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Anthropogenic ecosystem fragmentation drives shared and unique patterns of sexual signal divergence among three species of Bahamian mosquitofish

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Abstract
When confronted with similar environmental challenges, different organisms can exhibit dissimilar phenotypic responses. Therefore, understanding patterns of phenotypic divergence for closely related species requires considering distinct evolutionary histories. Here, we investigated how a common form of human-induced environmental alteration, habitat fragmentation, may drive phenotypic divergence among three closely related species of Bahamian mosquitofish (Gambusia spp.). Focusing on one phenotypic trait (male coloration), having a priori predictions of divergence, we tested whether populations persisting in fragmented habitats differed from those inhabiting unfragmented habitats and examined the consistency of the pattern across species. Species exhibited both shared and unique patterns of phenotypic divergence between the two types of habitats, with shared patterns representing the stronger effect. For all species, populations in fragmented habitats had fewer dorsal-fin spots. In contrast, the magnitude and trajectory of divergence in dorsal-fin color, a sexually selected trait, differed among species. We identified fragmentation-mediated increased turbidity as a possible driver of these trait shifts. These results suggest that even closely related species can exhibit diverse phenotypic responses when encountering similar human-mediated selection regimes. This element of unpredictability complicates forecasting the phenotypic responses of wild organisms faced with anthropogenic change – an important component of biological conservation and ecosystem management.

Introduction
Human-mediated ecological change represents the primary driver of contemporary evolutionary change (Palumbi 2001; Hendry et al. 2008). While we know that ecological alteration affects evolutionary futures of species and populations, we know less about the predictability or the consequences of these changes (Hendry et al. 2011; Sih et al. 2011; Barrett and Hendry 2012; Palkovacs et al. 2012). Given the striking and ongoing reduction of biodiversity caused by human activities (Dirzo and Raven 2003; Barnosky et al. 2011), these gaps in our knowledge require rapid attention if we are to devise new conservation approaches that employ evolutionary principles (Dawson et al. 2011; Carroll et al. 2014; Smith et al. 2014).

The predictability of evolutionary change has been repeatedly illustrated by parallel and convergent evolutionary patterns in a diversity of taxa (e.g., sticklebacks: Schluter and McPhail 1992; cichlids: Kocher et al. 1993; Anolis lizards: Losos et al. 1998; Mahler et al. 2013; guppies: Endler 1980; Reznick and Endler 1982; Iguanid lizards: Rosenblum 2006). Discovery of these shared adaptive responses to similar selection regimes has served to demonstrate natural selection as a major and predictable driver of evolutionary change (Schluter and McPhail 1993; Losos 2010). Yet, when distinct evolutionary histories or chance events interact with selection, evolutionary divergence may follow nonparallel (unique) trajectories (Gould and Lewontin 1979; Gould 1989; Taylor and McPhail 2000). Evolutionary histories can vary among lineages for reasons...
other than divergent selection regimes, including genetic drift (Fisher 1930; Lande 1981), founder effects (Mayr 1942; Nei et al. 1975), and mutation-order selection (Mani and Clarke 1990; Rundell and Price 2009; Schluter 2009). These different agents of evolutionary change can generate interpopulation differences in genetic architecture that in turn affect the magnitude and trajectory of evolution, even when selection pressures are similar (Travisano et al. 1995; Schluter 1996; Taylor and McPhail 2000; Langerhans and DeWitt 2004; Blount et al. 2008; Langerhans 2010; Rosenblum and Harmon 2011; Kaeuffer et al. 2012).

How predictable and repeatable are human-induced phenotypic shifts in the wild? Anthropogenic environmental alteration is pervasive and global in extent (Vitousek et al. 1997; Palumbi 2001), and as a result, humans are generating similar selection pressures in human-modified environments for multiple populations within species, and for multiple species across geographic regions. We currently know little about the predictability of phenotypic divergence in response to anthropogenic change, or the extent to which different groups of organisms (i.e., different populations or species) respond to the same human-induced stressor(s) in shared or unique ways. When faced with similar and severe human-induced impacts, do closely related species generally exhibit similar phenotypic responses, or do species-specific responses predominate? Answering this question is critical for understanding human-mediated evolutionary change in the wild and how to manage it (Hendry et al. 2011; Carroll et al. 2014; Smith et al. 2014).

