Hybridization between differentiated lineages can have many different consequences depending on fitness variation among hybrid offspring. When introduced organisms hybridize with natives, the ensuing evolutionary dynamics may substantially complicate conservation decisions. Understanding the fitness consequences of hybridization is an important first step in predicting its evolutionary outcome and conservation impact. Here, we measured natural selection caused by differential viability of hybrid larvae in wild populations where native California Tiger Salamanders (Ambystoma californiense) and introduced Barred Tiger Salamanders (Ambystoma tigrinum mavortium) have been hybridizing for 50–60 years. We found strong evidence of hybrid vigor; mixed-ancestry genotypes had higher survival rates than genotypes containing mostly native or mostly introduced alleles. Hybrid vigor may be caused by heterozygote advantage (overdominance) or recombinant hybrid vigor (due to epistasis or complementation). These genetic mechanisms are not mutually exclusive, and we find statistical support for both overdominant and recombinant contributions to hybrid vigor in larval tiger salamanders. Because recombinant homozygous genotypes can breed true, a single highly fit genotype with a mosaic of native and introduced alleles may eventually replace the historically pure California Tiger Salamander (listed as Threatened under the U.S. Endangered Species Act). The management implications of this outcome are complex: Genetically pure populations may not persist into the future, but average fitness and population viability of admixed California Tiger Salamanders may be enhanced. The ecological consequences for other native species are unknown.

Hybridization (interbreeding between differentiated lineages) occurs in almost all sexually reproducing groups of organisms (1–3). Consequences of hybridization include fusion of previously distinct lineages, extinction or local extirpation of one or both lineages, evolution of reproductive isolation via reinforcement, and production of novel, highly fit hybrid phenotypes. These outcomes depend on the distribution and inheritance of fitness among hybrids and often have important implications for both evolutionary and conservation biology.

In evolutionary biology, hybrid fitness and its genetic basis are important for understanding the evolution of reproductive isolation and the consequences of horizontal gene flow on the diversity and complexity of life (4–7). In conservation biology, hybrid fitness is a key to the mechanistic basis of inbreeding and outbreeding depression, the potential loss of biodiversity due to genetic swamping, and the evolution of invasiveness (8–11). These considerations become particularly important when introduced organisms successfully hybridize with natives.

Unlike cases of natural hybridization, some outcomes of human-mediated hybridization are generally considered less desirable than others. Low fitness of hybrids can reduce mean population fitness, rendering populations more vulnerable to extinction (8, 12). Novel hybrid genotypes may contribute to de novo evolution of invasiveness (9), and admixed populations may be considered less valuable than an authentic native gene pool (10). On the other hand, admixture may rescue declining native populations by alleviating inbreeding depression (13, 14) or facilitating adaptive evolution in modified or degraded habitats (15–17).

The long-term consequences of hybridization are strongly influenced by the genetic basis of hybrid fitness. In the case of hybrid vigor, genetic models fall into two classes: heterozygote advantage and recombinant hybrid vigor (18–20). Heterozygote advantage (overdominance) refers to beneficial interactions between heterospecific alleles of a single locus. Recombinant hybrid vigor depends on multilocus genotypes and may be caused by epistasis (beneficial interactions between heterospecific alleles from different loci) or by complementary effects of independent advantageous alleles from each parental population (19, 20). For example, if different loci have fixed deleterious alleles in each parental population, hybrids bearing the superior allele at each locus will have higher fitness than either parent.

Unlike heterozygote advantage, where the most fit genotype cannot breed true, recombinant hybrid vigor favors the fixation of a single recombinant genotype bringing together beneficial alleles from each parent. This new, true breeding genotype may be considered a modified version of one of the parental lineages or a new hybrid species. In conservation terms, fixation of a recombinant genotype may represent a minor evolutionary modification or total genetic “extinction” of a native form (this may depend more on one’s perspective than on the genetic details). Perhaps more important, the new form may have undesirable ecological effects on native communities (9).

