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THE INFLUENCE OF AN INNOVATIVE
LOCOMOTOR STRATEGY ON THE PHENOTYPIC
DIVERSIFICATION OF TRIGGERFISH
(FAMILY: BALISTIDAE)

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Innovations in locomotor morphology have been invoked as important drivers of vertebrate diversification, although the influence of novel locomotion strategies on marine fish diversification remains largely unexplored. Using triggerfish as a case study, we determine whether the evolution of the distinctive synchronization of enlarged dorsal and anal fins that triggerfish use to swim may have catalyzed the ecological diversification of the group. By adopting a comparative phylogenetic approach to quantify median fin and body shape integration and to assess the tempo of functional and morphological evolution in locomotor traits, we find that: (1) functional and morphological components of the locomotive system exhibit a strong signal of correlated evolution; (2) triggerfish partitioned locomotor morphological and functional spaces early in their history; and (3) there is no strong evidence that a pulse of lineage diversification accompanied the major episode of phenotypic diversification. Together these findings suggest that the acquisition of a distinctive mode of locomotion drove an early radiation of shape and function in triggerfish, but not an early radiation of species.

KEY WORDS: Adaptive radiation, balistiform, correlated evolution, disparity, generalized least squares, geometric morphometrics, locomotion, Tetraodontiformes.
Functional novelty is thought to spur diversification in eco-
morphological traits as a consequence of creating ecological
opportunity—the chance for a lineage to exploit new niches
(Schluter 2000; Gavrilets and Losos 2009; Yoder et al. 2010).
However, expectations of how functional innovations influence
the tempo of lineage, morphological, or functional diversification
are less clear. The ecological model of adaptive radiation predicts
a burst of morphological disparity and lineage diversification im-
mEDIATELY following functional innovation as lineages fill newly
available niches, then a slowing of diversification after niches fill
(Schluter 2000; Rabosky and Lovette 2006a; Rabosky and Lovette
2006a,b) or a consequence of reaching functional limits or hard
boundaries in morphospace (Blomberg et al. 2003; Freckleton and
Harvey, 2006; Harmon et al. 2010). However, some recent studies
have failed to find strong links between functional innovations and
phenotypic diversification (Slater et al. 2010) or species richness
(Alfaro et al. 2009a,b). Understanding why the consequences of
functional innovations vary so much among case studies repre-
sents a fundamental goal in macroevolutionary biology.

Within the more than 31,000 living fish species (Eschmeyer
2010) exists a tremendous diversity in locomotor morphology
and behavior including fascinating modes of swimming, gliding,
walking, crawling, and underwater flight. This variation presents
rich opportunities for the study of diversification following func-
tional innovation in locomotion strategies. However, despite an
explosion of interest in fish hydrodynamics enabled by increas-
ingly accessible three-dimensional (3D) flow visualization (e.g.,
Mittal et al. 2006; Tytell 2006; Lauder and Madden 2006; Dabiri
2009; Lauder 2010), there have been few comparative evolution-
ary studies of locomotor innovation and phenotypic diversifica-
tion. These studies have generally focused on the integration of
locomotion with other functional systems such as feeding (Rice
and Westneat 2005; Higham 2007; Collar et al. 2008) and have
found functional changes in locomotion to potentially influence
changes throughout the entire bauplan of a fish lineage. As lo-
comotion is central to a fish’s ecology, the question of whether
evolutionary change in locomotor structure following major func-
tional shifts follows the predictions of macroevolutionary theories
such as key innovation or the ecological theory of adaptive radia-
tion can be raised.

Triggerfish possess one of the more distinctive swimming
modes within fish, using coupled oscillation or undulation
of paired median fins to achieve forward thrust. This mode
of locomotion, termed balistiform swimming (Sfakiotakis et al.
1999), is found in several fish groups including flatfish
(Pleuronectiformes) and filefish (Monacanthidae), although the
greatest diversity in fin shape is found within triggerfish.
Shape diversity spans deep-bodied, large-finned species such as
Melichthys niger to shallow-bodied species with high aspect
ratio fins such as Canthideris maculata. Diversity in fin and
body shape is thought to partially reflect divergent locomotor
strategies (Lighthill and Blake 1990a; Wright 2000). Median
fin oscillators rely upon strongly tapering high aspect ratio fins
whereas undulators possess anteroposteriorly elongate fins of
more uniform depth (Lighthill and Blake 1990a,b,c,d). However,
triggerfish are not limited to this simple dichotomy of forms, and
a substantial diversity of intermediate morphologies are found
within triggerfish that have not been quantitatively examined in
hydrodynamic studies (see Wright 2000).

