Heritability of DNA methylation in threespine stickleback (Gasterosteus aculeatus)

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Abstract

Epigenetic mechanisms underlying phenotypic change are hypothesized to contribute to population persistence and adaptation in the face of environmental change. To date, few studies have explored the heritability of intergenerationally stable methylation levels in natural populations, and little is known about the relative contribution of cis- and trans-regulatory changes to methylation variation. Here, we explore the heritability of DNA methylation, and conduct methylation quantitative trait loci (meQTLs) analysis to investigate the genetic architecture underlying methylation variation between marine and freshwater ecotypes of threespine stickleback (Gasterosteus aculeatus). We quantitatively measured genome-wide DNA methylation in fin tissue using reduced representation bisulfite sequencing of F1 and F2 crosses, and their marine and freshwater source populations. We identified cytosines (CpG sites) that exhibited stable methylation levels across generations. We found that additive genetic variance explained an average of 24–35% of the methylation variance, with a number of CpG sites possibly autonomous from genetic control. We also detected both cis- and trans-meQTLs, with only trans-meQTLs overlapping with previously identified genomic regions of high differentiation between marine and freshwater ecotypes. Finally, we identified the genetic architecture underlying two key CpG sites that were differentially methylated between ecotypes. These findings demonstrate a potential role for DNA methylation in facilitating adaptation to divergent environments and improve our understanding of the heritable basis of population epigenomic variation.

Keywords: epigenetics; reduced representation bisulfite sequencing; intergenerationally stable methylation; meQTL; Gasterosteus aculeatus

Introduction

DNA methylation is a chemical modification to DNA that typically occurs at cytosines within CpG dinucleotides in animals (Suzuki and Bird 2008). It has been suggested that DNA methylation can play a number of biological roles, including gene expression regulation (expression, repression, alternative splicing, and spurious transcription prevention), cell-fate decision, and phenotypic evolution and adaptation to divergent environments (Bird 2007; Bossdorf et al. 2008; Maunakea et al. 2010; Feil and Fraga 2012; Jones 2012; Verhoeven et al. 2016; Neri et al. 2017; Richards et al. 2017). Recent genome-wide studies have revealed that DNA methylation variation is widely observed between closely related animal species and populations that have adapted to ecologically divergent environments (Massicotte et al. 2011; Liebl et al. 2013; Smith et al. 2015; Lea et al. 2016; Artemov et al. 2017; Le Luyer et al. 2017; Hu et al. 2018, 2019; Laporte et al. 2019; Heckwolf et al. 2020). In addition, methylation variation has been shown to have a substantial heritable component that selection can act on (Lim and Brunet 2013; Heard and Martienssen 2014; Taudt et al. 2016). Modification of the methylome may therefore be an important mechanism underlying phenotypic variation, adaptive evolution, and possibly ecological speciation (Jaenisch and Bird 2003; Turck and Coupland 2014; Verhoeven et al. 2016).

While theoretical studies have suggested that the evolutionary relevance of methylation variation is partially related to its heritability, experimental studies investigating heritable DNA methylation and its role in adaptive evolution are in their initial stages (Verhoeven et al. 2016; Hu and Barrett 2017; Richards et al. 2017). Although it is clear that DNA methylation levels can sometimes be intergenerationally stable (Jablonka and Raz 2009; Daxinger and Whitelaw 2012; Heard and Martienssen 2014), results have mainly come from plant studies, and the small number of animal studies have typically used isogenic lab lines (Morgan et al. 1999; Rakyan et al. 2003; but see Nätt et al. 2012; Weyrich et al. 2016, 2018; Heckwolf et al. 2020). The homogenous genetic backgrounds...
of these isogenic lines may mean that they are not representative of the methylation patterns occurring in more genetically heterogeneous populations (Herman et al. 2014; Verhoeven and Preite 2014). In addition, most studies in nonmodel species have so far been limited to describing broad patterns based on anonymous markers of DNA methylation (Schrey et al. 2013; Hu and Barrett 2017; Richards et al. 2017), which has hindered understanding of the functional relevance and genetic basis of stable methylation in these species.

Methylation variation is mainly under genetic control, which can be caused by DNA sequence variation in both cis- and trans-regulatory elements (Taudt et al. 2016; Hu and Barrett 2017). Recently, methylation quantitative trait loci (meQTLs) analysis has found both cis- and trans-acting genetic variants underlying methylation variation (Dubin et al. 2015; Orozco et al. 2015; Kawakatsu et al. 2016; Meng et al. 2016, Taudt et al. 2016). Cis-regulatory genetic variation typically affects methylation patterns of only one or a few nearby sites and is less pleiotropic, whereas genetic variants in trans-regulatory elements can simultaneously change the methylation levels of multiple sites (Taudt et al. 2016; Do et al. 2017; Schulz et al. 2017; Hannon et al. 2018; Gupta et al. 2019). However, with the exception of a few studies (Fan et al. 2019) almost all meQTL studies have been conducted in model species, and thus, the prevalence of cis- and/or trans-meQTLs, and their role in adaptive evolution in natural populations remains unclear.

To explore the stability of epigenetic modification between generations, and to study the genetic architecture of methylation variation between natural populations adapted to distinct environments, we used threespine stickleback (Gasterosteus aculeatus), an abundant fish species in both marine and freshwater habitats in the Northern Hemisphere. Since the end of the last ice age, marine stickleback colonized freshwater lake and stream habitats that were uplifted and landlocked, resulting in replicate freshwater populations that show repeated evolution of a suite of locally adapted traits (Bell and Foster 1994). The repeated adaptive divergence between marine and freshwater populations makes this a powerful system to study the ecology and genetic architecture of adaptation (Jones et al. 2012). In the last decade, a variety of genetic and genomic resources have been developed for this species (Baird et al. 2008; Hohenlohe et al. 2010; Jones et al. 2012; Ishikawa et al. 2017; Peichel and Marques 2017). In addition, genome-wide methylation variation between marine and freshwater populations (Smith et al. 2015) and between males and females (Metzger and Schulte 2018) have been characterized, as well as the demonstration of methylation responses to environmental change (Artemov et al. 2017; Metzger and Schulte 2017; Heckwolf et al. 2020). However, the intergenerational stability of methylation in stickleback, and the genetic architecture underlying methylation variation between marine and freshwater ecotypes, remain unclear.

We address these gaps by performing an epigenomic survey of fin tissue from sticklebacks under a common garden experimental design with controlled crosses. We first examined methylation divergence between marine and freshwater ecotypes. We then explored levels of methylation and its genetic basis across two generations of the marine-freshwater hybrid lines, and performed meQTL analysis with two F2 families to characterize the genetic architecture of methylation variation between ecotypes. We investigate four specific questions: (1) Is variation in DNA methylation stable between generations? (2) What is the genetic heritability of intergenerationally stable CpG sites? (3) What is the genetic architecture of DNA methylation differences between the stickleback ecotypes? (4) What are the relative contributions of cis- and trans-meQTLs to DNA methylation differences? Answering these questions will help to provide a baseline for understanding the heritability of methylation variation, and the role of methylation variation in facilitating population persistence and potentially local adaptation in natural populations.