In this study, we investigate patterns of phenotypic divergence within three species of Bahamian mosquitofish (Gambusia spp.) subjected to hydrologic fragmentation, a widespread form of anthropogenic change (Montague et al. 1987; Pringle 2001; Nilsson et al. 2005). Fragmentation of Bahamian tidal creeks, a primary habitat of Bahamian mosquitofish, results in consistent ecological change, especially the local extirpation of piscivorous fishes (Layman et al. 2004, 2007; Heinen-Kay et al. 2014; Chacin et al. 2015). Because piscivorous fishes often exert predictable negative selection pressure on conspicuous coloration in their prey (Endler 1980, 1983; Houde 1997; Martin et al. 2014; Giery and Layman 2015), we focused our investigation on divergence in mosquitofish coloration. Given reduced predation pressure in fragmented ecosystems, we predicted that populations inhabiting these tidal creek habitats would exhibit more conspicuous coloration. Because fragmentation also appeared to cause shifts in the spectral environment, a recognized agent of color divergence, we additionally examined the effects of turbidity (Seehausen et al. 1997; Dugas and Franssen 2011) and spectral composition (Endler 1991; Fuller 2002; Morrongiello et al. 2010) in driving phenotypic variation. To test our prediction, we applied a ‘shared and unique’ statistical approach (Langerhans and DeWitt 2004) that addresses two main questions: (i) Does fragmentation, a shared contemporary ecological disturbance, affect coloration in Bahamian Gambusia, and (ii) Are these effects consistent among three distinct Gambusia species?

**Methods**

We conducted our study in tidal, mangrove-dominated wetlands in the Bahamas, locally known as ‘tidal creeks’. Tidal creeks comprise shallow, tidally influenced wetlands, typically fringed by red mangrove (Rhizophora mangle). Hydrologic flux in tidal creeks derives from tidal exchange with the ocean, as the creeks have small watersheds with karstic bedrock. Consequently, intact tidal creeks have salinities around 35 ppt, and biotic communities similar to other mangrove-dominated systems (Layman et al. 2004; Valentine-Rose et al. 2007). Major predators of Bahamian mosquitofish in these systems include redfin needlefish (Strongylnura notata), snapper (Lutjanus spp.), and great barracuda (Sphyraena barracuda) (Layman et al. 2004; Araujo et al. 2014; Giery and Layman 2015).

Fragmentation of Bahamian tidal creeks results from road construction, usually near creek mouths. These road blockages often lack flow-conveyance structures, such as bridges or culverts, resulting in disruption of hydrologic connectivity. By blocking tidal exchange between landward sections of tidal creeks and the ocean, fragmentation precipitates a suite of environmental changes in blocked portions of tidal creeks including the extirpation of piscivorous fishes, lower dissolved oxygen daily minima, increased daily maximum temperature, altered salinity, higher turbidity, higher pH, and increased nutrient availability (Valentine-Rose et al. 2007; Allgeier et al. 2010; Heinen-Kay et al. 2014; Chacin et al. 2015; Fig. 1A). Therefore, fragmentation per se does not directly result in environmental change in blocked tidal creeks; rather, disruption of hydrologic connectivity induces an ecological cascade culminating in novel environmental conditions to which fragmentation-tolerant organisms, such as mosquitofish, may adapt. While the exact dates of each fragmentation event in this study are not known, most roads that bisect tidal creeks in this study were built between the 1950s and 1970s during an expansion of infrastructure associated with intensive logging and coastal development in The Bahamas.

Bahamian mosquitofish (Gambusia spp.) are small, live-bearing fish with short generation times (1–2 generations per year) found in a wide range of aquatic habitats in The Bahamas including tidal creeks, blue holes, and inland freshwater marshes (Downhower et al. 2000; Langerhans et al. 2005, 2007; Araujo et al. 2014; Giery and Layman 2015). Three species of Bahamian mosquitofish reside in...
Our study area, the northern Bahama Archipelago (Heinen-Kay et al. 2014). Gambusia hubbsi inhabits the Great Bahama Bank, including the islands of Andros and New Providence. Gambusia manni also inhabits several islands of the Great Bahama Bank (not known to occur sympatri- cally with G. hubbsi in any tidal creek), including the islands of Eleuthera and Long Island. A third, unnamed species, Gambusia sp., appears limited to the Little Bahama Bank, including the islands of Grand Bahama and Abaco. These three species, endemic to The Bahamas, form a monophyletic group of closely related species with similar natural histories and morphologies, and are collectively referred to as Bahamian mosquitofish. Current data suggest G. hubbsi and G. manni are sister taxa, with Gambusia sp. diverging from the clade of G. hubbsi and G. manni approximately 1.7–4.8 million years ago, and G. hubbsi and G. manni diverging from one another approximately 1.3–2.4 million years ago (Heinen-Kay et al. 2014).

Adult male Bahamian mosquitofish express heritable variation in dorsal-fin coloration that ranges from intensely reddish orange to pale yellow (Fig. 1B; Figure S1; Langerhans 2006; Martin et al. 2014; Giery and Layman 2015). Coupled with findings that show heritable female preference for brightly colored dorsal fins (Heinen-Kay et al. 2015), the conspicuous display of dorsal fins during courtship, exclusive presence in males, ontogenic elaboration, and condition dependence suggest that they are secondary sexual characters subject to sexual selection via female preference (Andersson 1994). Bahamian mosquitofish also exhibit numerous, small, black spots on their body, caudal and dorsal fins, and there is some evidence for a role of black spots in crypsis (Endler 1983) and signaling (Brooks and Caithness 1995; Horth 2003; Martin et al. 2014) in Poeciliid fishes. Previous work with G. hubbsi found reduced dorsal-fin coloration and body spotting in populations experiencing higher predation risk, suggesting that predation pressure can influence elaboration of these traits in Bahamian mosquitofish (Martin et al. 2014).

