

Genome-Wide Genotype-Expression Relationships Reveal Both Copy Number and Single Nucleotide Differentiation Contribute to Differential Gene Expression between Stickleback Ecotypes

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Abstract

Repeated and independent emergence of trait divergence that matches habitat differences is a sign of parallel evolution by natural selection. Yet, the molecular underpinnings that are targeted by adaptive evolution often remain elusive. We investigate this question by combining genome-wide analyses of copy number variants (CNVs), single nucleotide polymorphisms (SNPs), and gene expression across four pairs of lake and river populations of the three-spined stickleback (*Gasterosteus aculeatus*). We tested whether CNVs that span entire genes and SNPs occurring in putative *cis*-regulatory regions contribute to gene expression differences between sticklebacks from lake and river origins. We found 135 gene CNVs that showed a significant positive association between gene copy number and gene expression, suggesting that CNVs result in dosage effects that can fuel phenotypic variation and serve as substrates for habitat-specific selection. Copy number differentiation between lake and river sticklebacks also contributed to expression differences of two immune-related genes in immune tissues, *cathepsin A* and *GIMAP7*. In addition, we identified SNPs in *cis*-regulatory regions (eSNPs) associated with the expression of 1,865 genes, including one eSNP upstream of a carboxypeptidase gene where both the SNP alleles differentiated and the gene was differentially expressed between lake and river populations. Our study highlights two types of mutations as important sources of genetic variation involved in the evolution of gene expression and in potentially facilitating repeated adaptation to novel environments.

Key words: habitat-specific adaptation, CNV, copy number variation, eSNP, *cis*-regulatory regions, expression differentiation, three-spined stickleback.

Introduction

Uncovering the genetic mechanisms underlying adaptive evolution is a major research focus in evolutionary biology (Barrett and Hoekstra 2011). Adaptive phenotypes can result from changes in amino acid sequences that affect protein structure

and function (Hoekstra and Coyne 2007), as well as from alterations of gene expression patterns (Carroll 2008). Although gene expression can be plastic and respond to environmental stimuli (Gibson 2008), adaptive evolution of gene expression rests upon an inherited genetic basis.

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Gene expression differences between populations and species often carry a significant heritable component and impact fitness, contributing to adaptation (Stamatoyannopoulos 2004; Whitehead and Crawford 2006b; Pavey et al. 2010). A growing body of evidence has linked the acquisition of adaptive phenotypes in new environments to gene expression changes, including elongated beaks in cactus finches (Abzhanov et al. 2006), camouflage pigmentation in deer mice (Linnen et al. 2009; Mallarino et al. 2017), convergent thick lips in cichlids (Colombo et al. 2013), and repeated pelvic loss in three-spined sticklebacks (Chan et al. 2010). If the differentiation in expression confers an adaptive advantage across independent population clines, it may lead to parallel evolution at the gene expression level. The parallel evolution of expression patterns can be directly inferred when, for example, heritable gene expression variation correlates with an environmental cline rather than by ancestry (Whitehead and Crawford 2006a; Lenz 2015). Parallel gene expression has been observed in a few cases of diverging ecotypes or species of adaptive radiations (Derome et al. 2006; Pavey et al. 2010; Colombo et al. 2013; Manousaki et al. 2013; Stutz et al. 2015; Zhao et al. 2015; Hanson et al. 2017). Yet, the genetic variants associated with these gene expression patterns remain understudied.

Genomic studies of recurring ecotypes have revealed a major contribution of regulatory regions to parallel genomic divergence (Jones et al. 2012; Brawand et al. 2014). Combining gene expression surveys with genome-wide sequence analysis allows evaluating the role of genetic variants on the evolution of expression differences between ecotypes. The genetic basis of expression differences may reside in close physical proximity of a gene (in *cis*) or far away (in *trans*) (Gilad et al. 2008). Genetic mutations altering the sequence of *cis*-regulatory elements can affect the binding affinity of transcription factors, whose effects are mainly limited to expression variation levels of neighboring genes, whereas mutations that affect *trans*-regulatory elements typically encode transcription factors that regulate multiple downstream genes (Wittkopp and Kalay 2011). Due to the local effects of *cis*-regulatory elements that confer a lower extent of pleiotropy compared with *trans*-, *cis*-regulatory elements have been suggested to be more important than *trans*-regulatory elements in the expression divergence between species (Wittkopp et al. 2008).

In addition to sequence changes in its regulatory region, the number of copies of a particular gene can affect its expression. Gene copy number can differ among individuals due to genetic deletions and duplications, giving rise to copy number variations (CNVs), which natural selection can act upon (Nguyen et al. 2006; Katju and Bergthorsson 2013). Copy number is generally positively correlated with expression levels (Haraksingh and Snyder 2013; Gamazon and Stranger 2015), producing a gene dosage effect (Zhang et al. 2009). Gene dosage effects are often detrimental to fitness as they can

disrupt the stoichiometric balance in molecular networks (Papp et al. 2003; Veitia 2005; Birchler and Veitia 2012) and have been associated with diseases (Rice and McLysaght 2017). However, in some cases, dosage effects of CNVs have also been beneficial, such as the relationship observed between amylase gene copy numbers and starch diets in both humans and dogs (Perry et al. 2007; Axelsson et al. 2013), and the number of cytochrome P450 genes in insecticide-resistant populations of dengue mosquitoes (Faucon et al. 2015). Although variation in *cis*-regulatory elements and CNVs can both affect gene expression and contribute to adaptive phenotypes, their contribution to habitat-specific gene expression has not been systematically studied. Genotype-expression relationships become particularly interesting when divergence patterns across replicated populations independently adapted to different environments occur at both the genetic and expression levels, strongly suggesting a genetic basis underlying adaptive expression variation.

The three-spined stickleback (*Gasterosteus aculeatus*) is a powerful model species to investigate habitat-specific adaptation. After the last glaciation, marine three-spined sticklebacks repeatedly colonized different freshwater habitats, resulting in an adaptive radiation composed of habitat-specific ecotypes (McKinnon and Rundle 2002). In particular, recurrent adaptation to lakes and rivers (or streams) has given rise to distinct ecotypes across the northern hemisphere (Reusch et al. 2001), with morphological differences in body shapes and traits involved in foraging (Berner et al. 2008; Deagle et al. 2012; Kaeuffer et al. 2012; Ravinet et al. 2013; Lucek et al. 2014). Another profound difference between lake and river habitats is the distinct parasite community, in which lake fish generally suffer from a higher parasite burden than river fish, likely contributing to recurrent ecotype differences at both the phenotypic and genetic level (Kalbe et al. 2002; Eizaguirre et al. 2011; Feulner et al. 2015). Transcriptome analyses have revealed over a hundred genes with habitat-specific gene expression among wild-caught lake and river sticklebacks (Huang et al. 2016), some of which were also differentially expressed between lake and river sticklebacks in a laboratory-controlled parasite infection experiment (Lenz et al. 2013). Lake and stream sticklebacks raised in common garden conditions also exhibit parallel gene expression differences (Hanson et al. 2017). These results suggest a heritable component to habitat-specific gene expression, which is also supported by quantitative genetics analyses on pedigrees of sticklebacks (Leder et al. 2015). In sticklebacks, a greater contribution of *cis*-regulatory elements than *trans*-regulatory elements in expression variation and divergence between ecotypes has been suggested (Ishikawa et al. 2017; Pritchard et al. 2017; Verta and Jones 2019). However, unlike the parallel divergence observed between marine and freshwater sticklebacks at the sequence level (Jones et al. 2012) and in gene CNVs (Hirase et al. 2014), a

low degree of parallel genetic differentiation exists among repeatedly diverged lake and river ecotypes, both at the sequence level (Deagle et al. 2012; Roesti et al. 2012; Feulner et al. 2015; Stuart et al. 2017) and in copy numbers (Chain et al. 2014). This is despite habitat-specific patterns of gene expression (Huang et al. 2016; Hanson et al. 2017). Given the low degree of genomic parallelism, the genetic variation underlying the expression divergence between lake and river ecotypes remains elusive.

