neutral distribution in the Sashin Creek residents, suggesting there are loci on these chromosomes that also contribute to the genetic differentiation between migrants and residents. However, we were only able to map 1990 (25%) SNPs. It is therefore possible that we have missed regions of the genome that show signatures of selection.
QTL for different traits), and further support for the importance of these regions comes from QTL for growth (Wringe et al. 2010) and osmoregulation (Le Bras et al. 2011). Similarly, Nichols et al. (2008) found Omy5 and Omy10 being “hotspot QTL regions,” and additional studies have found loci on Omy5 to be associated with migratory related traits (O’Malley et al. 2007; Martinez et al. 2011) and timing of sexual maturity and development rate (Danzmann et al. 2005; Leder et al. 2006; Nichols et al. 2007; Miller et al. 2012). Although both Omy10 and Omy5 contain outlier $F_{ST}$ loci and a largely negative Tajima’s $D$ distribution in the Sashin Creek migrants, kernel smoothing suggests none of these chromosomes are enriched for such loci. However, it is important to keep in mind that only 25% of the total SNPs were matched to the $O$. mykiss genome. The relatively small number of loci confidently ordered in the genome is a prevailing issue that will be important in this and other nonmodel organisms. It is clear, not only that migration/residency is a complex trait involving many genes scattered through the genome, but also that there are striking differences in the number and location of such loci between populations. However, undoubtedly not all outlier loci are connected with the evolution of migration/residency. The separation of migrant and resident samples in the Sashin Creek dataset means other factors could also (due to different selection pressures, patterns of linkage disequilibrium or drift) cause segregation that overlaps with phenotype.

The evolution of marine and freshwater adaptation has probably been best studied in the three-spined stickleback (Gasterosteus aculeatus) and currently this is the only current comparative model for our study. Ecological genomic studies in sticklebacks have shown very convincingly that the evolution of multiple independent freshwater populations from a marine ancestor have involved parallel genetic changes (Colosimo et al. 2004; Cresko et al. 2004; Hohenlohe et al. 2010). However, the situation in freshwater rainbow trout is more complex. In this study, the population history of the two study sites is very different and demography has important implications for finding signatures of selection, but perhaps more importantly in the ability to reliably test for true associations between genotypes and phenotypes using GWAS (see Price et al. 2006; Nielsen et al. 2007). The Sashin Creek system includes barriers to migration that separate the two forms (Thrower et al. 2004). Although migrants from the upper watershed may migrate out to the ocean, large waterfalls prevent return migration to the upper sections of the Creek and Lake (Masuda et al. 2009). Similar patterns of strong differentiation (high $F_{ST}$ and a greater proportion of outliers) between migrants and residents above and below barriers have been found in other populations of $O$. mykiss (Narum et al. 2011; Limborg et al. 2012; Pearse et al. 2009; Martinez et al. 2011). In such systems there could be selection for above barrier fish to not produce migrant offspring as these would be lost from the population. Over time, this could result in purging of any alleles associated with anadromy. Such a scenario could produce patterns of genome variation similar to what we found in Sashin Creek with many loci associated with migration/residency, many outlier loci from $F_{ST}$ analysis, high LD, a largely negative departure from neutrality in the source population, and many private alleles being present in the migrants.

What followed in such systems can be thought of as a hard selection sweep in which alleles associated with residency were selected to a much greater frequency than the same alleles in the migrant population. Further support for selection against residents producing migrants comes from Thrower and Joyce (2004), who examined the survivability of migrant fish produced from crosses between resident fish and migrant fish. They found that migrant offspring produced by migrant fish had a greater proportion of returns than migrant offspring from resident fish, suggesting migrants are inheriting alleles that promote ocean survivability. In contrast, the Little Sheep Creek samples were collected from an area where migratory and resident life history types coexist providing greater opportunity for mating between the life history types (Bernston et al. 2011) as confirmed by the few loci associated with migration/residency, few $F_{ST}$ outliers, low levels of LD, largely neutral Tajima’s $D$ values, and low numbers of private alleles. Similar low levels of $F_{ST}$ and few outliers are found in other populations of $O$. mykiss where residents and migrants co-occur (Limborg et al. 2012). This finding suggests either an historic selective hard sweep that has been eroded by drift and interbreeding between resident and migrant forms, or a soft sweep in which selection pressure between forms is reduced.