We sampled 47 Gambusia populations comprising all three species across the Bahama Archipelago during March and April 2010 (Fig. 1C; Figure S2). Twenty-three of the collection sites were hydrologically fragmented by a road (Fig. 2A). At each site, we examined 3–8 adult males for color analysis. Fish were lightly anesthetized with eugenol (clove oil), laid on their left flank, and photographed on a Canon D10 digital camera. A ruler was included in each photo for measurement of body length (BL). Exposure and white balance were manually adjusted in Adobe Photoshop CS5 using a gray card exposure standard included in each photo. We sampled color using Adobe Photoshop CS5 in the RGB color space. RGB is a commonly employed color space for quantifying coloration in vertebrates (Stevens et al. 2007). The RGB color space defines color in a three-dimensional coordinate system where high values of R indicate red, G indicate green, and B indicate blue. We then generated an index measure of signal coloration ($R-G$)/($R+G$) following previous work by Endler (1990, 2012) and McKay (2013). This metric estimates color (hue) along an axis from red (+1) to yellow (0) and green (−1), and will be referred to as RG for the remainder of the paper. RGB measures were taken from 3 × 3 pixel samples from eight points on the dorsal fin and averaged for analysis. Four readings were located on the distal portion of the dorsal fin and four from the region near the fin insertion. The location of each measure was in areas of membrane between fin...
Color vision in fishes is taxonomically widespread and well developed (Loew and Lythgoe 1978; Lythgoe et al. 1994). Color vision in ecologically similar and closely related fishes suggest that Bahamian mosquitofish also possess well-developed color vision (e.g., Levine and MacNichol 1979; Archer et al. 1987; Körner et al. 2006; Watson et al. 2010). However, because RGB is not based on Gambusia visual systems it does not represent color as perceived by them or their predators. Determining whether the magnitude of variation detectible using our methodology is biologically relevant for our focal species would require additional information about their visual systems. Additionally, our methodology was not sensitive to UV light. However, UV coloration in Bahamian mosquitofish fins has been examined previously with spectrometry and has failed to reveal UV reflectance (Martin et al. 2014; Heinen-Kay et al. 2015).

We selected a set of environmental variables for measurement at each site based on their hypothesized roles as environmental agents likely to affect selection on coloration: predation pressure (Endler 1980, 1983; Horth 2004; Martin et al. 2014), turbidity (Seehausen et al. 1997; Dugas and Franssen 2011), and spectral environment (Endler 1991; Morrongiello et al. 2010; but see Schwartz and Hendry 2010). Fish communities were surveyed using roving diver surveys (Layman et al. 2004) and belt transects in areas where fish congregate such as mangrove fringes and pools (Faunce and Serafy 2007). Piscivore densities were calculated using the area sampled at each tidal creek, up to 1000 m² (Heinen-Kay et al. 2014). Fishes classified as piscivores included snappers (Lutjanus spp.), needlefishes (Strongylura spp.), groupers (Epinephelus spp.), great barracuda (S. barracuda), Atlantic tarpon (Megalops atlanticus), and jacks (Caranx spp.). Invertivores such as grunts (Haemulon spp.) and mojarra (Gerreidae spp.) were not classified as piscivores (Layman and Silliman 2002). Turbidity was measured in the field with an Oakton T100 turbimeter to the nearest nephelometric turbidity unit (NTU). To estimate the color of the aquatic habitat, we took underwater photos approximately 15 cm below the surface from a perspective perpendicular, and away from the shoreline at 1–4 locations within the study site and averaged for analysis (Fig. 2B). In each photo, the camera settings were maintained at 1/60th sec and F 5.6 exposure. The background coloration of the environment was estimated in Adobe Photoshop CS5 with a 3 × 3 pixel sample from the background of each photo. The dominant wavelength in each site was then measured in RGB color space. Color was estimated using two RGB color metrics: RG = (R−G)/(R+G) and GB = (G−B)/(G+B). As discussed above, RG measures the variation from red (1) to yellow (0) to green (−1), while GB measures variation from green (1) to cyan (0) and blue (−1). These two metrics allow us to capture major axes of environmental color variation observed in our study sites (Fig. 2).

The ‘snapshot’ nature of our sampling subjected our estimates to various extrinsic sources of environmental variation. We dealt with these sources of error in several ways. First, because radiance would be disproportionately affected by time of day or cloud cover, we restricted our analysis to parameters independent of total incident radiant flux (RGB ratios: Endler 2012). Second, we validated the temporal repeatability of our snapshot measures of water color using data available for a subset of sites (n = 13) following Lessells and Boag (1987). Underwater photographs from 2009, 2011, and 2012 show that our environmental color were temporally consistent (RG r = 0.45; GB r = 0.58; Table S1). Previous work has demonstrated temporal repeatability of piscivore density and turbidity in Bahamian tidal creeks (Heinen-Kay et al. 2014). Thus, overall our snapshot estimates of environmental parameters should provide meaningful estimates for comparison among sites.

Data analysis
Prior research suggests that fragmentation of tidal creeks largely results in consistent ecological changes (Layman...
et al. 2004; Araújo et al. 2014; Heinen-Kay et al. 2014). However, to be certain that fragmentation regime represented a consistent disturbance across the large spatial scale of this study, we evaluated the effect of fragmentation on the four putatively important drivers of variation in male coloration: water redness (hereafter $E_{RG}$), water blueness (hereafter $E_{GB}$), turbidity, and piscivore density across the study area. Turbidity and piscivore density were log-transformed for analysis (untransformed values in Table S2). For this analysis, we were primarily concerned with testing whether fragmentation resulted in consistent ecological change within the geographic range of each species (Figure S2). We used MANOVA in which environmental variables served as dependent variables, and fragmentation regime, species range, and the interaction between fragmentation and species range served as independent variables. If the effect of fragmentation (shared effects of fragmentation across the species ranges) is much greater than the effect of the interaction term (geography-dependent effects of fragmentation on environmental variables), then this suggests that each species has experienced a largely similar environmental shift subsequent to fragmentation, consistent with our assumption that fragmentation regime has a shared environmental effect across The Bahamas.

Our primary analysis of male coloration comprised an investigation into the shared and unique patterns of phenotypic divergence between fragmentation regimes across the three Gambusia species (Fig. 1D). To accomplish this, we conducted a mixed-model nested MANCOVA using five coloration measures as dependent variables: average dorsal-fin redness (hereafter $D_{RG}$), number of dorsal-fin spots, number of caudal-fin spots, and number of lateral body spots (all counts square-root transformed). Independent variables included BL (log-transformed size covariate), fragmentation regime (testing for a shared effect of fragmentation), species (testing for differences among species), and the interaction between fragmentation and species (to test for species specific, unique, effects of fragmentation), while island nested within species and population served as random effects.

The model was designed to test whether divergent species-level evolutionary histories of each population affected their response to shared ecological changes precipitated by ecosystem fragmentation. ‘Island’ and ‘population’ were random effects in our model to account for variation among replicate populations and because we wished to treat islands as random replicates for each species in order to focus on fragmentation effects across the three species. Random effects were included in MANCOVA using proc MIXED in SAS (SAS Institute, Cary, NC, USA). Details on this analytical method are found in Hassell et al. (2012) and Riesch et al. (2013). Significance tests for fragmentation, species, and their interaction were conducted with $F$-tests using restricted maximum likelihood and Kenward–Roger degrees of freedom adjustment (Kenward and Roger 1997). Remaining analyses were conducted in JMP (SAS) and Wilks’s partial $\eta^2$ was used to estimate the relative importance of model terms (see Langerhans and DeWitt 2004). We tested for heterogeneity of slopes among species, and between fragmentation regimes, by including BL in interaction terms with fixed main effects (Fragmentation and Species) and their interaction (Fragmentation $\times$ Species). Due to nonsignificance, we omitted these from our final MANCOVA analysis.

To interpret and visualize morphological divergence due to fragmentation, we (i) examined univariate patterns for each trait and (ii) derived canonical variates from independent variables in our MANCOVA model. First, we conducted separate analyses of variation for each phenotypic trait employing a univariate analog of the shared and unique model structure used in our MANCOVA analysis. Second, we examined patterns along canonical variates and inspected canonical loadings for each variate (pairwise correlations of traits and variates). These canonical loadings indicate the relative influence of each coloration variable on phenotypic divergence across factor levels (Fragmentation, Species, and Fragmentation $\times$ Species). Each canonical variate describes the multivariate linear combination of coloration variables that maximize differences between groups while minimizing differences within groups. To evaluate significant differences between fragmentation regimes within each species along the shared and unique phenotypic axes (canonical variates derived from ‘Fragmentation’ and ‘Fragmentation $\times$ Species’ terms in our MANCOVA, respectively), we conducted planned contrasts separately for each axis, using population as the unit of replication. Finally, to identify proximate relationships between environmental drivers and fragmentation-associated phenotypic divergence, we conducted a series of linear mixed models using population means of the shared and unique canonical variates derived from our mixed-model MANCOVA as dependent variables and environmental variables as independent variables. We tested for main effects of environmental color ($E_{RG}$ and $E_{GB}$), turbidity, and piscivore density on major axes of phenotypic variation. We explored pairwise interactions between piscivore density and other fixed terms in the model to examine possible predator-dependent effects of the spectral environment. However, all interactions were nonsignificant and therefore excluded from the final model. Species, and Island nested within Species (random term), were included to control for historical effects across species and islands.

**Results**

MANOVA analysis of the effects of fragmentation on putative environmental drivers of phenotypic variation showed that
tidal creek environments differed strongly according to fragmentation regime. We also detected environmental variation among the geographic ranges of each species (Table 1), although the effects of fragmentation on environmental variables were highly consistent across the three species ranges as indicated by the nonsignificant interaction term (Table 1). Wilks’s partial $\eta^2$ indicated that the majority of environmental variation was attributable to fragmentation regime rather than geography (i.e., a shared effect of fragmentation on environmental attributes within each species range; Table 1). Inspection of canonical variate loadings revealed that fragmentation was correlated with a reduction in piscivore density, a shift toward a more yellow/green spectral environment, and moderately increased turbidity (Table S3). Canonical loadings also suggest that the allopatric geographic ranges inhabited by each species differed in turbidity, environmental redisc, and piscivore density. However, further analysis suggested that these differences were minor. Pairwise multivariate contrasts between species revealed that only the ranges of Gambusia sp. and G. hubbsi differed in measured environmental factors (Wilks’s $= 0.59, F_{3,38} = 5.6, P = 0.001$) and follow-up univariate tests and pairwise comparisons (Tukey’s HSD) among species revealed that only turbidity, not environmental coloration or predator density, differed significantly among species ranges, with G. hubbsi inhabiting more turbid waters than Gambusia sp. (species means: Gambusia sp. = 0.48, G. hubbsi = 0.90, G. manni = 0.65).

In our analysis of male Gambusia coloration, we found all factors in our mixed-model MANCOVA exhibited highly significant effects (Table 2). Coloration was strongly affected by body size. The shared ecological disturbance, fragmentation, was clearly the most important term in our model other than body size, explaining a large portion of the observed phenotypic variation as reflected by partial $\eta^2$. Species, our historical variable, and the interaction between species and fragmentation, both explained less variance than the shared effect of fragmentation (Table 2). All species showed similar trajectories of phenotypic divergence along the shared axis; however, differences between fragmented and unfragmented populations were not significant for all species (Gambusia sp., $t = -1.5, P = 0.129$; G. hubbsi, $t = -0.5, P = 0.622$; G. manni, $t = -4.9, P < 0.001$).

| Effect                  | df  | F    | P    | Wilks’s $\lambda$ | Partial $\eta^2$ |
|-------------------------|-----|------|------|-------------------|------------------|
| Species range (SR)      | 8,76| 2.7  | 0.013| 0.61              | 0.22             |
| Fragmentation (F)       | 4,38| 3.5  | 0.015| 0.37              | 0.63             |
| F × SR                  | 8,76| 0.8  | 0.645| 0.86              | 0.07             |

Significant divergence along the first unique phenotypic axis also differed by species (Gambusia sp. $t = 3.4, P = 0.001$; G. hubbsi, $t = 1.1, P = 0.279$; G. manni, $t = -6.7, P < 0.001$). Finally, only one species differed along the second unique axis (Gambusia sp. $t = -2.9, P = 0.005$; G. hubbsi, $t = 0.0, P = 0.997$; G. manni, $t = -1.8, P = 0.074$).

Examination of canonical loadings revealed that all four traits ($D_{RG}$, caudal-fin spots, dorsal-fin spots, and lateral spots) were strongly correlated with one or more canonical axes (Table S4). Loadings for body size (BL) were positive and large for both dorsal-fin spots and caudal-fin spots (loadings = 0.63 and 0.94, respectively), indicating an overall increase in these trait values with the size of fish. The first canonical variate derived from the species term, the historical variable, was strongly correlated with dorsal-fin spots (loading = -0.81) and lateral spotting (loading = 0.50), while the second canonical variate was strongly correlated with dorsal-fin coloration (loading = 0.91). These results indicated that species differences in male coloration can be roughly characterized as: Gambusia sp. tend to have reddish-orange dorsal fins (high $D_{RG}$ values) with relatively few dorsal-fin spots, and high numbers of lateral spots; G. hubbsi have yellow-orange dorsal fins (low $D_{RG}$ values) with moderate numbers of dorsal-fin, and lateral spots; G. manni also has reddish-orange dorsal fins (high $D_{RG}$ values) with large numbers of dorsal-fin spots, and few lateral spots (Figure S3). Among shared and unique responses, fragmentation, the shared environmental factor, drove a strong shift in dorsal-fin spot number (loading = 0.79) reflecting a general pattern of reduced dorsal-fin spotting in fragmented populations, regardless of species identity (Fig. 3, Table S4). Unique axes of phenotypic divergence were correlated with fin coloration and dorsal-fin spots. The first unique canonical axis explained variation in fin coloration ($D_{RG}$ loading = 0.89). This reflected a pattern whereby species exhibited qualitatively different responses to fragmentation with respect to dorsal-fin color (Fig. 3). The second unique axis explained variation in coloration ($D_{RG}$ loading = -0.56) and dorsal-fin spotting (loading = 0.83); however, the amount of interaction variance explained by the second unique axis was small (<5%) and we do not consider it further.

