Differential DNA methylation in somatic and sperm cells of hatchery vs wild (natural-origin) steelhead trout populations

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Abstract

Environmental factors such as nutrition, stress, and toxicants can influence epigenetic programming and phenotypes of a wide variety of species from plants to humans. The current study was designed to investigate the impacts of hatchery spawning and rearing on steelhead trout (Oncorhynchus mykiss) vs the wild fish on a molecular level. Additionally, epigenetic differences between feeding practices that allow slow growth (2 years) and fast growth (1 year) hatchery trout were investigated. The sperm and red blood cells (RBC) from adult male slow growth/maturation hatchery steelhead, fast growth/maturation hatchery steelhead, and wild (natural-origin) steelhead were collected for DNA preparation to investigate potential alterations in differential DNA methylation regions (DMRs) and genetic mutations, involving copy number variations (CNVs). The sperm and RBC DNA both had a large number of DMRs when comparing the hatchery vs wild steelhead trout populations. The DMRs were cell type specific with negligible overlap. Slow growth/maturation compared to fast growth/maturation steelhead also had a larger number of DMRs in the RBC samples. A number of the DMRs had associated genes that were correlated to various biological processes and pathologies. Observations demonstrate a major epigenetic programming difference between the hatchery and wild natural-origin fish populations, but negligible genetic differences. Therefore, hatchery conditions and growth/maturation rate can alter the epigenetic developmental programming of the steelhead trout. Interestingly, epigenetic alterations in the sperm allow for potential epigenetic transgenerational inheritance of phenotypic variation to future generations. The impacts of hatchery exposures are not only important to consider on the fish exposed, but also on future generations and evolutionary trajectory of fish in the river populations.

Key words: steelhead; hatchery; rearing conditions; epigenetic inheritance; fish; genomics; phenotypic variation; sperm; generational

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**Introduction**

Epigenetics is a complementary mechanism with genetics for the molecular control of biology [1, 2]. A range of environmental factors including toxicants, stress, and nutrition can developmentally alter a variety of phenotypes in species from plants to humans through epigenetics [3]. Epigenetics is defined as “molecular factors and processes around DNA that regulate genome activity independent of DNA sequence, and are stable mitotically” [3, 4]. The ability of environmental factors to alter epigenetic programming, while not changing DNA sequence, provides a molecular mechanism for the environment to directly impact phenotypic variation and evolution [1, 3]. Although direct exposures of somatic cells to environmental factors can influence the individual exposed, epigenetic change in the germline (sperm or egg) can be transmitted to the next generation, and is termed “epigenetic inheritance” [2, 3]. In the event, the germline epigenetic alterations are transmitted to subsequent generations, in the absence of continued direct exposure, this is defined as “epigenetic transgenerational inheritance” [3, 5]. Therefore, environmental factors have the ability to developmentally impact epigenetic programming to influence the phenotype of the individual exposed. In addition, if the germline (e.g. sperm) is affected the potential for generational effects on phenotypic variation develop. The current study is designed to investigate the molecular effects of hatchery rearing on steelhead trout (*Oncorhynchus mykiss*).

Hatcheries involve both aquaculture facilities for food supply and operations for sustaining sufficient numbers of endangered fish such as the salmon populations in the Pacific Northwest, USA [6, 7]. Research shows that hatchery-reared fish differ from wild fish both in phenotype and in having reduced reproductive success in the wild [8–11]. In steelhead trout (*Oncorhynchus mykiss*), it has been shown that the offspring of fish experiencing even a single generation of hatchery rearing show marked phenotypic changes [8–22] (Table 1). This has also been observed in a variety of different hatchery salmon and trout species [12, 23–45] (Table 1). Pathologies observed include decreased fitness of hatchery-reared fish and offspring in the wild, and changes in age at spawning, morphology, growth rate, brain morphology, anti-predator behavior, and migration [8–45].

A model anadromous migratory salmonid fish species selected for the current study is the steelhead trout. A brood-stock hatchery operation (Winthrop National Fish Hatchery, WNFH, US Fish and Wildlife Service) on the Methow River in Washington State, USA was selected to compare wild natural-origin populations and hatchery populations of steelhead trout (Fig. 1). This hatchery used hatchery-reared steelhead crossbred with natural-origin fish from the same river as broodstock to produce offspring that were raised in hatcheries until the time of release as smolts, at which point they can migrate downstream to the ocean. Some steelhead were placed on a high level (plane) of nutrition and reached smolt stage within 1 year (fast growth/maturation), at which time they were released. Others were placed on a lower level (plane) of nutrition, and took 2 years to reach smolt stage (slow growth/maturation) prior to release. Natural-origin steelhead trout generally take 2 years to reach smolt stage in this river system. After returning from the ocean, fast growth/maturation (S1), slow growth/maturation (S2), and natural-origin (N) fish were collected and sampled.