**Materials and methods**

**Sampling and husbandry**

We collected adult threespine stickleback from one marine (Bamfield Inlet [BI], 48°49'12.69"N, 125° 8'57.90"W), and two freshwater (Hotel Lake [HL], 49°38'26.94"N, 124° 3'06.69"W, and Klein Lake [KL], 49°43'32.47"N, 123°58'7.83"W) locations in Southwestern British Columbia, Canada in May 2015 (Figure 1). We transported all fish to an aquatic facility at the University of Calgary, and separated them into population-specific 113 L glass aquaria. We maintained a common garden environment at a density of approximately 20 fish per aquarium, salinity of 4–6 ppt, water temperature of 15°C ± 2°C, and a photoperiod of 16L:8D for one year before making crossings. This period of time should minimize any effects of transportation and allow sufficient time for marine populations to acclimate to hypoosmotic conditions (McCairns and Bernatchez 2010; Morris et al. 2014; Wang et al. 2014; Artemov et al. 2017). We kept each aquarium as a closed system with its own filter, air pump, water supply, and temperature regulator. We fed all fish ad libitum once per day with thawed bloodworms (Hikari Bio-Pure Frozen Bloodworms).

**Crossing design**

Threespine stickleback are typically found in either marine or freshwater habitats, but distinct marine and freshwater ecotypes can hybridize, which can facilitate the detection of associations between genotype and phenotype (Ishikawa et al. 2017; Peichel and Marques 2017; Verta and Jones 2019). We generated
genetically heterogeneous marine-freshwater F1 families from wild-caught parents by collecting eggs from one marine female and extracting testes from one freshwater male per cross (Supplementary Figure S1). To generate a cross, we first equally distributed the eggs into a Petri dish containing fresh water. We then euthanized the male using an overdose of eugenol (>400 mg/L) and removed the testes. We crushed the testes in a Petri dish, with the water activating the released sperm and allowing fertilization. Fertilized eggs were left within the Petri dish for 20 min before being suspended in a well-aerated mesh-bottom container within 37°C glass aquaria, with an air stone for oxygenation and a sponge filter. In total, we produced one F1 family of BfxHL hybrids (hereafter referred to as HL_F1), and three F1 families of BfxKL hybrids (hereafter referred to as KL_F1). After hatching, the larval fish from the same family were reared in the same 37°C aquaria until reaching approximately 1 cm total length (TL), at which time the families were equally split into aquaria to maintain low densities. The fish fry were fed twice daily with live Artemia spp. nauplii. At approximately 2 cm TL, juvenile stickleback fish were transitioned to a diet of chopped thawed bloodworms once per day ad libitum. They were then transitioned to an adult diet of full thawed bloodworms gradually. We sampled caudal fin clips (hereafter referred to as “fin clips”) when individuals reached a 3.5 cm TL or more. In addition to the fish we used to generate the F1 crosses, we also sampled extra parental fish from the same marine or freshwater population. Fin clips were stored in 70% ethanol in microcentrifuge tubes at room temperature until extraction of genetic material.

To generate F2 families, we randomly selected and crossed one male and one female sibling within an F1 family from each hybrid line (HL or KL) using the same crossing methods. We produced one F2 family of HL hybrids (hereafter referred to as HL_F2) and one F2 family of KL hybrids (hereafter referred to as KL_F2). Fish were raised as described above. We randomly selected fish from HL_F2 and KL_F2 families, and sampled fin clips when an individual reached approximately 3.5 cm TL. We stored all fin clips as described above. In addition to the fish we used to make the F2 crosses, we also randomly sampled extra F1 fish from all F1 families. In total, we sampled 94 fish, including 11 parental fish (six marine females; two HL and three KL freshwater males), 19 F1 fish (7 HL_F1 and 12 KL_F1), and 64 F2 fish (28 HL_F2 and 36 KL_F2). Detailed information about sex and family is included in Supplementary Table S1. All sampling, crossing, and housing protocols were approved by the University of Calgary Life and Environmental Science Animal Care Committee (AC13-0040 and AC17-0050) following the ethical standards maintained by the Canadian Council for Animal Care.

Tissue choice
The choice of tissue used for genome-wide mapping of cytosines can influence the interpretation of methylation patterns (Stricker et al. 2017). We conducted our analyses using caudal fin tissue for several reasons. It has been shown that fin position, caudal fin depth, and caudal fin size are different between marine and freshwater stickleback, and that these phenotypic differences are heritable and associated with repeated adaptation to divergent marine and freshwater environments (Walker 1997; Jones et al. 2012). Because methylation is tissue-specific, choosing a tissue showing phenotypic differences between ecotypes increases the likelihood of finding meQTLs that contribute to this ecotype divergence. Caudal fins can also be dissected quickly and consistently, and the excision of fin tissue does not affect survival.

DNA extraction and sex determination
We extracted DNA from caudal fin using phenol: chloroform: isoamyl alcohol (25:24:1), and assessed the quality and quantity using Tecan Infinite® 200 NanoQuant and Quant-iT PicoGreen® dsDNA assay kit (ThermoFisher Scientific). We determined the sex of fish following Peichel et al. (2004).

Reduced representation bisulfite sequencing
To measure genome-wide DNA methylation levels, we used reduced representation bisulfite sequencing (RRBS) (Meissner et al. 2008; Gu et al. 2011), following Boyle et al. (2012) with some minor modifications. For each individual, we created a library from 120 ng of genomic DNA, and ligated the MspI-digested fragments in each library with unique Illumina TruSeq adapters. We targeted fragments of 160–340 bp (including ~120 bp of adapter sequence) using NaCl-PEG diluted SpeedBeads (Rohland and Reich 2012). We split the libraries into four pools (three pools of 24 libraries and one pool of 22 libraries), and treated the pools with sodium bisulfite (EpiTect, Qiagen) following a protocol for formalin-fixed paraffin-embedded samples (Gu et al. 2011). After two rounds of bisulfite treatment to ensure complete conversion of unmethylated cytosines, each pool was amplified with Illumina primers, and loaded in four lanes (100-bp single-end reads) of a HiSeq 2500 at the McGill University and Genome Quebec Innovation Centre. In total, we sequenced all 94 fish sampled across three generations (Supplementary Table S1). Each sample was sequenced to a mean depth (± SD) of 8.094 ± 2.532 million reads.

Read filtering and mapping
To remove adapter contamination, low-quality bases, and bases artificially introduced during library construction, we trimmed reads using Trim Galore! v0.4.4 (https://www.bioinformatics.babraham.ac.uk/projects/trim_galore/), with the “rbis” option. We then used the program Bowtie2 v2.2.9 (Langmead and Salzberg 2012), implemented in Bismark v0.17.0 (Krueger and Andrews 2012) to align trimmed reads for each sample to the stickleback genome (ENSEMBL version 98) with default settings, except for tolerating one non-bisulfite mismatch per read. We only included reads that mapped uniquely to the reference genome, and cytosines that had at least 10x coverage in downstream analyses. The average mapping efficiency (percentage of uniquely mapped reads among all reads) was 61.4 ± 4.7% (± SD). We quantified methylation at non-CpG motifs and found less than 1% non-CpG cytosines were methylated, suggesting a highly efficient bisulfite conversion. Only CpG context cytosine methylation was analyzed because CpG methylation is the most common functional methylation in vertebrates (Suzuki and Bird 2008).

