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Migration-related phenotypic divergence is associated with epigenetic modifications in rainbow trout

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Abstract

Migration is essential for the reproduction and survival of many animals, yet little is understood about its underlying molecular mechanisms. We used the salmonid *Oncorhynchus mykiss* to gain mechanistic insight into smoltification, which is a morphological, physiological, and behavioral transition undertaken by juveniles in preparation for seaward migration. *O. mykiss* is experimentally tractable and displays intra- and inter-population variation in migration propensity. Migratory individuals can produce non-migratory progeny and vice versa, indicating a high degree of phenotypic plasticity. One potential way that phenotypic plasticity might be linked to variation in migration-related life history tactics is through epigenetic regulation of gene expression. To explore this, we quantitatively measured genome-scale DNA methylation in fin tissue using reduced representation bisulfite sequencing of F₂ siblings produced from a cross between steelhead (migratory) and rainbow trout (non-migratory) lines. We identified 57 differentially methylated regions (DMRs) between smolt and resident *O. mykiss* juveniles. DMRs were high in magnitude, with up to 62% differential methylation between life history types, and over half of the

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Data Accessibility

Demultiplexed Illumina sequencing reads for the 18 analyzed individuals are available in the NCBI Gene Expression Omnibus, Accession No. GSE62195. This accession also contains raw DMC and DMR results as well as a readme file containing code for the bioinformatics analyses performed.

Author Contributions

Research was designed by MRB, MHM, MRS, RPN, KMHT, BM, and KMN. Laboratory experiments were conducted by AMG and computational and data analyses were performed by MRB. GHT provided the clonal line crosses. MRB wrote an initial manuscript draft, which was edited by all authors.

gene-associated DMRs were in transcriptional regulatory regions. Many of the DMRs encode proteins with activity relevant to migration-related transitions (e.g. circadian rhythm pathway, nervous system development, protein kinase activity). This study provides the first evidence of a relationship between epigenetic variation and life history divergence associated with migration-related traits in any species.

Keywords

epigenetics; life history variation; smoltification; *Oncorhynchus mykiss*; plasticity

Introduction

Migration is a complex phenomenon, with billions of animals annually responding to environmental heterogeneity by moving long distances (Bowlin *et al.* 2010). This allows them to exploit seasonally or geographically available resources (e.g. food, habitat, favorable climate or breeding conditions) or avoid unfavorable conditions (e.g. predators, disease; Alerstam *et al.* 2003). Migratory behavior occurs across all major branches of the animal kingdom, in groups as diverse as fish, insects, reptiles, crustaceans, birds, and mammals (Alerstam *et al.* 2003). It has evolved repeatedly within lineages (e.g. Chinook salmon *Oncorhynchus tshawytscha* (Waples *et al.* 2004), bats (Mammalia: Chiroptera) (Bisson *et al.* 2009)), and in some cases, rapidly (e.g. house finches *Carpodacus mexicanus* (Able & Belthoff 1998)). Changes in the propensity, timing, and patterns of migration may have large-scale implications for patterns of species connectivity and ecosystem health. Yet, despite human fascination with animal migrations (Nebel 2010), the molecular mechanisms that underlie migratory behavior are not well understood (Liedvogel *et al.* 2011). With rapidly changing environmental conditions and the associated challenges of managing species for future persistence, it is becoming critical to understand the mechanisms driving migration (Liedvogel *et al.* 2011).

Substantial phenotypic variation in migration trajectory, timing, and propensity exists across species and populations; this variability is attributed to a combination of environmental and genetic factors (Liedvogel *et al.* 2011; Dingle 2014). For example, many Pacific salmon species display variable migration timing during juvenile outmigration to the ocean, as well as timing in adult return to freshwater for spawning (Quinn & Myers 2005). These salmon species, and populations within them, also have variable migration trajectories, with some staying in coastal waters and others migrating to the open sea from rivers (Quinn & Myers 2005). Organisms respond to environmental cues (e.g. photoperiod, temperature, food availability), which can trigger internal cues (e.g. circadian rhythm, hormones, and fat deposits) to initiate migration (Folmar & Dickhoff 1980; McCormick *et al.* 1998). Heritability and genetic mapping studies across diverse taxa suggest there is genetic variation for the propensity to migrate (Berthold 1991; Thrower *et al.* 2004; Roff & Fairbairn 2007; Hecht *et al.* 2015). Additionally, phenotypic plasticity is believed to play a role in producing alternative migratory phenotypes (Hayes *et al.* 2012; Dodson *et al.* 2013). One potential way that phenotypic plasticity might be linked to variation in migratory tactics is through epigenetic regulation of gene expression (e.g. DNA methylation, histone tail

modifications, non-coding RNAs). Epigenetic modifications can mediate responses to environmental cues (Bastow *et al.* 2004; Petronis 2010; Feil & Fraga 2012), including behavioral (Champagne 2010, 2012; Mifsud *et al.* 2011) and circadian rhythm responses (DiTacchio *et al.* 2011; Fustin *et al.* 2013; Azzi *et al.* 2014). For example, DNA methylation is involved in transcriptional silencing, alternative splicing, and activating intragenic promoters (reviewed in Jones 2012) and acts as a reversible mechanism to drive circadian clock behavioral plasticity in mice (*Mus musculus*; Azzi *et al.* 2014).

Due to its experimental tractability and remarkable variability in migration propensity, the salmonid *Oncorhynchus mykiss* is an excellent model for studying mechanisms driving migration (Zimmerman & Reeves 2000; Riva-Rossi *et al.* 2007; Hayes *et al.* 2012; Courter *et al.* 2013). The species has two distinct migration life history strategies: migratory (anadromous steelhead) and non-migratory (resident rainbow trout). Both migrant and non-migrant individuals are born and rear in freshwater to the juvenile stage. Some juveniles then go through a process termed smoltification, during which they undergo morphological, physiological, and behavioral changes before migrating to the ocean (Hoar 1976); this migratory or anadromous form of *O. mykiss* is called a steelhead trout. Upon reaching reproductive maturity in the ocean, steelhead migrate back into freshwater to spawn, thus beginning the life cycle again. In contrast, the juveniles that do not undergo smoltification remain in freshwater throughout their lives; this resident form of *O. mykiss* is called a rainbow trout.

Migration-related life history traits are highly heritable within *O. mykiss* (Thrower *et al.* 2004; Hecht *et al.* 2015), but there is also substantial phenotypic variability among families and lineages. Genetic mapping studies have identified quantitative trait loci (QTL) associated with anadromy and residency across the genome. A linkage mapping study using the same F₂ family as this current study found QTL associated with smoltification-related traits across many chromosomes, including a major effect locus on chromosome Omy10 (Nichols *et al.* 2008). A genome scan of two closely related steelhead - rainbow trout populations found non-neutral outlier loci associated with smoltification across 19 chromosomes (Martínez *et al.* 2011). Both linkage mapping and genome-wide association studies (GWAS) have found many loci of small effect contributing to migration-related phenotypic differences across the genome using geographically diverse populations (Hecht *et al.* 2012a; b; Hale *et al.* 2013). However, there is evidence that some chromosomes (e.g. Omy5) contain strong associations with smoltification across many distinct populations (Pearse *et al.* 2014) and genetic mapping of this complex trait is on-going. Genes found proximal to identified QTL include those with functions pertaining to circadian rhythm, sexual maturation, development rate, and osmoregulation (Martínez *et al.* 2011; Hecht *et al.* 2012a; Hale *et al.* 2013). However, the specific genes triggering the developmental commitment to migration versus residency are not yet known.

