Population genomics reveals a possible history of backcrossing and recombination in the gynogenetic fish *Poecilia formosa*

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Unisexual sperm-dependent vertebrates are of hybrid origins, rare, and predicted to be short-lived as a result of several challenges arising from their mode of reproduction. In particular, because of a lack of recombination, clonal species are predicted to have a low potential to respond to natural selection. However, many unisexual sperm-dependent species persist, and assessing the genetic diversity present in these species is fundamental to understanding how they avoid extinction. We used population genomic methods to assess genotypic variation within the unisexual fish *Poecilia formosa*. Measures of admixture and population differentiation, as well as clustering analyses, indicate that the genomes of individuals of *P. formosa* are admixed and intermediate between *Poecilia latipinna* and *Poecilia mexicana*, consistent with the hypothesis of their hybrid origins. Bayesian genomic cline analyses indicate that about 12% of sampled loci exhibit patterns consistent with inheritance from only one parent. The estimation of observed heterozygosity clearly suggests that *P. formosa* is not comprised of direct descendants of a single nonrecombining asexual F1 hybrid individual. Additionally, the estimation of observed heterozygosity provides support for the hypothesis that the history of this unisexual species has included backcrossing with the parent species before the onset of gynogenesis. We also document high levels of variation among asexual individuals, which is attributable to recombination (historical or ongoing) and the accumulation of mutations. The high genetic variation suggests that this unisexual vertebrate has more potential to respond to natural selection than if they were frozen F1 hybrids.

Asexuality is common in many phyla (reviewed in ref. 7), but it is relatively rare in vertebrates (1). All known unisexual (all-female) vertebrates are products of hybridization events between sexually reproducing species (ref. 9 and references therein), constitute only 0.1% of extant vertebrate species (1, 9). One type of asexual reproduction found in unisexual vertebrates is gynogenesis, where females must mate with males of a closely related species (but refer to ref. 10 for exceptions), but the nonrecombinant embryos do not inherit any genetic information from the sperm donor (9). Because gynogens require sperm to initiate development of offspring, but no paternal genes are expressed, they are considered “sexual parasites” (11).

The maintenance of a gynogenetic species is paradoxical because gynogens face the costs of both sexual and asexual reproduction: the cost of finding a mate, exposure to diseases, accumulation of deleterious mutations, and lack of genetic recombination to facilitate adaptation. In addition, because male sperm donors do not gain a fitness advantage from mating with gynogens, selection should favor males that avoid mating with them.

Given the extensive and diverse list of challenges faced by gynogenetic species, they are predicted to be short-lived with a limited potential to respond to natural selection. Nevertheless, gynogenetic species persist, and some have origins in the distant past (12, 13). This suggests that gynogenetic species might be able to avoid or ameliorate some of the costs associated with their reproductive mode. One question that arises is how much genetic variation persists in gynogenetic species?

The Amazon molly (*Poecilia formosa*) is an excellent system to explore this question. *P. formosa* is the first vertebrate recognized as asexual (11) and is a gynogenetic species that uses *Poecilia mexicana* (Atlantic molly), *Poecilia latipinna* (sailfin molly), and *Poecilia latipunctata* (Tamesi molly) as sexual hosts (14). Like every other known unisexual vertebrate, *P. formosa* is thought to be a hybrid lineage (9, 13, 15–17). *P. mexicana* is recognized to be the maternal species of *P. formosa* (13, 15–17), whereas *P. latipinna* (or an extinct ancestor of *P. latipinna*) is the putative paternal species (17). *P. formosa* lives in sympathy with at least one of the two parent species throughout its range from the Tampico region in Mexico to the southeastern United States (Fig. 1A). Although recent studies suggest that *P. formosa* is a species that consists of “frozen” F1 hybrid clones (i.e., individuals with...
ancestry from *P. mexicana* and *P. latipinna* at all loci) (15–19), this result is based on limited genetic data. Additionally, it is still not clear whether *P. formosa* is the product of a single or multiple hybridization events, although a recent investigation supports the hypothesis of a single event possibly giving rise to several clonal lineages (19).