Triggerfish represent a good case study from which to explore
the diversification dynamics associated with functional innova-
tion for several reasons. Theoretical models assume symmetry
between dorsal and anal fins (Lighthill and Blake 1990a,b,c,d;
Wright 2000; Korsmeyer et al. 2002; Loofbourrow 2009), im-
plying that morphological evolution of the fins should be tightly
correlated. Because the ratio of fin to body depth has been demon-
strated to have a strong influence of the overall drag (Lighthill
and Blake 1990a,d), fin and some aspects of body shape evolu-
tion should also be highly correlated. These predictions about
morphological evolution in triggerfish have never been tested.
Moreover, the recent reconstruction of a phylogeny and chron-
ogram from this group (Alfaro et al. 2007; Dornburg et al. 2008)
provides the framework to ask how the evolution of a major func-
tional innovation (balistiform locomotion) influenced subsequent
cladogenesis and phenotypic diversification.

Here we use a suite of phylogenetic comparative methods
to ask two classes of questions about the influence of a novel
form of locomotion on the evolutionary dynamics of triggerfish
morphology: (1) Do triggerfish median fin and body shapes ex-
hibit strong patterns of correlated evolution as would be expected
if balistiform locomotion constrains their morphological evolu-
tion, and (2) is there evidence for a rapid or adaptive character to
the diversification of triggerfish lineages, morphology, and func-
tion as would be expected if the evolution of their distinctive
locomotor type catalyzed an adaptive radiation? To address those
questions, we characterize fin shape, aspect ratio, and body shape
in two-thirds of all extant triggerfish species and analyze patterns
of morphological and functional diversification in the context of
a time-calibrated molecular phylogeny.

**Methods**

To investigate patterns of correlated evolution between compo-
teents of the triggerfish locomotor system and infer the historical
processes that have helped to shape their standing biodiversity,
we gathered data on morphology, function, and phylogeny. We
measured morphological and functional diversity among species,
constructed morphospaces to identify the most important axes of
variation, and quantified the relationship between phylogeny and
fin and body shape morphospace to look for evidence of lineage
clustering within the phylomorphospace (Sidlauskas 2008).
QUANTIFYING PHENOTYPIC DIVERSITY OF TRIGGERFISH

We photographed 270 adult specimens comprising 26 species of triggerfish (Appendix S1) using 8–10 megapixel digital cameras. All specimens were photographed facing left, with individual photos taken of each fish’s body in addition to its dorsal and anal fins. Caudal fins could not be photographed consistently due to frequent damage and were not included in shape analysis. For each species, we aimed to digitize between 5 and 20 individuals, though for both the rare Red Sea endemic _Rhinecanthus assasi_ and the West African _Balistes punctatus_ only two individuals were available.

We used landmark-based geometric morphometrics (Bookstein 1991; Adams et al. 2004; Zelditch et al. 2004) to capture the shapes of the fins and body. We placed four total landmarks on the origin and insertion of the fin base and tips of the anterior and posterior rays and used sliding semi-landmarks (Bookstein 1997) to describe the curvature present along the distal margin (Fig. S1). The sliding semi-landmarks were placed by outlining each distal fin margin and resampling the curve to contain eight sliding semi-landmarks. We also placed one sliding semi-landmark along the fin base at the midpoint between the fin origin and insertion for a total of 13 fin landmarks (four fixed, nine sliding semi-landmarks). To quantify body shape, we used 27 homologous landmarks (Fig. S2) and positioned five semi-landmarks by eye to better define the curves of the body at the midpoint of the following pairs of landmarks: (1) 3 and 9, (2) 10 and 12, placed along the body, (3) 14 and 15, placed along the fin-ray insertion margin, (4) 17 and 18, placed along the body, and (5) 21 and 22. All of the landmarks were placed using TpsDIG2 (Rohlf 2006).

We quantified a functional property of each fin by calculating its aspect ratio (Lighthill and Blake 1990a; Wright 2000). Each fin was outlined in TpsDIG2 (Rohlf 2006) and the area and semispan were computed (Fig. S3). Wright (2000) defined the semispan as the length from the tip of the fin to the flapping axis, drawn perpendicular to the x-axis (Fig. S3). Following Wright (2000), we computed fin aspect ratios for each individual as the semispan squared, divided by the total fin area. We also repeated this analysis defining aspect ratio according to Walker and Westneat (2002) as two times the length of the fin’s leading edge squared, divided by the total fin area (Fig. S3). Mean aspect ratio values for each species were used in further analysis. The potential for biased error driven by allometry was assessed for both shape and mechanical data by testing for a correlation between fin or body centroid size and partial warp scores or aspect ratios using multivariate regression (Monteiro 1999) in TPSRegr (Rohlf 2003).