General methylation patterns
To identify general methylation patterns, we first performed a principal component analysis (PCA) on methylation levels in all samples using the prcomp function in R (R Core Team, 2018, v3.4.3). We ran the analysis by first identifying cytosines that were covered in all samples using the R package methylKit v1.4.1 (Akalin et al. 2012). Read coverage was then normalized between samples, using the median read coverage as the scaling factor. A minimum of ten reads in all samples was required at a CpG site for that site to be analyzed. We removed CpG sites that were in the 99.9th percentile of coverage from the analysis to account for potential PCR bias. We calculated the methylation levels by extracting the total amount of methylation-supporting reads,
and the total coverage of each CpG site, using the percMethylation function in the R package methylKit. To improve methylation estimates, we corrected for SNPs, which could have resulted in an incorrect methylation call if C-to-T and G-to-A SNPs were falsely interpreted as unmethylated cytosines, following Heckwolf et al. (2020). We first identified SNPs using the methylation value of each CpG site of all 11 parental individuals for input to Bis-SNP v0.82.2 (Liu et al. 2012) with the default parameters. Because Bis-SNP is sensitive to the directionality of the RRBS protocol (i.e. whether sequenced reads come from the original forward and reverse strands when calling C-to-T and G-to-A SNPs), we used a directional bisulfite-seq protocol that is similar to Krueger et al. (2012). We chose parental samples for identifying SNPs because they are the genetic source of the F1s and F2s, and are the most genetically heterogeneous samples. We used GATK’s VariantFiltration and SelectVariants to restrict variants to physical positions. We calculated pairwise weighted multiple-test corrected *P* values (Benjamini-Hochberg’s false discovery rate) < 0.1 and a minor allele frequency (MAF) greater than 0.0049 (Heckwolf et al. 2020), and more than 10% missing data across all 11 parental samples. We used MAF thresholds from 0.001 to 0.01 resulted in similar numbers of filtered SNPs. We calculated pairwise weighted *F*<sub>r</sub> between BI, HL and KL fish in the parental generation using VCFtools on filtered SNPs. We produced a list of positions (C-to-T and G-to-A SNPs) for correcting methylation estimates, using custom written Perl scripts from Heckwolf et al. (2020) and the R package GenomicRanges v1.30.3 (Lawrence et al. 2013). In addition, it has been suggested that sex specific methylation affects less than 0.1% of CpG sites on autosomal chromosomes, but more than 5% of CpGs on the sex chromosome in stickleback (Metzger and Schulte 2018). Therefore, to exclude a potential sex bias, we removed all CpGs located on the sex chromosome (group XIX). In total, we retained 52,940 CpG sites that passed the filtering step. To perform PCA, methylation levels at each CpG site were taken as input variables, whereas each point in multidimensional space represented a stickleback individual. Finally, to compare DNA methylation variation between F1 and F2 fish in each hybrid line, we calculated DNA methylation levels in 7,840 1-kb tiling windows (step = 1 kb; size = 1 kb) compiled from the same 52,940 CpG sites, and compared the standard deviations of methylation levels for each genomic window within each hybrid line.

**Analysis of methylation diversity between ecotypes**

To examine methylation diversity between ecotypes, we performed a differential methylation analysis between marine and freshwater populations from the parental generation, using the 52,940 CpG sites that passed the filtering step above. CpG sites were considered to be differentially methylated cytosines (DMCs) with a false discovery rate correction Q-value < 0.01 and a minimum required methylation difference of 15% between ecotypes, using the R package methylKit with sampling site (i.e. BI for the marine population, and KL and HL for the freshwater populations) as a covariate. We visualized differential methylation patterns across individuals and obtained clustering of samples and DMCs in heatmaps with the “complete” clustering method on Euclidian distances, using the R package heatmap version 1.0.8 (https://cran.r-project.org/web/packages/heatmapR/index.html). We clustered hyper- and hypo-methylated DMCs between ecotypes using relative percent DNA methylation, which is the normalized percent DNA methylation scaled for each DMC’s percent DNA methylation (median percent methylation as 0) of marine and freshwater fish in heatmaps. We also clustered individual fish based on overall methylation patterns across DMCs. We then analyzed the proportion of cytosines within genomic features [promoter/exon/intron/intergenic; promoters are defined as regions being upstream 1000 bp and downstream 1000 bp around the transcription start sites (TSSes)] for DMCs, using the R package ggaq v1.6.0 (Akalin et al. 2015). Because MspI restriction sites are not randomly distributed in the genome, we built a null distribution of genomic features based on all filtered CpG sites (i.e. 52,940 CpG sites). We chose precedence to promoters > exons > introns > intergenic regions when features overlapped (Smith et al. 2015; Hu et al. 2018). Finally, we annotated genes associated with DMCs, using the R packages biomaRt v2.34.2 (Durinck et al. 2005, 2009) and ChIPpeakAnno v5.12.3 (Zhu et al. 2010; Zhu 2013) on the stickleback reference genome from Ensembl 98 dataset, and performed gene ontology (GO) analysis on DMC-associated genes, using the R package topGO v2.28.0 (Alexa et al. 2006). Over-represented GO terms were those with multiple-test corrected *P*-values (Benjamini-Hochberg’s false discovery rate) below 0.1, based on a Fisher’s exact test. We compared DMC-associated genes with the genes associated with the 52,940 CpGs that passed the filtering step.

**Analysis of intergenerationally stable methylation**

We considered a CpG as intergenerationally stable when the CpG was not significantly differentially methylated between F1 and F2 generations of the same family within the same hybrid line (HL or KL) and fulfilled this criterion in both hybrid lines. Note that these “stable” sites are not necessarily “heritable” in the sense of methylation variation between individuals being due to additive genetic factors. DMCs between fish in F1 versus F2 generations were identified using the same method as described above for identifying DMCs between ecotypes, with sequencing lane as a covariate. We identified 137 and 82 DMCs within HL and KL hybrid lines, respectively. These sites were removed from the 52,940 CpG sites that passed the filtering step to provide a final dataset of 52,729 intergenerationally stable CpG sites for downstream analysis. We clustered fish based on the similarity of their DNA methylation profiles, with the “ward” clustering method on Pearson’s correlation distances, using the cluster function in the R package stats v2.15.1. We also compared the locations of DMCs between ecotypes with the locations of intergenerationally stable CpG sites to assess which of the sites involved in methylation divergence between ecotypes are stable across generations. Our criterion for identifying CpG sites with intergenerationally stable methylation is such that a type 2 error in our differential methylation test between generations (a false negative for differential methylation between F1 and F2 within a line) will lead to a false positive for stable methylation. To investigate the potential importance of this type of error in our data, we conducted a power analysis using a simulated data set in methylKit, following Wreczycka et al. (2017) with some minor modifications. To be conservative, our simulated dataset consisted of eight samples (four F1s and four F2s, matching the minimum number of fish in a line from each generation in the empirical dataset). We modeled the read coverage following a binomial distribution and defined the methylation levels following a beta distribution with parameters alpha = 0.4, beta = 0.5, and theta = 10. We ran simulations of differential methylation at 1% of 52,940 CpG sites, with effect sizes of 5%, 10%, 15%, 20% and 25% differential
methylation, respectively. After correcting for the covariate of sequencing lane (the same sequencing lane was assigned to two samples within F1 or F2 generation, for a total of four sequencing lanes), we adjusted P-values for multiple testing using the Q-value method (Storey and Tibshirani 2003), and considered CpGs to be DMCs with a false discovery rate correction Q-value < 0.01. Finally, we calculated the proportion of CpGs that were falsely identified as non-DMCs (false negatives) among all CpG sites under each effect size (5−25%) above.