The overall objective of this research was to examine genotypic variation in *P. formosa* at a genomic scale. We generated thousands of DNA sequence markers to address three main questions. (i) Is *P. formosa* the product of hybridization between *P. latipinna* and *P. mexicana*? This is the conclusion of several previous studies (11, 13, 15–19), and, here, we attempt to confirm this result with a genome-wide survey of *P. formosa* and its putative parent species. (ii) Is *P. formosa* composed of frozen F1 hybrid clones, or is there evidence of a more complicated history of nonclonal reproduction or recombination? Previous genetic analyses (13, 15–19), and the fact that no laboratory has been able to synthetize *P. formosa* from artificial hybridization experiments (15, 19–21), suggest that the genome of this species is more complicated than that of a simple F1 hybrid. If *P. formosa* is not composed of clones of F1 hybrid lineages, then, what is the genetic contribution of each parent species? (iii) How much genotypic diversity exists within *P. formosa* and is this genotypic diversity consistent with *P. formosa* having evolved from a single or multiple independently formed hybrid individuals? Quantifying genotypic variation will provide some estimate of this species potential to respond to selection.

Results and Discussion

We used a next-generation sequencing population genetics approach to collect information on variation from across the genomes of *P. formosa*, *P. latipinna*, and *P. mexicana* and to understand the genomic composition of the gynogenetic species. The methodology used herein allowed us to obtain genotype information from thousands of variable sites dispersed across the entire genomes of these fishes, with which we achieved a higher level of resolution of patterns of genetic variation than previous studies. We generated DNA-sequence data using the Illumina GAII platform for 192 fish: 41 *P. formosa* (5 localities where *P. formosa* is sympatric with *P. mexicana* and 6 localities where it is sympatric with *P. latipinna*), 82 *P. latipinna* (from 22 localities across Louisiana, Texas, and Mexico), and 69 *P. mexicana* (from 13 localities across Mexico and Honduras; Fig. 1A). We identified 26,313 single-nucleotide polymorphism (SNP) markers that were used for the analyses.

Our first goal was to confirm the hybrid origin of *P. formosa* and determine the genomic contribution from its parent species. Our results are in agreement with the findings of several previous studies: *P. formosa* is a hybrid between *P. mexicana* and *P. latipinna* (11, 14–19). A principal components analysis (PCA) of the genotype estimates of each individual at each locus (Fig. 1B–D), an admixture analysis performed in STRUCTURE with K = 2 (forcing individuals to be assigned to one of only two clusters) (Fig. 2A), calculation of genome-average pairwise GST (Fig. 3), and the hybrid index for each *P. formosa* (Fig. 3) all suggest that gynogenetic individuals have a genome that is intermediate between *P. latipinna* and *P. mexicana*. Principal component axes 1 (PC1) and 2 (PC2) collectively explained 63% of the genetic variation (Fig. 1B). PC1 appears to divide the three species into three distinct clusters, with *P. formosa* in an intermediate position between the parent species, consistent with the hypothesis of a hybrid origin for *P. formosa* (Fig. 1B). PC2 separates the asexual species from the sexual species and illustrates the variation in genotypes among individuals (Fig. 1B). Interestingly, the genotypic
variation within *P. formosa* was high: individuals of *P. formosa* did not all cluster together but were differentiated along the second principal component just as much as the sexual individuals. PC3 explained 5.7% of the variation and separated the populations of *P. mexicana* into three groups, which corresponded to three geographic regions (north, central, and south) (Fig. 1C). PC4 explained 4.0% of the variation and divided *P. latipinna* into two geographic regions: north plus central (the central group is comprised of populations in central Texas that were introduced from the north group, including population in Florida and the Gulf Coast of the United States) and south (Fig. 1D).

