Surprising spatiotemporal stability and frequency-independence across multiple fitness peaks driving adaptive radiation in the wild

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Abstract

The effect of the environment on fitness in natural populations is a fundamental question in evolutionary biology. However, most empirical field studies of fitness do not experimentally manipulate phenotypes or environmental conditions and rarely investigate more than a single species or population. Thus, the relative importance of the competitive environment versus intrinsic organismal performance in shaping the location, height, and fluidity of fitness peaks on the adaptive landscape remains largely unknown. We experimentally tested the effect of competitive environment on a multi-peak fitness landscape driving a microendemic adaptive radiation of generalist and trophic specialist pupfishes on San Salvador Island, Bahamas. We manipulated phenotypes, by generating lab-reared hybrid crosses, and competitive environment, by altering the frequency of rare phenotypes within field enclosures in their natural hypersaline lake environments on San Salvador. We tracked the growth and survival of 2,611 F4/F5 hybrids for 3 to 11 months in high- and low-frequency treatments replicated across two lake populations. We found strong evidence for frequency-dependent growth rates within and between enclosures, but no evidence for frequency-dependent survival differences. However, both fitness proxies supported a complex fitness landscape isolating generalist phenotypes on a local fitness peak separated by a small fitness valley from a higher fitness peak for molluscivores and a large fitness valley isolating the scale-eating phenotype in all trait dimensions. The striking consistency of this multi-peak fitness landscape across competitive environments, multivariate trait axes, and a previous field experiment provides experimental evidence for stasis, possibly due to fixed biomechanical constraints on organismal performance. These results challenge existing theory and highlight the interplay of organism and environment underlying the static and dynamic features of fitness landscapes.
Introduction

The adaptive landscape, the complex mapping of fitness onto phenotype or genotype, is both a central unifying concept in evolutionary biology and an empirical measurement [1–5] which links the microevolutionary processes of natural and sexual selection in wild populations with macroevolutionary patterns of speciation, novelty, and adaptive radiation [6–9].

Despite its central importance, it remains unclear what factors shape the fitness landscape across space and time. In classical views arising from both Wright’s [4] and Simpson’s [10] original conceptions of genotypic and phenotypic fitness landscapes, respectively, and Fisher’s geometric model [11], fitness optima are static on a high-dimensional fitness landscape. These static fitness peaks are due to epistasis within genotypic networks [12] or functional tradeoffs between different ecological niches or collections of similar niches known as adaptive zones [13]. However, a more recent paradigm originating from game theory [14–17] proposes that the fitness landscape is dynamic and resembles a trampoline: as the relative frequency of phenotypically similar individuals increases, their fitness decreases due to increased competition for resources, whereas rare phenotypes have a fitness advantage [5]. This is known as negative frequency-dependent disruptive selection and can lead to ecological speciation in sympatry, even when adapting to a unimodal resource distribution [18–22]. Laboratory and field studies of natural populations have provided extensive support for negative frequency-dependent disruptive selection [1,15,23–34] to the extent that some investigators assert its universality in all natural populations [35]. Due to its elegance and mathematical tractability, frequency-dependence has also been widely adopted by theorists as the sole mechanism for disruptive selection in most speciation models [5,9,36].
However, the relative contributions of the competitive environment (i.e. frequency-dependence) versus static fitness optima (i.e. functional tradeoffs) in shaping the broader topography of fitness landscapes across multiple species remains unknown. For example, although negative frequency-dependent disruptive selection may be ubiquitous within populations, the phenotypic scale of frequency-dependence is rarely measured (i.e. the competition kernel), particularly across multiple species, and can have major impacts on speciation \[18,37\]. Similarly, spatiotemporal shifts in the phenotypic location of a fitness optimum for a population due to environmental stochasticity is often assumed to follow a Brownian motion process \[38,39\] without accounting for hard boundaries imposed by functional tradeoffs or biophysical constraints (but see: \[40,41\]). Conversely, at broader macroevolutionary timescales the role of stable fitness optima in shaping trait diversification across a radiation is frequently tested and supported by fitting Ornstein-Uhlenbeck models of trait diversification using phylogenetic comparative methods \[42–46\] while ignoring the ecological effects of competition and environmental stochasticity (but see \[47,48\]).

To our knowledge, all previous field experimental tests of frequency-dependent selection involved only a single species or pair of ecomorphs \[15,24–26,29,31,34,49,50\] and there are still few studies spanning multiple habitats, traits, time periods, or species. In the few examples of studies at this larger scale, rare transgressive hybrid phenotypes appear to suffer a fitness cost, not an advantage \[51,52\]. For example, in a hybrid mesocosm experiment investigating male-male competition in stickleback, the rarest transgressive phenotypes experienced the lowest reproductive success \[52\], in contrast to predictions of sexual selection as a diversifying force \([53]\; \text{but see} \; [54,55])\). We also previously estimated multiple fitness peaks driving adaptive radiation in a clade of trophic specialist pupfish species endemic to San Salvador Island,
Bahamas by measuring the growth and survival of laboratory-reared F2 intercross and backcross hybrids placed in field enclosures [56]. Hybrid phenotypes resembling the widespread generalist species were isolated by a local fitness peak, separated by a fitness valley from a higher fitness peak corresponding to hybrid phenotypes resembling the molluscivore specialist, whereas hybrid phenotypes resembling the scale-eating specialist suffered the lowest growth and survival. Interestingly, rare phenotypes in this experiment did not experience a survival advantage and the scale of frequency-dependent survival appeared to operate only within the range of phenotypic diversity observed in the generalist population, not all three species [51]. These few experimental studies of multiple species suggest that frequency-dependent selection may have a limited phenotypic scale; however, experimental tests manipulating the frequency of rare hybrid phenotypes are needed.

Here we bridged these micro- and macroevolutionary scales by experimentally manipulating the frequency of rare hybrid phenotypes to test for an effect on the stability of multiple fitness optima in a tractable system for empirical measurements of the fitness landscape during nascent adaptive radiation. We created intermediate and transgressive hybrid phenotypes by multiple rounds of backcrossing and intercrossing between a generalist and two trophic specialist *Cyprinodon* pupfish species, manipulated the frequency of rare hybrid phenotypes between treatments in two independent lake populations on San Salvador Island, Bahamas, and then tracked individual hybrid survival and growth rates. We found negligible effects of competitor frequency on the survival of hybrids, but strong effects on growth. However, both fitness proxies supported a large and stable fitness valley isolating the novel scale-eating specialist across spatial and temporal environments. Our results lend strong empirical field support for stasis across multiple fitness peaks and valleys driving recent adaptive radiation.
Results

Field and laboratory survival of transgressive hybrid populations

We individually tagged and photographed 2,611 F4-F5 outbred juvenile hybrids resulting from crosses of all three species, generalist (*C. variegatus*), molluscivore (*C. brontotheroides*), and scale-eater (*C. desquamator*), from two different isolated lake populations on San Salvador Island, Bahamas before release into a high- and low-frequency field enclosure in each lake (Fig. 1; Table S1). The frequency-manipulation increased the frequency of transgressive hybrid phenotypes in the high-frequency treatment and generalist-type hybrids in the low-frequency treatment, resulting in a significant reduction in phenotypic variance on discriminant axis 2 (predominantly nasal protrusion) in lake 1 (Levene’s test, \( P < 0.0001 \)) and discriminant axis 1 (predominantly oral jaw size) in lake 2 (Levene’s test, \( P < 0.0001 \)) within the bivariate discriminant morphospace separating all three parental species (Fig. S1). The total density of hybrids was held approximately constant between high and low-frequency treatments (lake 1: high/low: 923/823 individuals; lake 2: high/low: 842/819 individuals; Table S1).