We distinguished between three categories of methylated sites that were stable between F1 and F2 generations: (1) constitutively hypermethylated sites, which are CpG sites with average DNA methylation levels greater than 0.9 in all samples, (2) constitutively hypomethylated sites, which are CpG sites with average DNA methylation levels less than 0.1 in all samples (Lam et al. 2012; Lea et al. 2016), and (3) methylated sites with average DNA methylation levels between 0.1 and 0.9 (hereafter referred to “variable sites”). Finally, we analyzed the proportion of cytosines within genomic features for CpG sites in each category and annotated genes associated with all intergenerationally stable CpG sites by testing for overlap between the locations of CpG sites and genomic regions of genes, following the same method as described above for ecotype DMCs.

Heritability of stable methylation

To determine the genetic heritability of the intergenerationally stable CpG sites across F1 and F2 generations, we first identified SNPs using the aligned reads of all F1 and F2 individuals for input to Bis-SNP, following the same SNP calling and filtering steps as described above with some minor modifications. We retained three sets of SNPs by filtering the 92,983 SNPs using a constant MAF cut-off (0.005) and three missing data cut-offs (10%, 30%, and 50%) across all F1 and F2 individuals. We used BCFTools (https://github.com/samtools/bcftools) to exclude sites that were in linkage disequilibrium (LD, pairwise $r^2 > 0.8$ within a window of 1 Mb) or on the sex chromosome. Finally, we used a linear mixed model implemented in PyLMM (http://genetics.cs.ucla.edu/pylmm/index.html) to test whether variation at SNPs is significantly associated with methylation levels at stable CpG sites in F1 and F2 generations, after correcting for sequencing lane variation and kinship based on the SNP data. We adjusted multiple-test P-values using Benjamini-Hochberg’s false discovery rate and considered an association to be significant when the corrected $P$-value < 0.05.

We estimated the narrow-sense heritability of DNA methylation levels for individual CpG sites of all F1 and F2 individuals using a linear mixed model approach (Yang et al. 2010) implemented in the R package lmmlite (https://github.com/kroman/lmmlite). We treated the methylation levels at individual CpG sites of all F1 and F2 individuals as phenotypes, and assumed each phenotype $y$ can be modeled as $y = \alpha + u + e$, where the random variable $u$ follows a normal distribution centered at zero with variance $\sigma_u^2$, and $e$ represents an independent noise component with variance $\sigma_e^2$. The matrix $K$ is the same kinship matrix as calculated above. For each trait we estimated $\sigma_u^2$ and $\sigma_e^2$ using the restricted maximum likelihood (REML) approach, with correction for the covariate of sequencing lane, and calculated the heritability as $h^2 = \sigma_u^2/(\sigma_u^2 + \sigma_e^2)$. Finally, we calculated the average heritability by taking the mean of heritability values of all CpG sites.

meQTL analysis

To identify the genetic architecture of methylation divergence between marine and freshwater stickleback, we performed meQTL mapping of the methylome in two F2 families of marine-freshwater hybrids. Due to the distinct methylation patterns that may be caused by genetic variation between the two hybrid lines (Supplementary Figure S2), we performed mapping separately for each hybrid line. We retained SNPs that were not located on the sex chromosome, had less than 10% missing data, and had low LD in HL_F2 or KL_F2 samples, using the same SNP filtering steps as described above. We then compiled a percentage methylation level matrix among HL_F2 or KL_F2 samples containing the 52,940 CpG sites that passed these filtering steps. Finally, we truncated these sites by the 10% minimum range of methylation variation across samples to reduce non-informative sites that could possibly inflate test statistics and create spurious SNP-CpG pairs. After filtering, we retained 525 SNPs and 27,614 CpGs in HL_F2, and 330 SNPs and 27,039 CpGs in KL_F2 for meQTL analysis, with no overlap between the retained SNPs in each line. We tested all genome-wide SNP-CpG pairs using the R package MatrixEQTL v2.2 (Shabalin 2012). This package enables rapid computation of QTLs by only retaining those that are significant at a pre-defined threshold. We fit an additive linear model to test if the number of alleles (coded as 0, 1, 2) predicted percentage DNA methylation levels (value ranging from 0 to 1) at each CpG site, including sequencing lane as a covariate. We used a Bonferroni-corrected multiple-test corrected threshold, set it to genome-wide significance for GWAS and divided by the number of CpG sites tested (i.e. HL_F2: $5 \times 10^{-7}/27,614 = 1.81 \times 10^{-12}$, KL_F2: $5 \times 10^{-9}/27,039 = 1.85 \times 10^{-12}$). We chose this stringent threshold to call meQTLs to minimize the possibility of false positives (Orozco et al. 2015). We calculated the distance between a SNP and a CpG site within a significant meQTL and defined a SNP as cis-acting if the SNP was located within 1 Mb from its associated CpG site or trans-acting if the SNP was located more than 1 Mb from its associated CpG site (Zhang et al. 2014). We then performed GO analysis on genes associated with unique SNPs within significant meQTLs and identified over-represented GO terms, using the same method as described above. The gene pools against which we compared the unique SNPs were the genes associated with the SNPs that passed the filtering step. Because previous studies have suggested that meQTLs and expression QTLs (eQTLs) are likely to co-occur in close genomic proximity, we compared locations of significant meQTLs in our study to significant eQTLs identified in Ishikawa et al. (2017), which also used a marine-freshwater hybrid design. Ishikawa et al. (2017) identified eQTLs under a range of salinities, but we only used eQTLs that they identified under 3.1 ppt, which is a similar salinity level to our experimental conditions. Finally, to investigate the role of meQTLs in adaptation to different habitats in stickleback, we compared locations of unique SNPs within significant meQTLs to previously documented regions of parallel genomic divergence between marine and freshwater sticklebacks (Hohenlohe et al. 2010; Jones et al. 2012; Terekhanova et al. 2014), and identified genetic architecture of DMCs between ecotypes by comparing the locations of DMCs and unique CpG sites associated with significant meQTLs. In addition to performing the meQTL analyses using MatrixEQTL as described above, we also validated meQTL results within each hybrid line using the R package R/qtl v.1.46-2 (Broman et al. 2003) with default settings, except for rescaling the basepair positions of SNPs by multiplying by a constant of $3.11 \times 10^{-6}$ due to the genome-wide recombination rate of 1%.
3.11cM/Mb in stickleback (Roesti et al. 2013). We calculated genome-wide logarithm of the odds (LOD) thresholds through 1000 permutations, using the n.perm function in the R package R/qtl and set the 95th percentile LOD score as the significance threshold (Höglund et al. 2020).