Migration is a threshold trait (Pulido *et al.* 1996; Dodson *et al.* 2013), with manifestation dependent on interactions between the environment and an organism's genetic architecture and development (e.g. body size (Jonsson & Jonsson 1993; Theriault & Dodson 2003)). For *O. mykiss*, environmental and landscape variables such as temperature, photoperiod, precipitation, elevation, and upstream distance can influence migration propensity (Zaugg &

Wagner 1973; Narum *et al.* 2008; Kendall *et al.* 2015). There is considerable phenotypic plasticity in life history diversity, with some populations containing individuals of a single life history strategy while other populations harbor individuals of both strategies (Jonsson & Jonsson 1993; Quinn & Myers 2005; Dodson *et al.* 2013). Each life history type can give rise to the other in both wild (Riva-Rossi *et al.* 2007; Courter *et al.* 2013) and hatchery (Viola & Schuck 1995; McMichael *et al.* 1997) settings. We hypothesize that the observed plasticity may be controlled by an epigenetic response threshold, which integrates parentally inherited factors, environmental factors, and developmental history (Yan *et al.* 2014). Therefore, divergent life history phenotypes may be a result of differential epigenetic modifications that link environmental factors and the genome to regulate internal cues (e.g. circadian rhythm). In this work we evaluate if epigenetic regulation (DNA methylation) is associated with smoltification, a migration-related phenotype, by comparing smolt and resident phenotypes from an *O. mykiss* F₂ intercross reared in a common environment.

Materials and Methods

Samples

The *O. mykiss* F₂ intercross used for this study was previously used for a QTL mapping study of smoltification wherein individual fish were quantitatively classified as smolts or residents (Nichols *et al.* 2008). In that study, steelhead (Clearwater River) and resident (Oregon State University) doubled haploid clonal lines (Nichols *et al.* 2007) were crossed to create F₁ hybrids. Gametes from an F₁ male then underwent androgenesis, to create F₂ progeny, each homozygous across loci, but genetically different from their siblings due to meiotic recombination. At the age of 2, when juveniles often begin to undergo smoltification, metrics strongly associated with smoltification (body size, condition, growth, morphology, osmoregulatory ability, and skin reflectance) were measured. The measurements from these phenotypes were compiled to identify a binary phenotype, classifying each individual as either a smolt or resident. Smolts were silver and streamlined in body shape, while residents had darker coloration, dark parr marks on the sides of their bodies, and greater body and caudal peduncle depths (Fig. 1; methods detailed in Nichols *et al.* 2008). Ten individuals (five males and five females) of each phenotype were selected for reduced representation bisulfite sequencing (RRBS). DNA was extracted from caudal fin using the Qiagen DNeasy kit. This tissue type is in intimate contact with the environment and undergoes phenotypic changes during smoltification, such as increased silvering and darkening, which are associated with metabolic changes that occur during the time frame leading up to migration (McCormick *et al.* 1998). We chose to use this doubled haploid cross in the first analysis of differential methylation between migration-related phenotypes because: 1) we could directly compare our differential methylation results with the location of QTL for smoltification by using fin tissue previously used for a genetic analysis of smoltification (Nichols *et al.* 2008) and 2) the doubled haploids come from a homogenous genetic background and were reared in a common controlled environment, reducing any potentially confounding noise from large amounts of genetic diversity or environmental variability.

Reduced representation bisulfite sequencing (RRBS)

RRBS libraries were constructed using a previously published protocol (Boyle *et al.* 2012). Briefly, DNA was quantified using the Qubit 2.0 fluorometer (Life Technologies) with the dsDNA BR assay kit. Genomic DNA (100 ng) for each individual was digested using the restriction enzyme *MspI* (New England Biolabs). DNA fragments were end-repaired, A-tailed, and ligated to the methylated Illumina TruSeq adapters. Ten samples were multiplexed per library, for a total of two libraries, with each sample barcoded with unique Illumina adapters to distinguish samples. DNA from fish representing the two life history types and sexes was equalized across libraries. Sodium bisulfite conversion was conducted using the EpiTect Bisulfite kit (Qiagen), following the protocol for formalin-fixed paraffin-embedded samples. After two rounds of bisulfite treatment to ensure complete conversion, the DNA was amplified using Illumina's TruSeq primers. The optimal number of PCR cycles (18) was determined empirically by comparing band intensities of PCR products on a 5% MetaPhor agarose gel (Lonza) for 10, 13, 16, 19, and 22 total cycles. Gel-free library size selection was performed with Agencourt AMPure XP beads (Beckman Coulter) as described by Boyle *et al.* (2012). Library DNA quantity and quality was assessed using a Qubit fluorometer and 5% MetaPhor agarose gel, respectively. After a quantitative PCR quality control check, the two libraries were sequenced using two lanes (50 bp single-end reads, 10 samples per lane) of an Illumina HiSeq 2000 at the Vincent J. Coates Genomics Sequencing Laboratory (University of California, Berkeley, USA). The RRBS protocol was repeated for nine samples with fewer than 10 million reads after quality filtering. These samples were sequenced as above, but with 100 bp single-end reads. Base calling was performed using Illumina's CASAVA 1.8.0. Initial read quality checks were done using FastQC (available at: <http://www.bioinformatics.bbsrc.ac.uk/projects/fastqc>). Trim Galore! (available at: http://www.bioinformatics.babraham.ac.uk/projects/trim_galore/) removed low quality and adapter contaminated sequences, using the "rrbs" option.

Read alignments to two *O. mykiss* genome versions

Recently, two versions of the *O. mykiss* genome (the French Genome Project (Berthelot *et al.* 2014) and the U.S. Genome Project (available at: <http://www.animalgenome.org/repository/aquaculture/>)), were released to facilitate large-scale genomic studies. Both assemblies used the same DNA source, but were produced using different methods. The French Genome Project (hereafter this Project's assembly will be referred to as the French Genome) used whole-genome shotgun sequencing approach, which enables all regions of the genome to be targeted, but has difficulty mapping reads from repetitive elements and paralogous sequences. This is a potential issue since paralogs exist throughout the salmonid genome due to an autotetraploidization event 25 - 100 million years ago (Allendorf & Thorgaard 1984). Contrastingly, the U.S. Genome Project (hereafter this Project's assembly will be referred to as the U.S. Genome) used a BAC library sequencing approach, which is better able to uniquely identify and map repetitive and paralogous sequences, but is limited to the regions of the genome contained within the BAC libraries. While these two *O. mykiss* genome assemblies have considerable sequence overlap, they also contain unique sequences. Conducting analyses with both genome assemblies in a comparative fashion is likely the most comprehensive approach until further genome development. Therefore, for the purposes of alignment, both genomes were *in silico* converted to bisulfite genomes using

Bismark (Krueger & Andrews 2011) in order to independently align reads to each genome. Although the French genome has longer N50 scaffolds, both genome assemblies contain hundreds to thousands of unassembled contigs. Therefore, to increase mapping efficiency (i.e. ability of reads to uniquely map to a single region in the genome), small scaffolds (< 50 kb for the French genome or < 10 kb for the U.S. genome) were removed prior to analysis. Bowtie (Langmead *et al.* 2009), implemented in Bismark, was used to align reads for each sample to the genome, tolerating one non-bisulfite mismatch per read, and producing SAM files for further analyses of differential methylation. Only reads that mapped uniquely to at least one of the reference genomes were included in downstream differentially methylated CpG site (DMC) and differentially methylated region (DMR) analyses. The percent of French genome CpG sites that are encompassed within an *MspI* recognition sequence ('CCGG') was *in silico* predicted using the SimRAD package (Lepais & Weir 2014) within the R computing environment (R Core Team 2014).