To sample from a broader range of environmental variability, we measured hybrid survival after 3 months in lake 1 (high-frequency: 77.1% survival; low-frequency: 75% survival) and after 11 months in lake 2 (high-frequency: 1.4% survival; low-frequency: 1.2% survival; Table S1), in the latter case spanning half the pupfish reproductive lifespan (approximately 2 years) but avoiding mortality due to senescence. There were no differences in survival probability between treatments in each lake (two-way logistic regression, treatment effect: \( P = 0.237 \)). For a control comparison, additional hybrids from each lake population (\( N = 199 \) total...
individuals) were simultaneously tagged, raised in laboratory aquaria, and their deaths and growth rates were tracked over one year.

Hybrid phenotypic diversity
Phenotypic similarity of each hybrid to the three parental species in each lake (n = 236) was calculated from 30 linear traits and angles measured from three pre-release photographs of each fish (Fig. S3). These traits were used to estimate two linear discriminant (LD) axes with major loadings (Table S2) of oral jaw size (LD axis 1) and nasal protrusion (LD axis 2), diagnostic traits of each specialist species and major axes of rapid trait diversification within this radiation [57,58]. Indeed, after correcting for standard length, residual jaw length variation within our hybrid populations exceeded the range of variation observed across allopatric Cyprinodon species and outgroup Cyprinodontidae species spanning over 20 million years since their most recent common ancestor (data from [57]; Fig. 2).

Strong directional and nonlinear selection for generalist and molluscivore phenotypes
We fit thin-plate splines using generalized cross-validation or restricted maximum likelihood to survival and growth rate data to visualize fitness landscapes across the discriminant morphospace [59]. We found evidence for directional selection on the survival of hybrid phenotypes most similar to generalist and molluscivore phenotypes in lake 1 (Fig. 3). We also found evidence of stabilizing selection on the growth rates of generalist and molluscivore phenotypes in this lake (Fig. S4). Despite low survival rates after 11 months in lake 2, we found strong evidence of nonlinear selection for hybrid phenotypes resembling the molluscivore. These landscapes were estimated from very few survivors (n = 22); however, the density of non-survivors in the
surrounding regions of morphospace and similar patterns in both field enclosures provides robust support for nonlinear divergent selection for the molluscivore phenotype in this lake. Patterns of survival in the wild were contrasted by strong directional selection for hybrids resembling the scale-eater in laboratory control populations from both lakes (Fig. 3).

We also used generalized projection pursuit regression to estimate the two multivariate linear phenotypic axes most strongly associated with survival across the 30-trait morphospace (Tables S3-S4) without making unfounded parametric assumptions about quadratic curvature as in canonical rotation analyses [60–62]. Visualization of survival fitness landscapes on the two major axes of selection indicated that the most transgressive hybrid phenotypes (i.e. least similar to parental phenotypes) suffered the lowest survival probability across treatments in both lakes (Fig. 4). In contrast, survival in laboratory control populations was shifted or opposite to the direction of selection in field enclosures along these two dominant axes of selection (Fig. 4).

We estimated the strength of multivariate selection gradients along these two major axes of selection and found significant evidence of directional selection ($P < 0.00001$) on ridge axis 1 in both lakes and marginal evidence of directional selection on ridge axis 2 in lake 2 (Table 1). The traits with the highest loadings on ridge axis 1 were a) lower jaw length and b) distance from the jaw joint to the orbit and on ridge axis 2 were a) angle between the premaxilla and orbit and b) distance from the premaxilla to the pectoral girdle (Tables S2-S3), further supporting strong selection on craniofacial trait diversification within this radiation [58].

No evidence of frequency-dependent survival differences between treatments

We found no evidence of significant treatment effects on either survival probability or the overall topography of the survival fitness landscape within the discriminant morphospace (Table 2). The
fixed effect of treatment did not improve the fit to the survival data in any of the generalized
additive models examined. Instead, models without the effect of frequency treatment were
strongly favored (Table 2; ΔAIC = 11). Similarly, we found no evidence for frequency-
dependent effects on survival between treatments on the two major axes of selection in either
lake estimated from generalized projection pursuit regression (Table 3). Models without the
effect of treatment provided a marginally better fit to the survival data on the two major axes of
selection in lake 1 (ΔAIC = 1) and were supported in lake 2 (Table 3; ΔAIC = 1.8).

Strong evidence of frequency-dependent growth rates between treatments
In contrast to survival, we found strong support for treatment effects on fitness landscapes
estimated from the independent fitness proxy of growth rate in lake 1 (Table 2; lake 2 was
excluded from all growth rate analyses due to the low number of survivors). Models including
the effect of frequency treatment on log-transformed growth rates in lake 1 were strongly favored
(ΔAIC = 68.6). This effect was robust to models including univariate splines and thin-plate
splines within the discriminant morphospace (Table 2).

Similarly, we found strong support for models including the effect of treatment for the
two major axes of selection across the entire morphospace in each lake estimated from
generalized projection pursuit regression (Table 3). The best supported model included the
effects of frequency treatment and univariate splines on the two major ridge axes of selection and
was strongly favored over a model without the effect of treatment (Table 3; ΔAIC = 46).

Phenotypic scale of frequency-dependence for growth rate but not survival
Within each enclosure we estimated the Mahalanobis distance for each hybrid individual, the
distance to the mean hybrid phenotype in the 30-dimensional morphospace correcting for trait
covariances, as an estimate of the rarity of each individual phenotype. We also calculated the
Euclidean nearest neighbor distance to the ten most similar hybrid phenotypes in 30-dimensional
morphospace for each hybrid as an estimate of the local frequency of competing phenotypes.
Overall, these measures estimate the frequency of similar hybrid phenotypes relative to each
hybrid to examine the phenotypic scale of frequency-dependence within each enclosure [51].
The frequency of similar phenotypes was not significantly associated with residual variation in
survival not explained by hybrid phenotype along the two discriminant axes (Fig. 5; Table 2).
Generalized additive models including the fixed effect of competitor frequency (distance to mean
phenotype) only marginally improved the fit to the survival data (Table 2; ΔAIC = 1; similar
results were found when substituting nearest neighbor Euclidean distance for Mahalanobis
distance: Fig. S1).

In contrast, generalized additive models including the fixed effect of competitor
frequency (both Mahalanobis and nearest neighbor distance) strongly improved the fit to the
growth data in lake 1 within the discriminant morphospace, even after accounting for the
treatment effect (Table 2; ΔAIC = 12). Similarly, for the two major axes of selection estimated
from generalized projection pursuit progression, models including the fixed effect of competitor
frequency substantially improved the fit to the growth data, even after accounting for the
treatment effect (Table 3; ΔAIC = 3.85).

Both fitness proxies support multiple fitness peaks on the adaptive landscape
Generalized additive modeling enables estimation and visualization of a joint fitness landscape after controlling for the effects of lake and treatment. The best supported models for survival included a fixed effect of lake and no effect of treatment, with either two univariate smoothing splines or a thin-plate plate and two smoothing splines modeling selection within the discriminant morphospace (Table 2). Models including spline terms were strongly supported over models including fixed linear effects of the discriminant axes (Table 2; ΔAIC = 18.7). The best supported models for growth rate in lake 1 included a fixed effect of treatment (lake 2 was omitted from these analyses due to the low number of survivors available for growth rate estimates) and two univariate smoothing splines within the discriminant morphospace (Table 2).

Strikingly, the best combined models for survival (across both treatments in both lakes) and growth rate (both lake 1 treatments) each independently supported an isolated fitness peak for hybrids resembling the generalist separated by a fitness valley from a region of higher fitness corresponding to the molluscivore phenotype (Fig. 6). This multi-peak landscape, consistent across both survival and growth rate fitness proxies and different exposure periods in each lake, was also striking similar to a previous independent field experiment using F2 hybrids in these same lakes [56]. This supports the surprising spatiotemporal stability of a complex fitness landscape topography spanning a recent adaptive radiation of trophic specialists across years, seasons, divergent lake environments, and manipulated frequencies of hybrid transgressive phenotypes.