Data accessibility
Raw Illumina sequencing reads for the 94 analyzed individuals can be downloaded from the NCBI Short Read Archive (BioProject ID: PRJNA587332). The cytosine coverage files (.cov) for the 94 analyzed individuals and codes used for analyses in this study are available through Github (https://github.com/barrettlabecoevogeno/Heritability_DNA_methylation_sticklebacks). Additional Supplemental material is available at figshare (https://doi.org/10.25386/genetics.13481877).

Results
General methylation patterns
To identify general methylation patterns, we performed PCA on the methylation levels of filtered CpG sites represented in all samples (Figure 2). PC1 reflected sequencing lane chemistry (Supplementary Figure S2) and so we included sequencing lane as a covariate in all downstream analyses. When analyzing all samples, PC2 (variance explained: 5.1%) clearly separated parental and F2 samples, with F1 samples filling the intermediate space between parental and F2 samples. Furthermore, the PCA separated the samples by sire (HL vs. KL) in F1 populations along PC3, which accounted for 3.2% of the variance observed in the data set (Figure 2A). In the parental fish, the PCA separated samples mainly by their habitat along PC3 (Figure 2B), whereas the F1 generation showed clustering based on family along PC2 (Figure 2C). The PCA also revealed some clustering between the HL and KL hybrid lines in the F2 generation, although there is no clear separation between lines (Figure 2D). Within families, we found significantly higher mean DNA methylation variance in the F2 generation than the F1 generation of families in the HL line ($W = 2.94 \times 10^{-7}$, $P = 5.61 \times 10^{-6}$), but not the KL line ($W = 3.08 \times 10^{-7}$, $P = 0.775$).

The average pairwise $F_{ST}$ calculated between populations was 0.03 (BI vs. HL), 0.04 (BI vs. KL), and 0.01 (HL vs. KL). These values are comparable to what has been reported between other marine and freshwater populations of stickleback using SNPs extracted from RAD-seq (e.g. Hohenlohe et al. 2010; Catchen et al. 2013; Lescak et al. 2015; Garcia-Elfring et al. Unpublished) and whole genome sequencing (Shanfelter et al. 2019), suggesting that our use of SNPs identified from RRBS should not bias estimates of genetic differentiation relative to other methods.

Methylation divergence between ecotypes in the parental generation
We identified 891 DMCs between parental fish sampled from marine versus freshwater habitats after false discovery rate correction. Based on Euclidean distances, individual fish clustered by their ecotypes, with the freshwater fish further clustered by their
sampling site (HL vs. KL, Figure 3A). When analyzing the methylation patterns of the 891 CpGs across generations, we found two major clusters, with the first cluster only including marine fish from the parental generation, and the second cluster including all freshwater fish from the parental generations and all F1 and F2 hybrids (Supplementary Figure S3). When comparing the mean methylation levels of the 891 ecotype DMCs in F1s versus parents, we found that in both hybrid F1 lines, there was a significantly greater number of CpG sites with methylation levels that were intermediate between the values of the parents than the number of CpG sites with methylation levels outside the values observed in the parents (KL_F1: G = 294, df = 1, P < 2.20 × 10⁻¹⁶; HL_F1: G = 124, df = 1, P < 2.20 × 10⁻¹⁶). In the F2 generation, we observed a greater proportion of sites showing methylation values outside the range observed in their F1 parents relative to the pattern between F1s and their wild parents (KL_F2: G = 241, df = 1, P < 2.20 × 10⁻¹⁶; HL_F2: G = 9.72, df = 1, P = 1.82 × 10⁻³). In addition, we found a marginally greater proportion of sites showing bias in methylation levels toward those of the mother in HL_F1 (G = 4.10, df = 1, P = 0.0428) but not KL_F1 (G = 0.576, df = 1, P = 0.448) when compared to the parental generation. However, sex is confounded with parental habitat in this comparison (marine fish are always female, and freshwater fish are always male). This confounding effect is not present in the F1 generation, where the same analysis found a marginally greater proportion of sites showing bias toward the F1 mother in KL_F2 (G = 3.85, df = 1, P = 0.0497) but not HL_F2 (G = 0.0627, df = 1, P = 0.802).

Identified DMCs between ecotypes showed no significant bias toward hyper- versus hypomethylation (430 hypermethylated and 461 hypomethylated DMCs; G = 1.21 × 10⁻³, df = 1, P = 0.972). However, we found significantly more DMCs within introns (G = 8.87, df = 1, P = 2.90 × 10⁻⁵) and significantly fewer within promoters (G = 24.5, df = 1, P = 7.33 × 10⁻⁷) when compared to the null distribution of all filtered sites (Figure 3B). In addition, we found no overlap between the locations of the 891 DMCs and the locations of sex-biased DMCs identified in Metzger and Schulte (2018), suggesting that removing CpGs located on the sex chromosome effectively minimized any potential sex-biased differential methylation. In total, DMCs mapped to 228 genes, with some genes having been shown to be associated with differential expression or methylation between ecotypes in recent studies (e.g. differentially expressed genes in gill: atp1a2a, g6pd, Artemov et al. 2017; differentially methylated genes in fillet: g6pd, chchd3a, Smith et al. 2015). GO analysis showed no significant GO term enrichment.

**Intergenerationally stable methylation**

We found that 99.6% of CpG sites (52,729 out of 52,940) were not differentially methylated across generations in both lines, suggesting the vast majority of sites show stable levels of methylation across generations. Our power analysis suggests that a small proportion of sites (<1%) are likely to have been falsely identified as non-DMCs (Supplementary Figure S4) across all effect size groups, suggesting the influence of type 2 error on our criterion for calling stable methylation would only affect a small number of sites. Based on Pearson’s correlation distance calculated from the 52,729 CpG sites, most individuals clustered by generation (F1 vs. F2) and by hybrid line (HL vs. KL) (Figure 4A).