Individual differentially methylated CpG (DMC) sites were identified between resident and smolt groups using logistic regression with the R package methylKit (Akalin *et al.* 2012). Read coverage was normalized between samples. A minimum of seven individuals per life history type, each with a minimum of ten reads, were required at a CpG site for that site to be analyzed. Default parameters were used (q-value < 0.01), with a minimum required methylation difference of 25% between smolt and resident phenotypes.

Differentially methylated regions (DMRs) were determined using the R package eDMR (Li *et al.* 2014) with default parameters. To be considered significant, a DMR needed to contain at least three CpG sites within an algorithm-specified genomic distance (see Li *et al.* 2014 for details), with at least one classified as a DMC (q-value < 0.01), and an absolute mean methylation difference greater than 20% when comparing smolt and resident phenotypes. Visualization of sequence reads within and across individuals, and compared to the reference genomes, was performed with Integrative Genomics Viewer (Robinson *et al.* 2011). Visualization of the differential methylation patterns across individuals was achieved using the heatmap2 function in the R package ggplot2 (Wickham 2009).

CpG island proximity, gene annotation, gene ontology, and pathway analysis

For CpG island prediction, EMBL-EBI's CpGplot (available at: <http://www.ebi.ac.uk/Tools/seqstats/>) was used with default parameters. DMRs were considered part of CpG shores if they were ≤ 2 kb away from an island (Irizarry *et al.* 2009) and part of CpG shelves if they were between 2 - 4 kb away from an island, as defined by Illumina.

We used published annotations of the French *O. mykiss* reference genome (available at: <http://www.genoscope.cns.fr/trout-ggb/cgi-bin/gbrowse/truite/>) for DMRs that aligned to this genome assembly. The U.S. *O. mykiss* genome has yet to be annotated so for DMRs that only aligned to this genome assembly, we used Bowtie to align the entire scaffold to the French *O. mykiss* genome to identify existing annotations. For the few reads that did not align, we used NCBI's megablast query to determine region-specific annotations for the DMR regions. For all annotations, scaffolds containing DMRs were trimmed to include 2.5 kb upstream and downstream of the DMR. An e-value threshold of $< 1e-50$ was required to assign a gene name.

We used Gene Cards (available at: <http://genecards.org/>) to identify gene ontology terms for the DMRs annotated as part of proximal regulatory regions (5' or 3' UTR) or gene bodies. Pathway analysis was conducted using Qiagen's Ingenuity Pathway Analysis (IPA), which identified functional associations among the annotated DMRs, such as particular canonical pathways and common gene networks found across the DMRs.

Results

Reduced representation bisulfite sequencing (RRBS) coverage and mapping

Total reads per library ranged from 135 - 147 million after quality filtering (Table S1). Total number of reads per individual ranged from 4 - 49 million (average: 14 million) after quality filtering. Two of the twenty samples did not have 10 million reads and did not undergo further analysis (samples 19 and 20 in Table S1). The bisulfite conversion was highly efficient, with all samples averaging >99.6% (median: 100%) conversion. One-third (33.2%) of the bisulfite converted reads mapped uniquely to the French version of the *O. mykiss* genome (i.e. at a single location; Table S1). One-quarter (25.9%) of the reads mapped to more than one genomic location. For the U.S. *O. mykiss* genome, 24.4% of the reads mapped uniquely while 35.7% mapped to multiple locations.

Identification of differentially methylated regions (DMRs)

A total of 117,082 and 158,381 CpG sites met the minimum coverage requirements for the U.S. and French *O. mykiss* genomes, respectively. For context, there were 22.8 million CpG sites in the French *O. mykiss* genome after removal of small contigs (refer to Materials and Methods). Therefore, less than 1% of all CpG sites were analyzed per genome assembly due to: 1) reads not uniquely mapping, 2) reads not being found in or near the *MspI* recognition sequence ('CCGG'), or 3) low sequence coverage. Single differentially methylated cytosines (DMCs) ranged from 113 (U.S.) to 521 (French). Based on these DMCs, we identified 57 DMRs after false discovery rate correction when comparing phenotypically classified resident and smolt life history types (Table 1). Given our lower limit differential methylation cutoff (20%), methylation differences between resident and smolt groups ranged from 20 - 62% per DMR. Individuals clustered primarily by life history type based on the 57 DMRs (Fig. 2). However, one smolt clustered with residents and two residents clustered with smolts. Because adult phenotypes (body morphology, sexual maturation) were recorded after these individuals reached age three, it is unlikely these individuals were phenotypically misclassified. Specifically, fish that were silvery and streamlined in their appearance were classified as smolts, and those that were dark colored, more deep-bodied, or producing gametes were classified as resident individuals. The DMRs displayed hyper- and hypo-methylation in nearly equal proportions (28 hyper-, 29 hypo-methylated when comparing residents to smolts). Each DMR contained 3 - 21 total CpG sites (average = 5), with 1 - 12 of these CpGs being differentially methylated (average = 3).

Thirteen of the DMRs were identified with both reference genomes, 31 with only the French genome, and 13 with only the U.S. genome, which has a lower N50 scaffold size in comparison to the French genome. Additional details about each of these DMRs, including contig and scaffold positions, can be found in Table S2.

Genomic features and gene annotation

We mapped 37 DMRs onto 12 chromosomes (Table S2). Chromosomes Omy10 and Omy19 contained the most DMRs, with 7 and 9, respectively (Fig. 4a; Table S2). Five other chromosomes (Omy3, Omy4, Omy7, Omy16, Omy20) contained two or more DMRs. Five chromosomes each contained a single DMR and twenty DMRs could not be mapped to a chromosome due to the incomplete nature of the *O. mykiss* reference genome(s).

Twenty-six of the DMRs localized within or proximal to annotated genes and a subset directly overlapped genomic regions with potential regulatory function. Over half of the DMRs for this study were found within CpG islands (CGIs), shores, or shelves (Fig. 4b). We found five of the annotated DMRs within or proximal to CGIs (i.e. CGI shores, shelves) located within putative promoter regions (Table S2). Six of the annotated DMRs were found in CGIs, shores, or shelves within gene bodies (Table S2).