Recent, human-mediated hybridization between native California Tiger Salamanders (Ambystoma californiense) and introduced Barred Tiger Salamanders, (Ambystoma tigrinum mavortium) constitutes an important case study of the evolutionary and conservation consequences of hybridization (21–23). Before the introduction 50–60 years ago (by bait dealers interested in selling salamander larvae to bass fishermen), the two lineages had been geographically separated for ≈5 my (24, 25). Despite such a long period of divergence, they hybridize readily and have formed a hybrid swarm in the Salinas Valley of California (21–23). Multiple introduction points, combined with subsequent dispersal and movements of salamanders across the landscape (23) have led to broad-scale admixture across 15–20% of the natural range of the native California Tiger Salamander. The recent listing of A. californiense as Threatened under the US Endangered Species Act (26) recognized hybridization as a major conservation concern.

In this study, we estimated viability of hybrid tiger salamander larvae over the first few weeks after hatching. This period of high mortality represents an enormous opportunity for natural se-
leation to operate on the diverse genotypes of a hybrid population (27). Observations were made in five wild breeding populations including the three major types of breeding habitats found in the Salinas Valley: natural vernal pools, seasonal cattle ponds, and perennial ponds (22). We categorized individual hybrids using multilocus genotypes based on nine physically unlinked molecular markers. Each individual was characterized as having native or introduced mtDNA and as being homozygous native, heterozygous, or homozygous introduced at each nuclear marker (22). Thus, our approach has the advantages of considering the fitness of genotypes defined more finely than simple pedigree categories (F1, backcross, etc.) and of estimating fitness in the wild across the range of larval habitats found within the hybrid swarm (28, 29). We focused on how early larval survival varies among hybrid genotypes in the wild. Specifically, is the pattern of survival consistent with hybrid dysfunction or hybrid vigor, and does fitness variation depend more on additive, dominant (heterozygous), or recombinant genetic effects?

Results

Generalized linear models relating survival probability to molecular marker-based hybrid indices clearly support hybrid vigor (Table 1 and Fig. 1). Survival was higher for individuals with high heterozygosity, \( h_k \), and intermediate ancestry, \( h_k^2 \) (see Methods). Negative coefficients for \( h_k^2 \) indicate that surviving larvae tended to be of less pure and more mixed ancestry. A more complex model (with \( h_k^2 \) and \( h_k \times h_k^2 \) terms) did not fit the data better than the reduced model shown in Table 1 [supporting information (SI) Table 2]. The significant effect of mixed ancestry over and above the effect of heterozygosity suggests that recombinant genotypes have enhanced survival that is not predicted by heterozygote advantage alone.

Details of the multivariate outcome differed among ponds (Table 1), but this is attributable to differences in initial genotype frequencies. Ponds that started highly introduced became more native and ponds that started highly native became more introduced (Fig. 1). This is consistent with selection favoring mixed ancestry over either pure native or pure introduced genotypes. Fig. 1 shows triangular regions of genotype space defined by the interdependence of ancestry and heterozygosity. Pure native or introduced genotypes, by definition, have no heterozygous loci. Individuals with all heterozygous loci must have evenly mixed ancestry, but mixed ancestry can also be achieved when some loci are homozygous for native alleles and others are homozygous for introduced alleles. Fig. 1 shows the density of individuals across this triangular genotype space before (hatchlings) and after (larvae) natural selection. In every pond, there is a clear shift toward a higher frequency of mixed-ancestry salamanders. That is, in all five ponds, the multilocus ancestry index shifts toward a more intermediate value and away from initially more extreme values. This was true regardless of whether the before-selection ancestry indices were fairly evenly distributed (Pond F) or skewed toward native (Pond G) or nonnative (BW1, CVP, JCL 2) ancestry values (Fig. 1).