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**Table 1. MANCOVA results for the effects of tidal creek fragmentation and geography on environmental conditions in 47 tidal creeks in The Bahamas.**

| Effect                  | df  | F    | P    | Wilks’s $\lambda$ | Partial $\eta^2$ |
|-------------------------|-----|------|------|-------------------|------------------|
| Species range (SR)      | 8,76| 2.7  | 0.013| 0.61              | 0.22             |
| Fragmentation (F)       | 4,38| 3.5  | 0.015| 0.37              | 0.63             |
| F × SR                  | 8,76| 0.8  | 0.645| 0.86              | 0.07             |

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**Table 2. Mixed-model MANCOVA results examining variation in coloration in male Gambusia collected from a series of fragmented and unfragmented tidal creeks in The Bahamas.**

| Effect                  | df  | F    | P    | Wilks’s $\lambda$ | Partial $\eta^2$ |
|-------------------------|-----|------|------|-------------------|------------------|
| Body length (BL)        | 3,457| 23.9 | <0.001| 0.28              | 0.72             |
| Species (S)             | 6,557| 17.3 | <0.001| 0.47              | 0.31             |
| Fragmentation (F)       | 3,457| 4.7  | 0.003| 0.13              | 0.87             |
| F × S                   | 6,557| 5.1  | <0.001| 0.73              | 0.15             |

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Univariate analyses mirrored our MANCOVA results; the effect of fragmentation on fin coloration depends on species identity while an overall effect of fragmentation on dorsal-fin spotting is evident although only nearly significant in the univariate analysis ($P = 0.074$; Tables S5 and S6). Overall, the three species exhibited the following phenotypic patterns across fragmentation regimes: *Gambusia* sp. exhibited little-to-no difference in dorsal-fin spotting and redder dorsal-fin coloration (high $D_{RG}$ values) in fragmented sites; *G. hubbsi* were similar in dorsal-fin spotting and fin coloration among fragmented and unfragmented sites; *G. manni* exhibited fewer dorsal-fin spots and more yellow dorsal-fin coloration (low $D_{RG}$ values) in fragmented sites.

Among environmental variables, turbidity and water coloration exhibited substantial evidence as drivers of phenotypic divergence (Table 3). Fin coloration showed evidence of a negative effect for turbidity and a positive association with environmental redness ($E_{RG}$). Turbidity was also negatively correlated with dorsal-fin spots although the relationship was merely suggestive ($P = 0.097$). Finally, environmental blueness ($E_{GB}$) and piscivore density failed to show any effect on the traits examined (Table 3).

**Discussion**

We document evidence for human-mediated phenotypic divergence among wild populations of Bahamian mosquitofish. This divergence exhibits evidence of both shared and unique responses among the three species. Below we discuss our evidence for parallel and nonparallel phenotypic divergence, putative mechanisms driving the observed divergence, and the implications of anthropogenically mediated change for the ecological and evolutionary futures of wild populations.

**Human-mediated divergence**

Hydrologic fragmentation of coastal wetlands across the Bahama Archipelago has driven replicated patterns of morphological divergence among three, closely related, species. While restricted to dorsal-fin spots, this shared, divergent response is consistent with our initial expectation that the drastic environmental changes associated with tidal creek fragmentation affects coloration consistently, among, and within species. Proffering an adaptive scenario for the reduction of melanic spotting of dorsal fins in mosquitofish occupying fragmented habitats is difficult without a better understanding of their function (crypsis, signaling, both, or other). However, given the negative correlation between dorsal-fin spots and turbidity suggested by our univariate analysis, reduction in fin spots due to fragmentation may reflect a consequence of reduced effectiveness of visual signals in turbid environments. Melanic patterns have been hypothesized to increase the conspicuity of color signals by increasing contrast between monochromatic backgrounds and the focal signal feature (Hailman 1979). Therefore, if turbidity relaxes positive selection on contrast, we might expect this signal modifier to fade or become less elaborate over time. Indeed, such a response has been documented in other study systems, lending support to this explanation (Victoria cichlids: Seehausen et al. 1997; Maan et al. 2010; blue-fin killifish: Fuller 2002). Regardless of the mechanism, which requires future investigation, we uncovered strong evidence for a shared decrease in dorsal-fin spotting.

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**Table 3.** Summary of univariate mixed-models examining the associations between population-level environmental variation and phenotypic divergence axes from the mixed-model MANCOVA (bold text emphasizing significant relationships).

| Divergence axis | Source | Estimate | df | F | P |
|-----------------|--------|----------|----|---|---|
| Shared $CV_s$ ($D_{RG}$) | $E_{RG}$ | -0.25 | 1,40 | 1.8 | 0.187 |
| | $E_{GB}$ | 0.06 | 1,40 | 0.6 | 0.435 |
| | Predator | -0.00 | 1,14 | 0.0 | 0.844 |
| | Turbidity | -0.05 | 1,40 | 2.9 | 0.097 |
| | Species | 2.2 | 26.0 | 0.022 |

| Unique $CV_u$1 ($D_{RG}$) | $E_{RG}$ | 0.59 | 1,38 | 5.0 | 0.031 |
| | $E_{GB}$ | 0.02 | 1,39 | 0.0 | 0.883 |
| | Predator | -0.00 | 1,40 | 0.0 | 0.827 |
| | Turbidity | -0.13 | 1,38 | 8.1 | 0.007 |
| | Species | 2.4 | 0.6 | 0.603 |
due to fragmentation in Bahamian mosquitofish (Fig. 3, Table 2).