DETERMINING THE MOST IMPORTANT AXES OF SHAPE VARIATION

To quantify the most important axes of shape variation for each fin or body shape dataset, we used a Procrustes fit (Rohlf and Slice 1990; see also Zelditch et al. 2004) to generate a mean shape for each species and remove variation due to scaling, rotation, and translation (Zelditch et al. 2000). We then used a second Procrustes fit of the 26 consensus configurations combined with a relative warps (RWs) analysis (Rohlf 1993) in TpsRelw version 1.46 (Rohlf 2007) to generate a morphospace (a series of orthogonal eigenvectors describing the major axes of shape variation). Because we set alpha to 0, the RW analysis was mathematically equivalent to a principal components analysis of the Procrustes coordinates (Rohlf 1993). In subsequent analysis, all RW scores were multiplied by one hundred to accommodate easier interpretation (following Sidlauskas 2008).

DIVERGENCE TIME ESTIMATION

To estimate the phylogeny and timing of evolutionary divergences in triggerfish, we assembled DNA sequence data for 28 balistid species (Table S1), representing 11 of 12 genera, and comprising approximately two-thirds of described triggerfish species. Most of these data derive from Dornburg et al. (2008), although two taxa, _Pseudobalistes naufragium_ and _Sufflamen verres_, were newly collected by MEA and sequenced, following the same protocols as Dornburg et al. (2008). Our study also includes three species of filefish (_Monacanthidae_), representing the closest sister group to the balistids (Santini and Tyler, 2003; Holcroft 2005; Alfaro et al. 2007; Dornburg et al. 2008; Yamanoue et al. 2008). We were unable to sample sufficient morphological data for adult _Sufflamen lunula_ and _Rhinecanthus verrucosus_, and these two taxa were subsequently pruned from our chronogram for analyses of morphometric data.

We conducted a Bayesian relaxed-clock time calibrated phylogenetic analysis (Drummond et al. 2006) of the triggerfish using two of the same calibration age priors as Dornburg et al. (2008). The first based the divergence of the lineages giving rise to extant families Balistidae and Monacanthidae on four fossil stem balistoids dated to 35 million years (MY): _Balistomorphus orbiculatus_, _B. ovalis_, _B. spinosus_, and _Oligobalistes robustus_ (Tyler and Santini 2002). We followed Alfaro et al. (2007) and assigned an upper bound of 70 MY to this calibration that reflects the appearance of several other tetraodontiform families in addition to the first stem tetraodontiforms in the fossil record. Divergence time analyses were repeated with the recently discovered Eocene taxa _Gornylistes prodigiosus_ used a calibration (Bannikov and Tyler 2008), although the change in the marginal posterior density of crown triggerfish ages was negligible. The second calibration age
prior placed a normally distributed prior age constraint on the age of crown balistids based on Alfaro et al.’s (2007) analysis. These prior age calibrations reflect the fossil record of balistids as no crown balistid fossils are known to be older than the middle Miocene (Schultz 2004), whereas stem balistids date back to at least 35 MY (Tyler and Santini 2002). We omitted one calibration used in Dornburg et al. (2008). Although the fossil *Balistes procapriscus* has been suggested to provide a minimum age on the split between *Balistes* and its sister group, *Pseudobalistes* (Santini and Tyler 2004), analyses of molecular datasets reveal *Pseudobalistes* to be polyphyletic (Dornburg et al. 2008), making assignment of this calibration ambiguous.

We estimated divergence times using the concatenated data under a model of uncorrelated but log-normally distributed rates using BEAST (Drummond et al. 2006), placing a birth–death prior on rates of cladogenesis. All fossil constraints incorporated soft upper bounds to avoid artificially truncating the posterior distribution of our divergence time estimates (e.g., Yang and Rannala 2006). Analyses were run with and without data to assess the influence of the prior on the posterior distribution of age estimates (Drummond et al. 2006). We used the nucleotide data partitioning strategy identified as having the highest Bayes factor support, with support being considered as Bayes factors greater than 10, resulting in partitioning our data by gene and codon (Kass and Raftery 1995; Brandley et al. 2005; Brown and Lemmon 2007) using the best-fit models of nucleotide substitution selected using AIC in jModelTest (Posada and Crandall 2000). The pool of candidate models included the pure birth (Yule) and birth–death model of speciation originally compared by Dornburg et al. (2008) as well as seven additional models. These included two fluctuating rate models (two rate Yule, and two rate birth death), a model allowing either speciation and extinction rates to vary through time, and two nested simpler models that held either the speciation or extinction rates constant while allowing the other parameter to vary through time. Finally the candidate model pool also included a log-normal and an exponential model of density-dependent lineage diversification that posit the lineage diversification rate to slow through time as the radiation progresses (Rabosky and Lovette 2006a,b). To test this hypothesis, we simultaneously compared the fit of several lineage diversification models using Akaike’s information criterion (Akaike 1973) in LASER. To test this hypothesis, we simultaneously compared the fit of several lineage diversification models using Akaike’s information criterion (Akaike 1973) in R using the LASER package (Rabosky 2006a,b). The pool of candidate models included the pure birth (Yule) and birth–death model of speciation originally compared by Dornburg et al. (2008) as well as seven additional models. These included two fluctuating rate models (two rate Yule, and two rate birth death), a model allowing either speciation and extinction rates to vary through time, and two nested simpler models that held either the speciation or extinction rates constant while allowing the other parameter to vary through time. Finally the candidate model pool also included a log-normal and an exponential model of density-dependent lineage diversification that posit the lineage diversification rate to slow through time as the radiation progresses (Rabosky and Lovette 2006a,b). To account for the potential effects of incomplete taxon sampling on these model fitting approaches to lineage diversification, we simulated 1000 random tree topologies using APE (Paradis et al. 2004) and Geiger (Harmon et al. 2007) under a pure-birth process using the empirical speciation rate inferred under a pure-birth process. This null distribution of trees was pruned to our level of taxon sampling (28 of 42 extant species) either randomly, or using one of three strategies that preferentially pruned younger taxa that had originated during the last 50%, 33%, or 25% of the time spanned by the phylogeny. These biased sampling strategies more accurately reflected our sampling of real triggerfish species, which was designed to sample at least one species from all major subclades and span reasonable levels of bias toward older bifurcations that may result from our empirical inclusion of all major triggerfish lineages. We assessed the fit of the lineage diversification models to this pool of pruned null trees and compared our empirical fit to the expected distribution of AIC score differences that result as a consequence of both random and nonrandom incomplete taxon sampling using the LASER package in R (Rabosky 2006b).