We found no significant enrichment of the stable sites in any of the genomic contexts when compared to the null distribution of all filtered sites (promoters: G = 4.53 × 10⁻⁴, df = 1, P = 0.983; exons: G = 4.94 × 10⁻⁵, df = 1, P = 0.994; introns: G = 3.49 × 10⁻⁵, df = 1, P = 0.995; intergenic regions: G = 1.01 × 10⁻⁴, P = 0.992; Figure 5A). Among the stable CpG sites, we found 6,462, 28,005, and 18,262 CpG sites that were constitutively hypermethylated, constitutively hypomethylated, and variable, respectively. When analyzing the genomic context of CpGs from these three
categories, we found a significantly biased genomic distribution, with constitutively hypermethylated sites enriched within exons ($G = 29.3, \text{df} = 1, P = 6.33 \times 10^{-4}$, Figure 5B), constitutively hypomethylated sites enriched within promoters ($G = 17.8, \text{df} = 1, P = 2.42 \times 10^{-5}$; Figure 5C), and variable sites enriched within introns ($G = 9.36, \text{df} = 1, P = 2.22 \times 10^{-3}$; Figure 5D) when compared to the null distribution of all filtered CpGs. We found that 94.8% (845 out of 891) of the DMCs between ecotypes were also identified as stable sites, a percentage not significantly different than the percentage of stable sites among all filtered sites ($G = 0.171, \text{df} = 1, P = 0.679$), suggesting that most sites involved in methylation divergence between ecotypes can be stable across generations, and could therefore be plausibly associated with adaptation to different habitats.

To calculate the genetic heritability of stable sites, we first identified 92,983 SNPs, and then filtered this dataset down to 350 SNPs that (1) had less than 10% missing data across all F1 and F2 individuals, (2) were not located on the sex chromosome, and (3) had low to no LD with each other. Six of these SNPs showed highly significant associations with the methylation values of F1 and F2 individuals ($Q < 0.05$), and 16,514 out of the 52,729 intergenerationally stable CpG sites had $h^2 > 0$ (Figure 4B). We also retained 3,007 and 4,203 SNPs after filtering the SNPs by 30% and 50% missing data, respectively, with 22 and 28 of these SNPs...
showing highly significant associations with the methylation values of F1 and F2 individuals, and 16,498 and 21,055 intergenerationally stable CpG sites having \( \hat{h}^2 > 0 \) (Supplementary Figure S5).

Finally, the kinship matrix estimated the narrow-sense heritability for CpG methylation levels at (on average) 24%, 32%, and 35% using the 350,3,007, and 4,203 post-filtering SNPs, respectively.

**Identification of meQTLs associated with methylation divergence between marine and freshwater ecotypes**

When analyzing meQTLs within each hybrid line, we identified 968 and 531 significant SNP-CpG pairs in HL_F2 and KL_F2 fish, respectively, corresponding to 335 unique SNPs and 75 unique CpG sites in HL_F2, and 201 unique SNPs and 72 unique CpG sites in KL_F2. We found that 85.0% (HL_F2: 823 of 968) and 94.4% (KL_F2: 501 out of 531) of the SNP-CpG pairs were also identified as significant SNP-CpG pairs when using R/qtl, and the \( P \)-value distributions showed no evidence of test statistic inflation (Supplementary Figure S6). A two-dimensional plot of meQTLs displayed significantly more trans- than cis-meQTLs (HL_F2: 965 trans- vs 3 cis-meQTLs, \( G = 1.30 \times 10^9 \), \( df = 1, P < 2.20 \times 10^{-16} \), KL_F2: 527 trans- vs 4 cis-meQTLs, \( G = 689 \), \( df = 1, P < 2.20 \times 10^{-16} \), with no significant GO enrichment for genes annotated with unique SNPs within significant meQTLs in either line. There was relatively low LD between SNPs and CpGs with significant cis associations (mean \( r^2 = 0.287 \)). We also found no overlap between CpGs associated with meQTLs and the DMCs identified between ecotypes in HL_F2, likely due to the small number of CpG sites associated with significant meQTLs. Alternatively, this could be due to the small amount (ranging from 24% to 35%) of methylation variation explained by additive genetic variation, suggesting that a significant proportion of CpGs are likely to be autonomous from additive genetic variation, and thus not detectable in a meQTL analysis. We found two CpGs associated with nine trans-meQTLs that overlapped with DMCs in KL_F2, with all nine SNP-CpG pairs verified by R/qtl. Although the two CpGs did not localize within any genes, they were in close genomic proximity (~15 kb) to zinc finger E-box binding homeobox 1b (zeb1b, Ensembl Gene ID ENSGACG0000001002) and centrosonal protein 76 (cep76, Ensembl Gene ID ENSGACG0000003686). The nine trans-meQTLs were annotated with four genes (Supplementary Table S2).

To assess the co-occurrence of meQTLs and eQTLs, we compared locations of meQTLs identified in our study to eQTL hotspots in Ishikawa et al. (2017). We found that an overall of 9.14% (HL_F2: 24 out of 335; KL_F2: 25 out of 201) of the unique SNPs overlapped with eQTL locations across HL_F2 and KL_F2 samples. This proportion does not suggest an excess of meQTL-eQTL overlap relative to null expectations that are built from all input SNPs for meQTL analyses (\( G = 0.245, df = 1, P = 0.621 \)). Finally, to investigate whether meQTLs might be associated with divergent selection in marine versus freshwater habitats, we examined whether the unique SNPs within meQTLs overlapped with genomic regions of high differentiation between the two stickleback ecotypes. We found a total of six (four in HL_F2 and two in KL_F2) unique SNPs within high differentiation regions, corresponding to 14 (11 and 3 in HL_F2 and KL_F2, respectively) trans-acting SNP-CpG pairs. The effect sizes (beta) of meQTLs ranged from 0.105 to 0.406 (median = 0.161). The six SNPs were annotated with four genes (Supplementary Table S3), which encode proteins likely to be relevant to marine-freshwater divergence in stickleback (e.g., sensing changes in osmoregulatory environment; see below).

**Discussion**

The role of DNA methylation in fundamental ecological and evolutionary processes has received increased attention in recent years (Metzger and Schulte 2016; Verhoeven et al. 2016; Hu and Barrett 2017). However, the extent to which variation in DNA methylation is stably transmitted across generations, and the prevalence of cis- and/or trans-acting genetic variants in contributing to methylome evolution remain poorly understood, particularly in natural animal populations. We used a quantitative, single-base-resolution technique (RRBS) to measure DNA methylation from fin tissue across two generations in threespine stickleback sampled from two distinct environment types. A large majority (99.6%) of CpG sites were identified as being intergenerationally stable, as indicated by consistent methylation levels across F1 and F2 generations in two hybrid lines. As a consequence, the subset of CpG sites that also showed significant divergence between marine and freshwater ecotypes in the grandparental generation, a large majority (94.8%) could be classified as being stable across generations. Epigenetic variation was associated with genetic variation to a certain extent, with a narrow-sense heritability of these stable sites ranging from 24% to 35%. These values are consistent with recent epigenome-wide association studies that have found that genetic variation can explain an average of 7–34% of methylation variation in animals (McRae et al. 2014; Orozco et al. 2015; Taudt et al. 2016; Carja et al. 2017). We found distinct patterns of genomic context between three categories of stable CpG sites: constitutively hypomethylated and hypermethylated sites were predominantly located within promoters and exons, respectively, whereas variable sites were enriched within introns. We also identified meQTLs in marine-freshwater F2 hybrid lines, with some meQTLs overlapping with genomic regions of high differentiation between marine and freshwater ecotypes in stickleback. Finally, we identified the genetic architecture underlying two DMCs between ecotypes that were also shown to have intergenerational stability in their methylation levels. Overall, our study provides the first investigation of the genetic basis of stable epigenetic variation in stickleback and identifies methylation differences that could be associated with local adaptation in marine and freshwater ecotypes.