The Methow River summer-run steelhead is part of an upper Columbia River evolutionary significant unit currently listed as threatened under the Endangered Species Act. For the past 60 years, the large number of hatchery fish released that breed with wild fish has had a significant influence on the wild steelhead population. Therefore, there is no distinct wild Methow River steelhead population without a molecular influence from hatchery impacts. The term natural-origin is used to refer to fish spawned in the wild and live their entire lives in the wild. Hatchery fish refers to fish generated through artificial rebreeding and crosses in the juvenile stages in the hatchery before being released into the wild with an adipose fin clip. In the current study, the wild (natural-origin) and hatchery (slow 2 years and fast 1-year juvenile growth/maturation) are compared. Natural and hatchery-origin adult steelhead returning in the Methow river in 2013 and 2014 and captured in winter and spring of 2014 and 2015 were used in this study (Supplementary Table S1).

The molecular effects of hatchery rearing and growth rate on somatic cells (red blood cells, RBC) and germline cells (sperm) were investigated. Genome-wide molecular analyses of differential DNA methylation regions (DMR) and genetic mutations (copy number variation, CNV) were performed. The hypothesis tested was that the hatchery spawning and rearing conditions alter the epigenetic programming of the steelhead somatic cells (RBC) and germline (sperm), such that later in life following return the adult steelhead will have altered epigenetics, with potential generational impacts through the sperm. This will result in an altered phenotypic variation, fitness, and evolutionary trajectory of the wild population.

**Results**

The steelhead trout was selected as a model salmonid fish species for the current study due to the migratory nature of the fish and availability of the trout genome sequence to facilitate molecular studies. The adult phenotypes and color after migration and return from the Pacific Ocean are generally similar between the hatchery and wild steelhead populations, as shown for the hatchery fish in Fig. 1. The hatchery was collected on the Methow River near Winthrop Washington, USA, during spring following the previous summer migration (Fig. 1a). A steelhead hatchery is located in Winthrop Washington that uses a broodstock population of adult river caught fish each year to spawn and rear the steelhead. An alternate fish hatchery also used was at Wells Dam below the mouth of the Methow River on the Columbia River (Fig. 1 and Supplementary Table S1). During collection, both hatchery (determined by a fin clip) and wild (natural-origin) steelhead were obtained. Since each cell type in the body has a unique epigenome (e.g. DNA methylation patterns), purified cell types are required for epigenetic analysis to allow unambiguous data interpretation. For the current study, both sperm and purified RBC (that contain nuclei in fish) samples were collected from the males. The DNA was isolated from the cell types obtained from adult male hatchery and wild steelhead (Fig. 1b). The isolated DNA from the RBC and sperm samples were then processed for the molecular studies.

The DMRs between the hatchery and wild male steelhead populations were identified in both the sperm and RBC separately. For each treatment group, equal amounts of DNA from three to five different males were pooled and three pools created for n = 9–15 animals per group. The treatment groups comprised fish from the Winthrop National Hatchery-fast growth/maturation (WNFH-S1; i.e. S1), Winthrop National Hatchery-slow growth/maturation (WNFH-S2; i.e. S2), and Winthrop National Hatchery standard hatchery (WNFH-H; i.e. H), and non-hatchery reared natural-origin wild fish (N). Information on the individual fish collections, characteristics, and labeling are presented.
in Supplementary Table S1. The information includes sample group, identification number, fly tag number, sex (males), date collection, natural or hatchery type, hatchery location, approximate age at collection, rearing (growth) strategy, and sample pool for each individual (Supplementary Table S1). Individual males used for the study had no difference in the length or age between the groups (Supplementary Table S1). Equal amounts of DNA from each individual upon collection was used in the hatchery or wild population pools, and then the DNA was fragmented through sonication and the methylated DNA immunoprecipitated (MeDIP) with an antibody to methylcytosine, as described [46] in the Methods. The MeDIP DNA was used to prepare libraries for next-generation sequencing (Seq) for an MeDIP-Seq analysis to identify the DMRs, see Methods. This genome-wide epigenetic analysis assesses >90% of the genome compared to other procedures such as reduced representation bisulfite sequencing (RRBS) that assesses <10% of the genome [47]. The sperm and RBC DNA samples were analyzed separately with MeDIP-Seq.

A comparison of the standard hatchery WNFH-H (H), fast 1-year growth/maturation WNFH-S1 (S1), slow 2-year growth/maturation WNFH-S2 (S2), and wild (natural-origin) (N) populations involved 2 years of collection with N1 in the first year collection for standard hatchery H vs N1 and a distinct collection of N2 in the second year collection for the S1 vs N2 and the S2 vs N2 comparisons. Therefore, these analyses were done separately. Summary of the individual fish and characteristics is presented in Supplementary Table S1. The MeDIP-Seq procedure was used to identify the DMRs between the group comparisons (Table 2). The analyses at several different statistically significant P values are shown for single sites (1000 bp window), and for multiple adjacent window sites (<2 1000 bp windows) with each being statistically significant with edgeR and false discovery rate (FDR), as described in the Methods. As the statistical threshold decreases (P-value) the number of identified DMRs decreases, as expected, and the edgeR P < 0.05 significance level was selected for subsequent use and data presentation. This generally correlated to an FDR of P < 0.05. Although the data analysis focused on the most stringently selected DMRs, the other DMRs at a lower statistical threshold are anticipated to also be important, but more variable between individuals. The DMR numbers for the sperm and RBC are presented in Table 2. For DMRs with multiple adjacent windows, the most predominant number of adjacent sites is 2 (1000 bp each) with the highest number of adjacent sites being 8. Therefore, the more stringently identified DMRs (all sites at P < 0.05) provide a reasonable set of DMRs (i.e. signature) to be used for further analysis. The lists of DMRs and genomic features comparing hatchery vs wild steelhead for the sperm are presented in Supplementary Table S2, and for the RBC in Supplementary Table S3.