Thirty-one of the 57 DMRs were found within the proximal putative regulatory or coding regions of annotated genes (Fig. 4c; Table S2). Specifically, 15 were found within gene bodies, 8 within 5 kb upstream of the transcription start site (TSS), 4 within 5-10 kb upstream of the TSS, and 4 within 1.5 kb downstream of the transcriptional stop codon. Twenty-six DMRs were located in or proximal to annotated *O. mykiss* genes (Table 1). Of the remaining 31 DMRs, 5 assigned as uncharacterized proteins (Table 1; Table S2). In addition to proximal regulatory region and gene body methylation differences, 26 of the DMRs were found within intergenic regions.

The most common gene ontology terms found across the 26 annotated DMRs included signal transduction and G-protein coupled receptor signaling pathway (biological process), protein binding (molecular function), and integral component of membrane, plasma membrane, and nucleus (cellular component; Table S3, Fig. S1).

Pathway analysis

Several molecular pathways associated with smoltification were identified via Ingenuity Pathway Analysis. Six of the top canonical pathways for the gene-associated DMRs were statistically significant (Table S4). We identified two gene networks, each with distinct functional roles as well as a common role relating to nervous system development and function (Table S4). These networks contained 11 - 15 of the 26 gene-associated DMRs (Fig. S2).

Discussion

Migration plays a central role in the life history of many animals but remains poorly understood at the molecular level, and the involvement of epigenetic mechanisms has not been previously explored for any species. We used a quantitative, high-resolution technique (RRBS) to measure DNA methylation across the *O. mykiss* genome and detected 57 highly significant differentially methylated regions (DMRs) relating to smoltification, a migration-related phenotype. Methylation differences of up to 62% per DMR were observed between resident and smolt fin tissue, a level that is on par with differences previously observed for other animal phenotypic changes associated with epigenetic mechanisms (e.g. Herb *et al.*

2012). Not only is the magnitude of methylation changes high, but some of the identified genes have relevance to migratory transitions. Hierarchical clustering based on methylation patterns alone differentiated the life history types, although there were a few individual exceptions (Fig. 2). Since it is unlikely these individuals were phenotypically misclassified, it appears that epigenetic heterogeneity is occurring. In these cases, we speculate that gene expression patterns resulting in alternative life history phenotypes may be due to 1) differential methylation at loci not captured in this study or 2) DNA methylation-independent mechanisms. Thus, distinct epigenetic profiles might underlie similar phenotypic outcomes.

Comparison to MS-AFLP approach and other salmonid epigenetic studies

The vast majority of ecological studies to date have used a methyl sensitive MS-AFLP approach (Cervera *et al.* 2002), which identifies global methylation changes, but does not provide the nucleotide-level resolution needed to derive functional genomic information and almost certainly would not have detected the small number of DMRs identified in this study. Previous salmonid studies that have used MS-AFLP have identified broad-scale methylation changes associated with seawater adaptation (Morán *et al.* 2013) and early maturation (Morán & Pérez-Figueroa 2011) in Atlantic salmon (*Salmo salar*). Global DNA methylation differences were not found in a study between wild and hatchery *O. mykiss* steelhead (Blouin *et al.* 2010); however, the authors acknowledge the potential for undetected gene-specific methylation changes. The RRBS approach used here has several advantages compared to the MS-AFLP approach (see review by Schrey *et al.* 2013 and citations therein), the most notable being its quantitative nature and its ability to identify particular genomic regions (as opposed to anonymous AFLP loci) that are differentially methylated. Identification of DMRs and their co-localization with genes and regulatory regions allows us to uncover gene annotation and gene ontology patterns as well as conduct pathway analysis so that results can be interpreted in a more integrative manner with other ecological and genetic studies for the trait of interest. The approach, however, does require a reference genome for alignment purposes. As more species are sequenced, it is our hope that the field of ecological epigenetics becomes more mainstream and used in tandem with other biological subfields (e.g. genetics, physiology, ecology) to gain more comprehensive and integrative knowledge on the molecular mechanisms underlying complex phenotypic diversity.

Paralogous sequences and tissue type affect interpretation of results

Several of the DMR-associated genes are members of gene families (e.g. CLDN10, MYOM3, PCDHGC5) and the *O. mykiss* genome, as previously mentioned, is highly duplicated. While our analyses exclude reads that aligned to more than one location in each of the genomes, we cannot be certain that all gene family members are contained in the current incompletely assembled versions of the genome. Therefore, it is possible that the specific gene family member with differential methylation may be misidentified in some cases. In the absence of bisulfite cloning to distinguish gene family members, results should be interpreted with caution. Given our requirement of unique mapping, it is likely that differential DNA methylation of many gene family members and other paralogous regions went undetected. Additionally, it should be noted that sampling choices, including tissue or

cell type analyzed and timing of collection, can greatly affect results for both epigenetic and gene expression studies.

Relating differential DNA methylation to genomic architecture

Over half of the annotated DMRs were found within or proximal to CpG islands (CGIs), strongly suggesting that epigenetic modifications are influencing transcriptional activity of associated genes. CGIs are CpG-rich regions that are predominantly unmethylated and serve as sites for transcription initiation (Deaton & Bird 2011). Approximately 70% of annotated promoters in vertebrates are associated with CGIs (Saxonov *et al.* 2006) and hypermethylation of these regions stabilizes gene silencing in promoter regions (Stein *et al.* 1982; Mohn *et al.* 2008; Payer & Lee 2008). Residents displayed a 32% increase in DNA methylation compared to smolts in a CGI shore of the *aryl hydrocarbon receptor 2 alpha* (*AHR2A*; Fig. 3) distal promoter (<10 kb from TSS), suggesting that *AHR2A* is more stably expressed in smolt than resident fin tissue. AHR is an environmentally-responsive transcription factor that mediates toxic responses to environmental pollutants (Hansson *et al.* 2004), circadian rhythm (Anderson *et al.* 2013), cellular differentiation (Abbott & Birnbaum 1989), apoptosis (Gu *et al.* 2000), and immune functioning (Nguyen *et al.* 2013).

In contrast to promoters, CGI methylation in gene bodies is more common and speculated to be associated with regulatory non-coding RNA (Dinger *et al.* 2008; Ørom *et al.* 2010), alternative splicing (Kornblihtt 2006), or intragenic promoter activity (Maunakea *et al.* 2010). An example of gene body differential methylation in this study is *nescient helix-loop-helix 2* (*NHLH2*), with residents having 38% higher DNA methylation levels than smolts within a gene body CGI shore (Fig. 3). *NHLH2* is a transcription factor important in modulating body weight and energy availability (Fox *et al.* 2007), which are both believed to be critical factors regulating the occurrence of salmonid smoltification (Thorpe & Metcalfe 1998; Theriault & Dodson 2003) and subsequent migration.

Almost half (46%) of the DMRs were not located within or proximal to known genes. This may be an indication of trans-acting regulatory elements (e.g. enhancers), which are quite commonly differentially methylated in the human genome (Ziller *et al.* 2014), or it may indicate the incomplete annotation of the *O. mykiss* genome precludes identification of additional genes and their proximal regulatory regions.