**Methylation divergence between ecotypes**

We found a similar number of differential methylation sites (891 DMCs) between marine and freshwater ecotypes of threespine stickleback to a recent study (737 DMCs in Smith et al. 2015). While we did not find any significantly enriched GO terms, some of these DMCs were annotated with genes that are likely to contribute to adaptation to marine and freshwater environments. For example, we found DMCs overlapped with genes related to osmoregulation (ion channel activity: trpc1, RYR3, gria3b, and sdr39u1), metabolic process (lipid and fatty acid metabolism: elou6f1, scap; glucose and carbohydrate metabolism: g6pd), immune response (hemopoiesis: kalma; myeloid cell and neutrophil differentiation: satb1b; erythrocyte maturation: klf3), and catalytic activity (alipl, phlp1, and sdr39u1). Because the osmotic environments, parasite communities, and migratory life cycles of marine and freshwater ecotypes differ (Smith et al. 2015; Huang et al. 2016; Artemov et al. 2017; Ishikawa et al. 2017), the
differential methylation of these genes suggests that the methylome could be associated with ecologically important phenotypic differentiation between ecotypes.

The genetic basis and functions of intergenerationally stable epigenetic variation

To understand how methylation divergence between ecotypes might be involved in the process of local adaptation, we next explored the stability of epigenetic variation across generations. While the approach of using experimental crosses to explore stable epigenetic variation and its underlying genetic basis has been widely applied in plant studies (e.g. Johannes et al. 2009; Roux et al. 2011; Li et al. 2014), few studies have used this type of experimental design in non-model animal populations (but see Nätt et al. 2012; Weyrich et al. 2016, 2018). Examination of the genetic basis of methylation sites is valuable for exploring the mechanisms that facilitate animal responses to novel environments, and for predicting the likelihood that populations will be able to evolve in response to environmental change (O’Dea et al. 2016). We identified 52,729 CpG sites that were not differentially methylated between F1 and F2 generations in both HL and KL hybrid lines (99.6% of all CpG sites). Similar to Heckwolf et al. (2020), this suggests that the majority of our analyzed CpG sites have stable levels of methylation across generations. These CpG sites are not necessarily heritable; it is possible that methylation levels are induced to similar levels across generations due to exposure to a similar environment. However, it is also possible that some methylation levels are heritable, either due to control by genetic variation (see below) or resistance to epigenetic reprogramming between generations. This pattern may also be due to non-global DNA methylation reprogramming during embryogenesis in fish, which can provide greater opportunity for transmitting DNA methylation from parents to the offspring (Schmitz et al. 2011; Skvortsova et al. 2018).

When assessing the contribution of the intergenerationally stable CpG sites to evolutionary processes, we found significant enrichment of constitutively hypomethylated CpG sites within promoters, and significant enrichment of constitutively hypermethylated CpG sites within gene bodies, suggesting that stable DNA methylation may directly regulate gene expression and facilitate alternative splicing, and thus contribute to genomic evolution by providing access to alternative promoter sites and increasing the number of transcriptional opportunities and phenotypes (Roberts and Gavery 2012). Consequently, mutations impacting intergenerationally stable methylation could accelerate the exploration of phenotypic space, and therefore allow populations to adapt to the changing environments more efficiently (Kliromonomos et al. 2013). Furthermore, the distinct distribution patterns of hyper- and hypo-methylated CpG sites are consistent with whole genome assessments of methylation in other fish species and in model animals and plants (Feng et al. 2010; Zemach et al. 2010; Long et al. 2013; Shao et al. 2014), suggesting a conserved role for constitutive hyper- and hypomethylation in a wide range of species.

We found a large proportion (~95%) of DMCs between marine and freshwater ecotypes in the grandparental generation also showed intergenerational stability in methylation levels, suggesting that the genes associated with these DMCs could play a role in facilitating adaptation to different environments. Theoretical work has suggested that environmentally responsive epigenetic changes that can be transmitted to the next generation might be beneficial when the effects of epigenetic variation increase both the parental and offspring fitness with low cost (Herman et al. 2014). As the functions of several DMC-associated genes identified here are relevant to responses to changes in aquatic environments such as salinity, parasites, and diet, our findings provide evidence for a possible adaptive mechanism in threespine stickleback whereby advantageous epigenetic changes that have been triggered by environmental stimuli are transmitted across generations.

There is substantial interest in biomedical and agricultural fields to understand the contribution of genetic variation to population epigenomic variation, with a number of recent genetic studies having quantified the heritable basis of population epigenetic variation in model animals (Taudt et al. 2016). When applying a stringent missing data cut-off (10%), we found a reasonably high average heritability of 24% for methylation levels across F1 and F2 generations. When applying more relaxed missing data cut-offs (30% and 50%), we find heritability estimates of 32% and 35%. In addition, we found that 31% to 40% of stable CpG sites had a measurable genetic component (narrow-sense heritability $h^2 > 0$), a percentage similar to previous findings in model species (Taudt et al. 2016). Together, our results suggest a plurality of mechanisms are likely contributing to stable levels of methylation variation across generations, including genetic control, epimutation, and exposure to past or current environmental factors.

Contribution of meQTL to methylation divergence between ecotypes

We characterized meQTLs in the F2 generation of two marine-freshwater hybrid lines, and detected two CpGs associated with significant meQTLs that overlapped with DMCs between marine and freshwater ecotypes, both in the KL_F2 line. The two CpGs were close to zeb1b and cep76, which are key genes involved in developmental processes. Zeb1b is a transcriptional factor regulating the expression of interleukin 2 (Williams et al. 1991), which is a cytokine gene involved in adaptive immune response in teleost fish (Zou and Secombes 2016). A recent study has also found that both interleukin 2 and genes involved in its signaling pathway are differentially expressed in kidney tissue between stickleback populations with different parasite prevalence (Fuess et al. 2020), suggesting a potential role for interleukin 2 and its regulators in facilitating stickleback responses to distinct parasite loads between marine and freshwater environments (Scharsack et al. 2016; Lugert et al. 2017). Cep76 is an important paralog of CC2D2A, which is a gene associated with development of the primary cilium, and is relevant to morphological differences between Pacific lamprey populations with distinct migratory behaviors (Hess et al. 2014). Because the morphology between marine and freshwater stickleback ecotypes differs significantly (Jones et al. 2012), the overlap between meQTL-associated CpGs and ecotype-DMCs suggests that these loci may be under divergent selection in marine versus freshwater habitats.