In this study, stickleback genomes and transcriptomes from the exact same individuals were used to study the molecular basis of habitat-specific adaptations between lake and river ecotypes. To identify candidate genes involved in adaptation to distinct parasite communities in lakes and rivers, we evaluated the relationships between gene expression variation in immune tissues and two types of variants, gene CNVs and single nucleotide polymorphisms (SNPs) in *cis*-regulatory regions. We tested for 1) associations between gene copy numbers or SNP genotypes and gene expression within and across individuals, 2) evidence of habitat-specific selection as inferred from different gene copy numbers between ecotypes or allele frequency differentiation of SNPs, and 3) differential gene expression between ecotypes. These serve as three pillars of evidence that genetic changes contribute to adaptive gene expression differences between ecotypes. In this way, we identified genetic variants that influence repeated differential expression between ecotypes, putatively contributing to habitat-specific adaptation.

Materials and Methods

Sampling Design

To study the genetic differentiation between lake and river stickleback ecotypes that underlie expression differentiation, we combined a whole genome and a whole transcriptome data set from a total of eight geographically widespread populations of three-spined sticklebacks that had been previously analyzed separately. The whole-genome sequence data set consisted of 48 fish from four parapatric population pairs; two independent drainages from Germany (G1 and G2), one from Norway (No), and one from Canada (Ca), with six individuals from each lake (_L) and each river (_R), respectively (Chain et al. 2014; Feulner et al. 2015; EBI Accession no: PRJEB5198; [supplementary fig. S1](#) and [table S1](#), [Supplementary Material](#) online). The average genomic coverage was 26-fold, and genotypes from the whole-genome sequencing were validated with 98% concordance by Illumina's Golden Gate assay (Feulner et al. 2015), yielding reliable SNP data reused in this study. The whole-transcriptome data set comprised gene expression data from a subset of the same individuals as referenced above (43 total fish, matched IDs indicated in [supplementary table S1](#), [Supplementary Material](#) online). These transcriptomes

were previously used to investigate habitat-specific gene expression between lake and river ecotypes (Huang et al. 2016). To understand the adaptation to distinct parasite environments between lake and river habitats, we focused on two immune tissues: we used 40 head kidney transcriptomes and 36 spleen transcriptomes (PRJEB8677). The average transcriptome library size was 6.5 million pair-end reads, which has limited power to detect genes with low expression but should be robust to quantify differences among medium to highly expressed genes (Tarazona et al. 2011; Ching et al. 2014).

Expression Profiling

Transcriptome libraries from the sampled populations were first analyzed following Huang et al. (2016; Dryad doi: 10.5061/dryad.hq50s). In short, transcriptome libraries from head kidneys and from spleens were analyzed separately. Weakly expressed genes with less than one read count per million in at least half of the respective tissue samples were removed and then libraries were normalized using the trimmed mean of *M*-value method (Robinson and Oshlack 2010) in EdgeR (Robinson et al. 2010). Expression levels were estimated as the log of normalized read count per million. The final set of expression profiles consisted of 12,105 genes from the head kidney and 12,451 genes from the spleen that were used in the analyses described below.

Identification of Gene eCNVs

CNV regions of the study populations were identified by Chain et al. (2014), where CNVs were assigned using consensus calls from the read depth approach implemented in the software CNVnator (Abyzov et al. 2011) and at least one other approach (paired-end and split-reads; for details see Chain et al. 2014). We identified genes with at least 95% length overlap with CNVs. Gene copy number was estimated using CNVnator and rounded to the closest integer. Genes showing no variation in estimated copy numbers among individuals of our study were excluded from copy number analyses. Genes with copy number estimates of zero but with detectable read depth >0 were removed from our analyses to avoid possible false deletion calls. A total of 832 autosomal protein-coding genes remained, referred herein as "gene CNVs."

Using gene copy numbers and the corresponding gene expression from the same fish, we evaluated the association between gene copy number and expression level for each gene CNV in each individual, and for each tissue type separately. Using a linear mixed effect model, gene copy number was set as a fixed effect, and the population and sex were set as random effects (expression levels \sim copy_number + (1|population) + (1|sex)). This approach makes use of the continuous nature of copy number genotypes and tests for dosage effects of CNVs, which is different

from the typical eQTL approach that associates expression variation to categorical genotypes. Benjamini–Hochberg’s multiple-test correction was applied to the P values of the fixed effect of copy number (Benjamini and Hochberg 1995). Genes with corrected P values <0.05 and positive correlation were considered as “gene eCNVs,” having statistically significant correlations between copy number and expression.

Identification of eSNPs

In addition to the evaluation of gene eCNVs, we mapped SNPs in *cis*-regulatory regions (eSNPs) to identify potential *cis*-regulatory elements that underlie gene expression variation. The eSNPs were determined for gene expression in head kidney and spleen separately, using SNPs within a 5 kb range of the transcription start sites (TSSs). We reasoned that the 5 kb upstream regions serve as a proxy for the location of potential *cis*-regulatory elements, based on empirical findings of *cis*-regulatory sequences in mouse (Shen et al. 2012). SNPs used in this study were extracted from a previous genome-wide survey (Feulner et al. 2015), excluding SNPs in CNV regions due to potential detection biases (Hartasánchez et al. 2018) and filtering SNPs for a minor allele frequency >0.05 in the four population pairs combined. Out of 12,105 and 12,451 genes expressed in the head kidney and in the spleen, respectively, 10,803 and 10,914 genes had a total of 815,341 and 841,063 SNPs, and jointly 870,917 SNPs that fulfilled our filtering criteria. For each expressed gene, we tested for a significant association between each SNP and expression levels in FastQTL v2.165 (Ongen et al. 2016) using the nominal pass and correcting for population stratification (population pairs and habitats) and sex. Two steps of multiple-testing correction (Benjamini–Hochberg) were applied on the P value for each SNP: the P values were first corrected for numbers of SNPs per gene and then for the total number of genes tested. SNPs with corrected P values <0.05 were considered as eSNPs.

Expression Differentiation between Stickleback Ecotypes

Differential expression (DE) analyses implemented in the package EdgeR was previously used to identify significantly differentially expressed genes between ecotypes, indicative of habitat-specific gene expression (Huang et al. 2016). To complement this binary categorization, we quantified the extent of expression differentiation in a continuous manner by computing the variable P_{CT} , which evaluates the relative variance in expression between groups (here lake vs. river ecotypes) compared with the variance within groups. We calculated P_{CT} between lake and river sticklebacks and accounted for expression variances among geographic population pairs and between sex using an ANOVA-based approach (methods adapted from Uebbing et al. 2016). P_{CT} as a measure of relative differentiation in gene expression between lake and river

ecotypes was calculated for each expressed gene and for the head kidney and spleen separately. Because the calculation of P_{CT} is conceptually equivalent to the calculation of copy number differentiation (V_{CT} , see below) and nucleotide differentiation (F_{CT} , see below), the evaluation of expression differentiation is made directly comparable to that of genetic differentiation. To determine how likely each P_{CT} value was obtained by chance, we recalculated P_{CT} 1000 times for each gene after random permutations of the ecotype labels. The p -values were calculated as the fraction of permuted values that exceeded the observed value and were corrected by the Benjamini-Hochberg method for the numbers of genes tested (Benjamini and Hochberg 1995). Genes with adjusted P values <0.05 for P_{CT} were considered significant. For candidate genes, we also calculated P_{CT} between each population pair separately, in the same way that P_{CT} was calculated for all populations combined but without population stratification in the ANOVA model.

Copy Number Differentiation of Gene eCNVs

In order to investigate the contribution of gene eCNVs in expression differentiation, we evaluated copy number differentiation between ecotypes across all population pairs together. For each gene eCNV, we calculated V_{CT} representing the relative variance in copy number between groups (here lake vs. river ecotypes) compared with the overall variance within groups, similarly to P_{CT} . V_{CT} was calculated using all individuals from the four population pairs with an ANOVA-based approach, where lake and river ecotypes were treated as two comparison groups, while accounting for variance between population pairs (copy_number \sim ecotypes * population_pair). As we exclude CNVs in the sex chromosome for our analyses, we did not include sex as a factor in the model. V_{CT} is different from V_{ST} , a measurement of copy number differentiation between populations without a nested structure (Redon et al. 2006), which was previously calculated on the same data set but between each lake and river pair separately in Chain et al. (2014). Including all population pairs together to estimate copy number differentiation (V_{CT}) increases sensitivity to detect differences between ecotypes, as it does not require differentiation signals to be extreme in each pair. We applied 1000 permutations following the methods for P_{CT} to identify gene eCNVs with significant V_{CT} , and p -values were corrected by the Benjamini-Hochberg method for the number of genes tested (Benjamini and Hochberg 1995). V_{CT} with corrected P values <0.05 were considered significantly differentiated between lake and river ecotypes. For candidate gene eCNVs, we also calculated V_{CT} between each population pair separately, in the same way as V_{CT} was calculated for all populations combined but without population stratification in the ANOVA model.

Allelic Differentiation of eSNPs

In addition to the evaluation of copy number differentiation, we calculated nucleotide differentiation between lake and river ecotypes for each SNP identified as eSNPs, evaluated as F_{CT} using the locus-by-locus AMOVA approach implemented in Arlequin (Excoffier and Lischer 2010). The F_{CT} was calculated as the percentage of variance between groups (lake vs. river ecotypes) relative to the total variance, using a hierarchical structure that groups lake and river ecotypes into four populations each. This AMOVA approach provides a more sensitive way to qualitatively evaluate habitat-specific patterns across replicated population pairs, compared with methods that scan for outlier regions in each population pair separately to identify parallel regions based on shared outliers (e.g., Feulner et al. 2015), for the same reason as mentioned above for V_{CT} . We used permutation tests implemented in Arlequin to determine the significance of the F_{CT} values and identify eSNPs with significant F_{CT} values ($P < 0.05$ from 1,023 permutations).

Identifying Correlations between Expression and Genetic Differentiation

A genome-wide correlation between gene expression differentiation (P_{CT}) and genetic differentiation (V_{CT} and F_{CT}) was performed on all expressed genes. For this analysis, V_{CT} was calculated for each of 350 gene CNVs that had expression (not gene eCNVs), and F_{CT} was calculated for each of 11,935 autosomal protein-coding genes that had expression (not only for eSNPs), excluding genes in CNV regions. F_{CT} was evaluated for each gene based on SNPs in the 5 kb upstream regions, using the AMOVA approach implemented in Arlequin (Excoffier and Lischer 2010). With the resulting matrixes of P_{CT} , V_{CT} , and F_{CT} of all genes expressed in the head kidney and/or spleen, the Spearman's rank correlation was used to test for correlation in each tissue 1) between P_{CT} and V_{CT} and 2) between P_{CT} and F_{CT} . All statistical analyses were carried out using the package R version 3.0.2 (R Development Core Team 2011) unless otherwise indicated.

Testing for Gene Ontology Enrichment in Genes with eSNPs and eCNVs

We tested for enrichment of gene ontology (GO) terms among the gene eCNVs, the genes with eSNPs, the gene eCNVs with significant V_{CT} , and the genes with eSNPs with significant F_{CT} . The enrichment tests were conducted with topGO (Alexa et al. 2006), based on Fisher's exact tests applying Benjamini-Hochberg's multiple-test correction. We used different background gene sets depending on the enrichment analysis: we compared gene eCNVs to all expressed genes in either tissue and to all gene CNVs that are expressed in either tissue; we compared genes with eSNPs to all genes that were included in the eSNP tests; we compared gene

eCNVs with significant V_{CT} to the set of gene eCNVs; we compared genes with eSNPs with significant F_{CT} to all genes with eSNPs. Overrepresented GO terms were those with corrected P values < 0.05 .

Results

We first evaluated genotype-expression relationships using CNVs and SNPs, and then investigated whether they contribute to expression divergence between ecotypes. Our overarching goal was to evaluate the relationship between genetic differentiation of the two variant types and gene expression differentiation between replicated pairs of lake and river three-spined stickleback ecotypes.

Gene Copy Numbers and Expression Levels Are Largely Positively Correlated

Out of a total of 19,782 protein-coding autosomal genes in stickleback genome, we identified 832 gene CNVs among our samples. Among these gene CNVs, 350 CNVs had available gene expression data, out of which 140 (40%) had a significant association between gene copy numbers and gene expression in at least one of the two immune tissues (corrected P values < 0.05). Five of these genes had a significant negative correlation between copy numbers and expression levels: *WBP1* (*VW domain binding protein 1*, ENSGACG00000000318), *slc47a1* (*solute carrier family 47, member 1*, ENSGACG00000020614) and two uncharacterized genes (ENSGACG00000020469 and ENSGACG00000012806) in head kidney samples, as well as *cyp3c1* (*cytochrome P450 family 3 subfamily A member 43*, ENSGACG00000010952) in spleen samples. The other 135 genes (39% of all expressed gene CNVs) had a positive correlation in at least one of the two tissues and were considered "gene eCNVs" (fig. 1 and supplementary table S2, Supplementary Material online). Among these 135 gene eCNVs, 13 were only expressed and had a positive correlation in one tissue (5 in head kidney and 8 in spleen), 62 were expressed in both tissues but correlated with copy number in one tissue, while 60 were expressed and correlated in both tissues. Among the genes that were expressed in either the head kidney or spleen tissues, gene eCNVs were enriched for antigen processing and presentation (GO: 0019882, with 4 out of 28 genes), immune response (GO: 0006955, with 5 out of 72 genes), major histocompatibility complex (MHC) protein complex (GO: 0042611 with 4 of 27 genes), and MHC class I protein complex (GO: 0042612 with 4 of 18 genes). MHC immune genes were among functional categories that were previously reported as enriched among all gene CNVs in sticklebacks (Chain et al. 2014). When comparing gene eCNVs against all gene CNVs that were expressed in either tissue, there was no GO term enrichment observed.

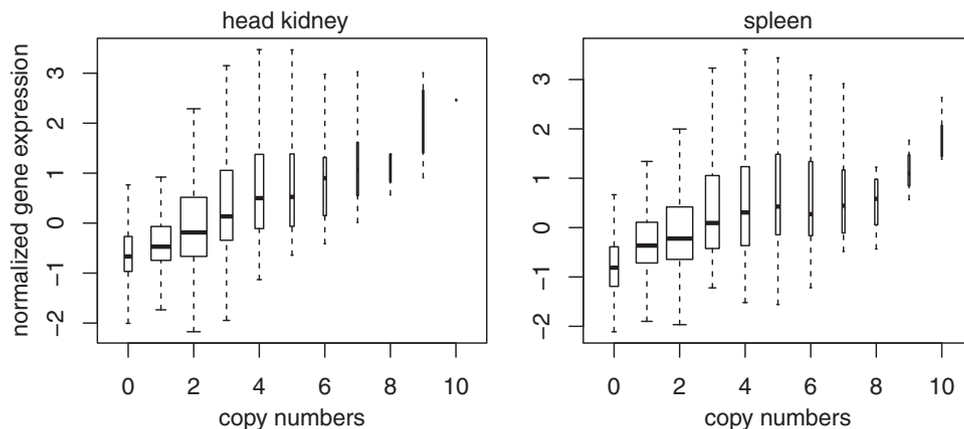


FIG. 1.—Normalized gene expression levels for a given gene copy number summarized across all gene eCNVs and individuals (n genes = 135; n individuals = 40 for head kidney, and n = 36 for spleen). Expression levels were evaluated in head kidney and spleen separately. Expression levels of each gene were centered to zero and scaled by the SDs. The widths of boxes represent the relative sample size (i.e., number of genes in each copy number category). Only copy numbers up to ten are shown.

Ten eCNVs Show Copy Number Differentiation between Ecotypes

As gene eCNVs are the putative genetic variants that affect gene expression, we evaluated differentiation in their gene copy numbers between ecotypes, which could contribute to gene expression divergence. We estimated V_{CT} for each gene eCNV, which is the relative variance in gene copy numbers between ecotypes compared with the variance within ecotypes. Out of the total 135 gene eCNVs, 10 (7.4%) have a significant V_{CT} (FDR < 0.05, permutation test), with V_{CT} values ranging from 0.144 to 0.578 (table 1). Of these ten genes, seven have higher average copy numbers in lake ecotypes than in river ecotypes, and three have higher copy numbers in river ecotypes. The 10 gene eCNVs with significant V_{CT} are distributed across 6 of 20 stickleback autosomes (fig. 2a). The GO annotations of the ten V_{CT} significant genes show that they are associated with various functions including ion binding, GTP binding, peptidase activity, diphosphatase activity, and transmembrane transport (table 1). But there was no functional enrichment of the ten gene eCNVs with significant V_{CT} compared with all gene eCNVs.

An Abundance of Genes with SNPs in *Cis* Is Associated with Expression

In addition to the CNVs associated with gene expression, we also investigated SNPs that are associated with gene expression. Out of a total of 870,917 SNPs within 5kb range of the TSSs of 11,360 genes expressed in either tissues, 8,353 SNPs were found associated with expression of 1,351 genes in the head kidney, 4,261 SNPs associated with expression of 746 genes in the spleen, including 1,336 SNPs associated with expression of 232 genes in both tissue types (corrected P values < 0.05, supplementary table S3, Supplementary

Material online). In total, 11,278 SNPs associated with 1,865 genes were determined as eSNPs that putatively contribute to gene expression differences among individuals. These eSNPs are symmetrically distributed across the 5 kb upstream and downstream range, with a slight peak within the 1kb range of the TSSs (supplementary table S3, Supplementary Material online). No GO term was enriched for the genes with eSNPs when compared with the joint set of 11,360 genes tested in the eSNPs analyses.

Fourteen eSNPs Show Allelic Differentiation between Ecotypes

For each eSNP, we evaluated the nucleotide differentiation, F_{CT} , between lake and river ecotypes. We found that 90.9% of eSNPs had negative or zero F_{CT} values, indicating no differentiation between lake and river fish populations. Out of the 1,112 eSNPs with a positive F_{CT} , 14 were significantly differentiated (P < 0.05, permutation test), with F_{CT} values ranging from 0.120 to 0.378 (fig. 2a). These 14 eSNPs were associated with expression of 14 different genes. These 14 genes are annotated with various functions spanning mRNA splicing, DNA binding, rRNA methylation, signal transduction, ATP binding, and GTP binding (table 2), with no significant enrichment of GO categories compared with the set of genes with eSNPs.

One eSNP and Two eCNVs Display Expression Differentiation between Ecotypes

The eSNPs and the gene eCNVs that are differentiated between ecotypes putatively contribute to expression differentiation. Among 12,105 genes expressed in the head kidney and 12,451 genes in the spleen, we identified 115 and 88 genes with significant P_{CT} , respectively (corrected P < 0.05, supplementary table S4, Supplementary Material online). Out of these genes, we found one gene with significant P_{CT} (0.217) in the

Table 1

Genes with Significant Differentiation in Gene Copy Numbers (V_{CT}) between Lake and River Ecotypes

Gene ID	Gene Name	GO Function			Tissue of eCNV	Higher Copy Number	V_{CT}	P_{CT}	
		Cellular Component	Molecular Function	Biological Process				HK	SP
ENSGACG00000008264	Novel Gene		Unknown		Both	River	0.245	-0.025	0.045
ENSGACG00000010952	Cytochrome P450 family 3 sub-family A member 43 (CYP3A43)	Membrane; integral component of membrane	Monoxygenase activity; iron ion binding; oxidoreductase activity; acting on paired donors, with incorporation or reduction of molecular oxygen; heme binding; metal ion binding	Oxidation-reduction process	SP	Lake	0.257	-0.024	-0.019
ENSGACG00000012073	Novel gene		Unknown		SP	Lake	0.278	NA	-0.021
ENSGACG00000015897	Cathepsin A	Unknown	Peptidase activity; serine-type carboxypeptidase activity; hydrolase activity	Proteolysis	Both	River	0.578	0.159	0.289*
ENSGACG00000016770	Deoxyuridine triphosphatase (dut)	Unknown	dUTP diphosphatase activity	dUTP metabolic process	Both	Lake	0.197	0.120	0.148
ENSGACG00000018877	GTPase, IMAP family member 7 (GIMAP7)	Unknown	GTP binding	Unknown	Both	Lake	0.348	0.245*	0.184
ENSGACG00000019933	si: dkey-85k7.12	Unknown	GTP binding	Unknown	Both	Lake	0.210	0.123	0.072
ENSGACG00000020614	Solute carrier family 47 (slc47a1)	Membrane; integral component of membrane	Drug transmembrane transporter activity; antiporter activity	Transmembrane transport	HK	Lake	0.178	0.032	-0.001
ENSGACG00000008242	Novel gene		Unknown		SP	River	0.144	0.082	0.002
ENSGACG00000009551	Ring finger protein 139 (rnf139)	Membrane; integral component of membrane	Zinc ion binding; metal ion binding,	Unknown	Both	Lake	0.215	0.186	0.136

NOTE.—NAs in P_{CT} , expression levels did not meet the filtering requirements and therefore P_{CT} were not calculated.

*Significant P_{CT} (Benjamini-Hochberg corrected $P < 0.05$).

head kidney that also had an eSNP with significant F_{CT} (fig. 2). The P_{CT} in the spleen was 0.142 (corrected $PR = 0.11$). The gene is *dehydrogenase/reductase (SDR family) member 13a, duplicate 3 (dhrs13a.3*, ENSGACG00000013614), a carboxypeptidase that catalyzes hydrolysis of peptide bonds (UniProt entry: G3PTQ4). The SNP residing 630 bp upstream of the TSS of this gene had a F_{CT} value of 0.204, and was significantly associated with gene expression in both tissues. We also found two genes with significant P_{CT} that exhibited both differentiation in copy numbers (significant V_{CT}) and significant correlations between gene copy numbers and gene expression (gene eCNVs) in both tissues. The gene *cathepsin A* (ENSGACG00000015897) had significant P_{CT} in spleen (0.289; P_{CT} of 0.159 in head kidney) and the highest V_{CT} (0.578) among all gene CNVs (fig. 2). The other gene, *GTPase, IMAP family member 7* (GIMAP7, ENSGACG00000018877), had significant P_{CT} identified in

head kidney (0.245; P_{CT} of 0.184 in spleen) and a V_{CT} of 0.348 (fig. 2).

eSNP Regulating Expression Differentiation in *dhrs13a.3*

Examining the differentiation signals within each population pair, the gene *dhrs13a.3* had higher expression levels in a subset of lake populations: in the head kidney of G1 ($P_{CT} = 0.648$) and G2 ($P_{CT} = 0.204$) but not in No (negative P_{CT}) and Ca ($P_{CT} = 0.076$) (fig. 3c); in the spleen of G1 ($P_{CT} = 0.305$) and No ($P_{CT} = 0.184$) but not in G2 ($P_{CT} = 0.076$) and Ca (negative P_{CT}). The genotypes of the eSNP residing 630 bp upstream of the TSS of *dhrs13a.3* were significantly correlated with gene expression levels across individuals in both tissue types (corrected $P < 0.001$, fig. 3a showed in head kidney). This SNP was differentiated between lake and river ecotypes and had consistently higher allele frequency of the

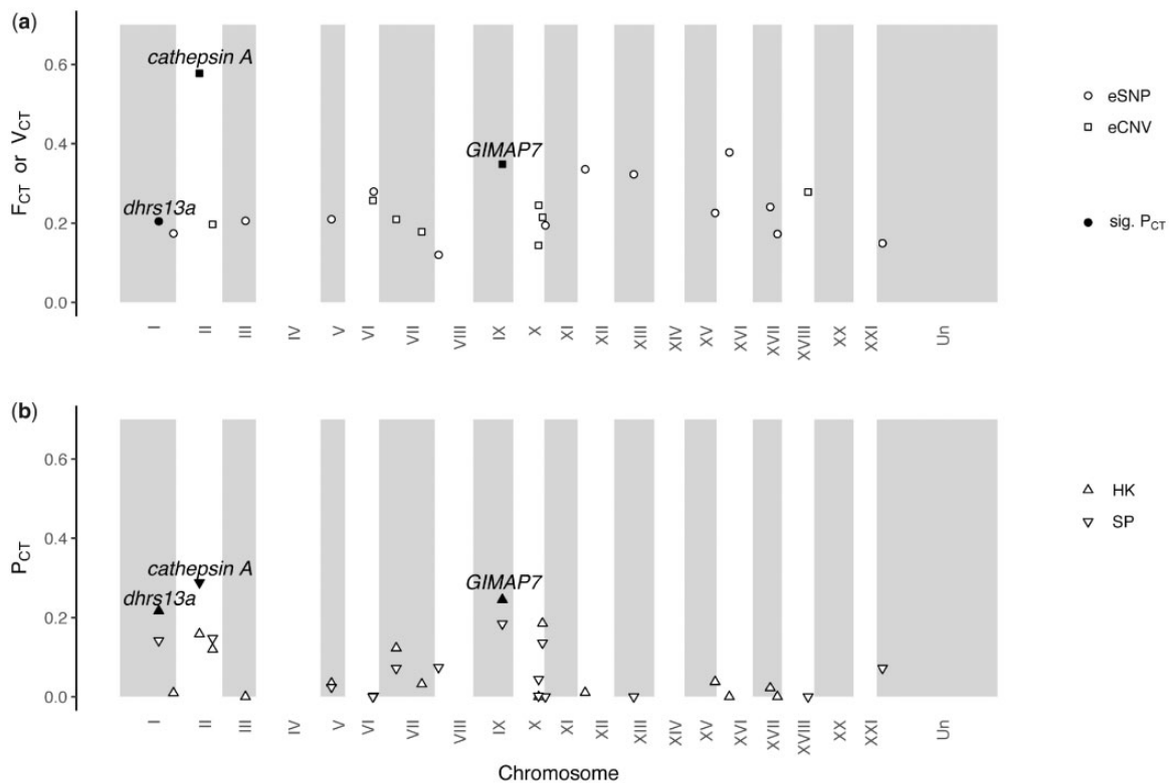


Fig. 2.—Genes with eSNPs with significant F_{CT} and eCNVs with significant V_{CT} between lake and river stickleback populations and P_{CT} of these same genes. (a) Genes with eSNPs with significant F_{CT} (circle) and gene eCNVs with significant V_{CT} (square) along the genome; and (b) P_{CT} of these same genes in the head kidney (triangle) and/or in the spleen (inverted triangle). Only the P_{CT} in the tissues where the eSNPs or the gene eCNVs were identified are shown. Genomic locations include 20 linkage groups of the stickleback genome representing autosomes (excluding the sex chromosome XIX), in addition to unplaced scaffolds (Un). The filled shapes indicate the three genes with significant P_{CT} .

allele G in the lake populations (fixed in G1_L and G2_L, and 83.3% in No_L and Ca_L) and higher allele frequency of T in the river populations (25% in G1_R, and 41.7% in G2_R, No_R, and Ca_R, [fig. 3b](#)). Both alleles occur in all four population pairs, and we confirmed that both were also present in an adjacent marine population from the North Sea ([Feulner et al. 2013](#)), with a low frequency of the T allele (8.3%). This suggests that the T allele derives from standing genetic variation in the ancestral marine populations, and repeatedly increased in frequency among river populations possibly due to positive selection. However, no selective sweep was found based on nucleotide diversity (π) in the 50-kb flanking region of the SNP, which did not differ between lake and river populations ([fig. 4a](#)). The gene region of *dhrs13a.3* harbors 51 SNPs across the four population pairs, with two synonymous and two nonsynonymous SNPs in the exons, and other SNPs in the introns. The nonsynonymous SNP, which substitutes a glycine with an arginine in the first exon, has the minor allele present in G1_L and G2_R with frequencies of 50% and 16.7%, respectively. The other nonsynonymous SNP, which substitutes a cysteine with a phenylalanine in the third exon, has the minor allele present in Ca_L with a frequency of 16.7%.

eCNV Regulating Expression Differentiation in *Cathepsin A*

The gene *cathepsin A* had higher expression levels in spleen among river sticklebacks in the two German population pairs G1 ($P_{CT} = 0.664$) and G2 ($P_{CT} = 0.409$; [fig. 3f](#)), but was not differentially expressed in No nor Ca (negative P_{CT} values). In head kidney tissues, this gene also had higher expression in river sticklebacks in the population pairs of G1 ($P_{CT} = 0.797$) and G2 ($P_{CT} = 0.190$) and Ca ($P_{CT} = 0.112$) whereas in No it had higher expression in the lake fish ($P_{CT} = 0.521$). The consistent DE in the two German population pairs was accompanied by copy number differentiation. This gene was the most differentiated gene CNV between lake and river sticklebacks in the two German population pairs (V_{CT} of 0.96 in G1 and 0.51 in G2) as previously reported ([Chain et al. 2014](#)), but not differentiated in No nor Ca ($V_{CT} = 0$) suggesting that the two German population pairs drive the overall habitat-specific signal ([fig. 3e](#)). We further identified *cathepsin A* as a gene eCNV, meaning that the gene copy numbers were significantly correlated with gene expression levels across individuals (corrected $P < 0.001$ in both tissue types, [fig. 3d](#)). To investigate whether the *cathepsin A* CNV is derived from standing genetic variation from an ancestral population, we searched

Table 2
Genes with Significant Differentiation in eSNPs (F_{CT}) between Lake and River Ecotypes

Gene ID	Gene Name	Cellular Component			GO Function			eSNP			F_{CT}		
		Cellular Component	Molecular Function	Biological Process	Position in Relation to TSS	Distance to TSS (bp)	Tissue	HK	SP	P_{CT}			
ENS5GACG00000000642	Novel Gene	Unknown	Unknown	Oxidation-reduction process	Up-stream	1,637	SP	NA	0.073				
ENS5GACG000000002647	alkB homolog 6 (alkbh6)	Unknown	Oxidoreductase activity	Oxidation-reduction process	Down-stream	3,811	HK	-0.021	-0.005				
ENS5GACG000000003827	si: ch73-14h10.2	P-body; nucleus; spliceosomal complex; U6 snRNP; U4/U6 x U5 tri-snRNP complex	mRNA splicing, via spliceosome; mRNA processing; RNA splicing	RNA binding	Up-stream	3,063	SP	NA	0.075				
ENS5GACG000000004256	Activity-dependent neuroprotector homoeobox b (adnppb)	Nucleus	Nucleic acid binding; DNA binding	Erythrocyte maturation	Down-stream	2,315	HK	0.010	-0.028				
ENS5GACG000000004442	Myocardin-related transcription factor Ba (mrtfba)	Unknown	Unknown	Unknown	Down-stream	1,161	Both	0.035	0.024				
ENS5GACG000000004844	RAS related (rras)	Membrane	Nucleotide binding; GTPase activity; GTP binding	Signal transduction; Notch signaling pathway; maintenance of epithelial cell apical/basal polarity	Down-stream	1,404	SP	0.001	-0.006				
ENS5GACG000000009164	Ly1 antibody reactive homolog (lyar)	Unknown	DNA binding	Unknown	Down-stream	1,080	HK	0.023	0.012				
ENS5GACG000000009941	DIM1 dimethyladenosine transferase 1-like (dimt1l)	Unknown	rRNA (adenine-N6, N6)-dimethyltransferase activity; RNA binding; methyltransferase activity; rRNA methyltransferase activity;	rRNA modification; rRNA processing; rRNA methylation; methylation	Down-stream	3,438	SP	-0.004	-0.028				
ENS5GACG000000011156	Sulfotransferase family, cytosolic, 6b, member 1 (sult6b1)	Unknown	Sulfotransferase activity; transferase activity	Cellular response to xenobiotic stimulus	Down-stream	2,520	SP	0.279	0.230				
ENS5GACG000000011426	si: ch73-267c23.10	Membrane; integral component of membrane	Unknown	Unknown	Down-stream	1,570	HK	0.173	-0.012				
ENS5GACG000000013118		None	Nucleic acid binding	Unknown	Up-stream	2,912	HK	0.226	0.039				

ENSGACG00000013614	B cell CLL/lymphoma 11B (BCL11B) Dehydrogenase/reductase (SDR family) member 13a, duplicate 3 (dhrs13a.3)	Unknown	Up-stream	630	Both	0.205	0.217 *	0.142
ENSGACG00000015279	Downstream neighbor of SON (DONSON)	Unknown	Up-stream	1,525	HK	0.174	0.009	0.268
ENSGACG00000016707	ATP-binding cassette, subfamily A (ABC1), member 4b (abca4b)	Membrane; integral component of membrane Nucleotide binding; ATP binding; ATPase activity; ATPase activity, coupled to transmembrane movement of substances	Down-stream	2,738	HK	0.206	-0.008	-0.002

NOTE.—NAs in P_{CT} , expression levels did not meet the filtering requirements and therefore P_{CT} were not calculated.

*Significant P_{CT} . (Benjamini-Hochberg corrected $P < 0.05$)

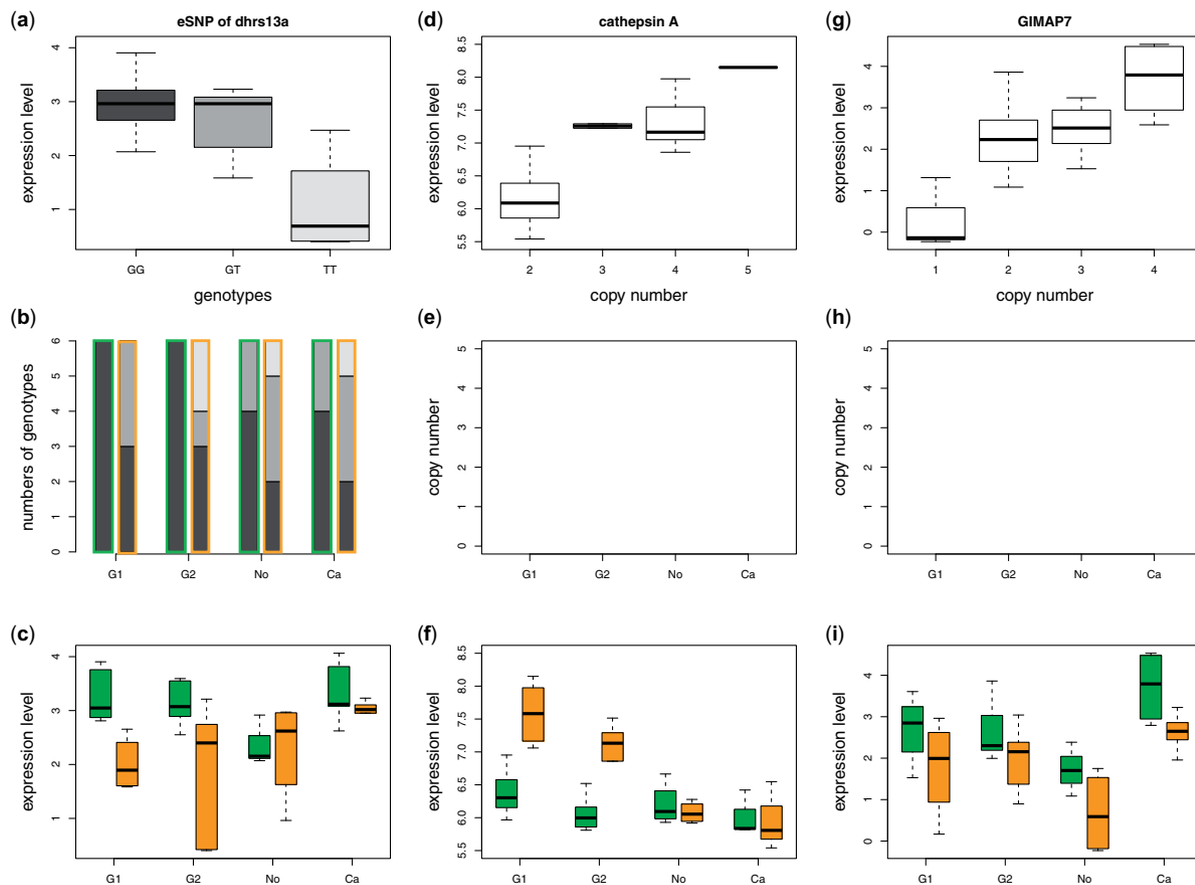


FIG. 3.—Gene *dhrs13a* with an eSNP with significant F_{CT} and gene eCNVs, *cathepsin A* and *GIMAP7*, with significant V_{CT} that also had significant P_{CT} between lake and river sticklebacks. (a) Association between eSNP genotypes and expression levels in the head kidney of *dhrs13a*, with y axes indicating expression levels of the different genotypes in boxplots summarizing normalized read counts across individuals. (b) Genotypes of the eSNP across four population pairs (G1: Germany 1, G2: Germany 2, No: Norway, Ca: Canada) where the bars with green border represent lake populations and the bars with orange border represent river populations. The colors for the genotypes are the same as in (a). (c) Expression differences in the head kidney across the same individuals where lake populations indicated in green and river populations in orange. (d and g) The association between gene copy numbers and gene expression in *cathepsin A* in the spleen (d) and *GIMAP7* in the head kidney (g). (e and h) Habitat-specific patterns of gene copy number of *cathepsin A* (e) and *GIMAP7* (h) across populations (dots represent lake and river individuals in green and orange, respectively). (f and i) the habitat-specific expression patterns of the same two genes, *cathepsin A* in the spleen (f) and *GIMAP7* in the head kidney (i) across populations.

for the presence of CNVs in the adjacent marine population from the North Sea (Feulner et al. 2013). The gene *cathepsin A* was not a CNV in the marine population, suggesting that the gene duplication occurred since the divergence of the freshwater populations (G1 and G2) from the marine population, or that the marine samples that were sequenced did not capture this variation. Note that the marine sampling only consists of six individuals, hence we lack power to detect variants at low frequency. A 5-kb region in the gene region of *cathepsin A* was depleted from SNPs in G1_R leading to a nucleotide diversity (π) of zero despite being duplicated compared to G1_L, suggesting a signature of background selection on the duplication (fig. 4b). In the other German populations, the gene harbors 23 SNPs, with two synonymous and one nonsynonymous SNP. The nonsynonymous SNP,

which substitutes a leucine by a phenylalanine in an alternatively spliced exon, has the minor allele present as heterozygous in three individuals in G1_L and in two individuals in G2_L, and as homozygous in one G2_L individual.

eCNV Regulating Expression Differentiation in *GIMAP7*

The gene *GIMAP7* had overall higher expression levels in the head kidney among lake ecotypes, and comparisons within population pairs found consistent directional differences across population pairs (fig. 3). P_{CT} in the population pairs ranged from 0.11 in G1, to 0.19 in G2, and 0.39 in Ca whereas expression levels did not meet filtering criteria in No. The expression in spleen tissues displayed the same direction of expression changes between lake and river

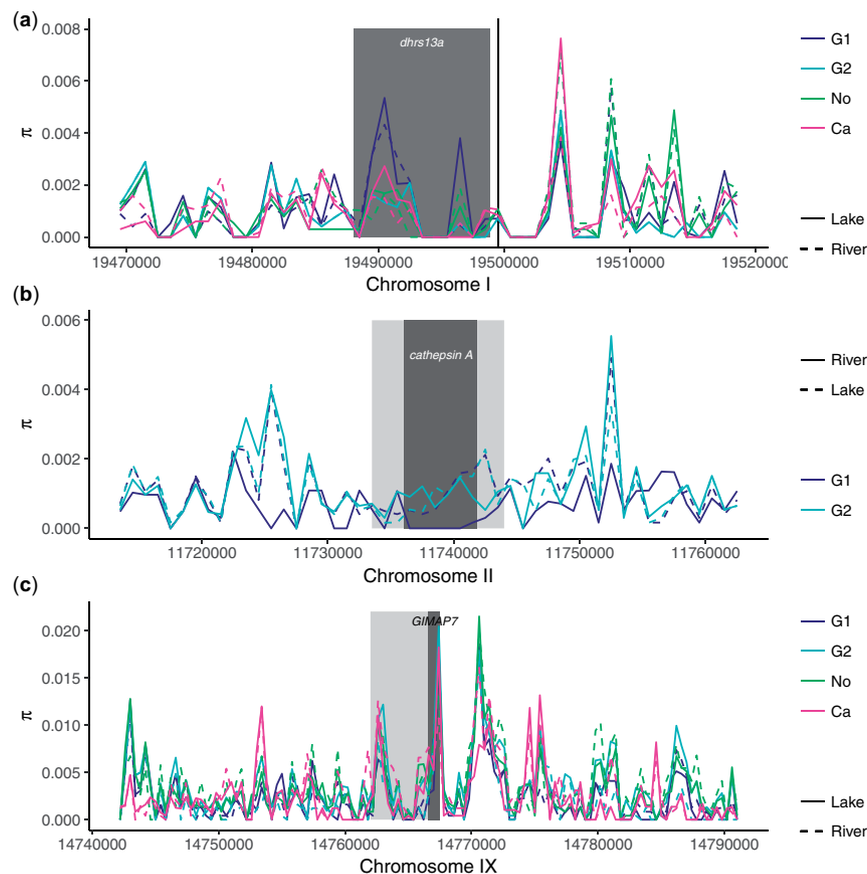


FIG. 4.—Nucleotide diversity (π) in the 50 kb flanking regions of the three candidate genes, the gene *dhrs13a* with an eSNP (a) and two eCNV genes, *cathepsin A* (b), and *GIMAP7* (c). In (a), the gene region is in dark gray and the eSNP denoted by a black vertical line. In (b) and (c), the gene regions are in dark gray and the CNV regions are in light gray. For the three genes, π was calculated for each population separately. For *dhrs13a* and *cathepsin A*, π was calculated for each 1 kb window and for *GIMAP7* π was calculated for 400 b window to adjust for SNP densities in each window. Solid lines represent populations with higher gene copy number (lake for *dhrs13a* and *GIMAP7* and river for *cathepsin A*) whereas dashed lines represent populations with lower gene copy number. For *cathepsin A*, we focused on G1 and G2 population pairs because CNVs were identified in only these two population pairs.

sticklebacks as in the head kidney, but differentiation was less pronounced: P_{CT} of 0.05 in G1, 0 in G2, 0.68 in No, and 0.07 in Ca. The V_{CT} values were reasonably high in at least three population pairs: 0.53 in G1, 0.64 in No, and 0.70 in Ca (fig. 3h). As with *cathepsin A*, *GIMAP7* was a gene eCNV (corrected $P=0.0074$ in head kidneys and corrected $P<0.001$ in spleen, fig. 3g). *GIMAP7* was not detected as a CNV in the North Sea marine population. This suggests independent duplication and deletion events in the freshwater populations since they diverged from the marine ancestor or that this variant is at low frequency in the marine population. In the genomic regions adjacent to *GIMAP7*, we found no differences in the levels of nucleotide diversity among the eight freshwater populations (fig. 4c). The gene region harbors a total of 38 SNPs across the four population pairs, 24 of which are nonsynonymous. This suggests that duplication and deletion of this gene might also contribute to the amino acid sequence diversification across population pairs.

Genome-Wide Correlation between Genetic Differentiation and Expression Differentiation

Genome-wide, F_{CT} in *cis*-regulatory regions did not significantly positively correlate with P_{CT} in either head kidney or spleen ($\rho=0.011$, $P=0.12$, $n=10,671$ in head kidney and $\rho=0.006$, $P=0.24$, $n=10,974$ in spleen; one-sided Spearman rank correlation). V_{CT} had a significant positive correlation with P_{CT} in spleen but not in the head kidney ($\rho=0.166$, $P<0.001$ for spleen; $\rho=0.064$, $P=0.064$ for head kidney; one-sided Spearman rank correlation).

Discussion

The genetic underpinnings of expression differentiation in adaptive evolution remain a focus of intense research. In this study, we combined genome-wide genetic variation and transcriptomic data from repeatedly evolved ecotypes of the three-spined stickleback to better understand their relationships in the process of adaptation to distinct habitats.

We first report a prevalent dosage effect of CNV genes on gene expression and numerous SNPs in *cis* associated with expression. The prevalent association between genetic variants and expression levels might provide phenotypic variation that promotes adaptation to distinct lake and river habitats. We describe one gene with a differentiated SNP that is associated with expression differentiation between lake and river populations, and two genes with significant associations between copy number differentiation and expression differentiation. These findings provide evidence that both SNPs and CNVs contribute to gene expression differentiation between recently diverged ecotypes.

Dosage Effects of CNVs Contribute to Expression Differentiation

CNVs reflect components of genome architectures that vary in the number of copies of a sequence and have been proposed to have a greater impact on gene expression compared with sequence modifications (Sudmant et al. 2015; Huddleston and Eichler 2016). We found that 39% (135) of all expressed gene CNVs have a positive association with expression in at least one of the two tissues sampled, with 60 gene CNVs showing significant positive association in both tissues. These results demonstrate prevalent dosage effects on gene expression across tissue types. Similar number of genes show associations between CNVs and expression changes in humans (e.g., 110 genes in Schlattl et al. [2011] and 44–96 genes in Stranger et al. [2007]) and a similar proportion (42%) of genes in *Drosophila* (Cardoso-Moreira et al. 2016). Recently, the Genotype-Tissue Expression (GTEx) Project also found large effect sizes of structural variations on gene expression in humans and highlighted the likely causality of many CNVs (Chiang et al. 2017). This is consistent with our findings of 135 gene eCNVs as putative causal variants for expression variation. While the 135 eCNVs are not enriched in any particular function compared with the expressed gene CNVs, they are enriched for functions of antigen processing and genes of the adaptive immune system (MHC genes) compared with the whole set of expressed genes. These two immune-related functional categories are a subset of enriched functions of gene CNVs overall (Chain et al. 2014), suggesting that the immune system might be amenable to expression differentiation via copy number changes. It is plausible that immune-related gene CNVs play an important role in adaptation to different parasite pressure in their natural environments, and contribute to observed divergences between lake and river ecotypes (Eizaguirre et al. 2009, 2011; Eizaguirre and Lenz 2010).

The integration of differentiation patterns of gene copy numbers and gene expression among gene CNVs in the same individuals enabled us to investigate the dosage effects of CNVs in the context of ecotype divergence. However, there was a weak genome-wide correlation between P_{CT} and V_{CT} . This is consistent with work performed on

Drosophila showing that the parallel differentiation of CNVs does not necessarily correlate with expression differentiation (Schridder et al. 2016). These together indicate that not all CNVs affect expression, at least not in all tissues, and that mechanisms other than linear dosage effects are also relevant. For example, some gene CNVs can be dosage insensitive (Zhou et al. 2011), and others can affect gene expression through compensatory mechanisms (Henrichsen et al. 2009). Although not all CNVs are expected to contribute to population differentiation, the ones where copy numbers and expression are differentiated between ecotypes are promising candidate genes involved in adaptation.

Genes underlying divergent adaptation should possess both high copy number differentiation (V_{CT}) and high expression differentiation (P_{CT}) between ecotypes, in addition to showing a positive correlation between copy numbers and gene expression levels (i.e., gene eCNVs). Here, we detected two genes, *cathepsin A* and *GIMAP7*, that fulfill both criteria and are therefore good candidates for being repeatedly driven by adaptive divergence between lake and river populations. The gene *cathepsin A* had the highest copy number differentiation among all gene CNVs and was present in more copies among the river ecotypes from the German populations than the German lake ecotypes, driving the overall differentiation signal. This gene encodes for a protein that plays an important role in processing endogenous bioactive peptides (Timur et al. 2016) and muscle metabolism (González-Prendes et al. 2017). Its isoforms CTS L and S have roles in MHC class II antigen presentation (Hsing and Rudensky 2005). More copies of the gene and therefore higher expression conceivably impact the immune response, whereas most of the gene region is depleted from variation despite the duplication in G1_R, suggesting background selection on the duplication. As river sticklebacks have lower MHC diversity compared with lake ecotypes (Eizaguirre et al. 2011), the higher copy number and expression of this gene potentially has a compensatory role and contributes to the defense against parasites specific to the river habitat. In contrast, lake ecotypes across population pairs were found to have higher copy numbers and higher expression of the gene *GIMAP7*, a GTPase that contains a domain AIG1-type G with immunity-associated functions (Krücken et al. 2004; Schwefel et al. 2010). The increase in *GIMAP7* copy number is associated with higher expression, possibly contributing to higher immune competence in lake individuals, as the parasite pressure is more intense in lake habitats (Scharsack et al. 2007; Eizaguirre et al. 2011). The matching habitat-specific expression patterns of *cathepsin A* and *GIMAP7* in immune tissues add to previous findings that CNVs are likely an important source of genetic variation that can help shape the host innate and adaptive immune response (Chain et al. 2014; Machado and Ottolini 2015). Our study on habitat-specific expression in immune tissues, which can potentially capture parasite-mediated

selection, has revealed two immune-related gene CNVs associated with expression differentiation, whereas other CNVs possibly contribute to habitat-specific adaptations in other tissues not sampled in our study. Previous investigation between marine and freshwater sticklebacks identified 24 gene CNVs consistent with parallel evolution, two of which were also found with DE between photoperiod treatments (*APOL2* and ENSGACG00000003408, Hirase et al. 2014). These two genes were also gene CNVs in our population system, with ENSGACG00000003408 also marginally differentiated between our lake and river populations ($V_{CT} = 0.124$, $FDR = 0.053$), but neither gene was expressed in our transcriptome data. In addition to Hirase et al. (2014), our findings of two gene eCNVs with significant V_{CT} and P_{CT} highlight an important role of gene CNVs in adaptation to new environments in sticklebacks.

eSNPs in *Cis* Also Contribute to Expression Variation

In addition to CNVs affecting gene expression, a total of 1,865 genes had SNPs in *cis*-regulatory regions identified as eSNPs putatively affecting gene expression. Though association tests between gene expression and SNPs do not necessarily reflect causal relationships, this result is consistent with previous studies that found abundant *cis*-eQTLs associated with expression divergence between stickleback ecotypes (Ishikawa et al. 2017; Pritchard et al. 2017; Kitano et al. 2019). Comparing marine and freshwater sticklebacks, Ishikawa et al. (2017) reported that about half of their local eQTLs resided in genomic regions of high divergence. Extending the comparison to multiple population pairs and between lake and river populations, we identified a gene differentiated between ecotypes both at the genetic level of an eSNP and in gene expression. The lake and river sticklebacks used in this study exhibit low parallel genomic divergence despite an isolation-by-adaptation signal (Feulner et al. 2015); genomic regions that most likely contribute to ecological divergence vary across different population pairs, suggesting the regulatory changes responsible for expression differentiation might also be population specific. As for *dhrs13a.3*, the homozygous T genotype of the eSNP 630 bp upstream of the TSS was associated with lower expression, and present in higher frequency in river populations where parasite abundance is generally much lower than in lakes (Scharsack et al. 2007; Eizaguirre et al. 2011). This allele is present in a detectable but low frequency (8.3%) in a source marine population (North Sea, Feulner et al. 2013) as well as in our lake populations, suggesting repeated increases in frequency in river habitats putatively due to habitat-specific adaptation.

Despite the abundance of genes with eSNPs, sequence differentiation of 5 kb upstream regions had an overall non-significant correlation with expression differentiation. This lack of genome-wide correlation between sequence-based differentiation in *cis*-regulatory regions and expression differentiation is consistent with other studies in whitefish,

flycatcher, and *Drosophila* (Renaut et al. 2012; Zhao et al. 2015; Uebbing et al. 2016), and can be at least partly explained by the narrow transcriptomic snapshot analyzed. Sequence differentiation might still impact expression differentiation in other tissues or at different developmental times not captured in our data. We also cannot exclude the impact that environmental plasticity might play in shaping expression differentiation. Although trans-regulatory changes may also contribute to expression divergence (e.g., Hart et al. 2018), we focused on *cis*-regulatory changes, which were found to account for large parts of parallel expression changes between marine and freshwater sticklebacks (Verta and Jones 2019). Taken together, our results highlight examples of SNPs and CNVs that contribute to expression differentiation linked to adaptive divergence.

Conclusion

By combining genome and transcriptome data from the same individuals across independently evolved population pairs, we describe generalities of the genetic basis of gene expression differentiation between lake and river sticklebacks. We revealed numerous changes of nucleotides in *cis*-regulatory elements that are associated with expression variation and prevalent dosage effects of CNVs on gene expression, providing variation that can foster rapid adaptation to different environments. We report one SNP in *cis* and two CNVs linked to gene expression differentiation that likely contribute to divergence between repeatedly evolved ecotypes. Our findings highlight both SNPs and CNVs as sources of genetic variation that promote repeated adaptation via *cis*-regulatory effect or dosage effect on gene expression.

Supplementary Material

Supplementary data are available at *Genome Biology and Evolution* online.

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Author Contributions

Y.H., F.J.J.C., and P.G.D.F. and E.B-B designed the analyses. Y.H. performed the analyses, and all authors contributed to discussions on research design and interpretation of the results. Y.H. drafted the article together with F.J.J.C. and P.G.D.F. All authors revised the article.

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