Comparison with previous genetic mapping studies

Our study demonstrates that examining smoltification from an epigenetic perspective uncovers genomic loci not previously associated with migration-related traits and also corroborates loci known to be associated with smoltification or migration propensity. The chromosome identified as having the second highest number of DMRs (Omy10, Fig. 4a) was previously found to have numerous QTL for traits associated anadromy/residency, including a main effect QTL for the binary smolt phenotype for this F₂ intercross (Nichols *et al.* 2008). This co-localization of genetic and epigenetic variation onto Omy10 may indicate that genetic and epigenetic factors can act in a coordinated fashion to modify migration-related phenotypes. Further work is warranted to determine if genetic and epigenetic variation overlap in particular chromosomal regions. This will require considerably higher marker

density (and genomic sequence data) than was possible in the original QTL mapping study (Nichols *et al.* 2008), which used microsatellite and amplified fragment length polymorphic markers.

Previous studies have identified QTL for smoltification and migration-related traits on many chromosomes in the *O. mykiss* genome, reflecting the highly complex and polygenic nature of the phenotype. It seems unlikely that all migration-related genetic variation is associated with epigenetic variation; therefore, it is not surprising that we did not identify DMRs on many of the chromosomes containing migration-related QTL. This lack of complete coordination suggests that genetic and epigenetic variation do not always affect the same loci. Additionally, there is considerable variation in the location of QTL regions across populations (Hecht *et al.* 2012a; b; Hale *et al.* 2013) and we expect the same may be true of epigenetic variation, which should be explored in future studies of additional populations. A study in Atlantic salmon further corroborates this idea, providing evidence that resident versus migratory life history types had a complex multigenic basis with little parallelism across populations (Perrier *et al.* 2013).

Comparison with candidate genes from GWAS and gene expression studies

We found similarities between genes associated with some DMRs and those identified in prior GWAS and gene expression studies. For instance, *neurogenic differentiation factor 4* (*NEUROD4*) and *metabotropic glutamate receptor 1* (*GRM1*) were identified as differentially methylated in our current study, both with 31% increased methylation in smolts, and as F_{ST} outliers in a previous *O. mykiss* migration-related GWAS (Hale *et al.* 2013). *NEUROD4* is a transcriptional activator that mediates neuronal differentiation (Bertrand *et al.* 2002) and *GRM1* is a G-protein coupled receptor for glutamate involved in central nervous system transmission and normal brain function (Frank *et al.* 2011; Yin & Niswender 2014). Glutamate receptors are involved in synaptic plasticity (Kano & Hashimoto 2009; Ohtani *et al.* 2014), which is extremely important for learning and memory.

Multiple DMR-associated genes were members of the same gene families found in a brain gene expression study comparing age two F_1 progeny of migrant and resident *O. mykiss* crosses from a wild Alaskan population (Hale *et al.*, submitted). Identification of these gene families (protocadherin, claudin, myomesin) in our current study suggests that neuronal modifications and tissue development may be epigenetically regulated in association with migration propensity. Specifically, *PCDHGC5* is involved in forming and maintaining neuronal connections in the brain (Chen *et al.* 2012) and has 49% increased methylation in residents, *CLDN10* can enhance paracellular permeability (Amasheh *et al.* 2010) and has 30% increased methylation in residents, and *MYOM3* is involved in muscle tissue formation and contraction (Schoenauer *et al.* 2005) and has 22% increased methylation in smolts. Myomesin was also found to be differentially expressed in Atlantic salmon brain and gill tissue during smoltification (Seear *et al.* 2009). This collection of genes, involved in neuronal and muscle development, may be quite important for migration-related transitions. Extensive neuronal modifications are required for these transitions (Herry *et al.* 2008; Durstewitz *et al.* 2010), including epigenetically-regulated modifications (Borrelli *et al.*

2008), while preparation for migration may facilitate increased muscle formation and body mass (Marsh 1984). Only *O. mykiss* that reach a certain size threshold typically transition to a migratory phenotype (Theriault & Dodson 2003; Páez *et al.* 2011). However, the functional link between neuronal and muscle development is less clear when considering the fin tissue used in our current study. It is quite possible in these cases that the genetic variation (which is primarily static across tissue types within an individual) is strongly influencing DNA methylation levels. A study comparing DNA methylation and genetic variation across human populations suggested that differentially methylated CpG sites are “evolutionarily established mediators between the genetic code and phenotypic variability” (Heyn *et al.* 2013). Therefore, the same epigenetic differences may be detected regardless of the tissue type chosen.

Another category of interest is immune response. The DMR with the greatest differential methylation between divergent life history phenotypes was *radical S-adenosyl methionine domain containing 2 (RSAD2)*, with smolts averaging 62% more methylation at 10 CpG sites in the putative 3' UTR (Fig. 3). *RSAD2* encodes for a protein with interferon-inducible antiviral and antibacterial properties (Fitzgerald 2011). A previous study found that residents overexpressed antiviral, including interferon-associated, transcripts in gills compared to migrants (Sutherland *et al.* 2014). This may indicate a potential link between immune response and migration propensity, which seems plausible since the immune system is partially controlled by circadian rhythms (Scheiermann *et al.* 2013). Brook charr (*Salvelinus fontinalis*) differentially express immune response genes in gills for alternative migratory tactics, although migrants have increased immune expression (Boulet *et al.* 2012). Since skin tissue, including the fin, is a primary site of initial pathogen entry in fish (e.g. Baerwald 2013), it also plays a critical role in immune response. Therefore, epigenetic changes associated with immune response genes in fin tissue are not surprising given these prior links between immune response and migration propensity studied in other tissues.

Potential functional significance of DMRs

Gene ontology, canonical pathway, and gene network analyses identified several molecular categories with genes capable of dramatically altering phenotypes, some with clear associations to behavioral phenotypes such as migration. These include protein kinases, G protein-coupled receptors (GPCRs), and genes linked to the circadian rhythm pathway. Three DMR-associated genes encode for proteins with serine/threonine and tyrosine kinase activity (*AATK*, *SBK1*, *STYK1*; Table S3). Kinase expression and activity is tightly regulated (Huse & Kuriyan 2002) and phosphorylation plays a key role in generating circadian rhythms (Reischl & Kramer 2011). GPCRs sense extracellular molecules to initiate signal transduction cascades (Marinissen & Gutkind 2001) and two DMR-associated genes encode proteins in this functional category (*ADORA3*, *GRM1*; Table S3). GPCR activation can induce tyrosine kinase activity as well as RAS (Luttrell *et al.* 1999), so several of the differentially methylated genes can be connected through common gene networks (see Fig. S2).