Despite the similarity of environmental shifts induced by fragmentation and the recent evolutionary differentiation of these species (1–5 my, Heinen-Kay et al. 2014), results also revealed an unexpectedly strong signature of species-contingent phenotypic divergence due to anthropogenic habitat alteration: Dorsal-fin coloration in Gambusia sp. and G. mannii differed between fragmentation regimes (in opposite manners), while G. hubbsi exhibited no such differentiation (Fig. 3). Below we discuss two questions that emerge from these results.

Why did fragmentation not influence dorsal-fin color in G. hubbsi? This finding was unexpected, especially in light of previous research from Andros Island blue holes showing that G. hubbsi fin coloration has adaptively diverged repeatedly between predation regimes (e.g., Langerhans et al. 2007; Langerhans 2010; Martin et al. 2014). One line of evidence points toward dorsal-fin color being a less important sexual signal within tidal creeks for G. hubbsi compared to the other two species. While the effect of fragmentation on tidal creek environments is highly consistent across the study area (Table 1), habitats within the range of G. hubbsi are relatively turbid, regardless of fragmentation regime (Table S2). Perhaps signal transmission of orange dorsal fins is generally poor in these turbid tidal creeks, with fragmentation having an effectively small influence on this signal's transmission environment. Future work is needed to test this hypothesis.

What generated dissimilar divergence trajectories in G. mannii and Gambusia sp. when selective environments appeared similar? Explanations of divergent responses to similar selection regimes typically rely on cryptic variation in selection regimes (e.g., sensory drive and female preference), differences in genetic architecture, or the chance in selection regimes (e.g., sensory drive and female preference). That is, one adaptive strategy to mitigate signal disruption is to increase signal investment (increased red color: $D_{\text{RC}}$), presumably increasing transmission distance and the probability of accurate reception (e.g., guppies: Endler and Houde 1995; perch: Kekäläinen et al. 2010; pygmy perch: Moringiello et al. 2010; rainbowfish: Kelley et al. 2012; shiners: Dugas and Fransson 2011; stickleback: Reinchen 1989). Such a compensatory response might resemble the increased fin redness of Gambusia sp. populations inhabiting fragmented, turbid habitats (Fig. 3). Another strategy of potentially equal adaptive value might be to reduce signaling investment in antagonistic environments to minimize costs of ineffective signals (Järvenpää and Lindström 2004; Heuschele et al. 2009). This would be expected if fitness benefits accrued through restraining signal investment are similar to or greater than that accrued through increasing signal effort. Redirecting energy or pigments to more effective signals or viability could adaptively recover negative fitness consequences of reduced signaling effectiveness.

Such a strategy might explain the general pattern of reduced dorsal-fin spotting in fragmented populations, the overall weakly colored, yellow dorsal fins in the turbid tidal creeks inhabited by G. hubbsi, and the reduced dorsal-fin redness within fragmented tidal creeks in G. mannii (Fig. 3). Evidence for such a strategy in the literature is not as strong; however, a few examples do illustrate a negative relationship between signal intensity and turbidity (Victoria cichlids: Seehausen et al. 1997; Maan et al. 2010; bluefin killifish: Fuller 2002).

Overall, both shared and unique patterns of divergent phenotypic responses among Gambusia species to fragmentation suggest that sexual signals are shifting due to changes to the spectral environment rather than predation risk. While compelling, these findings leave many questions unanswered. For example, what aspects of genome architecture differ among our study species? What are the contributions of genetic and environmental sources to phenotypic responses? And are the environmental shifts experienced by each species similar enough to be considered ‘shared’? We have simplified our analysis of environmental drivers by limiting the set of environmental factors to predation and spectral environment, yet fragmentation precipitates a range of ecological changes. One potentially important variable we do not address here is geographic variation in the effect of fragmentation on salinity. The Bahama Archipelago shows a strong north–south precipitation gradient (Figure S4) causing hydrologically isolated ecosystems, such as fragmented tidal creeks, to become hyposalinic in the north and hypersalinic in the south (Figure S5). We suspect this apparent interaction between geography and fragmentation might indirectly affect other factors important for signal production such as the types and quantity of carotenoids in mosquitofish prey (Cifuentes et al. 2001; Oren 2005). Such variation in carotenoid availability could underlie the unique fin coloration responses to fragmentation we observe in our dataset (Grether et al. 1999; Craig and Foote 2001; Grether 2005) and deserves further study.