A major hatchery condition that can be altered is the nutrition and amount of food intake that can promote a fast growth 1-year juvenile maturation or a more normal slow growth 2-year juvenile maturation. A group of WNFH fast (S1) and slow (S2) growth/maturation steelhead were compared with the wild natural-origin (N) groups (i.e. N1 and N2). Both the N2 vs S1 or S2 comparisons are presented in Table 2 for the identification of DMRs. Using a P < 1e−05 statistical threshold, similar numbers of DMRs were obtained for the wild natural-origin (N2),
hatchery-S1, or hatchery-S2 comparisons. An overlap of these DMR sets is shown in Fig. 2b for sperm and Fig. 3b for RBC. A comparison of the hatchery S1 vs hatchery S2 is also provided with a higher number of DMRs for RBC and lower number for sperm (Table 2). The Venn diagram overlaps of all the comparisons (Figs 2b and 3b) suggests each group comparison was primarily distinct with the largest overlaps between the N2 vs S1 and N2 vs S2, as well as the S1 vs S2 and N1 vs H. Therefore, at \( P < 1 \times 10^{-5} \), the DMRs are primarily distinct for each comparison.

The chromosomal locations of the DMRs for the sperm are presented in Fig. 2a and for the RBC in Fig. 3a. The DMRs are present on all of the annotated steelhead trout genome chromosomes for both sperm and RBC DMRs. The red arrowheads indicate the locations of the DMRs. Previous studies have demonstrated that DMRs can cluster on the genome to give statistically over-represented groups of DMRs [48]. These clusters of DMRs are shown in Figs 2a and 3a as black boxes on the chromosomes, and may function as epigenetic control regions [49]. Therefore, the DMRs are present throughout the genome and not isolated to specific regions or chromosomes. The RefSeq version of the Omyk 1.0 steelhead trout genome was used and obtained from NCBI. The read alignment rate was generally 76–83% for the steelhead sequencing with \(~55–100\) million reads per pool for comparison. The steelhead trout genome is now nearly fully assembled, but some of the sequence is still in unplaced contigs not localized to a specific chromosome. In Figs 2 and 3, chromosomes marked as \#1–\#29 and mitochondrial DNA (MT) all have DMRs mapped to chromosomes. The number of DMRs associated with unplaced contigs is listed in the figure legends. Therefore, the lack of a complete genome annotation provides an underestimate of the genome map findings, but is sufficient to allow interpretation of the majority of the data.

The chromosomal locations of the wild (N2) vs hatchery S1 (i.e. 1-year fast growth/maturation) or S2 (i.e. 2-year slow growth/maturation), and the hatchery S1 vs S2 are shown in Fig. 4a–d and in Supplementary Fig. S1. The lists of DMRs are presented in Supplementary Tables S2–S9 with genome location, statistical edgeR \( P \) values, log-fold change \([\text{[+]}]\) indicating an increase in

Figure 1: Hatchery locations, rivers, adjacent dams, and steelhead trout. (a) Map of Methow River and Columbia River confluence, and Winthrop hatchery. (b) Steelhead males (hatchery-origin)
| (a) Sperm wild (N1) vs Hatchery DMRs | P-value | All sites | Multiple windows |
|-------------------------------------|---------|-----------|------------------|
| 0.001                               | 4352    | 217       |
| 1e−04                               | 1328    | 83        |
| 1e−05                               | 577     | 34        |
| 1e−06                               | 296     | 14        |
| 1e−07                               | 167     | 8         |
| Number of significant sites         | 1       | 2         | 3                |
| Number of DMR                       | 543     | 32        | 2                |

| (b) Sperm Wild (N2) vs Hatchery S1 DMRs | P-value | All sites | Multiple windows |
|----------------------------------------|---------|-----------|------------------|
| 0.001                                  | 7332    | 323       |
| 1e−04                                  | 1689    | 71        |
| 1e−05                                  | 454     | 24        |
| 1e−06                                  | 149     | 13        |
| 1e−07                                  | 64      | 6         |
| Number of significant sites            | 1       | 2         | ≥4               |
| Number of DMR                         | 430     | 21        | 3                |

| (c) Sperm Wild (N2) vs Hatchery S2 DMRs | P-value | All sites | Multiple windows |
|----------------------------------------|---------|-----------|------------------|
| 0.001                                  | 6253    | 283       |
| 1e−04                                  | 1811    | 109       |
| 1e−05                                  | 693     | 56        |
| 1e−06                                  | 313     | 35        |
| 1e−07                                  | 167     | 22        |
| Number of significant sites            | 1       | 2         | ≥3               |
| Number of DMR                         | 637     | 46        | 10               |