Estimates of pairwise *G_{ST}* also confirmed the genomic intermediacy of *P. formosa* (Fig. S1 and Table 1). Differentiation between *P. latipinna* and *P. mexicana* was approximately twice that observed between *P. formosa* and either parent species (Table 1). Within-species (among-population) differentiation was an order of magnitude lower, and lowest in *P. formosa*, although distinctly nonzero. Estimation of hybrid indices for the asexual individuals was also consistent with the hypothesis of hybrid origin: for the 41 *P. formosa* examined, hybrid index ranged from 0.37 to 0.56 (mean, 0.49; Fig. 3). These estimates are in agreement with the estimates of admixture proportion calculated using STRUCTURE (Fig. 2a).

Given the patterns and diversity captured by both the estimates of hybrid index and the PCA, we calculated the probability of ancestry for each locus in *P. formosa* given the two parent populations and the individual’s hybrid index. This estimate is calculated using the Bayesian genomic cline model (22, 23) and is summarized in the α parameter. The α parameter is a population-level parameter that specified whether, on average, individuals are less or more likely to have either parent species’ ancestry given their hybrid index (which is a genome-wide measure of ancestry). The parameter α provides information about ancestry for a locus across individuals and not allelic state; thus, we used α to examine the organization of the genome of *P. formosa* with respect to ancestry. For frozen F1 hybrids obtained from parent species that are differentiated at all loci, we expect hybrid indexes of 0.5 (consistent with our mean estimates for *P. formosa*) and values of the cline parameter α to be zero across all loci, indicating no excess ancestry. However, our results suggest that *P. formosa* is either not a frozen F1 hybrid derived from a single F1 individual or substantial evolution has occurred in *P. latipinna* and *P. mexicana* since they gave rise to *P. formosa*. As shown in Fig. 3B, most loci in *P. formosa* have parameter α estimates not different from zero, indicating no excess ancestry for either parent species at those loci. However, about 12% of the loci have excess ancestry from one or the other parent species, consistent with the conclusion that genetic recombination might have occurred in *P. formosa*. An alternative explanation for these results could be that the allele frequencies of the parent species at the time of hybridization were much more similar than they are now. Interestingly 15% of the SNPs analyzed in this study are polymorphic in all three species, whereas 12.5% are shared only by *P. formosa* and *P. latipinna* and 21.6% are shared by *P. formosa* and *P. mexicana*. These results suggest that possibly *P. latipinna* and *P. mexicana* were more similar at the time of hybridization but have since diverged because of drift, mutations, and selection. In fact, only 1.8% of SNPs are now shared among species at the time of hybridization were much more similar than they are now. Interestingly 15% of the SNPs analyzed in this study are polymorphic in all three species, whereas 12.5% are shared only by *P. formosa* and *P. latipinna* and 21.6% are shared by *P. formosa* and *P. mexicana*. These results suggest that possibly *P. latipinna* and *P. mexicana* were more similar at the time of hybridization but have since diverged because of drift, mutations, and selection. In fact, only 1.8% of SNPs are now shared exclusively between the two sexual species, indicating that they are now clearly differentiated (as also suggested from the estimates of *G_{ST}*).