Changes in single-locus allele frequencies were idiosyncratic among markers and ponds (SI Table 3). However, most genotype frequencies (28/40 marker-pond combinations) showed a trend in the direction of increased frequency of heterozygotes after

Table 1. Multivariate analyses of the difference between hatchlings and survivors

<table>
<thead>
<tr>
<th>Pond</th>
<th>BW1</th>
<th>CVP</th>
<th>Pond F</th>
<th>Pond G</th>
<th>JCL2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Intercept</td>
<td>Coefficient</td>
<td>P</td>
<td>Coefficient</td>
<td>P</td>
<td>Coefficient</td>
</tr>
<tr>
<td>Intercept</td>
<td>1.151</td>
<td>0.088</td>
<td>1.756</td>
<td>0.002</td>
<td>7.388</td>
</tr>
<tr>
<td>( h_k )</td>
<td>1.911</td>
<td>0.002</td>
<td>13.744</td>
<td>0.000</td>
<td>4.246</td>
</tr>
<tr>
<td>( h_k^2 )</td>
<td>3.508</td>
<td>0.000</td>
<td>2.076</td>
<td>0.000</td>
<td>5.190</td>
</tr>
<tr>
<td>( h_k^2 )</td>
<td>1.756</td>
<td>0.088</td>
<td>-12.368</td>
<td>0.002</td>
<td>-6.276</td>
</tr>
</tbody>
</table>

A positive coefficient for \( h_k \) indicates selection favoring nonnative alleles, a positive coefficient for \( h_k^2 \) indicates selection favoring heterozygous hybrid genotypes, and negative coefficients for \( h_k^2 \) indicate selection favoring mixed genotypes regardless of heterozygosity. These heuristic interpretations of the coefficients are correct when all else is equal; refer to Fig. 1 to see actual changes in genotype distributions.

Fig. 1. Approximate densities of multilocus genotypes in hatchlings (before selection) and larvae (after selection) in five populations of hybrid tiger salamanders were estimated by using a two-dimensional density-estimation procedure with a bivariate normal kernel (function kde2d in R [30, 31]). Gray triangles represent potential genotype space described by ancestry (the fraction of introduced alleles in an individual’s marker genotype) and heterozygosity (fraction of markers heterozygous for native and introduced alleles). Darker areas are those with larger numbers of observed individuals.
selection, supporting the interpretation of hybrid vigor (SI Table 3). One possible exception is the GNAT1 marker, for which the dominance coefficient was negative (albeit not statistically significant) in all five ponds, suggesting low fitness of heterozygotes for this genomic region despite the overall pattern of hybrid vigor (SI Table 3).

Discussion

Our study of early-larval survival in hybrid tiger salamanders reveals higher fitness of hybrid genotypes with evenly mixed ancestry than genotypes composed predominantly of alleles from one parental lineage. Such strong evidence of hybrid vigor was unexpected based on previous studies suggesting constraints on admixture, inferred from high levels of linkage disequilibrium and habitat-dependent invasion success (21, 22). Our statistical analyses suggest that hybrid survival is influenced by both heterozygote advantage and recombinant hybrid vigor. Correct interpretation of the underlying genetics is crucial to predicting the evolutionary outcome of admixture between native California Tiger Salamanders and introduced Barred Tiger Salamanders. From a conservation perspective, hybrid vigor may be viewed as good or bad for the California Tiger Salamander, depending on whether one chooses to emphasize fitness of extant populations or genetic purity as a primary conservation goal.

The Genetic Basis of Hybrid Vigor. When considering the genetic basis of hybrid vigor, it is important to recall that our markers represent a very sparse sample of the genome. Our ability to derive information from this set of markers is enhanced by extensive linkage disequilibrium due to admixture in hybrid populations (22). Admixture linkage disequilibrium decreases the likelihood that a particular marker–trait association represents close physical linkage between the marker and a quantitative trait locus (32). By the same token, admixture linkage disequilibrium increases the accuracy of our marker-based estimates of genomic ancestry ($\theta_b$) and heterozygosity ($\theta_h$) because the sampled loci are correlated with many other regions of the genome (33, 34).