We identified multiple differentially methylated genes within the circadian rhythm pathway, which is a primary internal cue for migratory behavior. Circadian rhythms allow organisms

to anticipate and prepare for regular environmental changes. Although circadian rhythms are endogenous, they are adjusted or entrained to the local environment by exogenous environmental cues (Gwinner & Scheuerlein 1998; Lehmann *et al.* 2012). In addition to regular 24-hour clock changes, circadian rhythms impact seasonal physiological and behavioral changes, including migratory traits such as migration propensity (Gwinner 1996; Guerra *et al.* 2014). Circadian rhythm gene expression patterns and genotypes affect migration timing in diverse species (e.g. Williams 2001; Belden *et al.* 2007; Mukai *et al.* 2008). Furthermore, circadian rhythms are regulated by DNA methylation, histone modifications, and RNA-based mechanisms (DiTacchio *et al.* 2011; Fustin *et al.* 2013; Azzi *et al.* 2014), indicating that epigenetic mechanisms drive circadian clock plasticity. We found two DMR genes associated with circadian rhythm, *AHR2A* and *dexamethasone-induced RAS 1 (RASDI)*. AHR can sense environmental compounds and affect circadian rhythmicity and responsiveness to environmental light cues (Mukai *et al.* 2008; Anderson *et al.* 2013; Xu *et al.* 2013). *RASDI*, also abbreviated *DEXRASI*, plays a pivotal role in the circadian clock's responsiveness to external timing cues, including both photic and non-photoc stimuli (Williams 2001; Cheng & Obrietan 2006; Belden *et al.* 2007). Reduced *RASDI* expression can cause non-photoc stimuli to have more influence on clock phasing than photic stimuli (Cheng & Obrietan 2006). In our study, methylation was increased in residents in the putative promoter regions of *AHR2A* and *RASDI* by 32% and 22%, respectively. This may suggest higher expression of these genes in smolts, which then might improve their responsiveness to light cues for clock phasing compared to residents. However, we believe the methylation changes need to display similar patterns between fin tissue and other tissue types that respond to photic stimuli (e.g. brain or optic cells) for these functional associations to be biologically meaningful, which will require further study.

Epigenetically-influenced response threshold for triggering migration

Development into differentiated smolts or residents may be at least partially dependent on an epigenetic response threshold. Response thresholds may determine an individual's sensitivity to particular environmental stimuli, with interactions between genetic and epigenetic factors along with environmental cues causing a phenotypic transition when the threshold is achieved (Yan *et al.* 2014). In an ecological context, this may explain the observed plasticity for migration propensity (Kendall *et al.* 2015). Residents may give rise to migratory progeny, and vice versa, due to the complex interactions between the environment, genetics, and development leading to alternative phenotypic outcomes that are dependent upon these thresholds. Future research should focus on strengthening the connection between epigenetic and phenotypic diversity by integrating epigenetic results with genetic and gene expression datasets under different environmental conditions to examine the nature and extent of coordination between alternative sources of variation that ultimately drive complex phenotypes. For conservation purposes, the two life history types are often managed separately, which will require careful consideration of environmental and genetic variation, and how they interact to produce a phenotypic response.

Conclusions

The genome-scale methylation data presented here provide the first evidence of a relationship between epigenetic variation and diversity in migration-related traits. It also

demonstrates the power of large-scale epigenotyping in ecological studies for identifying loci associated with complex phenotypes. Epigenetic mechanisms link genetic and environmental factors to generate alternative life history strategies. As recently demonstrated by Sloat *et al.* (2014), flexible life history strategies “can be both heritable and subject to strong environmental influences” and our study indicates that differential DNA methylation at gene regulatory elements may be a critical molecular mechanism allowing interactions between an organism and its environment to balance phenotypic stability and plasticity, and ultimately modify migration-related phenotypes.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Fig. 1. Phenotypic differences between resident and smolt *O. mykiss* juveniles

Photographs of siblings with alternative life history types were taken on same date and age (June, age 2). Smolts are silvery and streamlined in body shape, while residents are darker in skin and fin coloration and have dark parr marks on the sides of their bodies.

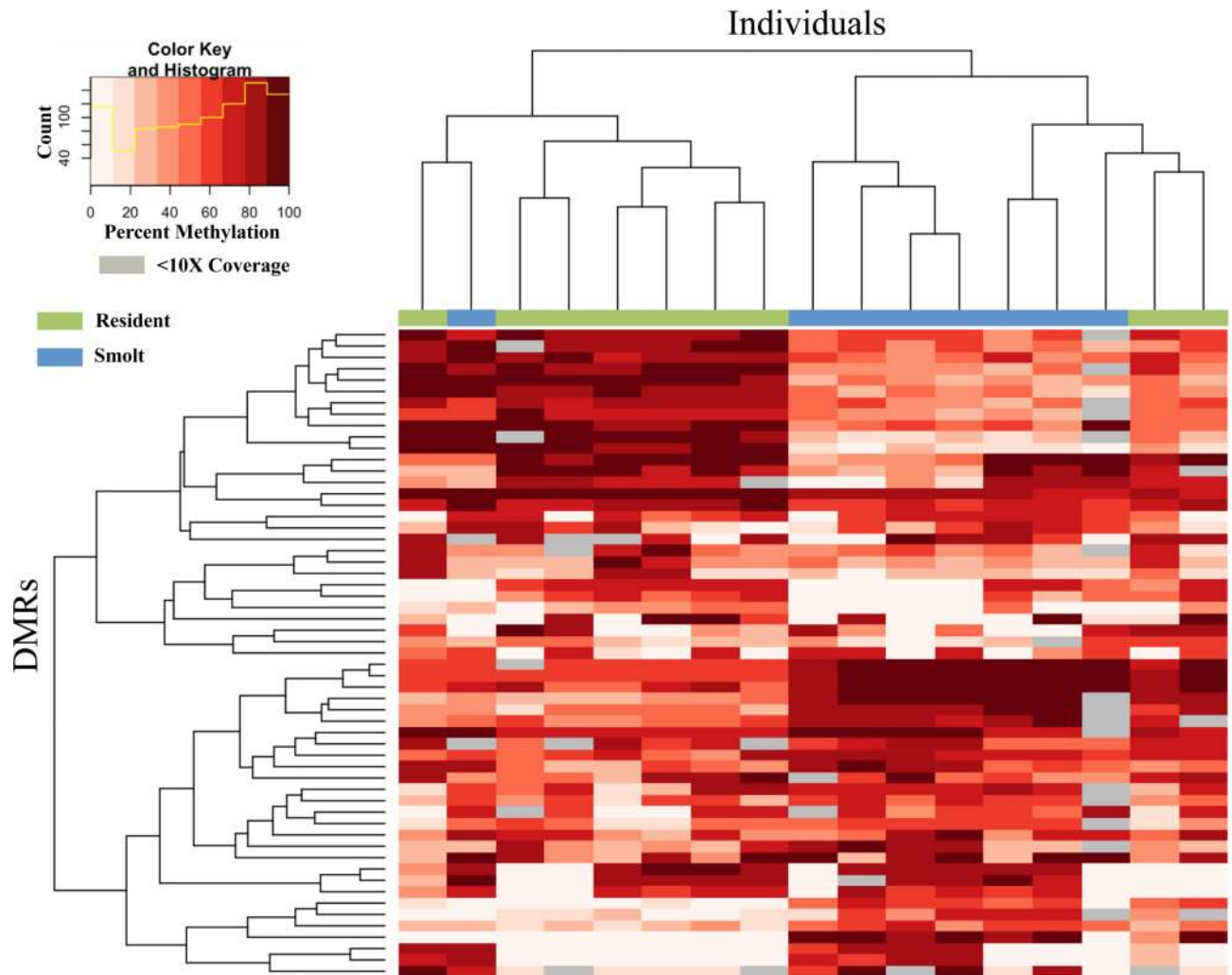


Fig. 2. Hierarchical clustering of hyper- and hypo-methylated DMRs for divergent migration-related phenotypes

This heatmap graphically displays the percent DNA methylation for each individual at the 57 DMRs. Each column represents a color-coded individual: green for residents and blue for smolts. Each row represents a differentially methylated region when comparing the two life history phenotypes. The darker the red, the more methylated that individual is for that DMR. Individual dendrogram positions are based on their overall methylation patterns across the 57 DMRs. DMRs below a minimum read depth of 10 were excluded and are represented as grey boxes.