Scale-eating trophic specialists are isolated by a highly stable fitness minimum

Across both treatments and lakes, hybrids resembling the scale-eater phenotype suffered the lowest survival and growth rates, except in laboratory control populations (Figs. 3-4, 6). This
pattern is consistent with previous observations of consistent low survival and growth rates across density treatments and lakes in scale-eater F2 hybrids; however, in the previous experiment few of the F2 hybrids fell within the phenotypic range of lab-reared scale-eaters [56]. In our current experiment, over 70 hybrids occurred within the 95% confidence ellipse of lab-reared F1 scale-eater phenotypes within the discriminant morphospace.

However, it is possible that some regions of the high-dimensional trait space may still connect scale-eater phenotypes to other regions of the morphospace through a fitness ridge [63,64]. To further explore the relative fitness of scale-eater hybrids, we visualized selection across all directions in the 30-trait morphospace by repeatedly sampling a random subset of 15 traits, calculating a discriminant axis for scale-eaters relative to generalists within this subspace, and estimating a survival spline for hybrid phenotypes on each arbitrary multivariate axis (Fig. 7). This results in a visualization of all possible fitness paths between generalist and scale-eater hybrid phenotypes for all subspaces within the 30-trait morphospace and, importantly, aligns these multivariate linear axes in the same direction from generalist to scale-eater phenotype for comparison of fitness curves across random subsets of the trait data. In three out of four field enclosures (with no relationship in the fourth), hybrids resembling scale-eaters suffered the lowest survival across nearly all visualized fitness paths, supporting their position in a high-dimensional fitness valley (Fig. 7). These analyses support a robust fitness minimum or ‘hole’ within the adaptive landscape isolating the scale-eater phenotype from other species which is consistent across different subsets of traits measured, lake environment, field exposure time, and frequency of competitors.

Discussion
No evidence of frequency-dependent survival in a multi-peak fitness landscape

We conducted an experimental field test of frequency-dependent selection in a nascent adaptive radiation of trophic specialist pupfishes. We found negligible evidence of frequency-dependent survival between treatments manipulating the frequency of rare hybrid phenotypes in two different lake environments nor any relationship between survival and the frequency of competitors (Figs. 3-5; Tables 2-3). In contrast, growth rate of survivors in each lake showed strong evidence of frequency-dependence between treatments and increased as the frequency of hybrids with similar phenotypes decreased in high-frequency enclosures (Tables 2-3, Fig. S6). These patterns were consistent across two important cross-sections of the 30-trait morphospace: the two discriminant axes separating the three parental species (Fig. 3) and the two strongest axes of nonlinear selection estimated from generalized projection pursuit regression (Fig. 4). The lack of any signal of frequency-dependent survival suggests that differences in survival among hybrid phenotypes are robust to competitive conditions and reflect intrinsic viability and performance constraints. These results complement most previous experimental studies of frequency-dependence which measured only growth rates [15,28,31].

Combined estimates of the fitness landscape for both survival and growth rate indicated a surprisingly consistent topography across space and time comprised of an isolated fitness peak corresponding to the generalist phenotype separated by a small fitness valley from a higher fitness peak corresponding to the molluscivore phenotype (Fig. 6). For survival, multiple peaks emerged from evidence of higher survival of generalist and molluscivore phenotypes after three months in lake 1 combined with evidence of much higher survival of the molluscivore phenotype after 11 months in lake 2 (Fig. 3). For growth rate in lake 1, multiple peaks emerged from higher growth rates of generalist phenotypes in both enclosures combined with moderate molluscivore
growth rates and very low scale-eater growth rates in the low-frequency enclosure (Fig. S4). These joint landscapes are admittedly reflective of combining different frequency treatments, exposure periods, lake environments, and potentially different selective regimes; however, comparison of general additive models provided no evidence of different selective regimes between treatments and only a fixed effect of lake environment, rather than a change in fitness landscape topography between lake environments (e.g. Table 2: very low support for models including a ‘by lake’ effect), thus supporting our inference of a single combined selective environment.

Across all four field enclosures and both fitness proxies the most prominent and consistent feature of fitness landscape topography was a large fitness valley isolating hybrids resembling the scale-eater, the most morphologically, ecologically, and genetically divergent specialist in the radiation [65–67], from all other hybrids across treatments, lake environments, field exposure periods, and across nearly all dimensions of the 30-trait hybrid morphospace (Figs. 3-4, 6-7). These empirical fitness landscape measurements were also consistent with observations of low fitness in hybrids partially resembling the scale-eater in a previous experiment [56].

Static features of a complex fitness landscape underlie the rare origins of trophic novelty

Overall, while there was strong selection against hybrid phenotypes driving species divergence in this radiation, hybrid survival showed minimal sensitivity to the frequency of competitors, supporting the classic view of static fitness peaks and valleys on the adaptive landscape. Indeed, the combined estimate of the fitness landscape across all four field enclosures in this study was strikingly similar to the original fitness landscape estimated for the high-density field enclosure
in lake 1 in our preview work [56]. This provides robust support for the spatiotemporal stability of an isolated generalist fitness peak separated by a large fitness valley from scale-eating and a smaller fitness valley and higher fitness peak for snail-eating. Our study also provides empirical field experimental data supporting a role for the stable fitness optima and minima frequently inferred from phylogenetic comparative studies [45,68] and observed in the fossil record [69,70].

This strikingly complex, frequency-independent, and persistent multi-peak fitness landscape within San Salvador Island’s hypersaline lakes provides an explanation for the rarity of trophic specialization across the Caribbean if outgroup generalist populations are under widespread stabilizing selection opposing trait diversification [56]. Indeed, Caribbean generalist populations show very little diversity in their trophic skeletal morphology or dietary diversity relative to San Salvador populations [58]. Scale-eating populations on San Salvador Island also show strong phylogenetic comparative evidence of adaptation to a new adaptive zone relative to generalist populations on San Salvador and neighboring islands [58] and stronger pre-mating isolation and genetic divergence from other species [71–73], consistent with faster rates of trait divergence and speciation driven by adaptation to this highly novel trophic niche.

What factors underlie spatiotemporally consistent fitness landscapes?

The frequency of similar hybrid phenotypes strongly affected growth rates both between and within enclosures; however, the ultimate survival of hybrids over both 3- and 11-month exposure periods was unrelated to their relative frequency within enclosures. In contrast to predictions of the theory of negative frequency-dependent disruptive selection [15,35,74], rare and transgressive hybrid phenotypes outside parental ranges exhibited the lowest survival rates within field enclosures. One possible explanation is that the intrinsic viability and basic
performance of survival tasks by these hybrids was impaired due to their mosaic hybrid genetic backgrounds. This may result in mismatched craniofacial traits leading to poor foraging performance or impaired suction-feeding or scale-biting performance necessary for successful foraging strikes. For example, the large oral jaws of scale-eaters appear to result from at least four moderate effect quantitative trait loci on different linkage groups that each increase jaw size [75], suggesting that the genetic basis of even this single trophic trait is moderately polygenic and may not be fully recovered within F4/F5 hybrids. Furthermore, F1 hybrid scale-eaters exhibit foraging kinematics during scale-biting strikes more similar to generalists which violates expectations of additivity, suggesting that kinematic behaviors may be non-additive and severely mismatched in more advanced hybrids [76]. Impaired foraging performance of hybrids similar to scale-eaters in field enclosures is also supported by the observation that hybrids with scale-eater morphologies in laboratory control aquaria showed the highest survival rates when fed only pellet foods (Figs. 3-4). This indicates that the field environment is contributing to the low survival of scale-eater hybrids.