Our analyses show a strong positive relationship between heterozygosity and probability of survival (Table 1 and Fig. 1). This result is consistent with heterozygote advantage (overdominance for fitness) contributing to the genetic basis of hybrid vigor. The coefficient on the “admixture” term ($\theta_b^2$) is consistently negative, implying that recombinant hybrid vigor also plays a role. Recombinant hybrid vigor may be caused by synergistic interactions between heterospecific alleles at different loci (epistasis) or complementary effects of superior introduced alleles at some loci and superior native alleles at other loci (19, 20). That is, $\theta_b^2$ is better viewed as an indicator of overall genomic admixture rather than additive $\times$ additive interaction per se.

Hybrid vigor in other systems has often been attributed to heterozygote advantage (35–39). However, heterozygote advantage is lost each generation because of segregation among gametes. If the most-fit genotype is heterozygous at many loci, it cannot breed true and will rarely, if ever, reappear in generations beyond the F1 (40). In contrast, if hybrid vigor is caused by complementary or epistatic genes, the most-fit genotype is homozygous for alleles derived from each lineage at different loci (19). Such recombinant homozygotes appear only in later generations, and they can breed true. True breeding recombinant genotypes can therefore establish new lineages with genomes that are mosaics of the two ancestral genomes (41–44).

If both heterozygote advantage and recombinant hybrid vigor are operating in the Ambystoma hybrid zone, then one likely outcome is a population of tiger salamanders that is fixed for native alleles at some loci, advantageous introduced alleles at other loci, and segregating native and introduced alleles as balanced polymorphisms at still other loci. However, the statistical pattern of high mean fitness of highly heterozygous genotypes may be transient, a temporary consequence of the current levels of admixture and linkage disequilibrium that may abate as the population genetic background changes from a highly variable hybrid swarm to a more stable mosaic of native and introduced alleles in different parts of the genome. As the population becomes fixed for a single admixed genetic background, the heterozygote advantage at a given locus may disappear (SI Text).

Variation, linkage disequilibrium, and divergent allele frequencies between habitats have been maintained over 12–24 generations of admixture between A. californiense and A. t. mavortium in the Salinas Valley (22, 23). However, hybrid vigor tends to maintain genetic variation, decrease linkage disequilibrium between conspecific alleles, and homogenize allele frequencies across habitats. All of these processes are illustrated in Fig. 1. Three processes are likely working against the tendency of hybrid vigor to bring about a stable distribution of mixed-ancestry genotypes. First, genetic drift contributes to variance among breeding ponds (45). Second, higher frequencies of introduced alleles are maintained in perennial than in seasonal breeding ponds (22). Finally, immigration from pure native populations outside of the hybrid swarm tends to increase native allele frequencies and contribute to a broad-scale geographic gradient in allele frequencies (23).

The high frequencies of introduced alleles in perennial ponds remains one of the strongest genetic patterns in the tiger salamander hybrid system; it may be maintained by habitat choice or selection in later life history stages. Fitzpatrick and Shaffer (22) hypothesized that introduced genotypes gain an advantage in perennial water bodies by extending their larval period or even breeding in the larval (paedomorphic) condition. Pure native California Tiger Salamanders must metamorphose to reproduce (46), but Barred Tiger Salamanders regularly forgo metamorphosis in perennial ponds and breed as paedomorphs (47, 48). Paedomorphs often reach sexual maturity earlier than metamorphs, produce larger clutches, and may breed earlier in a given season; any of these life-history factors may provide introduced genotypes with an advantage in perennial ponds (49, 50). The perennial pond populations studied here (BW1 and JCL2) show strong viability selection favoring hybrids during the early part of the larval period (Table 1 and Fig. 1). Antagonism between hybrid vigor for larval survival in all ponds and later selection favoring introduced genotypes in perennial ponds may explain the persistent pattern of high introduced allele frequencies in perennial ponds and relatively even native and introduced allele frequencies in seasonal ponds throughout the range of the hybrid swarm (23).