Although migration/residency in $O$. mykiss is undoubtedly a complex trait, the long-standing issue of how a “migration syndrome” is quantified or identified can have important implications on our synthesis of the results, especially when comparing between studies. In the study herein, all migrant fish were sampled as sexually mature adults returning to their natal streams, but the two resident populations were sampled at different time points during development. The Sashin Creek resident fish were tested for gamete expression before sampling only fish that produced gametes were included (sexual maturity precludes migration in salmonids, discussed below). In contrast, the Little Sheep Creek residents were sampled after fish reached a minimal size (fork length greater than 160 mm). Though the migrant phenotype is fairly easily scored (by the act of migration in downstream or upstream traps, distinctive body morphology and coloration), the resident phenotype may not always be definitively determined until reaching sexual maturation. Moreover, it is possible that some of the fish classified as residents in the Little Sheep Creek fish would have subsequently migrated. Similar complications arise when comparing our results with those from previous studies (Nichols et al. 2008; Hecht et al. 2012, 2013). We sampled migrants that had successfully returned from the ocean to spawn, whereas previous studies determined phenotype at 2 years of age (the point of smoltification, which occurs in the juvenile phase). This could be crucial, as the migrant fish in this study would have experienced additional selection and mortality in the ocean whereas previous studies did not. Therefore we are essentially measuring two different traits, the genetic basis of smoltification in the juvenile stage (Nichols et al. 2008; Hecht et al. 2012, 2013) and the genetic basis of successful migration (this study).

The annotation of genome scaffolds that are linked to candidate loci for directional selection produced associations to genes connected to development rate, osmoregulatory function, and reproductive maturity (Tables 1 and 2). Our BLAST results show that growth hormone 1 ($GH1$), a gene that is involved in growth and development (Dickhoff et al. 1997), is on the same contig as a tag that contains a marker that is associated with migration and certainly warrants further study. There are two BLASTn and four BLASTx hits to genes involved in ion exchange such as various potassium and calcium voltage-gated channels. Presumably such genes are involved in efficient osmoregulation and could be under heavy selection pressures in migratory individuals. Olfaction is also important in the ability of returning migrants to navigate to their natal streams, and an olfactory receptor (family C) is linked to a significant SNP therein (R08544, R03392).

Intriguingly, several genes involved in sexual differentiation and maturity were linked to significant SNPs in this study: gonadotropin-subunit beta-2 ($Gthb2$), SOX14, and sperm-associated antigen 16. Receptors for gonadotropin have been shown to be associated with return migration timing for spawning in chum salmon ($Oncorhyncus keta$; Onuma et al. 2010), a trait that could be under selection in all
migratory salmonids. Members of the SOX transcription factors have been shown to be involved in sex differentiation in *O. mykiss* (Baron et al. 2005) and other vertebrates. There appears to be a strong relationship between age of sexual maturity and sex, with the vast majority of sexual mature precocious fish being male in *O. mykiss* (e.g., Sharpe et al. 2011) and other species of salmonid (e.g., Larsen et al. 2010). This could have a direct effect on migratory behavior as smolification delays sexual maturity (Thorpe 1994). Confirming this, the Sashin Creek samples show a strong sex bias with 76% of the migrants being female compared to 29% of the residents. Of course, BLAST approaches like this are tenuous without a reference genome. We were only able to get BLAST information for a very small proportion (12%) of the significant SNPs, however they do contain some genes with functions connected to migration and warrant further analysis.

In this study, we have demonstrated the utility of RAD tag sequencing to find and genotype thousands of SNPs to understand possible adaptive divergence between migratory and resident ecotypes in natural populations. Although combining population genomic and association approaches is a powerful way to overcome some of the limitations of associations studies, GWAS in populations where phenotype is associated with strong population structure produces problems in separating true positives from loci that diverged simply as a function of demographic processes (see File S5). More generally, studies to dissect the genetic architecture of complex traits in non-model organisms will continue to be limited by: (1) confounding of traits with population demography, (2) the availability of a linkage map or genome sequence without which trends within and between chromosomes cannot be examined, (3) the amount of linkage disequilibrium within populations of study, (4) potentially confounding effects of environment on an individual’s phenotype, (5) the trait itself, and (6) the genetic architecture of the trait (simple, Mendelian or complex involving epistasis). With respect to the genetic basis of migration in *O. mykiss*, it is clear that many different regions of the genome are associated with this complex trait, both from this and other studies. Some of these regions seemed to be shared between different populations of *O. mykiss*, and some seem to be unique to individual populations, but note the differences in sampling returning adults in our study vs. smolting juveniles in other studies. We are still unsure whether this is a phenomenon unique to *O. mykiss*, salmonoids or whether it is found in many different taxa that have a genetic basis to migration and only future studies in other species will allow us to answer this question. In addition, more work needs to be done to determine which genes are involved and to fully understand the function of such genes. With the genome for *O. mykiss* soon to be completed, this will hopefully soon become a possibility.

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