An alternative non-mutually exclusive possibility is that intrinsic genetic incompatibilities within hybrids are contributing to their low survival rates, particularly if more transgressive hybrid phenotypes are associated with a greater number or more severe genetic incompatibilities. Although the San Salvador radiation only diverged approximately 10,000 years old, trophic specialists within the radiation contain ancient adaptive variants also found in outgroups that diverged over 5 million years ago (Richards et al. in prep.). Genetic incompatibility loci are known to segregate in wild populations ([77]; reviewed in [78]) and hundreds of genetic incompatibility loci have also been found between swordtail fish species of similar ages [79,80]. In support of this hypothesis, F1 hybrids of specialist species within the
San Salvador radiation show evidence of hybrid gene misregulation in approximately 10% of their differentially expressed genes, i.e. gene expression levels significantly different from parental expression levels, in whole larvae at 8 days post fertilization (dpf) and within craniofacial tissues at 17-20 dpf, respectively [81,82]. Although the fitness effects on hybrids are unknown, hybrid gene misregulation has been shown to affect hybrid viability and sterility in other systems [83–87] and misregulated genes in San Salvador Island pupfish species are enriched for developmental processes affecting ecological traits relevant to trophic specialization, including craniofacial morphology, muscle mass, and nitrogen metabolism [82]. However, the link between transgressive hybrid phenotypes and the extent of hybrid misexpression or genetic incompatibilities is still unknown.

**Does a large and stable fitness valley isolate scale-eating trophic specialization?**

Scale-eating (lepidophagy) is a particularly rare trophic niche among fishes and has evolved independently only 19 times across diverse marine, coastal, riverine, and lacustrine environments [65,88–90] and across ontogenetic stages from juveniles only to obligate scale-eating adults [91–94]. In particular, it is unusually rare among cyprinodontiform fishes: lepidophagy is only known to have evolved once within the San Salvador Island radiation and is thus separated by 168 million years of evolutionary time from the most closely related scale-eating specialists (within all three radiations of East African haplochromine cichlids [65]).

There are multiple scale-eating strategies and multiple hypotheses for the origins of scale-eating [88,90], supporting the observation that different fish lineages have different evolutionary potential to evolve scale-eating (e.g. relatively common in East African haplochromine cichlids, rare in cyprinodontiforms). For example, scale-biting pursuit predators, including the scale-
eating pupfish, wimple piranha (*Cataprin mento*), and cookie-cutter shark (*Isistius brasiliensis*), tear off scales and mucus from the side of their prey with perpendicular strikes using their entire oral jaws, whereas scale-rasping specialists, such as *Roeboides* tetras and khavalchor catfish (*Pachypterus khavalchor*), employ specialized externally protruding premaxillary teeth to dislodge scales from the sides of their prey [92,95,96]. However, in nearly all cases, scale-eaters are size-limited relative to their prey, unlike piscivorous fishes which generally grow much larger [97]. This difference suggests that the energetic payoff from scales is low relative the energetic demands of high-speed scale-eating or scale-rasping strikes. Scale-eating pupfish strike approximately once per minute in the wild, resulting in only a few scales and a mouthful of mucus per high-speed strike completed within 10 - 15 ms [98]. Overall, the rarity, high performance demands, and low caloric payoffs of scale-eating suggest that a wide and deep fitness valley isolates this niche and the necessary adaptive traits from all other ecological niches.

Consistent with this biomechanical prediction, our fitness data suggest that hybrids resembling scale-eaters suffered the highest fitness costs in growth and survival across all treatments, lakes, trait subsets, and time periods, but not in laboratory control aquaria fed only pellet foods. This provides an unexpected explanation for the rarity of scale-eating across cyprinodontiforms fishes if adapting to this specialized trophic niches requires multiple phenotypic traits that only provide fitness benefits in combination (i.e. fitness epistasis [12]). This is supported by existing population genomic and quantitative genetic evidence which suggests that adaptation to scale-eating is multifactorial and that multiple sources of adaptive variation from across the Caribbean contributed to trait diversification in this radiation [67,75,99]. The relevant adaptive alleles for scale-eating may be mismatched within hybrids,
such that all necessary morphological and behavioral traits never occurred within a single individual in our experimental hybrid populations. For example, aggressive and energetically demanding scale-eating behaviors may be highly deleterious in individuals with small oral jaws [76]. Although we included outbred backcrosses to scale-eaters in our hybrid populations, some generalist and molluscivore hybrid ancestry in these individuals may have impaired their scale-eating performance and behaviors so that scale-eating strikes were no longer energetically efficient. Conversely, algae-scraping with enlarged oral jaws or highly energetic strikes by hybrids resembling the generalist may also not provide a sufficient energy surplus for survival under field conditions.

Evidence for a holey adaptive landscape

One alternative metaphor to the fitness landscape is that the relationship between high-dimensional genotype or phenotype space and fitness more closely resembles a series of fitness ridges bypassing large regions of low fitness, or ‘holes’ [63,100]. The concept of holey adaptive landscapes emphasizes the possibility that two apparent fitness peaks in some low-dimensional phenotype or genotype space may be connected by a fitness ridge in higher-dimensional spaces. We tested this prediction by looking for fitness ridges connecting scale-eater phenotypes to generalist and molluscivore phenotypes across 500 random cross-sections in the 30-trait morphospace, rather than examining only a single two-dimensional cross-section. We found no evidence of a fitness ridge in any linear combination of these 30 traits connecting scale-eaters to other species in the radiation, instead nearly all survival curves appeared to decline in the region of the morphospace containing scale-eaters (Fig. 7). In one of the low-frequency treatments, survival curves showed no decline, but this treatment also included few hybrids closely
resembling the scale-eater phenotype and showed high survival overall (Fig. 7b). Thus, the entire volume of high-dimensional morphospace within our trait dataset containing the scale-eater phenotype appears to lead to a fitness minimum, or ‘hole’, isolating scale-eating phenotypes from the rest of the radiation. However, we failed to find evidence of any fitness ridges connecting these phenotypes to other species, one of the major predictions of the holey adaptive landscape hypothesis. Nonetheless, fitness ridges may still exist in genotype space or across unmeasured trait dimensions.

**Conclusion: on the origins of novelty during adaptive radiation on multiple fitness peaks**

The sensitivity of fitness landscape topography to the environment is rarely measured beyond a single population or fitness peak. Here we experimentally tested the effect of competitor frequency on a fitness surface spanning diverse hybrid phenotypes within a nascent adaptive radiation, comparable to phenotypic divergence spanning over 20 million years of Cyprinodontiform evolution (Fig. 1). Growth rate exhibited strong dependence on the frequency of similar phenotypes within and among field enclosures as predicted. However, hybrid survival showed no signal of frequency-dependence, challenging existing theory and previous experiments on a single population or species pair. Furthermore, major features of the fitness landscape, including generalist and molluscivore fitness peaks and a large fitness valley isolating scale-eaters, were strikingly consistent across lake environments, competitor frequency, and field exposure period. This challenges our existing view of empirical fitness surfaces as highly sensitive to environmental perturbation. Instead, multi-peak fitness landscapes spanning macroevolutionary levels of phenotypic disparity display both static and dynamic features across
space, time, and fitness proxy. These empirical results strengthen the connection between microevolutionary dynamics and static features of macroevolutionary fitness landscapes.

Materials and methods

Focal lakes

Laboratory breeding colonies of all three species were collected from two different focal lakes (lake 1: Crescent Pond; lake 2: Little Lake) in 2008, 2013, and 2014 using seine nets and hand nets while snorkeling. Crescent Pond is a 500 m x 60 m x 2 m deep hypersaline (40 ppt) lake containing a 0.1 – 0.3 m layer of fine silt on hard carbonate bedrock and is isolated from surrounding lakes by a small limestone ridge. Generalists and molluscivores are morphologically more similar in this lake than others due to a shorter nasal protrusion and exhibit a sister relationship across most of their genome [71,99]. There is only one other fish species in Crescent Pond, Gambusia hubbsi. Little Lake is a 2.3 km x 1 km x 4 m deep hypersaline (40 ppt) lake with a substrate consisting of Cerithium spp. snail shells and other molluscs and is connected to the larger interior Great Lake system through a sand bar. The lake contains two fish species Gambusia hubbsi and Atherinomorus stipes and the amphibious Kryptolebias marmoratus. Aquatic flora in both lakes is dominated by thick mats of three macroalgal species (Acetabularia crenulata, Batophora oerstedii, and Cladophora sp.) and wigeon grass (Ruppia maritima); only Little Lake is surrounded by red mangrove (Rhizophora mangle) forests [58].