Conservation Consequences of Hybrid Vigor. Hybridization between an introduced form (A. t. mavortium) and a declining native protected by the Endangered Species Act (A. californiense) raises several difficult conservation issues (8, 10, 23, 51). These include, (i) what are the consequences of hybridization for population viability? (ii) what are the consequences for other native species and the communities in which they occur? (iii) which individuals or populations of salamanders should be protected? and (iv) should any individuals or populations be eradicated? The answers to iii and iv depend, in part, on the answers to i and ii but may also depend on practical considerations and perspectives on the merit of genetic purity as a conservation goal. Our results have important implications for population viability and genetic purity of California Tiger Salamanders and for potential impacts on other organisms native to California ponds and grasslands. Hybrid vigor is probably not detrimental to population viability. However, some authors have raised the concern that negative effects of introgressive hybridization on fitness may not

Fitzpatrick and Shaffer

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be evident except during infrequent “ecological crunches” (10, 52). Such episodic changes in selection do occur (53–55), and may be widespread on the California landscape, given its tremendous among-year variance in rainfall and pond hydroperiod. Introgression could potentially contribute to loss of a crucial adaptation during an interlude between episodes of selection, leaving the population more vulnerable to extinction when the next crunch arrives. In addition, our characterization of hybrid fitness is incomplete, focusing only on early larval mortality. However, native and introduced tiger salamanders have been mixing in California for >50 years, and the hybrid swarm has survived and expanded (23) through droughts, El Niños, and substantial anthropogenic environmental change. The effect of admixture on population viability in California Tiger Salamanders remains an important open question, but it does not appear to us, at this time, to be a major threat.

Hybrid vigor implies that the native gene pool is not resistant to invasion of introduced alleles. Thus, a mixture of gene pools is expected, with fixation of advantageous introduced alleles and preservation of advantageous native alleles. This constitutes a loss of biodiversity, because alleles and genotypes unique to A. californiense are being lost. However undesirable it may be, this evolutionary transformation should not engender the same ethical concerns raised by true demographic extinction. In fact, the aesthetic appeal of a genetically authentic native gene pool may be in conflict with the best interests of the animals if hybrid vigor enhances population viability. Whether or not population viability is enhanced in this case depends on the relationship between average fitness and density regulation (56).

If hybrid vigor results in higher densities, larger body sizes, or more rapid growth of larvae in hybrid populations, the impacts of hybridization will almost certainly extend to other members of the ecological community. Tiger salamander larvae are voracious consumers of anuran tadpoles and aquatic invertebrates. Endangered, threatened, and sensitive prey that share habitats with tiger salamanders in the hybrid zone include California Red-Legged Frogs (Rana draytonii), Western Spadefoots (Spea hammondii), and Vernal Pool Fairy Shrimp (Branchinecta lynchi). Thus, hybrid vigor in tiger salamander larvae presents a credible but unproven threat to other pond-breeding organisms.

Our results support the view that hybridization can alter the evolutionary process by contributing novel genetic advantages to admixed populations (1, 15, 41). This view also suggests that hybridization between native and introduced lineages could accelerate the evolution of invasiveness and intensify the ecological impact of a biological invasion (9). Understanding the effect of hybridization on the ecological interactions of tiger salamander populations is a priority for future work in this system.

Methods
Natural selection among hybrid genotypes was estimated by using a cohort analysis in which we compared the distributions of genotypes in a single group of offspring at two different points in time. This provides a direct measure of the response to natural selection within a single generation (57). Furthermore, by focusing on diagnostic codominant molecular markers, we measured genetic changes that can be readily explained in terms of additive, dominant, and epistatic or complementary effects of differences between the native and introduced salamanders.

Sampling. We studied a subset of the breeding sites described in ref. 22. Pond F and CVP are natural vernal pools that fill during winter rains and dry by early to midsummer. They represent the natural, unmodified breeding habitat for native California Tiger Salamanders (46). Pond G is a man-made seasonal cattle pond with a hydroperiod similar to vernal pools. JCL2 and BW1 are perennial ponds that normally hold water through the summer, although both had dried completely during the summer before this study.