Conservation implications

At least half of the world’s major river drainages are dammed (Nilsson et al. 2005), and a large number of smaller drainages are heavily affected by hydrological fragmen-
tation. For example, in the United States alone, 2.6 million impoundments account for an estimated 20% of existing inland standing waters by area (Smith et al. 2002). Fragmentation impacts to coastal habitats are also frequent and heavy: even in the extensive marshlands of the southeastern United States, fragmentation directly affects approximately 84 000 ha (20%) of coastal wetlands (Montague et al. 1987). Damming is frequently cited as one of the most detrimental anthropogenic threats to freshwater faunal diversity, having extremely deleterious effects on endemic and anadromous species, especially fishes and invertebrates (March et al. 2003; Poff et al. 2007; Liermann et al. 2012; Cooney and Kwak 2013). Despite the global extent and huge ecological impacts of hydrologic disruption on biodiversity the number of studies investigating its impacts from an evolutionary perspective are remarkably few (e.g., Fransson 2011; Aguirre et al. 2013; Fransson et al. 2013; Cur- eton and Broughton 2014; Heinen-Kay et al. 2014; Santos and Araújo 2015). In each of these studies, species persisting in altered habitats show significant phenotypic change. Typically, trait shifts are related to viability (e.g., locomotion and foraging), but changes in sexual traits are also apparent (Heinen-Kay et al. 2014). In concert, these findings inform a general argument that hydrologic fragmentation and the novel ecosystems generated as a consequence drives widespread evolutionary divergence in a variety of wild fish populations.

Predicting the fate of wild populations subjected to such a pervasive ecological disturbance is perhaps the most pressing conservation challenge (Rice and Emery 2003; Stockwell et al. 2003; Carroll and Fox 2008; Hendry et al. 2011; Car- roll et al. 2014; Smith et al. 2014). Yet, as our results and others show (e.g., Fransson et al. 2013a), the direction and magnitude of phenotypic divergence may be difficult to predict, even when the focal species are closely related, ecologically similar, and subjected to seemingly similar selection regimes. These general conclusions, formed from this and other research exploring diverse responses to selection regimes (e.g., Kaeuffer et al. 2012) suggest a strong tendency for populations to diverge in parallel and nonparallel ways due to historically contingent factors. This fact complicates actions directed toward understanding and managing human-mediated evolutionary change in the wild (Carroll et al. 2014). Adequately capturing the predictably of phenotypic change required for evolutionarily informed conservation actions will require well-replicated studies of broad taxonomic, ecological, and geographic scope to uncover historically contingent influences on evolutionary trajectories.

Conclusion

Sexual signals play an important role in a diversity of biological functions and evolutionary processes (Smith and Grether 2008). Habitat alteration poses a significant threat to organisms because it can disrupt signal transmission during mate evaluation (e.g., Järvenpää and Lindström 2004; Engström-Öst and Candolin 2006; Candolin et al. 2007) and species recognition (e.g., Seehausen et al. 1997; Fisher et al. 2006; Walters et al. 2008). Consequently, habitat alteration represents an important threat to a diversity of species that implement chemical, auditory, electrical, and visual signals in their communication systems (van der Suijs et al. 2011; Rosenthal and Stuart-Fox 2012). Yet, the rapidity with which sexually selected traits can evolve (Endler 1986; Hoekstra et al. 2001; Kingsolver et al. 2001; Svensson and Gosden 2007), and the pervasiveness of anthropogenic environmental alteration, suggests that human-mediated divergence of sexual signaling systems is common among species that persist in altered ecosystems. Consequently, the future of wild populations may not depend on whether phenotypic change will occur. Rather, their fates will rely on whether the trajectory and magnitude of change is sufficient to ameliorate phenotype–environment mismatches and contribute to population persistence (Bell and Collins 2008; Barrett and Hendry 2012; Carroll et al. 2014; Zimova et al. 2014).

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Data archiving statement

Data for this study are available from the Dryad Digital Repository: http://dx.doi.org/10.5061/dryad.10b95.

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**Supporting Information**

Additional Supporting Information may be found in the online version of this article:

**Figure S1.** Male Bahamian mosquitofish (*Gambusia* spp.) show marked interpopulation variation in dorsal fin coloration, as well as spotting on fins and flanks.

**Figure S2.** Map of the upper Bahama Archipelago (inset) and its position in relation to the US and Greater Antilles.

**Figure S3.** Differences in male coloration among Bahamian mosquitofish species.

**Figure S4.** Latitudinal precipitation gradient in the Bahama Archipelago.

**Figure S5.** The relationships between latitude and salinity in fragmented and unfragmented tidal creeks.

**Table S1.** Repeatability analysis of environmental measures for a subset of fragmented and unfragmented tidal creeks on Abaco Island measured between 2009 and 2012.

**Table S2.** Untransformed means for *Gambusia* morphology and environmental variables among fragmented and unfragmented habitats.

**Table S3.** Loadings of environmental variables on canonical axes derived from MANOVA of effect of fragmentation and geography.

**Table S4.** Loadings of canonical variates derived from the Allometric (SL), Species (Historical), Fragmentation Regime (Shared), and Species × Fragmentation Regime (Unique) terms of the mixed-model MANCOVA examining variation in male Bahamian *Gambusia* coloration (important loadings bolded for emphasis).

**Table S5.** Summary results of shared and unique effects of fragmentation on phenotypic traits in Bahamian mosquitofish (*Gambusia* spp.).

**Table S6.** Least-squares means and standard errors for each significant term in our univariate analysis of shared and unique effects of fragmentation.