LINEAGE DIVERSIFICATION

If triggerfish experienced an ecological adaptive radiation (Schluter 2000), we predicted that species diversification rates would be highest early in the history of the clade and then slow through time as available niches filled (Schluter 2000; Rabosky et al. 2007). We initially tested this assumption using a modification of the Monte Carlo constant rates (MCCR) test (Pybus and Harvey 2000) that accounts for incomplete taxon sampling. Although the MCCR test was implemented in Dornburg et al. (2008), recent studies have found the gamma statistic of this test to be sensitive to biased, nonrandom, taxon sampling strategies, such as those employed by researchers attempting to sample all genera or functional groups (Cusimano and Renner 2010; Brock et al. 2011). As such, we implemented an extension of the MCCR test that accounts for nonrandom sampling of proportionally older splits while calculating the gamma statistic (Brock et al. 2011).

Although tests such as the MCCR test (Pybus and Harvey 2000) can detect early pulses of lineage accumulation, they cannot distinguish if this is a consequence of shifts in speciation, extinction, or alternate models such as density-dependent speciation (Rabosky and Lovette 2006a; Rabosky et al. 2007). For example, a higher than expected distribution of nodes toward the root of the tree might be the consequence of a variable rate of extinction, or could be explained by the expectations of a multirate birth–death model, and may not reflect a rapid initial radiation of species. To assess the best-fit model of lineage diversification for our data, we simultaneously compared all models using Akaike’s information criterion (Akaike 1973) in LASER. To test this hypothesis, we simultaneously compared the fit of several lineage diversification models using Akaike’s information criterion (Akaike 1973) in R using the LASER package (Rabosky 2006a,b). The pool of candidate models included the pure birth (Yule) and birth–death model of speciation originally compared by Dornburg et al. (2008) as well as seven additional models. These included two fluctuating rate models (two rate Yule, and two rate birth death), a model allowing either speciation and extinction rates to vary through time, and two nested simpler models that held either the speciation or extinction rates constant while allowing the other parameter to vary through time. Finally the candidate pool also included a log-normal and an exponential model of density-dependent lineage diversification that posit the lineage diversification rate to slow through time as the radiation progresses (Rabosky and Lovette 2006a,b). To account for the potential effects of incomplete taxon sampling on these model fitting approaches to lineage diversification, we simulated 1000 random tree topologies using APE (Paradis et al. 2004) and Geiger (Harmon et al. 2007) under a pure-birth process using the empirical speciation rate inferred under a pure-birth process. This null distribution of trees was pruned to our level of taxon sampling (28 of 42 extant species) either randomly, or using one of three strategies that preferentially pruned younger taxa that had originated during the last 50%, 33%, or 25% of the time spanned by the phylogeny. These biased sampling strategies more accurately reflected our sampling of real triggerfish species, which was designed to sample at least one species from all major subclades and span reasonable levels of bias toward older bifurcations that may result from our empirical inclusion of all major triggerfish lineages. We assessed the fit of the lineage diversification models to this pool of pruned null trees and compared our empirical fit to the expected distribution of AIC score differences that result as a consequence of both random and nonrandom incomplete taxon sampling using the LASER package in R (Rabosky 2006b).
CORRELATED EVOLUTION BETWEEN COMPONENTS OF BALISTIFORM Locomotor MORPHOLOGY

We tested for evidence of correlated evolution between the most important components of variation in our fin shape datasets while taking the expected covariance among traits due to phylogeny into account by using the phylogenetic generalized least squares (PGLS) method in the APE (Paradis et al. 2004) software package in R. As the model of evolution can affect this type of analysis, tests were conducted under both a Brownian (random walk) and OU (constrained) model of evolution, while comparing the AIC fit of each model (Table S2). To assess whether elements of body shape are integrated with the evolution of balistid locomotor morphology, we used a multiple regression using body shape as the dependent variable while taking the interactions of the fin shapes into account. Model fitting was conducted under both a BM and OU model of phenotypic change, with AIC values greater than four being used to select between model fits (Burnham and Anderson 2002).