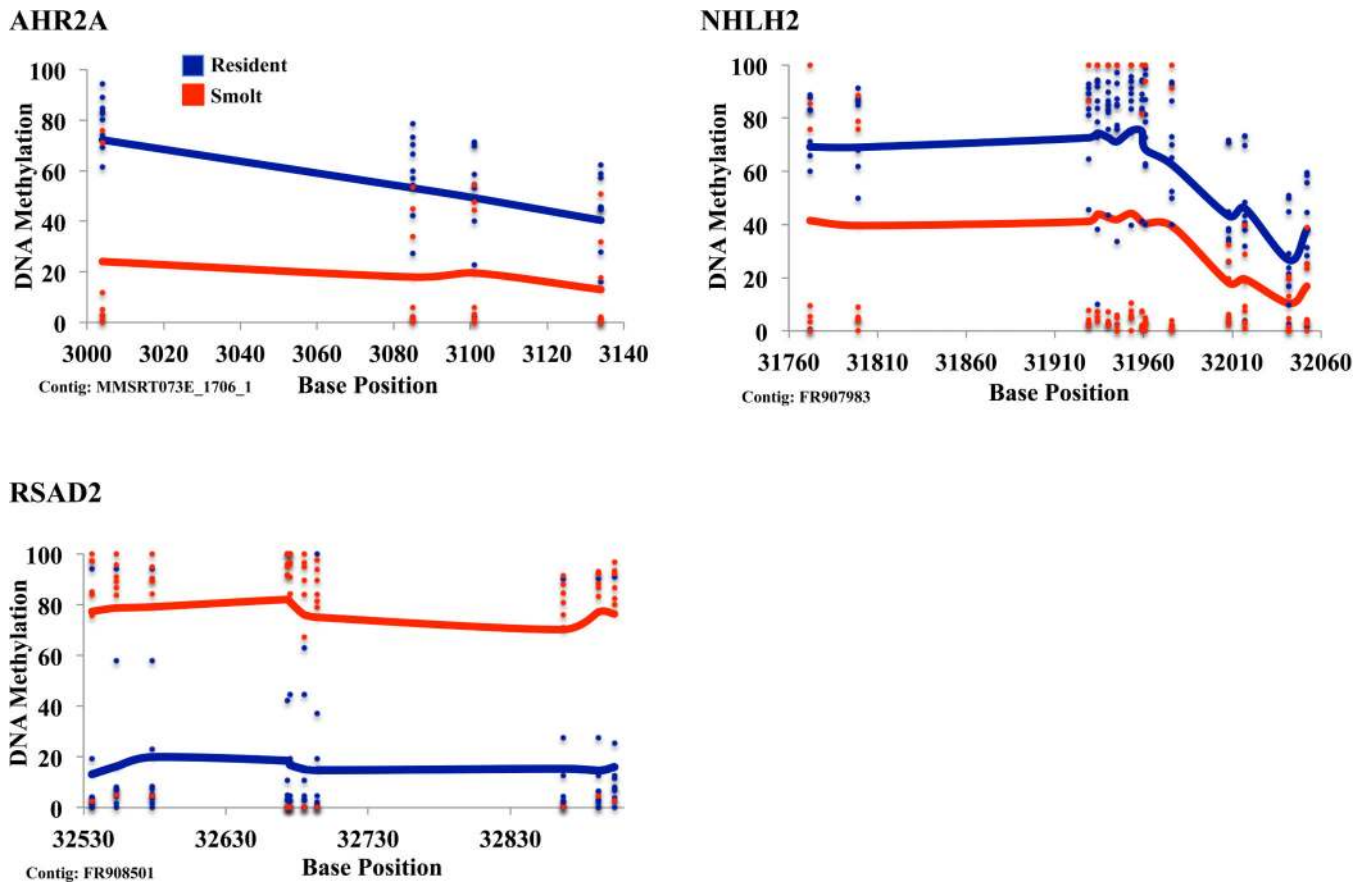


Fig. 3. Differential DNA methylation distinguishes residents and smolts

Three gene-associated DMR examples at the base-pair resolution. Points (blue=resident, red=smolt) represent DNA methylation levels of individual samples at every CpG site within the DMR and smoothed lines represent average DNA methylation levels for each life history type.