| (d) Sperm Hatchery S1 vs S2 DMRs      | P-value | All sites | Multiple windows |
|---------------------------------------|---------|-----------|------------------|
| 0.001                                 | 2560    | 56        |
| 1e−04                                 | 573     | 26        |
| 1e−05                                 | 194     | 11        |
| 1e−06                                 | 102     | 4         |
| 1e−07                                 | 63      | 3         |
| Number of significant sites           | 1       | 2         | 5                |
| Number of DMR                         | 183     | 10        | 1                |

| (e) RBC Wild (N1) vs Hatchery DMRs    | P-value | All sites | Multiple windows |
|--------------------------------------|---------|-----------|------------------|
| 0.001                                | 6795    | 609       |
| 1e−04                                | 1754    | 113       |
| 1e−05                                | 509     | 27        |
| 1e−06                                | 182     | 8         |
| 1e−07                                | 79      | 3         |
| Number of significant sites          | 1       | 2         | 3                |
| Number of DMR                        | 482     | 22        | 5                |

continued
DNA methylation, and (−) indicating a decrease in DNA methylation] between the comparisons, length (kb), CpG density, and gene associations. Although a mixture of DNA methylation increases and decreases at various DMRs were observed for all analyses, the sperm generally had a higher number of increases in DNA methylation at DMR (Supplementary Tables S2–S12).

Analysis of potential similarities in the natural (N1) vs hatchery (H) DMRs (WNFH-H vs N1) at $P < 1 \times 10^{-5}$ demonstrated only 25 DMRs with overlap. The list of these overlapped DMRs and associated genes is presented (Supplementary Table S10). The most common associated gene sites for both sperm and RBC were uncharacterized LOC, but genes involved in signaling and epigenetics were present. Although the DMR was primarily cell specific at the threshold of $P < 1 \times 10^{-5}$, an expanded sperm and RBC overlaps are presented in Fig. 5 to determine if greater overlap was present at a reduced statistical threshold. Interestingly, an extended overlap comparing the $P < 1 \times 10^{-5}$ DMRs with DMRs at $P < 0.05$ identified a greater degree of overlap among all the comparisons (Fig. 5c). Generally, a 15–30% overlap was observed unless the S1 and S2 populations were involved, which often had a 45–65% overlap. Therefore, using a lower statistical threshold allowed higher overlap of the DMRs between the comparisons. The altered S1 and S2 hatchery growth had an increased variation and allowed greater overlaps in DMRs. The N2 vs S1 and N2 vs S2 had an over 70% overlap in this comparison (Fig. 5c). The N1 vs H sperm had less overlap, but the RBC N1 vs H had ~50% overlap with the S1 and S2 RBC. Therefore, the hatchery growth/maturation comparisons indicated DMR had good overlap for the RBC, but were more distinct for the sperm DMRs.

Previous studies have demonstrated that MeDIP-Seq identified DMRs generally have low-density CpG content, and exist in CpG deserts [50]. Analysis of the CpG density of the DMRs identified in the current study also demonstrated the DMRs to have a low CpG density (Figs 2e and 3e and Supplementary Fig. S2). The predominant CpG density in the data sets was one or two CpG per 100 bp for both the sperm and RBC. In the CpG deserts of a few thousand bases, the CpG can cluster to presumably act as regulatory sites [50]. The size of the DMRs was found to be predominantly one or two thousand bases Figs 2f and 3f and Supplementary Fig. S2. A few DMRs were between 5 and 10 kb, but the majority were smaller. Similar observations were made with the other comparison of slow and fast growth (N2 vs S1, N2 vs S2, S1 vs S2) for ($P < 1 \times 10^{-5}$) DMRs, as presented in Supplementary Fig. S2. Therefore, the DMR genomic features identified were similar to DMRs previously characterized [3, 50, 51]. This is significant since the MeDIP-Seq procedure used in the current study optimally assessed lower density CpG deserts, while other procedures based on bisulfite (i.e. RRBS) assesses only a small percentage of the genome (<10%) and is optimal for high-density CpG sites.

A permutation analysis was performed to help verify the significance of the DMRs identified. This analysis involves randomly shuffling the samples between groups to obtain a null distribution for the number of DMRs expected due to random

| Table 2. (continued)   |
|------------------------|
| (f) RBC Wild (N2) vs Hatchery S1 DMRs |
| P-value | All sites | Multiple windows |
| 0.001 | 5596 | 206 |
| 1e−04 | 1268 | 47 |
| 1e−05 | 389 | 14 |
| 1e−06 | 175 | 6 |
| 1e−07 | 73 | 2 |
| Number of significant sites | 1 | 2 | 4 |
| Number of DMR | 375 | 12 | 2 |

| (g) RBC Wild (N2) vs Hatchery S2 DMRs |
| P-value | All sites | Multiple windows |
| 0.001 | 4978 | 129 |
| 1e−04 | 1191 | 46 |
| 1e−05 | 398 | 26 |
| 1e−06 | 170 | 17 |
| 1e−07 | 77 | 9 |
| Number of significant sites | 1 | 2 | ≥3 |
| Number of DMR | 372 | 23 | 3 |

| (h) RBC Hatchery S1 vs S2 DMRs |
| P-value | All sites | Multiple windows |
| 0.001 | 11611 | 665 |
| 1e−04 | 3556 | 188 |
| 1e−05 | 1097 | 61 |
| 1e−06 | 438 | 18 |
| 1e−07 | 193 | 9 |
| Number of significant sites | 1 | 2 | ≥3 |
| Number of DMR | 1036 | 54 | 7 |
chance. Internal population epigenetic variation is generally as-
sociated with hypervariable or metastable DMRs, as previously
described [52, 53]. There was generally a higher number of
DMRs in the full analysis when compared to the sperm and RBC
DMRs from the permutation comparisons of the hatchery and
wild populations (Figs 2d and 3d and Supplementary Fig. S3).
Two of the comparisons, sperm N2 vs S1 and sperm S1 vs S2,
did not show a higher number of DMRs with the full analysis
(Supplementary Fig. S3C and G). Although only a six-pool sub-
comparison is not ideal for a thorough permutation analysis,
the sperm with the S1 and S2 populations do show a higher de-
gree of variation. The DMR analysis was also supported with a
principal component analysis (PCA) of the DMRs, which demon-
strated good separation of the various groups for all compari-
sons (Figs 2c, 3c, and 4e–h, and Supplementary Fig. S4. Since
pools of animals were used, it is difficult to determine if the
Fig. 1 parameters such as age, size, hatchery, or collection data
create variability in the data, however, the PCA and permuta-
analyse indicate the DMR identified for the group compar-
isons is not a major factor. Within the specific groups, the
variation shown for each group in the PCA may be associated
with the variables shown in Supplementary Table S1.

The potential genetic variation between the hatchery and
wild steelhead populations was also investigated. A type of ge-
netic mutation previously shown to be one of the most
common and stable is CNVs of repeat elements [51, 54, 55].
Previously, we have shown the ability to use CNV as a mea-
sure of genetic variation in wild populations [56]. Sequencing
read depth data from the WNFH-H and N (i.e. N1) groups was
used for a CNV analysis (CNV-Seq). The CNV for the hatchery
and natural-origin wild populations for both the sperm and
RBC are presented in Supplementary Fig. S5. CNV analysis did
not show either hatchery or natural-origin samples to have
more CNVs than the other. The sperm had higher numbers of
CNVs than the RBC. None of the CNVs were common between
the different animal pools. Therefore, in contrast to the epige-
netic changes observed, negligible genetic changes between
the hatchery and wild populations were observed. A limitation
in this analysis is the read depth only allowed the identifica-
tion of large size CNV as presented. The negligible effect of
hatchery conditions on genetics has been previously reported
[57, 58].

The potential gene associations with the DMRs were investi-
gated using the steelhead trout genome annotations included
in the RefSeq genome. The DMR associated genes are presented
in Supplementary Tables S2–S9 for all the comparisons. The
majority of DMRs did not have known gene associations. The
cell-specific DMR associated gene functional categories are pre-
sented in Fig. 6a for sperm and Fig. 6b for RBC. The most pre-
dominant gene classification categories associated with all the

![Figure 2: sperm DMR N1 vs H chromosomal locations and analysis. (a) DMR chromosomal locations on the individual chromosomes vs size of chromosomes. All DMRs at a P-value threshold of $P < 1 \times 10^{-5}$ are shown with red arrowheads and clusters DMRs with black boxes. In addition, 212 DMRs were located on the unplaced concatenated scaffolds not shown. (b) Sperm comparisons DMR overlaps, $P < 1 \times 10^{-5}$. (c) Principal component analysis (PCA) of sperm DMRs N1 vs H. (d) Permutation analysis of sperm DMRs N1 vs H. (e) The number of sperm DMRs at different CpG densities. All DMRs at a P-value threshold of $1 \times 10^{-5}$ are shown. (f) The DMR lengths (kb) for sperm DMRs. All DMRs at a P-value threshold of $1 \times 10^{-5}$ are shown.](https://academic.oup.com/eep/article/7/1/dvab002/6274935)
group comparisons are transcription, metabolism, signaling, transport, development, protease, and epigenetics (Fig. 6). These gene categories are anticipated as they are the most abundant in most species and cell-type genomes. When the hatchery vs natural genes are put into a KEGG pathway analysis, the metabolic pathways (DCXR, ITPA, and PLCB3) and cytokine-cytokine receptor interactions (IL15, IL21) pathways were identified for the sperm, and proteasome (psme9a, psme1) for the RBC DMR associated genes (Fig. 6b). Limited numbers of DMR associated genes were present in the pathways, except for the S1 vs S2 comparison. Therefore, further analyses used a combination of all the comparison DMR associated genes.

The DMR associated genes were analyzed for physiological processes and pathology correlations using the steelhead annotations provided by NCBI. The predominant physiological processes were cell death, angiogenesis, and cell differentiation for sperm DMR associated genes (Fig. 7). Cell differentiation, cell death, and medulla development are predominant for RBC DMR associated genes (Fig. 8). The predominant sperm DMR gene-associated pathologies were intellectual disability, cutaneous discoloration, mitochondrial disease, and optic atrophy in sperm DMR associated genes (Fig. 7). Pathologies of hemoglobin disease, dystrophy, and anaplastic large cell lymphoma in RBC DMR associated genes were predominant (Fig. 8). Additional cellular processes and pathway correlations with statistical significance are presented in Supplementary Table S11 for the sperm DMR associated genes and in Supplementary Table S12 for the RBC DMR associated genes. The processes or pathologies are listed with the total number of genes, the overlap number of genes, and the $P$-value significance of the correlation is presented. Each of the different comparisons’ DMR associated genes are combined to identify the potential associated processes and pathologies correlated with the hatchery and rearing conditions.

**Discussion**

Observations indicate hatchery spawning and rearing induces epigenetic alterations when compared to a wild (natural-origin) populations of steelhead trout. In addition, feeding and growth rate (1-year fast growth S1 vs 2-year slow growth S2) of juveniles results in altered epigenetic programming in adulthood. The hatchery conditions impose nutritional, behavioral, or other types of stressors on the fish that can developmentally promote altered epigenetic programming and phenotypic variation. The exposure of somatic cells leading to epigenetic (e.g. DNA methylation) change may impact the exposed individuals’ health and phenotypes. Therefore, the hatchery populations can have reduced fertility, abnormal health, and survival when compared to the wild populations (Table 1). Since environmental exposures generally cannot directly alter genetic mutations, epigenetics provides a molecular mechanism for the physiological
changes observed. In the event, the hatchery induced epigenetic alterations (i.e. epimutations) that appear in the germline (egg or sperm), then the impacts may persist across generations. Previously, the transmission of epigenetic information through the germline to alter a variety of phenotypes in a number of different species, including fish, have been observed [3]. This epigenetic inheritance can influence the next generation’s phenotypes and health. If the germline epigenetic changes persist between generations in the absence of environmental exposures, then this is termed epigenetic transgenerational inheritance [3, 5]. Therefore, the hatchery impacts may not only be on the exposed individuals, but on subsequent generations. This has the potential to dramatically impact the wild natural-origin populations and their evolutionary trajectory. The current study demonstrates the ability of hatchery conditions to promote epigenetic alterations in both somatic cells (RBC) and germ cells (sperm), so both directly exposed fish populations and future generations need to be considered.

Previous studies have demonstrated hatchery impacts on fish phenotypic alterations and health (Table 1). Hatchery rearing has been shown to decrease reproductive success in steelhead trout by about 40% per captive-reared generation in the first two generations when fish are moved back to natural environments [8, 13]. Hatchery rearing of salmonids affects lateral line and auditory structures [11]. Phenotypic changes and decreases in fitness can occur even when wild fish are incorporated as broodstock in hatchery operations [9]. Although the relationships of the molecular epigenetic alterations and DMRs found in the current study cannot currently be causally related directly to the phenotypes, hatchery-induced epigenetic alterations (i.e. epimutations) provide a potential molecular mechanism for the phenotypic variation and health effects previously observed.

The hatchery fish are exposed to hatchery conditions and reared through development to the smolt stage, so all early development periods are impacted. Primordial germ cells (PGCs) in fish species, including salmonids, are specified by the localization of germplasm components into certain cells early in embryonic development [59]. These germplasm components include the products of the vasa and nanos genes [59]. PGCs migrate to the gonadal anlagen, then undergo mitotic proliferation and differentiate to become either spermatogonia or oogonia [59]. The PGC is a critical stage for the male germline when dramatic epigenetic reprogramming occurs. Puberty in male salmonids is associated with growth and size, photoperiod, and hormones including 11-ketotestosterone [60]. Once spermatogenesis is initiated, spermatogonia are surrounded by Sertoli cells to form cyst-like structures (spermatocysts) inside which the germ cells undergo mitotic divisions, produce spermatocytes that undergo meiotic divisions to form spermatids, which then undergo spermiogenesis to produce spermatozoa. The spermatozoa are released from the spermatocysts into the testis tubules [61]. The earlier stages of spermatogenesis are regulated by follicle-stimulating hormone (FSH) from the pituitary, while the later stages are regulated by luteinizing hormone (LH) [62]. Any of these developmental stages may be impacted by environmental factors that result in epigenetic changes to the germ cells.

There is evidence that epigenetic shifts during early life play a significant role in directing the life history and phenotypes of fish during adulthood. This is clearly demonstrated in cases of temperature-dependent sex determination, as seen with sea
bass [63]. In Atlantic salmon, the temperature of embryonic development affects growth rate of the smolt stage, and is associated with larval myogenin expression and DNA methylation levels [64]. Similarly, stressors during embryonic and larval stages for salmon induce changes in the immune response at 4 months of age that are associated with changes in DNA methylation in gill tissue [65]. These early developmental impacts on later-life health and disease involve the developmental origin of health and disease (DOHAD) mechanisms observed in many species, including humans. Epigenetics is one of the primary molecular mechanisms involved in this phenomenon.

In the current study, the epigenetic alterations observed were DMRs identified in the hatchery steelhead somatic and germline cells, compared to natural-origin populations. The presence of a reproducible DMR is termed an epimutation [1, 3, 4]. The molecular procedure used to identify the DMRs was methylated DNA immunoprecipitation (MeDIP) followed by Seq for an MeDIP-Seq analysis [66]. The MeDIP procedure is biased to low-density CpG analysis (<10% CpG) [67], which constitutes ~95% of the genome [68]. In contrast, due to the informatics and alignment issues, a bisulfite procedure (RRBS) can be biased to higher density CpG analysis (>10%), which constitutes ~5% of the genome [47]. Therefore, the current study used MeDIP-Seq to capture the majority of the genome, while previous studies have used bisulfite (i.e. RRBS) procedures [58, 69]. The DMRs identified in the current study had low CpG content, and exist primarily in CpG deserts [50]. Evolutionarily, CpG methylation has been shown to increase susceptibility for C to T conversions, and leads to regions of the genome with low CpG content. Small clusters of CpG in these CpG deserts were likely conserved due to regulatory roles for these CpG clusters [50]. The genomic locations of the DMRs also demonstrate a genome-wide distribution on most chromosomes. Although the assembly of the steelhead trout genome is not complete, it has been significantly improved recently and the most recent reference genome (Oncorhynchus mykiss, Omyk_1.0, accession GCF_002163495.1, https://www.ncbi.nlm.nih.gov/assembly/GCF_002163495.1), was used in the current study. The comparison of the hatchery and wild (natural-origin) steelhead populations identified a large number of single-site DMRs and a smaller set of DMRs using a more stringent selection of adjacent site DMRs. Therefore, the epimutations identified in this study appear to be due to alterations in epigenetic programming between the hatchery and wild natural-origin steelhead populations. Although variables such as age, size, and hatchery conditions will impact an individual’s epigenome, these variables were controlled for in this study to the extent possible. The permutation analyses and PCA plots did not identify any obvious sources of unexplained variability.
In the current study, comparisons were also made between hatchery-reared steelhead fed at a high plane of nutrition and reaching smolt stage within 1 year (fast-growth; WNFH-S1), and hatchery-reared steelhead fed less and reaching smolt stage within 2 years (slow-growth; WNFS-S2), or non-hatchery reared wild natural-origin steelhead that typically take 2 years to mature to the smolt stage (N). Similar to what was found when comparing hatchery-reared to wild steelhead trout, the DMRs identified occurred primarily in CpG deserts, and a large proportion of them was present in intergenic regions. A comparison of the slow and fast growth/maturation hatchery conditions also demonstrated similar genomic features in the DMRs identified. Therefore, the epimutation characteristics found in the current study are similar to those previously identified in other species following environmental exposures [1, 3, 70].

Genetic analysis in the current study used an evaluation of CNVs to estimate gene sequence similarity between treatment groups. Observations indicate there was negligible genetic variation between these populations. Negligible genetic mutations have been previously observed between the hatchery and wild populations compared [57, 58, 71]. This is to be expected in the Methow River fisheries system examined, as hatchery-reared broodstock are crossed with stream-raised “wild” natural-origin steelhead at each generation. In addition, hatchery-reared fish can spawn with wild fish in the river. This blends the genetic backgrounds of the populations examined. These observations suggest hatchery or feeding rate-induced epigenetic alterations may have a significant role in the phenotype and health differences between the populations. However, as with any biological system, the integration of genetics and epigenetics will be required to fully realize the phenotypic variation and

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**Figure 6**: DMR associated gene categories. The gene classification is listed and correlated to the number of DMR associated genes within the specific classification category for (a) sperm DMR associated gene categories and (b) RBC DMR associated gene categories. (c) Pathways and processes with multiple genes.
environmentally induced health effects. The initial environmentally induced epigenetic alterations are anticipated to generationally lead to genetic changes, as previously described [72], such that an integration of both molecular mechanisms is involved.

The DMR associated genes indicate the epigenetic alterations observed may have major effects on genome activity, potentially correlating to the phenotypic alterations known to occur (Table 1). The majority of DMRs did not have gene associations, suggesting distal regulation of genomic activity may be involved, as previously suggested for other species [48]. For those DMRs with associated genes, a large number of gene classification categories were implicated in the hatchery vs natural-origin steelhead studies including transcription, development, protease, transport, immune, epigenetic, metabolism, and hormone (Fig. 6). Specific cellular pathways were also identified, but only 2–3 genes were represented in each pathway. A large number of cellular processes and pathologies were correlated with the DMR associated genes for both the sperm and RBC (Supplementary Tables S11 and S12 and Figs 6–8). Observations suggest a number of different physiological processes and pathologies may potentially be affected by the epigenetic DNA methylation alterations observed. These potential alterations in genome activity are speculated to correlate to the phenotypic variation and health impacts of the hatchery steelhead populations, but future research is needed to identify causal relationships.

Accelerated growth and maturation rates in the hatchery promoted the highest level of epigenetic changes in the mature fish. The 1-year S1 matured smolt as an adult had the highest level of epigenetic change compared to the 2-year S2 matured smolt as an adult which was similar to the wild natural-origin population. However, the S2 matured population also had a high level of epigenetic change compared to the wild natural-origin population. This suggests other hatchery environmental factors than growth in the S2 population also exist and

Figure 7: Sperm DMR associated gene correlations. Cellular localization of associated genes with processes and pathologies in box
contribute to the hatchery impacts on the epigenetic programming. Clearly, moving to a more normal hatchery condition to mimic the natural-origin wild population [73] will help improve the physiology and health of the steelhead trout population and reduce the abnormal phenotypes observed in Table 1. In addition to the early developmental rearing and nutrition parameters, a large number of other parameters can impact the migration and adult fish development [7, 74]. This can include the impacts of fin clips [75], migration impacts of Dams [76], behavioral impacts from predators [77], water temperature and quality [78], and ocean conditions [79]. Therefore, the epigenetic analysis of the adult fish in the current study will incorporate not only the hatchery rearing impacts, but these other variables as well. This needs to be considered in the data interpretation.

The current study supports a potential role for hatchery and growth rate-induced epigenetic change in steelhead trout. The observations correlate with the phenotypic variation between hatchery and wild natural-origin fish populations. Although the programming and signatures of DMRs were found to be associated with specific cell types, the functional impacts of these DMRs need to be further investigated. A recent study has also examined the genetic and epigenetic differences between hatchery-reared and natural steelhead trout from the Methow River system [58]. In this study, Restriction Site Associated DNA Sequencing (RAD-Seq) was used to assess genetic similarity by identifying single nucleotide polymorphisms (SNPs). Similar to the current study, negligible genetic differences were found. Gavery et al. [58] used RBBS to evaluate DNA methylation, and again similar to the current study, found epigenetic differences between hatchery-raised and natural fish. The current study provides a more genome-wide analysis due to the MeDIP-Seq focus on lower density genome regions, and a CNV analysis that also reinforces the previously published genetic results. Therefore, the current hatchery conditions used can impact
phenotypic variation through environmentally induced epigenetic transgenerational inheritance.

Conclusions
Clearly, the current study supports a critical role for epigenetics when considering the molecular source for hatchery impacts, however, it will be an integration of epigenetics and genetics that will influence the molecular control of phenotypic variation. Since the sperm were found to have epigenetic alterations, the generational impacts through epigenetic inheritance also need to be considered. The potential that these epigenetic germline effects can be transmitted in the absence of direct exposure through epigenetic transgenerational inheritance now needs to be assessed. In hatchery operations involving food production, such as aquaculture, or when the hatchery fish are not allowed to breed with the wild fish population, impacts on the wild population will be reduced. However, in the event a hatchery population can breed with a wild fish population, the long-term impacts on the wild population and the environment could be dramatic and alter the future trajectory of the wild population. Further research into environmental epigenetic impacts on hatchery and wild fish populations is now needed. Interestingly, the epigenetic alterations observed could be used as biomarkers to further identify hatchery impacts on the fish, as well as correlate with phenotypic alterations observed in the future.

Methods
Animal Studies
As described in the Supplementary Methods [58], adult steelhead trout, Oncorhynchus mykiss, were collected from the Methow River and Winthrop National Fish Hatchery in Winthrop, WA, USA. The purified sperm and RBCs were collected for DNA preparations and epigenetic analysis. The sample collections were obtained by the WNFH staff and the staff of Dr Penny Swanson’s laboratory at the Northwest Fisheries Science Center, and the National Marine Fisheries Service (NMFS, NOAA) in Seattle WA, who had the appropriate vertebrate animal approvals for the study. All fish were reared and sampled according to the WNFH and University of Washington Institutional Animal Care and Use Committee (Dr Penny Swanson’s laboratory at the Northwest Fisheries Science Center, and the National Marine Fisheries Service (NMFS, NOAA) in Seattle WA, who had the appropriate vertebrate animal approvals for the study. All fish were reared and sampled according to the WNFH and University of Washington Institutional Animal Care and Use Committee (Dr Penny Swanson, Protocol #2313-90). Samples were shipped on dry ice and stored at −80°C for analysis. The initial natural and hatchery origin adult steelhead returning to the Methow River in summer 2013 and captured spring 2014 were used as designated N1 and H. Subsequent years were collected in a similar manner for the N2 and S1 or S2 populations (Supplementary Table S1).

Epigenetic Analysis, Statistics, and Bioinformatics
DNA was isolated from sperm and RBCs as described in the Supplementary Methods [56]. Methylated DNA immunoprecipitation (MeDIP), followed by Seq (MeDIP-Seq) was performed. MeDIP-Seq, sequencing libraries, Seq, and bioinformatics analysis were performed as described in the Supplementary Methods [66]. All molecular data has been deposited into the public database at NCBI (GEO # GSE145887), and R code computational tools are available at GitHub (https://github.com/skinnerlab/MeDIP-seq) and www.skinner.wsu.edu.

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Data Availability
All sequence data have been deposited to NCBI, GEO # GSE145887.

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Conflict of interest statement
The authors have declared that no competing interests exist.

Supplementary data
Supplementary data are available at EnvEpig online.

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