We used partial least squares (PLS) (Rohlf and Corti 2000) in TPSPLS (Rohlf 2005) and Morphof (Klingenberg 2008) to identify pairs of axes that explain the maximum covariance between the datasets, in a manner mathematically similar to a principal components analysis emphasizing covariance as opposed to variance (Bookstein and Rohlf 2004; Zelditch et al. 2004). This method can only assess covariance between pairs of datasets so we conducted three pairs of PLS analyses for our shape data, one for each possible combination of the fin and body shape matrices. To test whether the shared covariance of the datasets was robust to the influence of phylogeny, we also extracted pairs of covarying PLS axes and tested them for significant correlated evolution using PGLS.

MORPHOLOGICAL AND FUNCTIONAL DIVERSIFICATION

Clades that have undergone an ecological adaptive radiation (Schluter 2000) are expected to partition more ecofunctional trait variation among rather than within subclades (Harmon et al. 2003). We used phylomorphospace visualization (Sidlauskas 2008) and analyses of disparity through time (Harmon et al. 2003) to investigate whether triggerfish exhibited this pattern. Phylomorphospaces combine morphometric and phylogenetic datasets to provide a visual assessment of how lineages partition available morphospace and how closely phylogenetic proximity predicts morphological similarity. We plotted the three RWs explaining the greatest percentage of total variance for each dataset (body, dorsal fin, and anal fin) in 3D morphospaces, and projected the phylogeny linking these species into the same space by reconstructing the morphological position of the internal nodes using weighted squared change parsimony. All phylomorphospaces were plotted using the Rhetenor module (Dyreson and Maddison 2003) in Mesquite (Maddison and Maddison 2008).

Assessing the relative subclade disparity among lineages has become an integral part of macroevolutionary studies focusing on how lineages occupy a morphospace (e.g., Foote 1997; Eble 2000; Valentine and Jablonski 2003; Villier and Eble 2004; Jablonski 2005). If the novelty of the triggerfish locomotive system has driven a rapid early pulse of diversification in their shape morphology, triggerfish lineages should achieve higher relative subclade disparity early in their history than would be expected under a random model of evolution. We tested this hypothesis by calculating the relative subclade disparity through time for fin aspect ratios and the most important axes of fin and body shape (Harmon et al. 2003). To assess whether triggerfish diversity patterns differed from a null model of Brownian evolution, we simulated the evolution of fin and body shape variable on the triggerfish topology using the empirical variance for each trait 1000 times and calculated the morphological disparity index (MDI) using the Geiger package (Harmon et al. 2007) in R. Negative MDI values indicate that subclades vary strongly from one another whereas positive values indicate that subclades have converged. To account for our level of incomplete taxon sampling of tipward taxa, we followed Harmon et al. (2003) and restricted our analysis of MDI values to the first 60% of the time spanned by the phylogeny.

**Results**

**DETERMINING THE MAJOR AXES OF MORPHOLOGICAL VARIATION**

Two RW axes explained approximately 76% of the total body shape change variation for triggerfish (Table 1). The first axis of body shape change described an elongation of the overall body coupled with an anteroposterior compression of the cranium (Fig. 1). The second RW axis for body shape change described a dorsoventral compression of body depth coupled with an anteroposterior elongation of the body and cranium (Fig. 1). The combined body shape data also provide evidence for substantial restructuring of the triggerfish cranium, with the dorsal slope of the cranium becoming steeper and elongating.

For the dorsal and anal fins, the first three RWs explained 90% and 93% of the overall variance for each fin shape, respectively (Table 1). For both the dorsal and anal fins, the first RW

| Table 1. Percent of variance explained by the first three relative warps. |
|-----------------------|----------|----------|----------|
|                       | Anal fin | Dorsal fin | Body shape |
| Relative Warp 1      | 57.18    | 59.72     | 51.94     |
| Relative Warp 2      | 20.62    | 19.49     | 23.4      |
| Relative Warp 3      | 12.69    | 13.86     | 7.5       |
TRIGGERFISH DIVERSIFICATION

**Figure 1.** Dorsal fin, body shape, and anal fin shape change quantified by the first two relative warps. Positive and negative extremes represent the maximum observed phenotypic divergence on either end of the respective RW axis.

(explaining 57–60% variance) described the change in the length of the anterior (leading edge) fin rays (Fig. 1). The second RW in both fins (explaining 19–21% variance) described variation in the anteroposterior length of the fin coupled with a dorsal to ventral elongation of fin rays posterior to the leading edge (Fig. 1). The third RW axis (explaining 13–14% of the variance in shape change) described a change in curvature along the distal margin of the fin (image not shown).

**TRIGGERFISH PHYLOGENETICS AND DIVERGENCE TIME ESTIMATION**

Our inferred phylogeny contains six major clades: (1) *Canthidermis*; (2) *Sufflamen*; (3) *Rhinecanthus*; (4) *Abalistes*; (5) *Balistes*; and (6) all other balistids, providing strong support for the paraphyly and polyphyly of multiple genera congruent with the findings of Dornburg et al. (2008) (Fig. 2). Our results mirror Dornburg et al. (2008), as we found *Balistes* to be paraphyletic, with strong support present for nested placement of *Pseudobalistes fuscus* and *P. naufragium* within this clade. There was strong support for a sister relationship between *Balistoides viridescens* and *P. flavimarginatus*, suggesting these genera are also not monophyletic. Our analysis revealed high support values (PP > 0.95) for most nodes in the tree, with the exception of the three most basal divergences (Fig. 2).

We estimated a crown age of balistids of approximately 10 MY, with the 95% highest posterior density (HPD) interval spanning approximately 7–14 MY, suggesting Balistidae to be relatively young (Fig. 2). The chronogram also suggests that the stem lineages of the six major clades identified above began to diversify in the Late Miocene. *Abalistes* and *Canthidermis* appear to have split relatively rapidly from the lineage that gives rise to clades 2 and 3 (~9 MY). The confidence intervals of our age estimates are all well within the bounds of the ages estimated by Dornburg et al. (2008), suggesting that the placement of the additional calibration age prior based on the fossil *B. procapriscus* had a minimal influence on the analysis (Table S2).

**PATTERNS OF TRIGGERFISH LINEAGE DIVERSIFICATION**

The rate of lineage diversification ($\lambda_G$) for the triggerfish was estimated at approximately 0.266 lineages per million years, based on a pure-birth (Yule) model. Similar log likelihood scores for the fit of the Yule ($\sim$4.1363) and birth–death ($\sim$4.1360) models of lineage diversification prevented us from being able to reject a pure-birth model as the underlying process for triggerfish lineage diversification. When the candidate pool of models was expanded to include seven additional models of lineage diversification, a density-dependent model of speciation was inferred to be the best fit, suggesting rates of lineage diversification to have declined over the history of triggerfish evolution. However, this pattern of declining rates of cladogenesis most likely reflects a methodological bias as our simulations designed to mimic various taxon sampling strategies demonstrated that this result is within the range of the model-fit expectations that would occur based on incomplete and nonrandom taxon sampling (Fig. 3). Similarly, accounting for nonrandom taxon
Figure 2. Consensus chronogram of triggerfish divergence times. Bars around nodes represent the 95% HPD. Shaded dark boxes at nodes indicate posterior probabilities (PP) >0.95, whereas numbered nodes correlate with Appendix 2. White numbered nodes indicate the presence of prior age constraints. Nodes with PP less than 0.5 do not contain a 95% HPD on the age estimate.

CORRELATED EVOLUTION AMONG COMPONENTS OF BALISTIFORM LOCOMOTION

PGLS analysis provided strong evidence for correlated evolution between the two most important RWs of the dorsal fin and each of the reciprocal RWs of the anal fin (Table 2). This suggests that the heightening, elongation, or skewing of one of these fins is linked to a similar change in the other (0.00001 < P < 0.05). These results were identical under both a Brownian and constrained (OU) model of phenotypic evolution. We observed congruent patterns when analyzing pairs of PLS axes while accounting for the expected covariance due to shared phylogenetic history using PGLS (Table 3). Further, plotting dorsal fin shape RW1 against anal fin shape RW1 reveals a striking pattern of tight correlation between the two RWs, confirming that the two traits to have undergone correlated evolution (Fig. S5).

Multiple regression analysis provided strong evidence for a pattern of integrative evolution between the interacting fin shapes and the first major body shape warp under both a BM and OU model (r^2 ~ 0.71). Conversely, we obtained no significant correlation between the second RW of body shape and the interactions of the anal and dorsal fin shapes (r^2 ~ 0.3), supporting a decoupling of the evolution of dorsalventral compression of the body from the elongation and lengthening of fin shapes.
TRIGGERFISH DIVERSIFICATION

Figure 3. Distribution of ΔAIC differences (Δ1AIC) generated by fitting a Yule and exponential density-dependent models of speciation (DDX) to pruned trees simulated under a pure birth process. Positive Δ1AIC indicate preferred fits for the DDX model. Dark arrows represent empirical fit of triggerfish data. (A) Expected Δ1AIC distribution generated by randomly sampling Yule trees (P ∼ 0.01); (B) Expected Δ1AIC distribution generated by preferentially pruning lineages from the more recent 50% of each tree's length (P ∼ 0.07); (C) Expected Δ1AIC distribution generated by preferentially pruning lineages from the more recent 33% of each tree's length (P ∼ 0.09); and (D) Expected Δ1AIC distribution generated by preferentially pruning from the more recent 25% of each tree's length (P ∼ 0.09).

MORPHOLOGICAL AND FUNCTIONAL DIVERSIFICATION

We find evidence of phylogenetic clustering for both the fin and body-shape data, with subclades occupying distinct regions of morphospace (Fig. 4A–C). Additionally, plotting the body-shape RW1 against either the dorsal or anal fin RW1 (Fig. 4D and E) reveals that nearly every major clade occupies a position in morphospace that is divergent from that of its sister clade. The only major qualitative difference between these figures is the wide separation of members of lineage 5, identified as "Balistes" sensu Dornburg et al. (2008) and the genus Canthidermis in the anal-fin shape/body shape phylomorphospace (Fig. 4E).

Table 2. Testing for correlated evolution between fin shapes.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Axes</th>
<th>P-value/β-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>First major warp vs. Second major warp</td>
<td>dorsal fin vs. dorsal fin</td>
<td>0.057/0.001</td>
</tr>
<tr>
<td>First major warp vs. Second major warp</td>
<td>dorsal fin vs. anal fin</td>
<td>0.16/0.015</td>
</tr>
<tr>
<td>Second major warp vs. Second major warp</td>
<td>dorsal fin vs. dorsal fin</td>
<td>0.015/0.005</td>
</tr>
<tr>
<td>Second major warp vs. Second major warp</td>
<td>dorsal fin vs. anal fin</td>
<td>0.16/0.015</td>
</tr>
</tbody>
</table>

Results from the phylogenetic generalized least squares test for correlation between fin shapes. Results in bold indicate significant evolution between these traits. Bold P-values indicate significant results.

We find evidence of phylogenetic clustering for both the fin and body-shape data, with subclades occupying distinct regions of morphospace (Fig. 4A–C). Additionally, plotting the body-shape RW1 against either the dorsal or anal fin RW1 (Fig. 4D) reveals that nearly every major clade occupies a position in morphospace that is divergent from that of its sister clade. The only major qualitative difference between these figures is the wide separation of members of lineage 5, identified as "Balistes" sensu Dornburg et al. (2008) and the genus Canthidermis in the anal-fin shape/body shape phylomorphospace (Fig. 4E).

Table 3. Testing for correlated evolution among principal PLS axes.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Axes</th>
<th>P-value/β-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dorsal fin vs. body shape first vs. second</td>
<td>0.0009/0.223</td>
<td></td>
</tr>
<tr>
<td>Dorsal fin vs. body shape second vs. first</td>
<td>0.0002/0.144</td>
<td></td>
</tr>
<tr>
<td>Anal fin vs. body shape first vs. second</td>
<td>0.0001/0.141</td>
<td></td>
</tr>
<tr>
<td>Anal fin vs. body shape second vs. first</td>
<td>0.0016/0.314</td>
<td></td>
</tr>
</tbody>
</table>

Results from the phylogenetic generalized least squares test for correlated evolution between morphospace axes of variation from the partial least squares analysis. Bold P-values indicate significant results.
Figure 4. Phylomorphospace visualizations for the major relative warps. Comparisons of how lineages occupy different morphospaces: (A) The first and second dorsal fin RWs; (B) the first and second anal fin RWs; (C) the first and second body shape RWs; (D) comparison between the first dorsal fin RW and the first body shape RW; (E) the first anal fin and the first body shape RW; (F) the aspect ratio of the dorsal fin and the aspect ratio of the anal fin. *Genus names follow the classification proposed in Dornburg et al. (2008).
Discussion

Three lines of evidence suggest that the functional demands of balistiform locomotion have influenced triggerfish shape evolution. First, the evolution of the most important axes of anal and dorsal fin shapes is highly integrated. Second, key aspects of median fin and body shape have coevolved in triggerfish. Third, triggerfish appear to have explored the limits of their modern phenotypic and functional diversity early in their history. These results support the idea that the majority of shape diversity observed in modern triggerfish reflects an underlying early diversification of function and are consistent with the expectation that evolutionary novelty spurs functional diversification. However, our analysis of lineage diversification fails to find a corresponding early pulse of speciation, suggesting morphological and lineage diversification to have been decoupled during triggerfish evolution. Thus, the history of triggerfish diversification does not conform to the expectations of the classical adaptive radiation model.

Patterns of Correlated Shape Evolution

The results of every test for correlated evolution that we performed (Tables 2 and 3) as well as the observed linear relationship between the RWs of the median fins in phylomorphospace (Fig. S7) strongly suggest that the two major locomotor structures of triggerfish, the dorsal and anal fins, evolved in lockstep. Further, the phylomorphospace visualizations of aspect ratio (Fig. 4F) show that mechanical evolution in triggerfish median fins closely reflects patterns of morphological evolution (Fig. 4A and B). The visualizations also imply ancestral crown triggerfish to have had a medium aspect ratio (Fig. 4F), with a subsequent history of most lineages converging on low and medium aspect ratios (Upper left corner, Fig. 4F). Low aspect ratio fins are correlated with increased maneuverability and are associated with fish that remain in close proximity to reefs, whereas higher aspect ratios are associated with increased water column usage and also shallower habitats with more wave energy (Fulton et al. 2005). These expectations suggest two functional hypotheses to explain the tightly correlated evolution of dorsal and anal fin shape and function: (1) balistiform swimmers may be functionally constrained to achieve propulsive forces using coupled symmetrical fins, or (2) there may be a developmental constraint in which the dorsal and anal fin belong to the same module. These hypotheses are not mutually exclusive. The potential lift generated by the dorsal and anal fins may constrain these fins to evolve in tandem for efficient maneuverability in the water column, whereas developmental modules could also shape the development of the fish’s underlying musculature and skeletal elements (e.g., Mabee et al. 2002). Sorenson (2007) recently found a startling degree of symmetry between the underlying dorsal and anal fin structural elements in Rhinecanthus rectangularis, supporting the idea of a developmental module underlying the primary locomotor components of triggerfish.

Our analyses also suggest that major features of body shape evolve in tandem with fin shape. For example, we find that elongate bodies are coupled with rounder, less-sigmoidal fins (Table 3, Fig. S6 left panel). Fish with this body plan tend to be more reef associated (Lieske and Myers 2001; Bean et al. 2002) suggesting this to be an efficient body plan for maneuvering complex 3D environments. Triggerfish lineages associated with pelagic and open environments are also characterized by high aspect ratio fins and bulbous reduced crania (Fig. S6). This suggests modifications of the cranial morphology to be a potentially important and understudied aspect of the hydrodynamics of balistiform locomotion.

What Drove Functional and Morphological Diversification in Triggerfish?

The morphological disparity indices and the phylomorphospace visualizations show that triggerfish colonized nearly all of their presently occupied morphospace early in their radiation. All axes of shape and functional change are partitioned among rather than within lineages early in the history of the group (Fig. 4), and the accumulation of disparity among lineages appears to occur rapidly, mostly during the first 1–2 MY of the group’s history. This pattern is consistent with recent adaptive radiation models in which early lineages invade disparate regions of morphospace whereas later lineages subpartition initially colonized regions (Schluter 2000; Harmon et al. 2003). However, triggerfish depart from these models in one important sense: they do not show the expected pattern of initially rapid lineage diversification.

Although we recover a pattern of elevated early cladogenesis while assuming random taxon sampling, the results of simulations that account for nonrandom taxon sampling clearly demonstrate the sensitivity of these test statistics to violations of this assumption (e.g., Cusimano and Renner 2010; Brock et al. 2011). Two scenarios might explain the apparent patterns of uncoupled
Figure 5. Disparity through time plots for anal fin shape RW1 (A), Dorsal fin shape RW1 (B), body shape RW1 (C), body shape RW2 (D), anal fin aspect ratio (E), dorsal fin aspect ratio (E). The solid curve represents the empirically inferred pattern subclade disparity through time. The dashed curve represents the median of the Brownian simulations. All plots were generated using the Geiger (Harmon et al. 2007) package in R.
lineage diversification and morphological evolution in triggerfish. It is possible that these processes were in fact coupled, but the elevated rates of extinction eroded the signature of early rapid lineage diversification (Quental and Marshall 2009; Rabosky 2009). We regard this scenario as unlikely for two reasons. First it requires morphologically disparate subclades to resist total elimination even while extinction acts within them to reduce richness. This is not a frequent outcome of standard models of high lineage turnover, which tend to eliminate early branching lineages entirely and generate very young crown clades (Raup 1985; Sidlauskas 2007). Second, some simple violations of the assumption of random extinction, such as the heritability of extinction rates, are expected to bias methods for testing diversification rates toward inferring an early pulse of diversification even when the true diversification rate remains constant (Rabosky 2009). In spite of that potential bias, we fail to recover evidence of initially rapid diversification. Thus, although it cannot be ruled out completely, we believe it unlikely that triggerfish experienced an invisible burst of speciation shortly after evolving balistiform locomotion.

A second possibility is that triggerfish adaptive diversification reflects a process where phenotypic diversification is decoupled from cladogenesis. Although the most prevalent model of ecological adaptive radiation (e.g., Schluter 2000) predicts that these processes will be linked, the most cited examples of this phenomenon are clades with restricted geographic distributions such as islands or lakes (Day and Wilkinson 2006; Seehausen 2006, Losos 2009; Johnson et al. 2009), quite unlike the broad species ranges that characterize most triggerfish. In widespread marine species, long-distance larval dispersal (e.g., