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Title: Population genomics of natural and experimental populations of guppies (*Poecilia reticulata*)

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Running Title: Population genomics of guppy populations

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Abstract

Convergent evolution represents one of the best lines of evidence for adaptation, but few cases of phenotypic convergence are understood at the genetic level. Guppies inhabiting the Northern Mountain Range of Trinidad provide a classic example of phenotypic convergent evolution, where adaptation to low or high predation environments has been found for a variety of traits. A major advantage of this system is the possibility of long-term experimental

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studies in nature, including translocation from high to low predation sites. We used genome scans of guppies from three natural high and low predation populations and from two experimentally established populations and their sources to examine whether phenotypic convergent evolution leaves footprints at the genome level. We used population genetic modeling approaches to reconstruct the demographic history and migration among sampled populations. Naturally colonized low predation populations had signatures of increased effective population size since colonization, while introduction populations had signatures of decreased effective population size. Only a small number of regions across the genome had signatures of selection in all natural populations. However, the two experimental populations shared many genomic regions under apparent selection, more than expected by chance. This overlap coupled with a population decrease since introduction provides evidence for convergent selection occurring in the two introduction populations. The lack of genetic convergence in the natural populations suggests that convergent evolution is lacking in these populations or that the effects of selection become difficult to detect after a long time period.

Key words: Natural selection; convergent evolution; genome-scan; *Poecilia reticulata*; long-term field experiments

Introduction

Convergent evolution, where similar adaptive phenotypes have independently arisen across similar environmental contrasts or clines, provides some of the strongest evidence for natural selection in the wild (Arendt & Reznick 2008). Classic examples include similar mouth morphology of cichlids in different lakes in East Africa, similar body size and limb morphology in Anolis lizards on different Caribbean islands, and latitudinal clines of temperature tolerance in *Drosophila melanogaster* ((Hoffmann *et al.* 2002), reviewed in (Schluter 2000)). With few exceptions, e.g. melanism in rock and beach mice to match dark backgrounds (Hoekstra 2006), we are ignorant about the genes that underlie such convergence because of the difficulties associated with identifying these genes by forward genetics. The situation is currently changing, as the facile generation of large DNA variation data sets supports genome-wide scans for candidate loci that are undergoing selection in similar environments. Genome-wide scans in multiple pairs of populations adapted to alternative environments have been used to identify convergent evolution in a variety of systems, e.g. threespine stickleback (*Gasterosteus aculeatus*, (Hohenlohe *et al.* 2010; Jones *et al.* 2012)), groundsels (*Senecio spp.*, (Roda *et al.* 2013)), lake whitefish (*Coregonus*

clupeiformis (Renaut *et al.* 2012)), sunflowers (*Helianthus spp.*(Renaut *et al.* 2014)), and stick insects (*Timema cristinae* (Soria-Carrasco *et al.* 2014)).

It remains to be seen how common genetic convergent evolution is and what factors allow or hinder this kind of repeated evolution. Many factors could influence the probability that the same genetic changes are selected for independently, including population size, gene flow, and similarity of starting genetic variation (Rosenblum *et al.* 2014). Rarely, however, do we know how or if these factors vary within a single system allowing researchers to test these predictions. Methods are now available to reconstruct the demographic history of populations at the whole genome level (e.g. (Gutenkunst *et al.* 2009; Li & Durbin 2011)), allowing researchers to formally compare populations with different demographic and population histories. Timescale is also important in detecting convergent genetic evolution because linkage disequilibrium between adaptive and non-adaptive loci is expected to decrease with time since the selective sweep (Barrett & Schluter 2008). This is especially important when only a small portion of the genome is surveyed, e.g. RAD-seq. Here, we present the first study to use a genome scan approach in the guppy, *Poecilia reticulata*, one of the prime models for microevolution and convergent evolution in vertebrates. The guppy system offers a unique resource, long-term experimental populations, which can provide direct insight into the importance of population history and timescale in detecting selection at the genetic level.

Guppies inhabiting the rivers in the Northern Range Mountains of Trinidad display convergent adaptations to life in different predator regimes. Rivers traversing the mountain range are frequently punctuated by steep waterfalls that exclude large piscivorous fishes from upper reaches of the rivers. Only a single species of small predatory fish is found above many such waterfalls. Because of the mountainous barriers between river systems, guppy populations in headwater streams of different rivers must have evolved independently from one another, and there appears to be repeated and independent adaptation to reduced predation in the tributaries of each river. The rivers show population-genetic separation (Barson *et al.* 2009; Suk & Neff 2009; Willing *et al.* 2010), presumably because of limited gene flow between them. The rivers draining the southern slopes of these mountains reside in two major drainages: the Caroni, which drains to the west, and the Oropuche, which drains to the east. Populations from the Caroni and Oropuche drainage have been shown to be diverging for some 600,000-1.2 million years (Carvalho *et al.* 1991; Fajen & Breden 1992;

Alexander *et al.* 2006) and are sufficiently different from one another to have been named different species (Schories *et al.* 2009).

Investigators have documented several instances of phenotypic convergent evolution in guppies from low versus high predation populations from different rivers. Males in the low predation environments of upper reaches tend to be more colourful than high predation populations in the lower reaches, bearing a larger number of spots and larger individual spots (Endler 1980). Populations in opposing predation regimes also differ in several life history traits; in low predation populations, both males and females are larger and mature later, and females give birth to larger and fewer young (Reznick 1982; Reznick *et al.* 1996). Finally, behaviour and swimming performance differ between predation regimes; in low predation populations, both sexes are less wary of predators (Kelley & Magurran 2003), shoal less (Seghers 1974; Seghers & Magurran 1995), and males rely more on courtship than sneaking when attempting to mate (Houde 1997). Guppies from high predation localities are less likely to be eaten by predators in experimental trials and have faster escape responses than guppies from low predation localities (O'Steen *et al.* 2002; Ghalambor *et al.* 2004). Many of these phenotypes have been shown to be highly heritable in laboratory studies, e.g. (Reznick 1982; Reznick & Bryga 1996). Therefore, the guppy is an ideal system to explore how phenotypic convergence is reflected at the genetic level.

A major advantage in the guppy system is the availability of long-term experimental populations, where researchers have transferred fish from high predation localities to streams, isolated by waterfalls, that were previously free of both guppies and most predators (e.g. Endler 1980). Long-term monitoring of these experimental populations showed that guppies have evolved towards phenotypes typical of low predation environments; they mature later, invest less in reproduction, and are more colourful than their ancestral populations from high predation localities downstream. In some cases, adaptation after experimental introduction has been shown to be quite rapid, occurring within four years (approximately eight generations) (Endler 1980; Reznick & Bryga 1987, 1996; Reznick *et al.* 1990; O'Steen *et al.* 2002). These experimental populations, therefore, provide researchers an opportunity to test predictions concerning genetic convergent evolution by comparing them to naturally colonized populations that differ in their population history and time since colonization.

Here we present the first genome scan to examine convergent evolution in the guppy, using a RAD-seq approach (Miller *et al.* 2007). This study builds on recent examples of convergent evolution at the genetic level by using the unique attributes found in the guppy system, i.e. largely independent replicates in pairs of high and low predation environments in separate rivers of known ancestral relationships and *in situ* experimental populations. By comparing putative signatures of selection among many river pairs we have the power to delineate drift from selection. Additionally, we estimate the demographic history and relatedness of each natural and experimental high and low predation pair to determine whether the likelihood of genetic convergent evolution is related to these factors.

Materials and Methods

Sampling

Fish were collected in February 2012 (Table 1, Figure 1). Fish were caught with butterfly nets and euthanized with MS222 and stored in 95% ethanol. This study was performed in accordance to the Max Planck Institute guidelines for treatment of animals. The three natural population comparisons were chosen because the differences in adaptive anti-predator traits have been well documented. Differences in amount of colour (Endler 1978, 1980), life history characteristics (Reznick & Endler 1982; Reznick *et al.* 1996), and behaviour (O'Steen *et al.* 2002) have been shown for each river pair. Also, the populations have shown differences in predator communities (Endler 1978; Reznick & Endler 1982).

Details about the Aripo and El Cedro introduction experiments can be found in Endler (1980) and Reznick and Bryga (1987), respectively. Briefly, approximately 200 individuals were introduced into the Aripo Introduction site in 1976 from a high predation Aripo source, and 100 individuals were introduced into the El Cedro Introduction site in 1981 from a high predation El Cedro source, both with equal amounts of males and females. Because female guppies can store sperm (as many as 9 sires per brood has been reported (Neff *et al.* 2008)) the estimated founding census population size is likely to be higher than the introduced number. Both introduction sites were checked multiple times before the introduction of high predation fish to determine that they were free of guppies and therefore migration from natural populations is unlikely. Guppies in the Aripo Introduction population were found to show an increase in the number and size of orange, black, and iridescent colour spots (Endler 1980), while guppies in the El Cedro Introduction population were found to show an increase in iridescent colouration (Kemp *et al.* 2009). Both populations showed differences in life

histories with males and females maturing later and at larger sizes than their source populations (Reznick & Bryga 1987; Reznick *et al.* 1990, 1997). In addition, the Aripo Introduction site fish evolved the production of larger offspring and smaller litter sizes early in life.

Sequencing

Genomic DNA was extracted from caudal peduncle tissue using the DNeasy Blood and Tissue Kit (Qiagen). We genotyped each individual using a RAD-seq method adapted from Poland and colleagues (Poland *et al.* 2012) (Miller *et al.* 2007; Baird *et al.* 2008). Briefly, double-digested genomic DNA (enzymes: *PstI* and *MseI*) was annealed with cut-site specific sequencing adaptors bearing individual barcodes. Barcoded samples were amplified via PCR (12 cycles, with barcoded samples multiplexed). Genomic libraries were size selected for 250-500 bp by gel extraction (Qiagen). Each individual had a unique barcode, using 159 of the original barcodes from Poland and colleagues (Poland *et al.* 2012). A total of five multiplexed libraries were sequenced, four with 159 barcoded samples, and one with 96 barcoded samples. Of the 153 individuals used in the final analysis, 68 samples were sequenced three times and 85 samples were sequenced twice in separate libraries to account for variation in barcoding efficiency and low coverage. The resulting fragments were sequenced on an Illumina HiSeq 2000, with 100 bp single-end reads.

Read Mapping and SNP Calling

As a reference sequence, we used the guppy genome produced from multiple paired-end libraries from a female from a high predation Guanapo population (located between the Arima and El Cedro rivers) that had been inbred for five generations in the lab. The genome assembly consists of 3,028 scaffolds and has an L50 of 5.27 Mb (NCBI: GCF_000633615.1 DDBJ/EMBL/GenBank: accession AZHG00000000).

We used SHORE v0.8 (Ossowski *et al.* 2008) to import the data, assign reads to barcodes, filter poor quality reads, and convert reads to fastq files. When assigning reads to a barcode, we allowed for 2 mismatches in the barcode and restriction enzyme sequence. Quality filtering was done with default cutoffs in SHORE (chastity violation 57, quality violation 3, quality cutoff read trimming of 5). Chastity filters were used to remove reads that resulted from overlapping clusters. Resulting high quality reads were mapped to the reference genome using STAMPY v1.0.21 (Lunter & Goodson 2011) with default parameters and assuming an

expected divergence of 4% from the reference. Reads were realigned around indels with IndelRealigner in GATK v1.6-9 (DePristo *et al.* 2011). Resulting sequences are available on NCBI's short read archive (SRA, BioProject ID PRJNA248075).

A total of 967 million reads were generated, of which 8.5% were discarded due to poor quality, and a further 9.5% could not be assigned to a barcode, leaving approximately 792 million high quality (HQ) reads. Individual samples with fewer than 100,000 reads were removed from subsequent analyses. After filtering, the average number of reads per individual was 3,177,282.1 (min = 400,733, max = 13,707,656). Overall, 465,063,290 million reads could be mapped to the reference, with a mean of 3,019,892 (96%) per individual (min = 168,921, and max = 13,175,144).

The average number of HQ reads per individual varied among populations (Table 1), with the high predation Aripo having the highest number, and the low predation Oropuche the lowest. This pattern was the same for mapped reads (Table 1). Even though there remains a high amount of variation in coverage per population the mapping performed equally well on all populations (Table 1). As a test of mapping we examined the relationship between scaffold size and number of reads for a random sample of 10 individuals. Number of mapped reads and scaffold size was highly correlated (Spearman's rho = 0.72, N = 1436, p < 0.0001, for scaffolds larger than 10 kb Spearman's rho = 0.90, N=675, p < 0.0001).

SNPs were called with GATK v1.6-9 (DePristo *et al.* 2011) using the following settings: stand_call_conf = 30 and stand_emit_conf = 30 (the minimum phred-scaled confidence threshold at which variants should be called and emitted, respectively). Resulting SNPs were then filtered using a minimum depth of 8, maximum depth of 200, and minimum genotype quality of 20 (corresponding to a false positive rate of 0.0004). Only biallelic SNPs were considered because the GATK algorithm works best with biallelic SNPs (DePristo *et al.* 2011). The number of multiallelic sites accounted for approximately 9% of SNPs called with high quality and coverage. Within populations, we removed possible paralogs by finding 100 bp regions with mean observed heterozygosity above 60% (mean of 93% percentile cutoff across all populations). Finally, at least 70% of individuals within each population needed to be genotyped at a given SNP and a minor allele frequency of above 0.01 within river to include it in subsequent analysis. Filtering and summary statistics were calculated using VCFTOOLS v0.1.9 (Danecek *et al.* 2011) or custom R (v3.0.0) or PERL (v5.12.3) scripts.

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Population Structure Analysis

We used STRUCTURE v2.3 (Pritchard *et al.* 2000) to estimate the population genetic clusters in our data. For this analysis, we considered only SNPs represented in all populations by at least 10 individuals and a minor allele frequency of 0.01 within river, and were separated by 10 kb to be ‘unlinked’, when SNPs were within 10kb of each other the most polymorphic SNP was chosen. We used a burn-in of 10,000, followed by another 10,000 MCMC steps. We assumed an admixture model and correlated allele frequencies with no prior information. We estimated lambda, with the maximum cluster (K) equal to 1 (lambda = 0.46). Then we estimated the most likely number of clusters by estimating the maximum likelihood for K = 2 - 9. We ran each cluster option five times. The most likely number of clusters was determined by finding the region where the likelihood reached a plateau and did not vary among replicate runs, e.g. (Evanno *et al.* 2005). The resulting graphical display was created in DISTRUCT v1.1 (Rosenberg 2004). A neighbour-joining unrooted tree based on Nei’s distance and a PCA was calculated to summarize population structure using the R package ADEGENET v1.3-9 on the same SNP set (Jombart 2008). Here missing data was replaced with the mean SNP frequency (Jombart 2008).

Demographic Analysis

We inferred within river population history using the joint allele frequency spectrum (AFS) diffusion based approach implemented in $\partial a \partial i$ (Gutenkunst *et al.* 2009). For each pairwise comparison we projected the dataset to 10 (averaging over all possible re-samplings of the larger sample size) for each population pair. This was to minimize missing data in the allele frequency matrix but maximize the number of segregating sites (Gutenkunst *et al.* 2009). The number of segregating sites ranged from 933.5 – 2429 (mean = 1635.6) depending on the pair analysed. We used the isolation migration model provided by $\partial a \partial i$, with the slight modification that founding populations were estimated independently, instead of estimating the size of population 1 as ‘s’ and the size of population 2 as ‘s-1’. This modification was necessary because low predation populations had extremely small sizes compared to high predation populations causing these parameters to be estimated at their boundaries of 1 and 0, which is problematic for exploring likelihood space. The parameters estimated were; founding effective population size of the high and low predation populations, the current effective population size of both populations, time at which the populations split, and migration between both populations. Times are given in $2N_{\text{ref}}$ generations, migration in $2N_{\text{ref}} M_{ij}$, and sizes are relative to N_{ref} .

We used a nested-model to determine whether migration between populations was likely, first fixing both migration estimates to 0, then migration of high to low predation to 0 (upstream migration is considered to be less likely), and then allowing migration between both sites. Significance of model fit was made with log-likelihood ratio tests. We used the log_{fmin} optimizer, bounds of parameters were at least 10 times higher than what was estimated, and the maxiter was set to 20. Optimization runs always began by first perturbing the starting parameters by 1 fold to ensure that independent runs converged on similar values, and grid sizes were much higher than sample sizes (grid sizes = 40,60,80). To estimate parameters we used conventional bootstrapping (fitting 100 datasets resampled over the loci), we report the median and 5 and 95% percentiles.

We decided that for two river analyses, the El Cedro and Arima, a 2-population model did not adequately reflect the population history because of admixture from outside rivers as determined in the STRUCTURE analysis. For these two cases we used a similar population model but with a 3rd ‘ghost’ or unsampled population to represent the entire meta-population of the Northern Mountain Range guppies. Here the parameter estimates were similar to the 2-population model but with a prior time of split between the high predation and ghost population, and migration between the ghost and high predation population.

To compare among models we converted time at split into years and effective population size of the low predation population into numbers of individuals. To do so, we obtained N_{ref} from theta ($\theta = 4N_{\text{ref}}\mu L_{\text{eff}}$, where μ is the mutation rate and L_{eff} is the effective length of the genomic region used). We used a mutation rate of 4.89×10^{-8} (estimated through genotyping parental and F1 SNPs, Künstner et al. unpubl. data), effective sequence length of 233,013 bp (corresponding to 3377 RAD-seq tags of 69 bp), and 2 generations per year (Reznick *et al.* 1997).

Outlier Analysis

We used an F_{ST} window outlier approach to detect regions undergoing divergent selection between paired high and low predation populations. Here each pairwise comparison within river was considered (9 populations, 5 pairwise comparisons). We also measured overall F_{ST} by comparing pooled groups (all natural low predation populations versus all high predation populations, e.g. (Axelsson *et al.* 2013)). We adapted the methods of Hohenlohe and colleagues (Hohenlohe *et al.* 2010) to detect outliers in F_{ST} , with the modification of

removing the kernel-smoothing algorithm. This was to avoid non-independence among windows, needed to use a χ^2 test used to test significance of overlap between comparisons. We used Weir and Cockerham's F_{ST} estimate (Weir & Cockerham 1984). Significance was determined by using a bootstrapping approach, where SNPs were bootstrapped 10,000 times and outliers were identified if their F_{ST} occurred within the upper 95% or 99% of bootstrapped values. We report analyses that employed a 150 kb window size. Different window sizes were assessed and we found that the divergent regions of the genome did not vary with these numbers within population comparison. We found that a window size of 150 kb limited the variation of windows assayed and SNP density among different river comparisons. Significant difference in allele frequency within river at individual SNPs was determined using a G-statistic test, with an FDR p-value of less than 0.05 (Hohenlohe *et al.* 2010).

We tested for independence among outlier lists using χ^2 tests. For the multidimensional χ^2 test we only evaluated windows that were covered in all populations. A Venn diagram was created using the program R package gplots. We next compared outlier lists to test specific predictions about convergent evolution. If similarity in standing genetic variation were related to the repeatability of evolution, we expected that the overlap among outlier lists would reflect the similarity across the entire genome and geographic proximity. If, however, population history or timescale are more important in detecting convergent evolution, we expected different degrees of overlap between introduction and natural populations. We tested for independence between any two outlier lists using a χ^2 test and using all assayed windows within the comparison. Methods concerning the distribution and genomic attributes of the outlier windows can be found in the supplementary materials. We also evaluated individual SNP F_{ST} , extracting the set of SNPs above the 95% F_{ST} quantile for each river pair. Then the overlap of these SNPs was evaluated using a χ^2 test as with the window analysis.

To investigate whether allele frequencies at any individual SNP loci correlated with predation environment we used the program BAYENV2, which accounts for overall population relationships (Günther & Coop 2013). We chose this approach over other F_{ST} outlier methods because it has been shown to be robust to false positives in populations with recent range expansions (Lotterhos & Whitlock 2014). The covariance matrix was estimated from the 'non-linked' SNP set used in the STRUCTURE analysis and was calculated by averaging over 10 random iterations sampled after an initial 50,000 iterations. The individual SNP matrices

were then correlated to a standardized environmental matrix (populations with high predation coded as 1 and those with low predation as 0) with 100,000 iterations. Outlier SNPs were identified as those with a Bayes Factor of above 20, rho of above 0.4, and minor allele frequency of above 0.01 within river pair.

Annotations

Annotations were obtained for regions of interest from gene model predictions from guppy reference genome (Künstner et al. unpubl. data, NCBI ascension: GCF_000633615.1). We evaluated enrichment of functional classes in specific contrasts using Gene Ontology classification. GO IDs for gene models in the reference database were derived using Blast2Go (Conesa & Götz 2008). We evaluated over-representation of biological process ontologies using the *topGo* package. To account for correlation in the GO graph topology, we used the *elim* algorithm, which eliminates genes from ancestor terms if they are significantly enriched in a child term. We report those with a p-value of less than 0.05 and enrichment of at least 3.

Results

We used RAD-seq for genome-wide genotyping of 9 populations found in the Northern Range Mountains of Trinidad (Figure 1, Table 1), with roughly equal numbers of males and females from each population, for a total of 153 individuals. After SNP filtering, we were able to evaluate 7,757 SNP loci from at least 70% of individuals in all populations (corresponding to 3,801 loci more than 10kb apart). As expected (Lynch 2007), there were more transitions (5,066) than transversions (2,691). For within-river pairwise population comparisons, we had information on an average of 45,535 SNPs (min = 13,285 in Oropuche and max = 83,066 in Aripo, Table S1).

Population Structure

High predation populations had a significantly higher expected heterozygosity (H_E) than their low predation counterparts (Table 1, paired t-tests across loci for all pairs of populations; all p-values < 0.0001). The largest difference in H_E was found between the Arima populations, likely due to admixture from other populations found in the high predation Arima (discussed below). The highest diversity was found in the high predation population found in the Oropuche ($H_E = 0.21$), and the lowest in the El Cedro introduction ($H_E = 0.05$). Populations within a river were more related than populations between rivers (Table S2). Populations

from different rivers and from the Caroni vs. Oropuche drainages were more distantly related to one another than those from within the same river (Table S2).

STRUCTURE analysis revealed that the most likely number of clusters was four (Figure 2a, Figure S1). In general, clusters were in agreement with geography. Three of the clusters are from the Caroni drainage and the other one from the Oropuche drainage. Within the Caroni drainage, the three clusters distinguish between the El Cedro, Arima, and Aripo River populations. High predation populations, found in the lower stretches of each river, showed more evidence of admixture than low predation populations, as evident by individuals with a probability of less than 90% of belonging to a single cluster (Table S3). The highest amount of admixture was found in the high predation Arima, where evidence of admixture is visible in all individuals from many different clusters. We also found four individuals from the El Cedro high predation population to have evidence of admixture with the Arima populations (Table S3). Similar groupings were obtained with the neighbour-joining tree (Figure 2b) and PCA (Figure S2).

Demographic Analysis

We estimated demographic parameters within the five river pairs using a joint Allele Frequency Spectrum (AFS) diffusion approach (Table 2a). First, we used a nested model approach to determine whether migration between high predation and low predation populations was likely (Table S4). No migration was likely between the populations in the Oropuche River, nor between the Aripo and El Cedro Introduction populations with their source sites. Migration downstream (low predation to high predation) was likely within the Arima River and migration both upstream and downstream was likely in the Aripo River. In the Aripo River, the estimate for low predation to high predation migration was five times higher than high predation to low predation migration.

We estimated population parameters by bootstrapping the data over all loci (Table 2a). Among the naturally colonized populations, the Aripo populations had the oldest divergence time, while the Oropuche populations had the youngest. Estimates of population split in the two introduced population were slightly shorter than the actual times of the events (estimate of 10-24 years ago for the Aripo Introduction [actual 36 years], and 8-30 years ago for El Cedro Introduction [actual 31 years]).

The ancestral and current effective population sizes were approximately the same in the Oropuche and Aripo high predation sites since they split with their respective low predation populations (Table 2a, specifically $HP_{Anc} < HP$ in 52%, 42% of the bootstrap replicates for Aripo, and Oropuche, respectively). The Arima and El Cedro high predation populations seemed to have increased in size using the two-population model ($HP_{Anc} < HP$ in 100%, 96% for Arima and El Cedro respectively). Because these two populations also showed evidence of substantial admixture in the STRUCTURE analysis, we tested a three-population model (Table 2b). While the estimated time of population split between high and low predation populations was not different from the two-population model, the estimated current population sizes of the high predation populations were smaller and were comparable to the estimates for the other high predation sites, from the Aripo and Oropuche in the two-population model (Table 2b).

All naturally colonized low predation populations have grown in size since the population split (Table 2, specifically $LP_{Anc} < LP$ 100% of the bootstrap replicates for all natural low predation populations). Conversely, the two introduction populations showed evidence of population decrease after their introduction (Table 2, $LP_{Anc} > LP$ 94% of replicates in El Cedro and 100% of replicates Aripo Introduction). The direction of population size change in the Arima low predation population and El Cedro Introduction in the two-population model was not different from the estimates derived from the three-population model.

Outlier Window Analysis

We used an F_{ST} outlier window approach to probe for evidence of selection between paired high and low predation populations within river (Table 3). A median of 3,901 windows of 150kb length was assayed for each river pair (Table 3). 2,126 windows were common to all five comparisons (three between natural high-low predation pairs and two between introduction and ancestral sites). These included 955 stretches of immediately adjacent windows on 157 scaffolds, with the greatest distance between windows being 1,950 kb.

Overall, 9 to 13% of windows were detected as outliers at a 95% threshold and 3 to 7% were detected as outliers at a 99% threshold. Among the comparisons between high-low predation sites, the Arima and El Cedro comparison had the highest percentage of outlier windows and the Oropuche comparison had the fewest. We observed similar patterns when we probed for F_{ST} outliers at individual SNPs (G-statistic; Table S1). The number of outlier windows was

not correlated with the total number of windows assayed, not considering the Oropuche comparison. Indeed, when we reran the analyses on just the subset of SNPs genotyped in all populations, the general patterns among river analyses remained the same (Table S5).

We refined our evaluation of genomic regions under selection by combining immediately adjacent outlier windows to produce outlier intervals (comparable to ‘peaks’ in (Hohenlohe *et al.* 2010)). The El Cedro and Arima comparisons had the highest number of outlier intervals and the Oropuche comparison had the fewest. The Arima comparison had many outlier windows, but these combined into relatively few, long intervals in comparison to the other rivers (Table 3). We found that both the Aripo and El Cedro Introduction analyses and the Arima analysis had non-random distributions of outlier windows on linkage groups (Table S6). Both the Arima and El Cedro comparisons had an enrichment of outlier windows on linkage group 12 (LG12) and LG15 (Table S6). The other enriched LGs were unique to each river. The enrichment found in El Cedro remained the same after removing the ‘admixed’ individuals found in the high predation El Cedro population. Outlier windows are unlikely to be caused by differences in genetic content; there were no differences in percentage of Ns between outlier and non-outlier windows and only the Aripo comparison had a significantly higher GC content in outlier windows and even here the effect size was quite small (GC content in non-outliers = 0.36, in outliers = 0.37, $p < .002$; Table S7). Indeed, outlier windows were significantly more diverged than non-outlier windows, using the absolute divergence measure Dxy (Cruickshank & Hahn 2014) (Table S7). To determine whether similar types of genes were within outlier windows in different river analyses, we analyzed Gene Ontology (GO) enrichment in Biological Processes (BP) annotations for gene models in outlier windows for each river analysis separately. No GO category was found to be significantly enriched in more than one river comparison (data not shown).

The difference between linkage disequilibrium (LD) within outlier windows compared to all windows varied between rivers. The low predation Arima, El Cedro Introduction, Aripo Introduction, and high predation El Cedro (only before removing admixed individuals) all showed significantly higher LD in outlier windows than non-outlier windows. No difference was found in the other populations (Table S7). Autocorrelation of F_{ST} outliers along LGs was found in four LGs in the El Cedro, two in Arima, and one in the Aripo (Table S8). Similar to the clustering analysis, we found an autocorrelation of outliers in El Cedro on LG12 and in

Arima on LG15. Linkage disequilibrium was not higher across LGs between outlier and non-outlier windows in any of the river comparisons (Table S7).

Overlap of Outlier Windows

We used a multi-dimensional χ^2 test to quantify the extent of overlap in outlier windows among rivers. When we compared all five river pairs, the outlier lists were significantly dependent on one another ($\chi^2 = 43.35$, $df = 1$, $p < 0.001$). However, no windows were found to overlap in all five comparisons, two overlapped in four river comparisons (expected 2), 33 overlapped in three comparisons (expected 28), 238 overlapped in two river comparisons (expected 208), and 690 were unique to the river (Figure 3). The two windows that overlapped in four of the river comparisons were on scaffold 56 (LG22) and scaffold 9 (LG19). Neither showed a significant difference in F_{ST} when all populations were pooled, but both showed a significant decrease in heterozygosity when LP populations were pooled (Table S9, Table S10). Nine gene models were found in these two windows (Table S11). There were a few genes that may underlie adaptive phenotypes related to growth and life history on scaffold 56. The *SLC2A2* gene is involved in sugar transport and homeostasis and may be related to sugar transport to yolk and developing embryos in zebrafish (Castillo *et al.* 2009) and *GHSR* is related to somatic growth and metabolism in fish (Jönsson 2013). A similar lack of overlap in all river comparisons was found when individual SNPs were queried, where no outlier SNPs overlapped in four or five river comparisons and only 4 SNPs overlapped in three comparisons (Table S12 and Table S13).

We next compared outlier lists to test whether different factors influenced the probability of genetic convergent evolution. The outlier lists did not overlap significantly when we considered only the naturally colonized river pairs (Arima, Aripo, and Oropuche: $\chi^2 = 1.48$, $df = 1$, $p = 0.22$). Three windows overlapped in all three rivers (expected 4), 90 overlapped in two rivers (expected 82), and the remaining 584 were unique to a single river pair. Of the three windows that overlapped in the natural river pairs, two of the windows were outliers in the El Cedro Introduction and one was an outlier in the Aripo Introduction. Two were also significant when all high and low predation populations were pooled and F_{ST} calculated (Table S9). None of these windows were found to show a consistent decrease in heterozygosity across either high or low predation populations but two showed a significant decrease in heterozygosity in the low predation populations when all low predation populations were pooled (Table S10). We analysed the gene content of the windows that

overlapped in all natural population comparisons. These regions contained 21 gene models (Table S11).

We next examined lists of outliers in each pairwise comparison to test whether some outlier lists overlapped more than others, using a χ^2 method (Table 4). None of the comparisons among natural high-low predation pairs overlapped with each other more than expected by chance (Table 4). The Aripo high-low predation and Aripo Introduction-high predation outlier lists overlapped significantly (Table 4), but the Aripo high predation population was common to each outlier list so these pairs are not independent of each other. The El Cedro Introduction-high predation outlier list had a significant amount of overlap with the Arima outlier list (Table 4). After removing the four individuals from the high predation El Cedro showing admixture with the Arima population, the overlap between Arima and El Cedro became non-significant. Finally, the two introduction experiment comparisons (El Cedro and Aripo Introductions) overlapped more than expected by chance (Table 4).

Overall the overlap between introduced populations occurred more than the naturally colonized populations. The 63 outlier windows shared in both experimental introduction population comparisons collapsed to 59 intervals distributed across 40 scaffolds and 16 LGs. Figure 4 shows the distribution of this overlap in the four linkage groups with the most overlap in introduced populations compared to the overlap found in natural populations. The outlier regions were not distributed randomly. There was a significant enrichment of shared outlier windows on LG8, with 8 windows (2 expected; $\chi^2 = 11.46$, $df = 1$, $p < 0.0007$, Figure 4) and LG23 with 6 windows (2 expected; $\chi^2 = 13.79$ $df = 1$, $p = 0.0002$, Figure 4). LG8 was also enriched with outliers in the El Cedro Introduction (Table S6). GC content did not differ between overlapping outlier windows and the rest of the genome (outliers = 0.36 ± 0.005 , all = 0.36 ± 0.0005 , $t = 0.36$, $df = 63.41$, $p = 0.72$), nor percentage of Ns (outliers = 0.085 ± 0.011 , all = 0.077 ± 0.001 , $t = 0.72$, $df = 63.64$, $p = 0.48$) and the absolute divergence was higher in outlier windows than non-outlier windows (IE outliers = 0.0015 ± 0.00007 , non-outliers = 0.0011 ± 0.000007 , $t = 5.5$, $df = 112263$, $p < 0.0001$, IA outliers = 0.0018 ± 0.0001 , IA non-outliers = 0.0010 ± 0.000009 , $t = 7.9$, $df = 77237$, $p < 0.0001$). Of the outlier windows 22 had low heterozygosity in pooled introduction populations, while only one had low heterozygosity in the pooled high predation populations (Table S14). We also annotated the gene models found in the overlap regions between the two introduction populations; 396 gene models mapped to this region. Of these gene models 168 could be annotated with biological

process in a GO analysis (Table S15). Many enriched developmental, reproduction, and gamete generation terms were found among them. We report the annotation for gene models in these overlapping windows (Table S16). There were a few interesting candidate genes, for example genes involved in growth and metabolism (CTSF is involved in fat metabolism (Russo *et al.* 2004) and CASQ1 in diabetes susceptibility (Das & Elbein 2006)), and in ovarian development and resource provisioning in zebrafish (inhibin-alpha (Wu *et al.* 2000)).

Individual SNP Analysis

We identified five SNPs that were significantly correlated with predation environment using BAYENV2. As with the F_{ST} outlier approach, no SNP showed allele frequency differences in all 5 pairwise comparisons (Table 5, Figure 5, Table S17).

Two of the outlier SNPs, “scaffold_50_4557695” and “scaffold_50_4575198”, were within ~17.5 kb of each other on LG15. These SNPs were fixed for the minor allele in the Arima low predation, Aripo Introduction, and El Cedro Introduction populations. There was no difference in allele frequency between the Aripo low and high predation populations for either SNP. A difference in allele frequency was seen for only one SNP in the Oropuche. These two SNPs were also in a region of strong autocorrelation and enrichment of outlier F_{ST} in the Arima and El Cedro paired comparisons on LG15 (Table S6, Table S8). One of these SNPs (scaffold_50_4557695) was also found to be above the 95% quantile of F_{ST} values in three river comparisons (Table S13).

The other three outlier SNPs were distributed across different scaffolds and LGs (Table 5). SNP ‘scaffold_5_11523021’ differed in allele frequency in the comparison of Aripo populations and was fixed for the minor allele in the Aripo and El Cedro Introductions. SNP ‘scaffold_148_1123620’ was fixed for the minor allele in the El Cedro Introduction, and allele frequencies differed between the Arima populations. However, no difference was seen in the Oropuche, the Aripo nor Aripo Introduction populations. Finally, SNP ‘scaffold_67_757135’ was fixed for the minor allele in the Arima low predation and El Cedro Introduction and showed a difference in allele frequencies between the Aripo populations but not in the Aripo Introduction or Oropuche populations. This SNP was also found to be above the 95% quantile of F_{ST} values in three river comparisons (Table S13). We annotated the closest gene models to these outlier SNPs, which provides candidates for investigating allelic differentiation between groups (Table 5). We also annotated genes within 50kb of these SNPs

(Table S18). Results for all major analyses were similar with low coverage individuals removed (Tables S19-S22).

Discussion

Here we investigated convergent evolution in natural and experimental guppy populations using genome-wide selection scans. While we detected perhaps a surprisingly high number of F_{ST} outlier windows differentiating each matched high and low predation population, we did not find evidence for convergent evolution in all five pairwise comparisons. Importantly, there were only two outlier windows that were common to at least four comparisons; this was equal to what was expected by chance. Similarly, using individual SNP analysis, we found loci differing in a maximum of four of the five high and low predation comparisons. This may seem surprising given the considerable evidence for phenotypic convergence among guppy populations, e.g. (Endler 1980; Reznick 1982; Kelley & Magurran 2003). We examined all pairwise combinations of overlapping windows to test specific hypotheses concerning the importance of similar standing genetic variation, timescale, and population history in detecting convergent evolution. The sharing of outlier windows in the natural populations was lower than what would be expected by chance. This was different for the El Cedro and Arima contrasts but the overlap between them was apparently caused by recent admixture from the Arima to the El Cedro. The two recent experimental introduction populations also shared a significant number of outliers. Our demographic analysis revealed evidence of effective population size decrease in the introductions. In contrast, all naturally colonized low predation populations showed an increase in effective population size since colonization (Table 2). While, this decline in population size could be caused by random mortality, our results suggests instead that convergent evolution has occurred in these introduced populations due to the high amount of overlap in outlier windows in these two populations.

There were other indications of recent, convergent directional selection in the two introductions. The overlap of outlier windows occurred over many LGs and the affected regions were enriched with gene annotations associated with reproduction and growth (Table S15 and S16). LD was higher in the introduced populations outlier windows than in non-outlier windows. Many of the windows also had low heterozygosity in both introduced populations, a pattern not seen in their source populations, as is expected if there was a recent selective sweep. Together with previous evidence of rapid phenotypic change, we now have

strong evidence that introduced populations underwent recent selective events. Reznick and colleagues (Reznick & Bryga 1987; Reznick *et al.* 1997) reported a change in male size and maturation age after only four years (approximately eight generations) and in both male and female size at maturity after seven years (approximately 14 generations) in the El Cedro Introduction population. Similarly, male and female age and size at maturity, as well as number of offspring in the first litter and offspring size had all significantly changed after only 11 years in the Aripo population (Reznick *et al.* 1990). While there is no published census population size for these introduction populations, from personal observation it is in the hundreds in the Aripo Introduction and the thousands in the El Cedro Introduction, i.e. much higher than the estimated current effective population sizes (Table 2). The population dynamics of transferring fish from a high predation population, where they have short generation times and high fecundity, into a low predation environment, where they have reduced risk of mortality, have been simulated (Reznick *et al.* 2004). The simulations suggested rapid population growth, with populations exceeding 1,000 individuals in less than one year. Rapid population growth can lead to fast adaptation because it is permissive of natural selection without local extinction. It is likely that this signal of genetic convergent evolution found in the introduced populations and not in natural populations is due to this shared demographic history or recent timescale of convergence.

Population structure

As reported before (Crispo *et al.* 2006; Barson *et al.* 2009; Willing *et al.* 2010), guppies from high predation populations had a higher amount of heterozygosity and more admixture from other rivers when compared to their low predation population counterparts. The high predation Arima population had a large and consistent amount of admixture from the Oropuche and El Cedro rivers. Each individual genotyped was assigned to both the Oropuche and El Cedro cluster with approximately 20% probability. The possible basis for this admixture is that the high predation Arima site is adjacent to many research facilities visited by guppy biologists. We know that fish from multiple populations were accidentally released in the Arima in September 2001 (F.H. Rodd personal communication). It is possible that other similar accidental introductions have occurred (Magurran 2005). The high predation El Cedro also showed evidence of admixture with the Arima, likely because these rivers lie in close proximity to one another and there are no major barriers between the two populations. The other rivers are strongly differentiated from one another, with only five individuals (only one from a low predation population) assigned to its river cluster with less than 90%

probability. Differences in heterozygosities within populations likely did not affect the F_{ST} outlier tests, since absolute divergence differs significantly between outlier windows and non-outlier windows (Table S7 (Cruickshank & Hahn 2014)). Our observed genetic differentiation among populations confirms inferences from allozyme and low-density genotyping studies (Carvalho *et al.* 1991; Alexander *et al.* 2006; Suk & Neff 2009; Willing *et al.* 2010).

Our estimates of effective population size and migration rates are high but not outside of the range previously reported for guppies. The overall patterns of relative population size are comparable to those observed by Barson *et al.* (2009). Low predation populations had smaller or similar effective population sizes as their high predation counterparts, except for low predation Aripo (this discrepancy was also reported in (Barson *et al.* 2009)). Migration was higher downstream (low to high predation) than upstream (high to low predation). To these data we add estimates of founding effective population sizes and colonization times. We estimated that some low predation populations were colonized by very few individuals and the largest ancestor population size was 112 individuals. It is well documented that guppies are strong colonizers; they have successfully invaded at least 69 countries outside of their natural range, where they were initially introduced for mosquito control (Deacon *et al.* 2011). Also, Deacon and colleagues (Deacon *et al.* 2011) have shown that a single pregnant female could establish a population of up to 60 individuals and that the population remained viable for 2 years (the duration of the experiment). The evidence for inbreeding depression and inbreeding avoidance in natural guppy populations is mixed (Pitcher *et al.* 2008; Johnson *et al.* 2010).

Detecting convergent evolution

One possible explanation for the differences in the signatures of selection between natural and introduced populations is that they were caused by differences in the amount of initial genetic variation in the founders. Our results do not fully support this alternative. The estimated ancestral effective population size in the introduced populations was not larger than in all natural low predation populations. However, a limitation of these genetic data is that they alone cannot distinguish multiple natural colonization events occurring in quick succession from one larger colonization event. Furthermore, high migration and reduced divergence affects the allele frequency spectrum similarly, both showing high correlation of allele frequencies between the populations. Shared low-frequency alleles will distinguish the

two models but these may have been filtered or missed in our data (Gutenkunst *et al.* 2009). The Aripo River analysis yielded the longest divergence time, smallest low predation ancestral population, and was also the only river to show evidence for migration upstream. The Oropuche population analysis yielded the youngest divergence time and no migration was found to be likely between the populations. It is also possible that low predation populations are older than estimated here, but that they have gone through repeated genetic bottlenecks due to flooding and multiple colonization (van Oosterhout *et al.* 2007). Also, generation time (used to estimate time in years) itself is under selection and may change over time (Reznick & Endler 1982). Therefore our migration and time of divergence estimates should be interpreted with caution. Another possible reason for the strong overlap in introduced populations is that introduced populations are more similar in selective forces other than predation compared to naturally colonized populations. Future studies measuring pathogen pressures and diet of these populations would help clarify the interactions and importance of different selective pressures in this system.

The number F_{ST} outlier windows, up to 13% at the 95% confidence threshold, within individual rivers may appear surprisingly high. Outlier windows occurring in a single river pair should be interpreted with caution because they cannot be distinguished from the effects of drift and bottlenecks. The possibility of drift over-shadowing selection is likely to be common in the guppy system because of the limited number of founders and limited migration into the upper reaches of rivers (van Oosterhout *et al.* 2006; Crispo *et al.* 2006). Additionally, populations in fractal landscapes like rivers are more likely to have false positive F_{ST} outliers due to correlated ancestry and inflated variance in F_{ST} measures (Fourcade *et al.* 2013). These factors could also be responsible for the lack of genetic convergent evolution found in other similar study systems (e.g. Atlantic salmon (*Salmo salar*) (Perrier *et al.* 2013) and Rainbow trout (*Oncorhynchus mykiss*) (Hecht *et al.* 2013)). However, these factors are unlikely to generate false-positives in the same genomic regions in multiple comparisons because they would affect randomly distributed regions of the genome. Overlap in outlier regions could be caused by other neutral factors such as reduced recombination around centromeres and isochores or differences in mutation rate. Indeed, Roesti *et al.* (Roesti *et al.* 2012) found a large bias of F_{ST} outliers in non-recombining centromeres using a similar approach in lake-stream stickleback. However, we have no evidence that outlier regions had different genetic properties than the rest of the genome

(Table S7). Therefore, overlapping regions showing evidence of genetic divergence in multiple populations are more likely signatures of selection.

Our analyses revealed little evidence of genetic convergence in guppy evolution. Only two windows were outliers in at least four river pairs and only three outliers were found in all natural paired comparisons, not more than expected by chance alone. Similarly an analysis detecting correlation between environment and allele frequencies, after accounting for overall covariance among populations, showed a difference in allele frequency in a maximum of four river pairs. We may have missed the genetic regions selected in all populations because we used a RAD-seq approach, which sequences only a small fraction of the genome. Increased recombination between selected loci and neutral loci would further decrease the chance of detecting selected regions. LD was significantly higher in outlier windows than non-outlier windows in the Arima low predation and the two introduction sites. There were no such differences in the natural Oropuche and Aripo populations. The Arima and introduction populations also had the strongest evidence for convergent evolution in the form of shared F_{ST} outlier windows. The differences in the presence of LD are thus suggestive of recombination having erased the evidence of selection in the paired comparisons of LP and HP populations in natural rivers.

Another intriguing possibility is that phenotypic convergent evolution was mediated by different genes in different rivers. A recent review of genome scans to detect outlier loci revealed that only a small fraction of loci (1 - 12%) were found to be outliers in more than one divergent pair within the same system, suggesting that convergent phenotypic evolution cannot be easily equated with convergent genetic evolution (Nosil *et al.* 2009). Similarly, a meta analysis has shown that the likelihood of the same gene undergoing selection in independent systems decreases with increasing age of ancestral node (Conte *et al.* 2012). Evolution in which similar phenotypes are the product of different genes has been found in a variety of systems. For example, adaptive melanism in different populations of rock pocket mice has apparently distinct genetic causes (Nachman *et al.* 2003). Likewise, different genes could be responsible for similar adaptive changes in the guppy. Many adaptive traits in this system, like body size, body shape, behaviour, swimming performance, colouration, and life histories, are likely to be polygenic. Additionally, if different enhancers affecting the same gene were selected, we would not be able to detect a signal of convergent evolution. While phenotypic differences between the populations may be examples of adaptive plasticity

(Ghalambor 2007), many of these phenotypic differences remain heritable in common lab conditions (Reznick 1982; Reznick & Bryga 1996), indicating that they are caused by genetic differences. Differences in ecological variables have been shown to have a large influence on adaptive phenotypes in the guppy. For example, substrate size relates to male colour spot size (Endler 1980) and canopy openness influences primary productivity and food availability, which in turn affect life history traits and male colour (Endler 1995; Grether *et al.* 2001; Reznick *et al.* 2001). Clearly, convergent evolution at the phenotypic level in the guppy is nuanced in a way that could reduce the odds of correlated genetic convergence.

Sticklebacks adapted to freshwater and marine environments present an iconic example of aligned phenotypic and genetic convergent evolution, where the derived allele at the *Eda* locus, which is associated with reduced lateral plates (Colosimo *et al.* 2005), is found segregating at low frequencies in marine populations and has been fixed repeatedly in freshwater populations (Hohenlohe *et al.* 2010; Jones *et al.* 2012). Why would the results for sticklebacks be so much more deterministic than those for guppies? One possibility is the nature of the founding populations. We know of no estimate for founding population sizes for lake stickleback from marine populations but it is most likely large, to account for high amounts of standing variation (Hohenlohe *et al.* 2010). Also, continuous migration between the founding freshwater and marine populations is likely (Bell & Foster 1994). Our estimates of population histories in guppies paint a very different picture. We inferred that the natural low predation populations were founded with relatively few individuals, approximately 10 – 100, and even the introduced populations had low ancestral effective population sizes. We also have strong evidence that migration upstream is very limited. Therefore the starting genetic variation in low predation populations was most likely low, decreasing the likelihood of selection acting on similar standing genetic variation in natural populations (Rosenblum *et al.* 2014). Our results also imply that the founders of guppy populations in different low predation environments may have been genetically distinct at the outset. In contrast, the founders of the freshwater stickleback populations in the Pacific Northwest were all derived from a large, panmictic marine population. Therefore the guppy represents a system with equally striking phenotypic convergent evolution as the stickleback system but with a very different population history, one that would decrease the likelihood of genetic convergent evolution (Rosenblum *et al.* 2014).

Limitations of RAD-seq analyses

Several studies have examined the biases and limitations of reduced-complexity genotyping, such as RAD-seq in selection studies. Due to differences in restriction enzyme recognition sites, RAD-seq may underestimate diversity through allele drop out, although F_{ST} appears to be fairly robust to this problem (Arnold *et al.* 2013). It has been suggested including low read coverage worsens allele drop out effects (Gautier *et al.* 2012). We therefore used minimum coverage thresholds. Restricting the analysis to loci with complete information for all individuals may be helpful as well, but is rarely sensible, as this would eliminate the majority of loci when many individuals are studied. As a compromise, we only considered loci with information from at least 70% of individuals from each population. We found roughly equal heterozygosity for four populations and even higher heterozygosity for two populations than what had been found with conventional genotyping methods (Willing *et al.* 2010), the opposite of what would be expected with allele drop out. Another concern with RAD-seq is ascertainment bias, as genotypes closer to the reference genome are more easily determined (Sousa & Hey 2013). Again reassuringly, we were able to map a similar fraction of reads for all populations, arguing against ascertainment bias. While we found fewer SNPs in the most distantly related populations from the reference (the reference genome was created with a fish from the Guanapo River found in the Caroni watershed between the Arima and El Cedro rivers), these also had the lowest sequence coverage.

Conclusions

Results from high-density genotyping of paired high/low-predation natural and experimental populations have revealed several notable patterns of selection. First, using population genetic modeling approaches to reconstruct the demographic history and migration among sampled populations, we found that naturally colonized low predation populations experienced population growth since colonization, while introduction populations shrank. Second, we detected only very few regions across the genome with signatures of selection common to all populations. In contrast, the two experimental populations shared many such regions. Together our results suggest convergent genetic evolution in introduced populations but not natural populations. Our results emphasize that detecting the signature of convergent evolution at the genome level requires sampling at the appropriate timescale or population history; otherwise neutral processes or population-specific selective forces may obscure or decrease the likelihood of convergent genetic evolution.

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Data Accessibility

All demultiplexed read data used for genotyping submitted to NCBI SRA: BioProject ID PRJNA248075. The reference genome used for mapping and annotation is available on NCBI assembly accession: GCF_000633615.1.

A full SNP dataset is available on dryad. Dryad doi:10.5061/dryad.n35j4

Author Contributions

BAF, DNR, CD, DW designed the research. BAF and DNR collected the samples. BAF did the molecular work. BAF and AK conducted the bioinformatics analysis. BAF analysed the data. BAF wrote the manuscript with input from the other authors.

Figure 1: Sampling sites in Northern Trinidad.

LAM = Low predation Arima, HAM= High predation Arima, HE= high predation El Cedro, IE= Introduction El Cedro, LA = Low predation Aripo, HA = High predation Aripo, IA = Introduction Aripo, LO = Low predation Oropuche, HO = High predation Oropuche. Dashed circles indicate drainages (red = Caroni drainage, blue = Oropuche drainage).

Figure 2: Population structure among sampled populations.

(a) Structure analysis with $K=4$ clusters. Each line represents one individual and the colour distinguishes clusters. The population is indicated on the x-axis. (b) Neighbour-joining tree based on Nei's distance between sampled populations. Population names can be found in Table 1.

Figure 3: Overlap in outlier windows for all river pairs

A Venn diagram indicating the degree of overlap in F_{ST} outlier windows found in all 5 river pairs using the set of windows that were assayed in all 5 rivers.

Figure 4: Overlap in outlier windows in introduced populations compared to naturally colonized populations

F_{ST} for windows for each river pair, Arima (pink), Aripo (orange), Oropuche (blue), Aripo Introduction (red), and El Cedro Introduction (yellow). Outliers within river pair is indicated by a grey triangle, while outliers shared by any two naturally colonized pair is indicated by blue triangle, and outliers shared by both introduced populations by a red triangle. Shown are the four linkage groups with the most overlap between introduced populations.

Figure 5: Loci with strong correlation between allele frequency and predation regime.

Shown are the frequency of major allele (in black) and minor allele (in grey) for each population grouped by river pair for each outlier SNP.

Table 1: Summary of population samples.

For each population we report the river, population ID (H=high predation, L= low predation, I= introduction to low predation habitat), N = Number of individuals (after filtering for low coverage individuals), grid reference, (UTM 20 P), average number of High Quality Reads (HQ) per individual, average number of mapped reads per individual, and expected heterozygosity (H_E).

River	Population	N	Grid Reference	Avg. HQ reads	Avg. mapped reads	H_E
Arima	HAM	14	PS 82000 686800	2,288,761	2,211,423	0.16
	LAM	19	PS 1181500 687300	2,376,598	2,366,832	0.08
El Cedro	HE	16	PS 1178900 689500	4,105,864	3,910,696	0.10
	IE	20	PS 1179900 689500	3,495,089	3,328,987	0.05
Aripo	HA	19	PS 1179500 6937000	4,782,516	4,564,046	0.14
	LA	19	PS 1181900 692 900	4,130,695	3,936,416	0.11
	IA	13	PS 1179900 693700	2,169,641	2,067,852	0.09
Oropuche	HO	18	PS 1180900 697000	3,079,306	2,959,967	0.21
	LO	15	QS 1178800 704100	1,128,363	1,082,971	0.18

Table 2: Demographic analysis of population river pairs.

- (a) Parameter estimates (medians and 95% CI) for a 2-population isolation with migration model. S is the number of segregating sites, HP is the high predation, LP is the low predation, and ‘anc’ the ancestral population at time of split, T is the time since population split ($2N_{ref}$ generations), migration between populations (M_{HP-LP} M_{LP-HP}) are in units of $2N_{ref}m_{ij}$, $\Theta = 4N_{ref}\mu$, and ll is the long-likelihood of the model. To make comparisons among river pairs LP_{ANC} and LP were translated to number of individuals ($LP_{ANC} N_E$ and $LP N_E$) and T to years (T_{years}) based on θ . Migration estimates were fixed at zero depending on the nested likelihood models (see supplement). Parameter medians and confidence intervals were estimated by conventional bootstraps. (b) Similar population parameters were estimated using a 3-population isolation with migration model. The third population (G) represents a ghost or unsampled population. T is time is the split between the HP and G population, while T_2 is the time of split of the HP and LP population.

(a) 2-Population Model

	Arima	Aripo	Oropuche	Aripo Introduction	El Cedro Introduction
S	1732.5	1629.5	2429.0	1453.4	933.5
HP_{Anc}	2.80 1.35 – 4.49	1.54 0.83 – 2.85	0.11 0.05 – 0.18	1.25 0.80 – 2.79	1.50 0.77 – 2.76
LP_{Anc}	0.0099 0.0062 – 0.017	0.0017 0.0007 – 0.0037	0.0045 0.0028 – 0.0072	0.010 0.005 – 0.018	0.0084 0.0044 – 0.014
$LP_{Anc} N_E$	112.03 70.98 – 181.93	16.06 6.67 – 35.89	74.19 45.39 – 117.42	103.29 53.50 -178.83	55.66 29.48 – 93.68
HP	70.53 35.07 – 93.85	1.48 0.81 – 2.61	0.12 0.07 – 0.19	1.54 0.83 – 3.04	4.24 2.26 – 8.62
LP	0.20 0.10 – 0.34	2.92 1.75 – 5.38	0.51 0.26 – 0.96	0.0040 0.0022 – 0.0080	0.0026 0.0013 – 0.0040
$LP N_E$	2207.82 1199.95 – 3838.86	27917.53 15946.72 - 54020.57	8408.33 4292.79 – 15820.25	39.69 22.31 – 80.16	17.41 8.33 – 26.40
T	0.039 0.027 – 0.051	0.15 0.10 – 0.24	0.0036 0.0024 – 0.0048	0.0021 0.0013 – 0.0030	0.0026 0.0017 – 0.0037
T_{years}	439.56 314.51 -562.02	1461.88 947.45 – 2250.28	58.27 40.46 – 78.70	21.56 21.84 – 69.41	16.95 11.58 – 24.19
M_{HP-LP}	0	1.45 0.82 – 2.67	0	0	0
M_{LP-HP}	7.63 5.26 – 12.32	8.22 5.83 – 11.75	0	0	0
$Theta$	512.0 499.6 – 523.3	431.8 395.7 – 468.4	746.5 743.0 – 753.5	456.8 455.9 – 457.6	301.8 301.0 – 302.3
Ll	-219.6 -225.4 – -218.0	-160.0 -167.6 – -156.7	-159.4 -161.0 – -158.5	-168.5 -168.9 – -167.8	-181.9 -182.4 – -181.7

(b) 3-Population Model

	Arima	El Cedro Introduction
HP_{Anc}	2.78 1.56 – 5.57	1.56 0.82 – 2.87
G_{Anc}	0.95 0.55 – 1.82	1.03 0.54 – 1.81
LP_{Anc}	0.02 0.01 – 0.04	0.008 0.005 – 0.017
$LP_{Anc} N_E$	140.94 68.66 – 306.35	64.49 29.03 – 114.09
HP	0.62 0.31 – 1.02	1.02 0.54 – 1.79
G	1.54 0.79 – 2.63	0.96 0.53 – 1.86
LP	0.07 0.04 – 0.11	0.0039 0.0021 – 0.0061
$LP N_E$	504.22 280.38 – 778.15	24.67 13.21 – 50.14

<i>T</i>	1.91 1.10 – 3.57	0.58 0.29 – 1.30
<i>T2</i>	0.05 0.03 – 0.08	0.0024 0.0013 – 0.0037
<i>T2years</i>	379.95 200.04 – 578.87	15.91 8.04 – 30.30
<i>M_{HP-G}</i>	10.41 6.12 – 17.56	1.06 0.54 – 1.84
<i>M_{G-HP}</i>	10.41 5.90 – 18.46	4.33 2.12 – 7.68
<i>M_{HP-LP}</i>	0	0
<i>M_{LP-HP}</i>	11.06 6.42 – 17.97	0
<i>Theta</i>	327.0 244.2 – 421.9	299 229 – 463
<i>Ll</i>	-178.6 -208.4 – -162.2	-182 -189 – -180

Table 3: Summary of F_{ST} outlier analysis.

Using a window size of 150 kb, F_{ST} was calculated between paired high and low predation populations within a river. Outlier windows were determined with a bootstrapping approach for F_{ST} at both the 95 and 99% confidence level. Outlier windows were then collapsed into intervals. We reran the same analysis on El Cedro with the four ‘admixed’ individuals removed (El Cedro NoAd).

River	mean SNP density (per window)	mean F_{ST} (min-max)	Total # of windows	# outliers windows 95% (%)	# outlier windows 99% (%)	# of outlier intervals	mean outlier interval length kb (min-max)
Arima	9.74	0.11 (0-0.92)	3990	480 (12%)	230 (6%)	329	217 (150 – 1350)
Aripo	15.73	0.05 (0-0.5)	4114	475 (11%)	243 (6%)	362	196 (150 – 750)
Oropuche	3.88	0.06 (0 – 0.53)	2354	222 (9%)	74 (3%)	202	164 (150 – 450)
Aripo Introduction	6.80	0.06 (0 – 0.83)	3812	348 (9%)	154 (4%)	290	179 (150 – 750)
El Cedro Introduction	10.57	0.10 (0-0.59)	3980	503 (12%)	274 (7%)	364	206 (150 – 900)
El Cedro Introduction NoAd	7.91	0.11 (0-0.70)	3821	492 (13%)	273 (7%)	350	210 (150 – 1350)

Table 4: Tests of independence of outlier windows between rivers.

Shown above the diagonal are individual χ^2 values with associated significance. Below the diagonal is the number of overlapping outlier windows at a 95% confidence level. The El Cedro set without admixed individuals is given in brackets for the El Cedro and Arima comparisons.

	Arima	Aripo	Oropuche	Aripo Introduction	El Cedro Introduction
Arima		0.58	0.05	3.6	9.97*(0.96)
Aripo	62		0.61	15.3**	0.43
Oropuche	30	32		2.6	0.06
Aripo Introduction	53	64	15		9.98*
El Cedro Introduction	82 (67)	64	30	63	

** p-value < 0.001

* p-value < 0.01

Table 5: Correlation of individual loci to predation regime

Loci with a strong correlation between allele frequency matrix and environmental matrix after correcting for overall population covariance using a Bayesian approach. For each SNP we report the ID (which is its scaffold and position concatenated), the linkage group (LG), Bayes factor (BF), Spearman's rho, and annotation of the closest gene.

ID	LG	BF	Rho	closest Gene
scaffold_50_4557695	15	568.66	0.60	B-cadherin-like
scaffold_50_4575198	15	99.72	0.50	B-cadherin-like
scaffold_5_11523021	16	32.03	0.41	pleckstrin homology domain containing, family G (with RhoGef domain) member*
scaffold_148_1123620	11	27.22	0.44	MHC class I related gene
scaffold_67_757135	21	24.27	0.43	protein-like uridine-cytidine kinase-like 1

* outlier SNP was in the gene model