We quantified multilocus genotypes during two early life-history stages. Our before-selection samples consisted of eggs collected during the laying season, whereas our after-selection samples were drawn early in the larval stage. Each pond was outfitted with artificial oviposition sites consisting of 25 bamboo stakes (1 m long and 0.5 cm in diameter) planted into the pond bottom in water 30 cm deep and four 1.2-m-long coils of 12-gauge galvanized steel wire in water 40–50 cm deep. Ponds were visited weekly from December 2002 through February 2003. On each visit, all salamander eggs (if present) were removed from all stakes and wires to guarantee that all eggs collected were laid during the preceding week. Eggs were counted and placed in a bucket, from which a random sample of 50 eggs was collected. Remaining eggs were scattered in shallow water where they could readily adhere to submerged vegetation and rocks (natural oviposition sites). Collected eggs were held in cups containing 20% Holtfreter’s solution (58) and allowed to hatch. Hatchlings were killed and stored at −80°C.

Young larvae were captured by haphazardly seining ponds. Collections were made on February 6 and March 26, 2003. Reproduction in Pond F did not begin until the week of February 28, when there was a single bout of egg laying. The artificial oviposition sites in Pond G were destroyed in early February by cattle. Therefore, these two ponds are represented by single samples of larvae. In the other three ponds, we found no statistically significant (α = 0.05) differences between the two samples of larvae (data not shown) and therefore pooled them for all hatching vs. larvae comparisons below.

Molecular Methods. Each individual was assayed for the nine molecular markers used by Fitzpatrick and Shaffer (22). These are single nucleotide differences that distinguish native A. californiense from introduced A. t. mavorium alleles at mtDNA and eight mapped nuclear genes. Each individual was characterized as having native or introduced mtDNA and as being homozygous native, heterozygous, or homozygous introduced at each nuclear marker. Primer sequences and PCR conditions can be found in ref. 22 (also see refs. 21 and 59). Diagnostic alleles were identified by restriction enzyme digestion and agarose electrophoresis of PCR products, as described by Fitzpatrick and Shaffer (22).

Data Analyses. We used both single-marker and multilocus statistical analyses to evaluate variation in survival among genotypes. For single locus selection analyses, we used standard marker-trait regressions (60, 61):

$$\log\left(\frac{P}{1-P}\right) = \text{intercept} + b_A X_A + b_H X_H + \text{error}, \quad [1]$$

where P is the probability that a data point is a surviving larva, \(X_A = -1, 0, \) or 1 for homozygous native, heterozygous, and homozygous introduced genotypes, respectively, and \(X_H = 0 \) or 1 for homozygous and heterozygous genotypes, respectively. The fitted coefficients \(b_A \) and \(b_H \) represent additive and dominance effects. Regressions were fitted in the program R 2.4.1 (31) by using the glm() function with a binomial error distribution and observations weighted by stratum (see below). We assume fitted models represent indirect responses to selection on loci with unknown linkage relationships to our markers. The residual error includes variance due to incomplete linkage between markers and selected sites as well as environmental and sampling variance (37).

For multilocus selection analyses, we used the quantitative genetics approach described by Lynch (18). This model identifies
additive, dominance, and epistatic effects on hybrid fitness as terms in a linear statistical model. In our case,

\[
\log\left( \frac{P}{1-P} \right) = \text{intercept} + \alpha_1 S_3 + \delta_1 \delta H_2 + \alpha_2 S_3^2 + \delta_2 \delta H_2^2 + (\alpha \delta_1) \delta H_2 + \text{error},
\]

where individual genotypes are described by \( \theta_3 \), a linear ancestry index ranging from \(-1\) (all native alleles) to \(+1\) (all introduced alleles) and \( \delta H_2 \), a heterozygosity index ranging from \(-1\) (all markers homozygous) to \(+1\) (all markers heterozygous). The fitted coefficients \( \alpha_1 \) and \( \delta_1 \) describe simple additive and dominance effects, whereas \( \alpha_2 \) and \( \delta_2 \), and \( (\alpha \delta) \) describe additive \( \times \) dominance, dominance \( \times \) dominance, and additive \( \times \) dominance epistasis, respectively \((18)\). This model was fitted to the data from each pond by using the glm() function with a binomial error distribution and observations weighted by stratum, as for the single locus case (see below).