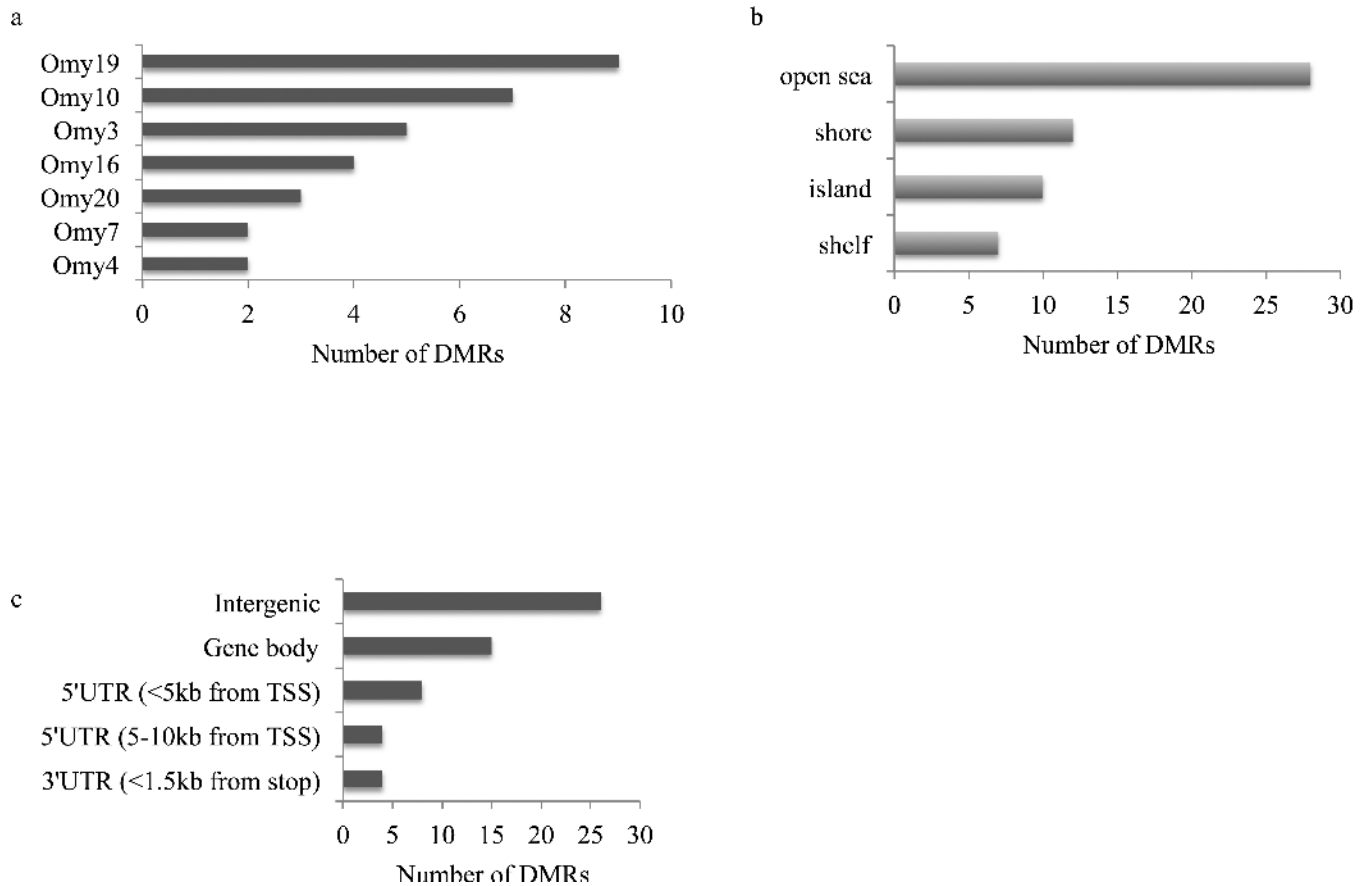


Fig. 4. Genomic features of the 57 migration-related DMRs for *O. mykiss*

(a) Seven chromosomes contain two or more DMRs. (b) Approximately half (29) of the DMRs are found within or proximal to CpG islands. (c) Over half (31) of the DMRs are found within gene bodies or proximal regulatory elements.

Table 1Differentially methylated regions (DMRs) when comparing resident and smolt juvenile *O. mykiss* phenotypes.

Symbol	Entrez gene name	Meth diff ¹	p-value	q-value
AATK	apoptosis-associated tyrosine kinase	-20.7	1.1E-21	1.4E-21
ADHFE1	Alcohol Dehydrogenase, Iron Containing, 1	-38.2	6.2E-29	9.9E-29
ADORA3	adenosine A3 receptor	-23.5	5.4E-45	2.4E-44
AHR2A	aryl hydrocarbon receptor 2 alpha	31.9	1.1E-43	3.6E-43
CCDC93	coiled-coil domain containing 93	20.7	6.8E-18	8.6E-18
CD247	CD247 molecule	22.8	3.2E-34	6.3E-34
CLDN10	claudin 10	29.5	3.0E-36	6.8E-36
CRYBB3	crystallin, beta B3	21.3	1.0E-35	2.1E-35
DDX10	DEAD (Asp-Glu-Ala-Asp) box polypeptide 10	29.5	1.2E-31	2.2E-31
GEMIN2	gem (nuclear organelle) associated protein 2	-34.1	2.5E-30	4.1E-30
GRM1	glutamate receptor, metabotropic 1	-31.3	1.2E-53	1.0E-52
HES6	hes family bHLH transcription factor 6	-20.7	2.2E-14	2.6E-14
KIAA1467	KIAA1467	23.2	1.8E-17	2.2E-17
MYOM3	myomesin 3	-21.7	7.9E-10	8.0E-10
NEUROD4	neurogenic differentiation 4	-31.4	8.3E-36	1.8E-35
NHLH2	nescient helix loop helix 2	38.0	3.2E-76	1.4E-74
PALLD	palladin, cytoskeletal associated protein	24.4	7.2E-62	7.9E-61
PCDHGC5	protocadherin gamma subfamily C, 5	48.8	3.1E-49	1.7E-48
RASD1	RAS, dexamethasone-induced 1	21.5	1.8E-16	2.2E-16
RGL2	ral guanine nucleotide dissociation stimulator-like 2	21.2	1.2E-15	1.3E-15
RSAD2	radical S-adenosyl methionine domain containing 2	-61.8	2.7E-69	5.9E-68
SBK1	serine/threonine-protein kinase SBK1-like	-28.6	7.0E-86	1.9E-84
SETD3	SET domain containing 3	42.5	5.4E-53	4.0E-52
STYK1	serine/threonine/tyrosine kinase 1	28.3	3.5E-40	1.3E-39
TMEM150A	transmembrane protein 150A	-23.6	6.4E-24	8.7E-24
TUBA1A	tubulin, alpha 1a	-26.9	9.1E-34	1.7E-33
-	Uncharacterized protein	42.0	1.2E-30	2.1E-30
-	Uncharacterized protein	-21.1	1.3E-10	1.3E-10
-	Uncharacterized protein	-21.7	2.7E-11	3.0E-11
-	Uncharacterized protein	-31.7	8.3E-36	1.8E-35
-	Uncharacterized protein	-32.3	1.4E-22	1.7E-22
-	Unknown	-21.0	7.9E-29	1.2E-28
-	Unknown	-33.8	1.2E-43	3.6E-43
-	Unknown	20.6	1.1E-09	1.1E-09
-	Unknown	23.7	2.0E-28	3.2E-28
-	Unknown	26.2	2.1E-38	7.1E-38
-	Unknown	-30.0	2.2E-42	8.9E-42
-	Unknown	31.9	8.3E-36	1.8E-35
-	Unknown	29.1	5.3E-46	2.6E-45

Symbol	Entrez gene name	Meth diff ¹	p-value	q-value
-	Unknown	21.3	3.9E-23	5.5E-23
-	Unknown	-20.2	1.4E-31	2.4E-31
-	Unknown	-22.3	6.7E-37	2.0E-36
-	Unknown	-23.7	2.8E-49	1.7E-48
-	Unknown	-24.0	6.8E-38	2.1E-37
-	Unknown	-20.5	1.9E-11	2.1E-11
-	Unknown	30.5	5.3E-23	7.3E-23
-	Unknown	20.6	4.6E-11	4.9E-11
-	Unknown	-21.6	2.1E-23	3.0E-23
-	Unknown	24.4	8.3E-36	1.8E-35
-	Unknown	26.6	6.1E-28	9.3E-28
-	Unknown	-31.3	1.5E-65	2.2E-64
-	Unknown	-31.0	8.3E-36	1.8E-35
-	Unknown	20.1	6.9E-37	1.7E-36
-	Unknown	-23.1	6.1E-33	1.1E-32
-	Unknown	35.4	8.7E-60	1.2E-58
-	Unknown	22.0	8.7E-27	1.2E-26
-	Unknown	-26.3	7.5E-22	1.0E-21

¹Percent methylation differences averaged from all CpG sites within the defined region. The comparison is residents compared to smolts with positive values representing increased methylation for residents and negative values representing increased methylation for smolts.

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