Experimental hybrid populations

Independently for each focal lake, laboratories colonies of all three species were crossed in all directions resulting in both outbred F2 intercrosses and backcrosses (as originally described in
Hybrid populations from each lake were raised for an additional one to two generations in a common laboratory environment before backcrossing to independently sampled colonies of each parental species from each lake (collected in 2013 and 2014), resulting in an outbred population of F4 and F5 intercross and backcross hybrids for each lake, each originating from colonies of 10 – 20 wild-caught individuals from all three species.

Over six weeks, approximately 6,000 F4 and F5 hybrids were bred and raised in recirculating 151-liter aquaria at 5-10 ppt salinity, 25 – 29° C, on a diet of newly hatched brine shrimp, commercial pellet foods, frozen seafood, and dried seaweed. 95% water changes were performed every other day. Juvenile hybrids were size-sorted using 1/8” mesh bags identical to the field enclosure material resulting in a range of SL from 11.2 – 22.0 mm. Hybrids were packed in oxygen-permeable shipping bags (Kordon, Inc.) and shipped to San Salvador Island, Bahamas in packing crates (Rubbermaid ActionPacker Storage Box) as checked airline baggage.

Field enclosure experiments

Hybrids were temporarily held in concrete holding tanks with flow-through seawater at the Gerace Research Centre after arrival. Each individual was again size-sorted through 1/8” mesh bags, anesthetized in a solution of buffered MS-222 (Finquel, Inc.), photographed on the left, right, and dorsal sides using a Canon EOS 60D with an EF-S 60 mm f/2.8 USM macro lens mounted on a tripod with external flash, and finally injected subcutaneously in the left dorsal musculature with a 1 mm x 0.1 mm stainless steel sequential coded wire tag using a handheld multi-shot injector (Northwest Marine Technologies, Inc.). A fin-clip from the caudal fin of each fish was removed and stored with the archival tag in 100% ethanol at -20° C. Tagged hybrids were allowed to recover for at least 4 days in flow-through holding tanks at the Gerace Research
Centre to fully regrow their caudal fins while fed on a diet of newly hatched brine shrimp and commercial pellet foods.

Field enclosures were 3.6 m x 4.9 m rectangular fully-enclosed bags with a mesh size of 0.318 cm (Christiansen’s Net Company, Inc.) secured to either PVC pipe set in concrete (lake 1) or iron rebar hammered into the substrate (lake 2; Fig. 1). Two enclosures were deployed in the littoral zone of each lake after removing any debris, then the bottom mesh wall was weighted down with rocks and logs covered in macroalgae from the surrounding area and filled with benthic substrate, macroalgae, and wigeon grass from surrounding areas. Care was taken to avoid introduction of any adult fishes, but smaller pupfish and mosquitofish could still enter the enclosure through the mesh.

One enclosure in each lake was randomly selected as the high frequency treatment and the second enclosure was the low frequency treatment. Hybrids were individually selected for each treatment by eye, selecting more divergent phenotypes for the high-frequency treatments and selecting the most generalist-like hybrids for the low-frequency treatments. This resulted in reduced phenotypic variance and morphospace occupation within the low-frequency treatment within each lake. This also effectively reduced the frequency of scale-eater hybrids falling within the 95% confidence interval of parental scale-eater phenotypes. Hybrid densities within enclosures approximated the natural densities of 0.9% and 3% scale-eaters and 6 and 5% molluscivores in Crescent Pond and Little Lake, respectively [65].

In Crescent Pond (lake 1), tagged hybrids were released in large batches into the high-frequency enclosure on May 15th, 19th, and 28th and low-frequency enclosure on May 21st, 26th, and 28th, 2014, respectively. In Little Lake (lake 2), all tagged hybrids were released into the high-frequency enclosure on May 18th and into the low-frequency enclosure on May 25th and
Surviving hybrids were recovered from lake 1 on August 26th and 27th, 2014 after 3 months by carefully removing the substrate and sequentially lifting the entire mesh bottom, then photographed laterally and stored in 100% ethanol. To sample from a wider range of seasonal environments and recover the full time to reproductive maturity, surviving hybrids were recovered from lake 2 on April 28th and 29th, 2015 after 11 months in field enclosures. Tags were dissected from all survivors after preservation, read using a 100x tag-reading scope from Northwest Marine Technologies, Inc, and matched with archival tags to identify the survival status (0 or 1) of each tagged hybrid.

**Laboratory control**

Additional hybrids from each population (n = 199, Table S1) were raised in two 151-liter laboratory aquaria concurrent with the field experiment for 11 months. Control hybrids were raised on a diet of only commercial pellet foods to provide a uniform resource offering no advantages for specialized trophic morphology. Hybrids were fed once daily an amount of food that could be consumed in five minutes but not *ad libitum* and raised at 26 – 27°C in 5-10 ppt salinity (Instant Ocean) with weekly 95% water changes. Laboratory hybrids grew faster than fish placed in field enclosures and high densities within each aquarium population led to intense competition for food and high mortality rates, whereas survivors collected from field enclosures never reached maximum adult sizes or approached senescence. The day of each laboratory death was recorded, followed by removal of the tag to identify the pre-release photograph of that individual. Laboratory deaths were tracked from July 1st, 2014 until May 21st, 2015.

**Morphometrics**
Each hybrid used in field experimental \((n = 2,611)\) and laboratory control populations \((n = 199)\) was measured for 21 landmarks (Fig. S3) on both left and right lateral sides and 10 landmarks on the dorsal surface of the head plus a 4 mm size-standard grid for calibration of each image using TpsDig2 [101]. F1 lab-reared individuals of each parental species from each lake \((n = 236)\) were also measured in the same way. 27 linear distances and three angles were calculated from these landmarks and then averaged for both lateral sides, resulting in 30 trait measurements plus standard length. Traits were selected to capture phenotypic divergence in craniofacial morphology and body shape (Fig. S3).

Linear trait measurements and angles were then imported into R (R Development Core Team 2018) and size-corrected by taking the residuals from a standard major axis regression relative to log-transformed standard length (SL) for each trait in the entire hybrid pool and F1 lab-reared parental individuals from both lakes \((n = 110 ~\text{generalists}, 45 ~\text{molluscivores}, 81 ~\text{scale-eaters})\) using the sma function in the smatr package [103] in R. Standard major axis regression is appropriate when trait measurement error is present on the x-axis and the y-axis and is equivalent to the first principal component of phenotypic variance between these two axes [103]. Initial inspection of size-correction plots indicated that ordinary least squares (OLS) regression tended to overestimate the slope of the regression line, particularly for highly variable traits among parental and hybrid populations such as oral jaw length. However, our results were robust to OLS size-correction. No allometric scaling was observed among different species except for nasal protrusion distance and nasal protrusion angle, which exhibited no association with log-transformed SL and were not size-corrected. All size-corrected trait residuals and uncorrected nasal protrusion distance and angle were standardized to a standard deviation of one and mean of zero for comparisons across traits. There was no effect of standard length at introduction on
survival of hybrids (GLM logistic regression with effects of field enclosure and log-transformed SL: $P = 0.709$)

Visualization of fitness landscapes

Fitness landscapes were visualized in each enclosure by fitting thin-plate splines to the survival (binomial) or growth rate (normal) data for each hybrid using generalized cross-validation (GCV), which minimizes residual prediction error of the spline surface. Splines were estimated using the Fields package [59] in R. When over-fitting was apparent, restricted estimation of maximum likelihood (REML) was used to estimate the curvature of the spline instead of GCV (used for both survival landscapes in lake 2; REML estimation of splines was identical to GCV surfaces in lake 1). Growth rate was only examined in lake 1 due to the low number of survivors with growth rate data from lake 2 (Table S1).

We focused on two different cross-sections of the 30-dimensional hybrid morphospace. First, we examined selection on the two-dimensional linear discriminant morphospace maximizing phenotypic separation among the three parental species from both lakes using the lda function in the MASS package in R [104]. This morphospace provides a simple index of hybrid similarity to each of the three parental species, comparable to multivariate hybrid indices from similar studies of hybrid fitness between two species (e.g. [28,105]). Second, we examined the two major axes of nonlinear selection within the 30-dimensional morphospace using generalized projection pursuit regression for binomial data using the gppr function in the gsg package in R [62,106], following the original recommendations of [61]. This approach avoids the problematic quadratic assumptions of canonical rotation [107,108], particularly for highly nonlinear data such as ours, and enables visualization of the strongest axes of nonlinear selection.
within the dataset. We calculated the first two ridge axes using the gppr function with a binomial family of response distributions and then projected hybrid phenotypes onto each ridge axis by matrix multiplication. We performed generalized projection pursuit regression separately for lake 1 and lake 2 hybrid populations due to the large differences in survival (Table S1).

Generalized additive modeling

We formally tested for experimental treatment effects on fitness landscapes using generalized additive modeling in the mgcv package [109] in R. This modeling framework enables incorporation of spline terms into generalized linear models and comparisons of models containing spline, fixed, and random effect terms using AIC. For the survival data, we compared models with the fixed effects of treatment and lake and all combinations of univariate smoothing splines and thin-plate splines on both discriminant axes (Table 2) or both major ridge axes of selection estimated from generalized projection pursuit regression (Table 3). We also explored models allowing the thin-plate spline surface (i.e. the fitness landscape) to vary between lake environments using the ‘by’ term within the thin-plate function. Finally, we included models with a covariates including log-transformed standard length and distance measures of competitor frequency within each enclosure based on either Mahalanobis distance or nearest-neighbor Euclidean distance (see below). Models were compared using AIC. We examined a similar range of models for the growth rate data, calculated from the difference in log-transformed SL between pre-release photographs and surviving fish. However, we excluded lake 2 from all growth rate analyses due to the low number of survivors in this lake.

Analyses of frequency-dependent selection within enclosures
We used two approaches to measure the frequency of competitors within each enclosure. First, we calculated the Mahalanobis distance from each hybrid phenotype to the mean hybrid phenotype in the full 30-trait morphospace using the mahalanobis function in R. This distance estimates the disparity of each hybrid relative to the most abundant hybrid phenotypes while accounting for trait correlations. We also measured the frequency of competitors in the local region of morphospace surrounding each hybrid by calculating the sum of the Euclidean distance to the ten nearest neighbors in the full 30-trait morphospace, following the approach in [51]. We used the knn.dist function in the FNN package [110] to calculate the matrix of distances among all hybrid pairs.

Analyses of a fitness valley for scale-eater phenotypes

To evaluate the stability of a survival fitness valley near scale-eaters across all dimensions in our 30-trait morphospace, we estimated and visualized smoothing splines for survival from random subsets of the trait dataset. We first estimated a smoothing spline for survival relative to the discriminant axis (LD1 in Fig. 3) for all 30 traits separating parental scale-eater phenotypes from generalist phenotypes. We then randomly drew 15 traits from the dataset and recalculated the generalist-scale-eater discriminant axis and a new survival spline for 500 trait subsets using a custom script in R. This enabled alignment of each discriminant vector from generalist to scale-eater regardless of the subset of traits sampled. Each of the 500 discriminant vectors was then rescaled to the mean parental scale-eater phenotype, so that each hybrid could be scored on the same scale proportional to their phenotypic similarity to the scale-eater (i.e. 1 = full match). Survival splines were estimated separately for each field enclosure and plotted on a shared,
rescaled generalist-scale-eater discriminant axis to visualize the overall evidence for a fitness valley in the scale-eater region.

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Table 1. Directional selection gradients ($\beta$) and matrix of quadratic and correlational selection gradients ($\gamma$) on the two main ridge axes of selection estimated using generalized projection pursuit regression on the 30-trait morphological dataset for each lake (frequency treatments pooled based on model selection evidence). $\beta$ and $\gamma$ were estimated in separate regressions. Directional selection gradients in bold were significant in a one-way logistic regression model with fixed effects of ridge terms. Quadratic coefficients from the multiple regression model were doubled to estimate quadratic selection gradients [111]. However, please note that a quadratic approximation is inappropriate to model these highly nonlinear survival data [60].

| Lake 1 (Crescent Pond) ppr ridge axis | $\gamma$ |
|--------------------------------------|----------|
| $N = 1756$ A1 A2 |          |
Table 2. Model selection comparison of general additive models for survival and growth rate of hybrids placed in field enclosures in both lakes. The best supported model is indicated first. Notation is adopted from the mgcv package in R: $s(LD1)$ indicates a smoothing spline fit to discriminant axis one; $tps(LD1, LD2)$ indicates a thin-plate spline fit to the two discriminant axes; competitor distance indicates the Mahalanobis distance from each hybrid phenotype to the mean phenotype within the 30-trait morphospace while accounting for trait correlations. Growth rate models were only analyzed for lake 1 due to the low survival rates in lake 2.

| model                                                                 | AIC       | ΔAIC |
|-----------------------------------------------------------------------|-----------|------|
| survival ~ s(LD1) + s(LD2) + lake                                      | 2033.095  | -    |
| survival ~ tps(LD1, LD2) + s(LD1) + s(LD2) + lake                     | 2033.096  | 0.001|
| survival ~ tps(LD1, LD2) + lake                                       | 2042.247  | 9    |
| survival ~ tps(LD1, LD2) + lake + competitor distance                 | 2043.28   | 10   |
| survival ~ tps(LD1, LD2) + lake + treatment + logSL                   | 2044.308  | 11   |
| survival ~ tps(LD1, LD2) + lake + logSL                               | 2044.120  | 11   |
Table 3. Model selection comparison of general additive models for survival and growth rate of hybrids placed in field enclosures in both lakes for the first two major axes of selection (A1 and A2) estimated using generalized projection pursuit projection. The best supported model is indicated first. Notation is adopted from the mgcv package in R: s(A1) indicates a smoothing spline fit to ridge axis one; tps (A1, A2) indicates a thin-plate spline fit to the two ridges; competitor distance indicates the Mahalanobis distance from each hybrid phenotype to the mean phenotype within the 30-trait morphospace while accounting for trait correlations.
distance
log(growth) ~ s(A1) + s(A1) + logSL + treatment -3226.259 3.8
log(growth) ~ s(A1) + s(A1) + logSL + competitor distance -3185.980 44
log(growth) ~ s(A1) + s(A1) + logSL -3180.214 50

| lake 2          | survival ~ s(A1) + s(A2) | 108.7232 |
|-----------------|------------------------|---------|
| (Little Lake)   | survival ~ s(A1) + s(A2) + treatment | 110.4928 |
|                 | survival ~ s(A1) + s(A2) + tps(A1,A2) + treatment | 110.4925 |
|                 | survival ~ tps(A1,A2) + treatment | 110.4927 |
|                 | survival ~ A1 + A2 + treatment | 110.4921 |
|                 | survival ~ s(A1) + s(A2) + treatment + competitor distance | 110.6482 |

![Fig. 1](image)
Fig. 1 High- and low-frequency field enclosures and the associated benthic macroalgae communities inside each enclosure typical of surrounding littoral zone habitats in lake 1 (a,c: Crescent Pond) and lake 2 (b,d: Little Lake) after 3 month and 11 month field exposure periods, respectively.

Fig. 2
Fig. 2 Residual jaw length variation in a) allopatric Cyprinodontidae species (grey) diverging over 20 million years ago ([112]; data from [57]) and b) lab-reared F4/F5 hybrid populations (orange) measured from the field experiment in this study. The minimum and maximum residual upper jaw lengths across allopatric stickleback populations in the Pacific Northwest (PNW; data from Fig. 1 reported in [113]) are also included for comparison. Each a) species mean or b) individual F4/F5 hybrid is represented as tick marks on the x-axis plus density plots as an estimate of residual variation. Residuals were calculated from a linear regression of log-transformed lower jaw length on log-transformed standard length without mean and variance standardization for comparison on the same absolute scale across all three studies.

Fig. 3
Fig. 3 Survival fitness landscapes for hybrid populations in high- (first column) and low-frequency (second column) field enclosures and laboratory controls (third column). Thin-plate splines predict the probability of survival (heat color) across a single linear discriminant morphospace separating generalist and scale-eater phenotypes (x-axis: LD1) and generalist and molluscivore phenotypes (y-axis: LD2). Survivors in field enclosures are depicted in black relative to deaths over the 3-month and 1-year exposure periods, respectively. Laboratory control points are proportional to the number of days each hybrid survived within 151-liter aquaria. All hybrids are plotted within a shared linear discriminant morphospace calculated from lab-reared F1 individuals of parental populations in both lakes (first row: lake 1; second row: lake 2). 95%
confidence ellipses for each parental population in each lake are shown for generalists (I. grey), molluscivores (II. purple), and scale-eaters (III. red).
Fig. 4 Survival fitness landscapes for hybrid populations in high- (first column) and low-frequency (second column) field enclosures and laboratory controls (third column) across the two major ridge axes of selection within the 30-trait morphospace. Thin-plate splines predict the probability of survival (heat color) across the two major ridge axes associated with survival (Table S3-4), estimated separately for lake 1 (first row) and lake 2 (second row) hybrid populations using generalized projection pursuit regression. Survivors in field enclosures are depicted in black relative to deaths over the 3-month (first row) and 1-year (second row) exposure periods, respectively. Laboratory control points (third column) are proportional to the number of days each hybrid survived within 151-liter aquaria. 95% confidence ellipses for each
parental population in each lake are shown for generalists (I. grey), molluscivores (II. purple), and scale-eaters (III. red).
Fig. 5 Residual survival probability relative to the density of similar hybrid phenotypes in high- (first column) and low-frequency (second column) field enclosures in each lake. Residuals calculated from the survival probability not explained by the thin-plate splines estimated for hybrid phenotype in the discriminant morphospace (Fig. 2). The frequency of similar hybrid phenotypes was calculated for each hybrid from Mahalanobis distance to the mean hybrid phenotype within the full 30-trait hybrid morphospace account for trait correlations. Similar patterns were found using sum of the nearest-neighbor Euclidean distances to the ten most similar hybrid phenotypes in the full 30-trait morphospace (Fig. S5).
Fig. 6 Joint survival (first column) and growth (second column) fitness landscapes estimated across treatments and lake environments using generalized additive modeling. Thin-plate splines estimate the probability of \( a \) survival controlling for lake and treatment effects across all four enclosures and \( b \) growth rate controlling for treatment effects in lake 1 (lake 2 was excluded due to low survival rates). Thin-plate splines and smoothing splines (± 1 SE) are depicted within the linear discriminant morphospace separating generalist and scale-eater phenotypes (x-axis: LD1) and generalist and molluscivore phenotypes (y-axis: LD2) calculated from laboratory-reared individuals of parental populations in both lakes. 95% confidence ellipses show the location of generalist (grey), molluscivore (purple), and scale-eater (red) parental populations from lake 1 (small dashed line) and lake 2 (large dashed line). All hybrids are represented by points or tick marks on the x-axis and parental individuals are represented by tick marks on the upper margin.
Fig. 7 Spaghetti plots illustrate all possible fitness paths between generalist and scale-eater hybrid phenotypes in high- (first column) and low-frequency (second column) field enclosures. Each orange line depicts the relationship between survival and a random discriminant axis separating generalist and scale-eater phenotypes estimated from generalized cross-validation of a smoothing spline for 500 random subsets of 15 size-corrected traits (out of 30); the black line illustrates the smoothing spline estimated for a discriminant axis from all 30 traits, the grey lines illustrate the smoothing spline estimated for each subset. Each subsampled discriminant axis was rescaled to the mean parental scale-eater phenotype = 1 (red arrows).
Parental phenotypes are illustrated as black (generalist), purple (molluscivore), and red (scale-eater) tick marks on the x-axis.
**Table S1.** Sample sizes for high-frequency and low-frequency field enclosures and laboratory controls for each lake hybrid population.

| lake       | exposure period | high-frequency treatment survivors / total | low-frequency treatment survivors / total | control |
|------------|-----------------|--------------------------------------------|------------------------------------------|---------|
| Crescent Pond | 3 months        | 712 / 923; 77.1%                            | 662 / 883; 75.0%                          | 69      |
| Little Lake | 11 months       | 12 / 842; 1.4%                              | 10 / 819; 1.2%                           | 130     |
Table S2. Trait loadings on the two linear discriminant axes maximizing phenotypic separation among F1 lab-reared individuals of the three parental species from both lakes. Numbered landmarks correspond to illustrations of linear distances and angles in Fig. S3.

| trait         | trait LD1 | trait LD2 |
|---------------|-----------|-----------|
| cranialwidth  | -0.10     | 0.00      |
| innereyetosnout| -0.33     | 0.28      |
| suspensorium  | 0.74      | 0.37      |
| dorsalsnoutlen| 0.11      | -0.37     |
| adductorht    | 0.22      | -0.23     |
| jawlen        | 1.07      | -0.19     |
| ad2pect      | -0.07     | 0.16      |
| pmxlen        | 0.52      | -0.38     |
| foreeyewidth  | 0.17      | -0.42     |
| bodydepth     | 0.10      | -0.27     |
| dorsaltocaudal| -0.12     | 0.01      |
| headht        | 0.55      | -0.53     |
| analtocaudal  | -0.29     | -0.07     |
| caudalpedht   | -0.19     | 0.29      |
| pmx2add       | -0.76     | 0.49      |
| jaw2pect     | -0.10     | -0.07     |
| snoutlen      | 0.69      | -0.19     |
| foresnout     | -0.64     | 0.26      |
| eyewidth      | 0.01      | 0.04      |
| eyetosnout    | 0.29      | 0.71      |
| headwidth     | -0.11     | 0.23      |
| nose          | -1.06     | -1.61     |
| hindeyewidth  | -0.12     | 0.03      |
| eyeht         | -0.76     | -0.11     |
| topeyeangle   | 0.35      | 0.43      |
| lowereyeangle | -0.23     | 0.68      |
| nasalangle    | 0.49      | 0.01      |
| headlen       | 0.02      | -1.07     |
| bellylen      | -0.44     | 0.3       |
| pectinsertion | -0.19     | -0.19     |
| buccalwidth   | 0.16      | 0.3       |
Table S3. Trait loadings on the two ridge axes most strongly associated with survival probability within the Crescent Pond high and low-frequency field enclosures estimated using generalized projection pursuit regression. Numbered landmarks correspond to illustrations of linear distances and angles in Fig. S3.

| Trait                              | Trait                          | Ridge axis 1 (A1) | Ridge axis 2 (A2) |
|------------------------------------|--------------------------------|-------------------|-------------------|
| 1. nasal protrusion                | 1. nose                        | -0.02             | 0.06              |
| 2. interorbital width              | 2. cranialwidth                | 0.2               | 0.02              |
| 3. orbit to premaxilla             | 3. innereyetosnout             | 0.27              | 0.22              |
| 4. suspensorium length             | 4. suspensorium                | -0.24             | 0.23              |
| 5. dorsal facial length            | 5. dorsalsnoutlen              | -0.26             | -0.22             |
| 6. adductor height                 | 6. adductorht                  | 0.26              | -0.03             |
| 7. lower jaw length                | 7. jawlen                      | 0.21              | 0.42              |
| 8. subopercle to pectoral girdle   | 8. ad2pect                     | -0.11             | -0.26             |
| 9. premaxilla length               | pmxlen                         | -0.25             | -0.31             |
| 10. jaw joint to orbit             | foreeyewidth                   | 0.36              | 0.04              |
| 11. body depth                     | bodydepth                      | 0.15              | 0.09              |
| 12. dorsal to caudal distance      | dorsaltocaudal                 | -0.18             | 0.01              |
| 13. head height                    | headht                         | 0.17              | -0.35             |
| 14. anal to caudal distance        | analtocaudal                   | -0.02             | 0.1               |
| 15. caudal peduncle height         | caudalpedht                    | -0.11             | 0.17              |
| 16. lateral skull length           | pmx2add                        | -0.13             | -0.17             |
| 17. upper jaw to pectoral girdle   | jaw2pect                       | -0.05             | 0.15              |
| 18. lateral facial length          | snoutlen                       | 0.05              | -0.2              |
| 19. nasal length                   | foresnout                      | -0.02             | -0.06             |
| 20. horizontal orbit diameter      | eyewidth                       | -0.07             | -0.08             |
| 21. adductor to premaxilla         | eyetosnout                     | -0.09             | 0.14              |
| 22. max. neurocranium width        | headwidth                      | 0.05              | 0.13              |
| 23. orbital neurocranium width     | hindeyewidth                   | -0.12             | -0.02             |
| 24. vertical orbit diameter        | eyeht                          | 0.13              | 0.23              |
| 25. premaxilla to orbit angle      | topeyeangle                    | -0.23             | 0.14              |
| 26. premaxilla to adductor angle   | lowereyeangle                  | -0.16             | 0.24              |
| 27. nasal protrusion angle         | nasalangle                     | 0.06              | 0.18              |
| 28. neurocranium to premaxilla     | headlen                        | -0.2              | -0.15             |
| 29. orbit to anal fin insertion    | bellylen                       | -0.22             | 0.03              |
| 30. pectoral fin insertion width   | pectinsertion                  | -0.25             | -0.04             |
| 31. gape width                     | buccalwidth                    | -0.23             | -0.12             |
Table S4. Trait loadings on the two ridge axes most strongly associated with survival probability within the Little Lake (lake 2) high and low-frequency field enclosures estimated using generalized projection pursuit regression. Numbered landmarks correspond to illustrations of linear distances and angles in Fig. S3.

| trait                              | trait              | ridge axis 1 (A1) | ridge axis 2 (A2) |
|------------------------------------|--------------------|-------------------|-------------------|
| 1. nasal protrusion                | nose               | 0.18              | -0.02             |
| 2. interorbital width              | cranialwidth       | -0.07             | 0.14              |
| 3. orbit to premaxilla             | innereyetosnout    | -0.08             | 0.24              |
| 4. suspensorium length             | suspensorium       | -0.19             | 0.12              |
| 5. dorsal facial length            | dorsalsnoutlen     | 0.2               | -0.33             |
| 6. adductor height                 | adductorht         | 0                 | 0.01              |
| 7. lower mandible length           | jawlen             | -0.26             | 0.11              |
| 8. subopercle to pectoral girdle   | ad2pect            | -0.13             | 0.02              |
| 9. premaxillla length              | pmxlen             | 0.13              | -0.04             |
| 10. jaw joint to orbit             | foreeyewidth       | -0.05             | 0.19              |
| 11. body depth                     | bodydepth          | -0.03             | 0.15              |
| 12. dorsal to caudal distance      | dorsaltocaudal     | 0.03              | -0.17             |
| 13. head height                    | headht             | 0.04              | -0.07             |
| 14. anal to caudal distance        | analtdocaudal      | -0.07             | -0.18             |
| 15. caudal peduncle height         | caudalpedht        | -0.05             | -0.15             |
| 16. lateral skull length           | pmx2add            | -0.23             | 0.22              |
| 17. upper jaw to pectoral girdle   | jaw2pect           | 0.34              | -0.43             |
| 18. lateral facial length          | snoutlen           | -0.17             | 0.19              |
| 19. nasal length                   | foresnout          | 0.03              | -0.05             |
| 20. horizontal orbit diameter      | eyewidth           | 0                 | 0.08              |
| 21. adductor to premaxilla         | eyetosnout         | -0.23             | 0.22              |
| 22. max. neurocranium width        | headwidth          | 0.08              | -0.08             |
| 23. orbital neurocranium width     | hindeyewidth       | 0.15              | -0.12             |
| 24. vertical orbit diameter        | eyeht              | 0.25              | -0.19             |
| 25. premaxillla to orbit angle     | topeyeangle        | -0.51             | 0.47              |
| 26. premaxillla to adductor angle  | lowereyeangle      | -0.26             | 0.13              |
| 27. nasal protrusion anagle        | nasalangle         | 0.02              | 0.01              |
| 28. neurocranium to premaxilla     | headlen            | 0.21              | 0.02              |
| 29. orbit to anal fin insertion    | bellylen           | 0.03              | -0.12             |
| 30. pectoral fin insertion width   | pectinsertion      | 0.12              | 0.04              |
| 31. gape width                     | buccalwidth        | -0.19             | 0.01              |
Fig. S1 Log-transformed lower jaw length versus log-transformed standard length for

a) allopatric Cyprinodontidae species (black) and b) hybrid populations (one color per treatment) used in this study. The minimum and maximum upper jaw lengths of allopatric stickleback populations in the Pacific Northwest are also shown for reference (brown). Analyses of data published in [57] and Fig. 1 reported in [113].
Fig. S2 Histograms depicting the phenotypic variance of hybrid populations in high- (gray bars) and low-frequency (orange/blue) treatments in lake 1 (first row) and lake 2 (second row) on the first and second discriminant axes (LD1 and LD2 from Fig. 3).
Fig. S3 Morphometric landmarks indicating the 28 linear distances, 3 angles (25-27), and standard length (SL) for a) lateral, b) close-up of the craniofacial region, c) dorsal view, and d) close-up of the injected coded wire tag in the dorsal musculature including injection site (note different hybrid image used here for clarity). Numbers correspond to named traits in Tables 2-4. Lateral measurements were collected from pre-release photographs of both the left and right sides of each hybrid and averaged for all analyses. Dorsal view anteroposterior measurements (2,3,5,23,28) were also measured on each side and averaged for analyses. The 2 mm grid boxes in each image were used for calibration.
Fig. S4 Growth rate fitness landscapes for a) high-frequency and b) low-frequency treatments in lake 1 (Crescent Pond). Thin-plate splines predict the growth rate (heat color) across a single linear discriminant morphospace separating generalist and scale-eater phenotypes (x-axis: LD1) and generalist and molluscivore phenotypes (y-axis: LD2). Survivors in field enclosures are depicted in black relative to deaths over the 3-month exposure period. All hybrids are plotted within a shared linear discriminant morphospace calculated from lab-reared F1 individuals of parental populations in both lakes. 95% confidence ellipses indicate generalist (I. grey), molluscivore (II. purple), and scale-eater (III. red) regions of the morphospace.
Fig. S5 Residual survival probability relative to the density of similar hybrid phenotypes in high- (first column) and low-frequency (second column) field enclosures. Residuals calculated from the excess survival probability not explained by the thin-plate splines estimated for hybrid phenotype in the discriminant morphospace (Fig. 3). The frequency of similar hybrid phenotypes was calculated for each hybrid from the nearest-neighbor Euclidean distances to the ten most similar hybrid phenotypes in the full morphospace.
Fig. S6 Residual growth rate relative to the density of similar hybrid phenotypes in high- (first column) and low-frequency (second column) field enclosures. Residuals calculated from the growth rate variance not explained by the thin-plate splines estimated for hybrid phenotype in the linear discriminant morphospace (Fig. S5). The frequency of similar hybrid phenotypes was calculated for each hybrid from \( a-b \) the Mahalanobis distance to the mean hybrid phenotype within each treatment and \( c-d \) the sum of nearest-neighbor Euclidean distances to the ten most similar hybrid phenotypes in the full 30-trait morphospace.
Fig. S7 Histograms depicting the frequency of competitors in high- (gray bars) and low-frequency (orange/blue) treatments in lake 1 (first row) and lake 2 (second row) for both Mahalanobis distance (first column) and nearest neighbor distance to the ten nearest phenotypes (second column).