Two additional issues needed to be addressed in our statistical analyses. First, an interesting problem was created by the temporally extended breeding season in three of our study ponds (BW1, G, and JCL2). In these ponds, breeding consisted of three to five major episodes corresponding to significant rain storms; each episode potentially involved a different set of adults with different genotype frequencies. Thus, our before-selection samples were stratified simple random samples. Each weekly sample (stratum sample) was weighted according to stratum size \((\text{stratum size} / \text{total sample size})\) and those weights passed to the glm() function. Under this weighting scheme, the after-selection individuals each get a weight of 1.0 because they come from a single stratum, and the averages of the individual weights before and after selection are equal \((62)\).

Second, traditional estimates of confidence intervals for regression coefficients do not account for error in the independent variables. However, \( \theta_3 \) and \( \delta H_2 \) were estimated with error because our markers are samples of the genome \((\theta_3 = 2P_T - 1, \delta H_2 = 2P_{CT} - 1, \text{where } P_T \text{ is the fraction of an individual's marker alleles derived from } A. \ t. \text{maavoritum, and } P_{CT} \text{ is the fraction of markers heterozygous for native and introduced alleles})\). Therefore, we used a bootstrap and randomization procedure to incorporate this sampling variance into hypothesis tests regarding the fitted coefficients described in Eq. 2. For each of 10,000 replicates, we performed a two-step Monte Carlo procedure.

First, we bootstrapped across nuclear markers. Bootstrapping across markers addresses error in individual estimates of \( \theta_3 \) and \( \delta H_2 \) due to sampling only a fraction of each salamander’s genome. This is in the same spirit as bootstrapping across characters in phylogenetic analysis \((63)\) and results in each individual having associated with it a distribution rather than a single point estimate of each \( \theta_3 \). MtDNA was treated as a fixed effect; it was included once and only once in every replicate, because there is only a single mitochondrial allele to sample. Excluding mtDNA slightly affected parameter estimates but not their signs or \( P \) values. Including mtDNA in the pool of markers to resample is inappropriate because it does not contribute to \( \theta_3 \). The second step in each replicate was to randomize observations to simulate the null hypothesis that hatchlings and surviving larvae were random samples from a single distribution of genotypes. We recorded the fraction of replicates giving coefficients larger \((f_{\text{larger}})\) and smaller \((f_{\text{smaller}})\) than the estimate from the original data and estimated two-tailed \( P \) values as \( 2 \times \min(f_{\text{larger}}, f_{\text{smaller}}) \). The procedure was applied separately to each of our five study ponds.

The full Lynch \((18)\) model (Eq. 2) may overfit some data sets (particularly those with highly skewed allele frequencies), leading to problems with collinearity of variables and high variances around coefficient estimates. Therefore, we also fitted a reduced model:

\[
\log\left( \frac{P}{1-P} \right) = \text{intercept} + \alpha_1 S_3 + \delta_1 \delta H_2 + \alpha_2 S_3^2 + \text{error}.
\]

This model focuses on fitness variation as a function of individual ancestry \((\theta_3)\), individual heterozygosity \((\delta H_2)\), and as a function of individual admixture \((\theta_3^2)\) decoupled from ancestry and heterozygosity \((\delta H_2^2 = 1)\) in both pure native and pure introduced genotypes, and it equals 0 in individuals with evenly mixed ancestry, i.e., half native and half introduced alleles regardless of heterozygosity). The coefficient \( \alpha_2 \) represents effects of mixed ancestry not depending on heterozygosity but does not distinguish true epistasis from complementary effects. We compared the likelihoods of the full and reduced models by means of Akaike’s Information Criterion \((AIC)\) \((64)\).

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