Although methylation variation between marine and freshwater ecotypes can be caused by both cis- and trans-regulatory changes, we found only trans-meQTLs within genomic regions of high differentiation between ecotypes. This is interesting because we expected to detect a bias toward cis-meQTLs due to the close genomic proximity of the SNPs and CpGs from the same RRBS fragments. Moreover, a number of recent studies have shown greater contribution of cis-regulatory than trans-regulatory genetic variants in gene expression divergence in the gill, brain and liver tissues of stickleback (Ishikawa et al. 2017; Pritchard et al. 2013). In addition, the distinct distribution of DMCs between marine and freshwater sticklebacks (Zou and Secombes 2016) suggests a contribution of epigenetic variation to the differentiation between marine and freshwater stickleback ecotypes.
2017, Verta and Jones 2019). However, predominantly trans-regulatory changes in gene expression have also been found in the tooth plate of stickleback and Drosophila (McManus et al. 2010; Osada et al. 2017; Hart et al. 2018). These contrasting results have been attributed to a number of factors such as inter-versus intra-specific comparison and tissue heterogeneity, where trans-regulatory effects dominate in intraspecific comparisons and in more heterogeneous tissue (Hart et al. 2018). Our findings fit with these explanations in that we conducted an intraspecific comparison using heterogeneous caudal fin tissue that consists of epidermis, osteoblasts, dermal fibroblasts, and vascular endothelium (Tu and Johnson 2011).

Although selection may initially favor master regulator genes that regulate distant genes through trans-acting mechanisms during rapid adaptation, it has been suggested that different evolutionary scenarios and selective contexts may alternatively favor trans- and cis-acting mechanisms during intraspecific adaptive divergence (Cooper et al. 2003; Lemos et al. 2008; Stern and Orgogozo 2009; Hart et al. 2018). In the case of marine stickleback invading freshwater environments, local adaptation must often occur in the presence of ongoing gene flow (Nelson and Cresko 2018). This scenario may initially favor trans-acting mechanisms that are less susceptible to being eroded through recombination. However, as the population reaches later stages of adaptation to the local environment, the advantage of responses mediated by trans-regulatory genes may shift to favor cis-regulatory mechanisms, where co-evolved mutations are more closely linked to each other and the genes they regulate (Verta and Jones 2019).

Our functional analysis identified multiple genes associated with meQTLs that are located in genomic regions that have been shown to have significant differentiation between marine and freshwater populations, and could therefore be relevant for local adaptation (Hohenlohe et al. 2010; Jones et al. 2012; Terekhanova et al. 2014) (Supplementary Table S3). For example, we found genes annotated with osmosis and electrolyte transport (KCNB2), and skeletal and fibroblast growth (5ko5a1a), which have also been found in previous stickleback studies investigating differential gene expression or methylation in gill or fillet tissue between marine and freshwater ecotypes (Smith et al. 2015; Artemov et al. 2017). These results suggest that this collection of genes might be important for facilitating adaptation to these divergent environments in stickleback, although the direction of causality between DNA methylation variation and gene expression remains elusive. Interestingly, it has been shown that genomic regions that are not significantly differentiated between ecotypes can still play an important role in adaptation to novel aquatic environments in stickleback (DeFaveri et al. 2011; Leinonen et al. 2012; Ellis et al. 2015; Erickson et al. 2016; Ferchaud and Hansen 2016). Our findings suggest that the genetic architecture underlying methylation divergence and physiological adaptation to different aquatic environments in stickleback is complicated and could include SNPs from genomic regions that experience either neutral or selective processes.

Limitations

Our study has a number of caveats that should be noted. First, an intrinsic problem of in vivo studies using next-generation sequencing techniques such as RRBS is the heterogeneity of analyzed tissues. Fin tissues consist of many different cell types including epidermis, osteoblasts, dermal fibroblasts, and vascular endothelium (Tu and Johnson 2011). Therefore, various proportions of different cell types could introduce biases in measures of methylation levels (Kratochwil and Meyer 2015). In addition, we only used fin tissues from a single developmental stage of sticklebacks, whereas methylation and gene expression patterns are known to be development-related and tissue-specific (Wang et al. 2009; Feil and Fraga 2012). Thus, overlap between the locations of meQTLs identified using fin tissue in this study and the eQTLs identified using brain tissue in Ishikawa et al. (2017) should be interpreted with caution. Further studies extending our work to a broader range of tissues and developmental stages will be helpful for a more comprehensive characterization of methylation variation, and its role in gene regulation and development.

Second, the reduced representation genome sequencing method used here can only cover a small proportion of all possible methylation patterns in these populations. Thus, we are inevitably missing a large number of stable CpG sites and SNPs located outside of the regions of the genome represented here. In addition, because the accuracy of SNP calls from bisulfite sequencing data can be affected by the conversion rate of unmethylated cytosines (Barturen et al. 2014), the SNPs identified in our study could be different than those that would be obtained using a sequencing method that produces independent SNP data (e.g. restriction-site associated DNA sequencing, RAD-seq; Baird et al. 2008). Furthermore, while we mainly focused methylation patterns in promoters and gene bodies, other regulatory elements such as enhancers and transposons, although less well annotated in stickleback, are also important drivers of regulatory and phenotypic evolution (Wittkopp and Kalay 2012) and thus warrant further research.

Third, the number of individuals and families used in our study is limited. Thus, including additional samples from more families would provide additional information on family-level variation, as well as more loci associated with methylation variation that would increase the power of our heritability and meQTL analyses. In addition, we generated marine-freshwater F1 families by crossing marine females and freshwater males, and recent studies in zebrafish have suggested that epigenetic patterns at early developmental stages can often reprogram to reflect the paternal state (Jiang et al. 2013; Potok et al. 2013). Whether such reprogramming is common to teleosts remains unclear (Skvortsova et al. 2018), but additional reciprocal crosses using marine males and freshwater females, as well as pure crosses within marine or freshwater populations, would allow a more comprehensive understanding of parental effects on epigenetic inheritance (Laporte et al. 2019).

Finally, although we have corrected for the possibility of falsely interpreting C-to-T and G-to-A SNPs as epigenetic variation by excluding them from methylation estimates, it is possible that some SNPs were miscalled. Thus, our results provide a necessarily coarse map of the genetic architecture underlying stable methylation and methylation divergence between marine and freshwater stickleback populations. A wider investigation of regulatory elements in combination with genome-wide sequencing of chromatin modifications (e.g. chromatin immunoprecipitation followed by sequencing (ChIP-seq); Park 2009; Furey 2012) and whole-genome resequencing (e.g. Le Luyer et al. 2017) would provide a more comprehensive and precise understanding of the relationship between genetics and DNA methylation, and the role that epigenetic responses may play in facilitating evolutionary change.
Conclusions

Here, we provide the first insights into the genetic architecture of DNA methylation in threespine stickleback. Our genome-wide methylation data reveals that the vast majority of CpG sites have stable methylation levels across generations, including the sites that show significant divergence in methylation levels between marine and freshwater ecotypes. Some of these sites show evidence of genetic control, while others are likely to be autonomous from genetic variation. We also explored the genomic distribution of methylation in marine-freshwater hybrid populations and found meQTLs that overlap with previously identified genomic regions of high differentiation between marine and freshwater populations. In addition, our data demonstrates different contributions of cis- and trans-meQTLs to methylene divergence in stickleback. Our study adds to the few studies using non-model, outbred vertebrates to test for the genetic basis of intergenerational stability over the short-term, as well as population persistence and adaptation over longer evolutionary time scales.

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Conflicts of interest

None declared.

Literature cited