The possibility that our results are attributable only to the fact that the parent populations were more similar at the time of hybridization and differentiated since is not a trivial one. To estimate both the hybrid index and α parameter we must assign
putative parent species to *P. formosa*. We chose populations of *P. mexicana* found in the northern part of the species’ range and populations of *P. latipinna* found in the southern part of its range because previous work has suggested that the region of Tampico, Mexico, is where *P. formosa* has originated (13, 17). However, this ad hoc decision might have influenced the estimates of α, given that the allele frequencies of the sexual individuals found in those populations now might be significantly different from the allele frequencies during the time of hybridization. To address this possibility, we also estimated α for all of the other parent population combinations and found that the locus-specific α estimates did not correlate between different parent population combinations (Table S1 and Fig. S2). The patterns found during the estimation of α across the genome can be explained by (i) recombination in the gynogenetic species coupled with differentiation between the parent populations or (ii) high differentiation in the parent species (and populations within each parent species) after the hybridization event. The estimation of linkage and Hardy–Weinberg (HW) disequilibria (Fig. S3) provide evidence to support the hypothesis that some recombination has occurred in *P. formosa*. In fact, calculations of a Burrow’s composite measure of linkage disequilibrium demonstrate that *P. formosa* exhibits less disequilibrium than expected for an asexual lineage of hybrid origin descending from a single F1 individual, which suggests that recombination has reduced the linkage disequilibria created by admixture (SI Results and Discussion and Fig. S3).

To rule out the hypothesis that the excess ancestry recorded was attributable to recent gene flow with either of the parent species, we compared the estimations of the α parameter for the northern and southern populations of *P. formosa*, where *P. formosa* is sympatric with *P. latipinna* and *P. mexicana*, respectively. This analysis revealed no difference in the pattern of parent contribution (Fig. 3C). Thus, the excess ancestry for ~12% of the loci surveyed does not vary with geography across *P. formosa* and does not appear to be a function of recent gene flow from the parent species. Although previous studies have found that microchromosomes might be inherited by *P. formosa* (24), we found no evidence to support their results. However, given that we only used SNPs that had a minimum coverage of five reads per population, and microchromosome inheritance has been recorded only in a handful of populations not sampled extensively in this study, it is possible that microchromosomes were not included in our analysis at all because they are not present in every population.

Together with investigating the ancestry of *P. formosa*, we tried to shed light on the allelic state of the loci used for the analyses. The estimation of observed heterozygosity across loci revealed patterns that are inconsistent with the hypothesis that *P. formosa* is a frozen F1 hybrid descended from a single F1 individual (Fig. S4). Specifically, loci with observed heterozygosities

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**Table 1. Summary of genome-wide G_{ST} calculation**

<table>
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<tr>
<th>Pairwise comparisons</th>
<th>n</th>
<th>Mean G_{ST}</th>
<th>Maximum G_{ST}</th>
<th>Minimum G_{ST}</th>
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<tr>
<td><em>P. latipinna</em> vs. <em>P. latipinna</em></td>
<td>3</td>
<td>0.06 ± 0.02</td>
<td>0.043</td>
<td>0.081</td>
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<tr>
<td><em>P. mexicana</em> vs. <em>P. mexicana</em></td>
<td>3</td>
<td>0.096 ± 0.02</td>
<td>0.077</td>
<td>0.125</td>
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<td>1</td>
<td>0.028</td>
<td>Upper bound: 0.0278</td>
<td>Lower bound: 0.0288</td>
</tr>
<tr>
<td>Interspecific comparisons</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>P. latipinna</em> vs. <em>P. mexicana</em></td>
<td>9</td>
<td>0.361 ± 0.01</td>
<td>0.341</td>
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<td><em>P. mexicana</em> vs. <em>P. formosa</em></td>
<td>6</td>
<td>0.155 ± 0.02</td>
<td>0.132</td>
<td>0.180</td>
</tr>
<tr>
<td><em>P. formosa</em> vs. <em>P. latipinna</em></td>
<td>6</td>
<td>0.163 ± 0.01</td>
<td>0.144</td>
<td>0.170</td>
</tr>
</tbody>
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Mean and range of G_{ST} for comparisons between populations of *P. latipinna* and *P. mexicana* and for comparison within species among populations of *P. latipinna* (North, Central, and South; Fig. 1) and *P. mexicana* (North, Central, and South; Fig. 1). Mean and upper and lower bounds provided for the comparison between the two regions (North and South) of *P. formosa*. |
different from 0 (for loci at which the parents are not differentiated) or 1 (for loci at which the parents are differentiated) are not expected in F2 hybrids descended from a single F1 individual (which would be consistent with the hypothesis of a single hybridization forming the mother of all P. formosa). The variation in heterozygosity across loci (Fig. S4) is, instead, consistent with a short history of backcrossing, perhaps before the onset of gynogenesis or with the hypothesis that P. formosa is comprised of descendants of multiple, independently formed F1 gynogenetic individuals. However, the data gathered for the present study and work performed in other laboratories suggest that a short history of recombination is more likely than the hypothesis of independent origins of clonally reproducing F1 hybrids. The ancestral P. formosa might have been a sexually reproducing hybrid for some time before becoming gynogenetic, as hypothesized by Turner et al. (16), and later by Stock et al. (19). Unsexual sperm-dependent organisms might be rare because certain genetic compositions (and possibly specific epistatic interactions) are necessary for the evolution of gynogenesis. Stock et al. (19) referred to this idea as the “recombinant hypothesis.” Perhaps the specific combination of alleles required for gynogenesis only occurred after some cycles of recombination and independent assortment of alleles. This possibility could explain why no one has been able to reproduce P. formosa in the laboratory, even after extensive attempts to do so (13-16, 19-21).

A recent study also found some loci in P. formosa that were homozygous for one of the parent species, and the authors suggested that mitotic gene conversion (or mitotic recombination) during gamete formation might explain the pattern (18, 19). Our results are consistent with the hypothesis of mitotic gene conversion. When this particular type of recombination occurs, some loci become homozygous for one of the alleles (25), causing a loss of heterozygosity and an increase in linkage disequilibrium. The probability of the occurrence and success of gene conversion varies across the genome (26). This mechanism causes genotypes to vary among individuals and causes a decay of admixture linkage disequilibrium because recombination within admixed individuals and between chromosomes of different ancestry occurs. Thus, mitotic gene conversion could potentially explain the observed variation in ancestry across loci in P. formosa and the variation among individuals of P. formosa.

The last goal of our study was to determine the amount of genotypic variation present within P. formosa. The GST calculations and the PCA suggest relatively high genotypic diversity in P. formosa. This observation is in agreement with previously published results (17, 27, 28), which all found that P. formosa was genotypically variable. Stock et al. (19) suggest that the high genetic diversity in P. formosa is attributable to high mutation rates because the phylogenetic analyses of mtDNA variation suggested a monophyletic origin of P. formosa (19). Turner et al. (27) also suggested that high mutation rates are more probable than multiple hybrid origins based on allozyme data. However, some of the P. formosa studied by Turner et al. (27) were collected in the population where triploid individuals are present, and, therefore, the high clonal diversity found in this population in the Rio Purification might have been caused by the presence of triploids. We did not include triploids in this study. To address the possibility suggested in previous studies, that high mutation rates within P. formosa contribute to high genetic variation, we calculated the percentage of variable SNPs private to P. formosa and found that 3% of the SNPs used in this study are only found in the gynogenetic species. Thus, mutation accumulation within P. formosa has been moderate and does not fully explain the genotypic diversity in this species.

Our analyses using more than 25,000 SNPs from across the genome of P. formosa document considerable genotypic variation within this gynogenetic species. Given the complexity of the genomic patterns across loci and among individuals, it is not currently possible to make definitive inferences about the number of clonal lineages or the number of hybridization events involved in the origin of P. formosa. We can conclude, however, that our results are consistent with the hypothesis that the formation of P. formosa as a unisexual species is the result of hybridization and possible subsequent recombination. Recombination might have occurred following hybridization and the resulting genotypic variation has obscured our ability to discern distinct lineages. Results from calculation of genetic distances between individuals across loci illustrates that individual P. formosa are not identical to one another (Fig. S5). However, differentiation between P. formosa from different geographic regions appears to be less than among populations of either parent species (Fig. S5). Recombination following hybridization (i.e., recombination via sexual reproduction) could have occurred before the onset of the gynogenetic reproductive mode, resulting in numerous genotypes (as suggested from the calculation of observed heterozygosity). Alternatively, some form of asexual recombination, most likely mitotic gene conversion or autodixis (29, 30, 31), might have occurred (or might still be occurring). Similar mechanisms for the production of genetic diversity have been proposed for the unisexual lizards in the genus Darevskia [formerly Lacerta, Lacertidae (32, 33)] and have been shown to possibly play a significant role in the maintenance of asexual species (8).

Regardless of the origins of genetic variation in P. formosa, it is clear that this variation could contribute to the persistence of this species. Coexistence between a unisexual sperm-dependent species, and its host can be achieved and maintained if genetic variation is present in a population because natural selection can select against the clones that overlap extensively in resource use with their host species (frozen niche variation) (34, 35). Intriguingly, some form of asexual recombination, such as mitotic gene conversion or automixis, might also facilitate a reduction in the rate of accumulation of deleterious mutations (2) and increase the longevity of P. formosa beyond what is predicted by theoretical models (36).

Materials and Methods

In-depth descriptions of the protocols, models, and calculations used are in SI Materials and Methods. This work was performed under Institutional Animal Care and Use Committee no. 0818_0325_18.

Next-generation DNA sequence data from 192 fish were generated with the Illumina GAII platform following recently developed methods (for more details see, refs. 37 and 38 and SI Materials and Methods). A total of 32,492 variable sites were identified using custom Perl scripts together with samtools and balsam versions. Because of the low numbers of individuals sampled from each locality, we pooled individuals across localities into eight geographical regions to obtain adequate sample sizes to perform all of our analyses (Fig. 1A).

We trimmed data to only those SNPs with a minimum of five reads per marker per region (population grouping), which produced 26,313 SNPs, and then used Bayesian hierarchical models to estimate allele frequencies for each locus based on the observed data by using the allele frequency Bayesian model presented in Gompert and Buerkle (37). We summarized population genetic structure at the individual level using both a PCA and STRUCTURE 2.2 (40, 41). We also summarized population genetic structure at the population level by calculating pairwise GST statistics (42) for all combinations of regional groupings (Fig. 3).

To investigate the genomic composition of P. formosa, we used a Bayesian approach to estimate hybrid index for all P. formosa individuals and to assess ancestry (relative to the putative parental species) at all SNP loci for all individuals. To obtain a clearer picture about the robustness of the pattern obtained using the putative parent populations (P. latipinna and P. mexicana in northern Mexico), we repeated the estimation of both the hybrid index (Fig. S2) and α for all combinations of possible parent populations and calculated the correlation coefficient between these new estimates vs. the estimates obtained with the putative parent populations (Table S1).

Results from the genomic clines analysis (Results and Discussion) provided possible evidence of a history of recombination in P. formosa. This was unexpected given the hybrid species (P. formosa) and the proposed lack of recombination in this asexual species. Consequently, we predicted substantially higher linkage disequilibrium in this species compared with the parental species. We, therefore, calculated Burrow's composite measure of...
linkage disequilibrium (Δ) between all pairs of variable loci (43, 44) and performed a set of simulations to untangle the effects of linkage disequilibrium and HW disequilibrium on Δ.

To further investigate the results of the linkage disequilibrium estimation and to better understand the allelic state of the loci analyzed, we calculated the observed heterozygosity for each locus in each population (Fig. S4).

To address the question of whether mutation accumulation only can explain the variation present in *P. formosa* (as suggested by previous studies), we calculated the proportion of variable SNPs private to *P. formosa*, *P. mexicana*, and *P. latipinna*, as well as the proportion of SNPs shared by all species and by only two species.

As an alternative means of illustrating the genotypic variation observed within *P. formosa* (Fig. 1B), we calculated the "genotypic distance" between each pair of individuals at each locus as a measure of genotypic dissimilarity among individuals (Fig. S5).

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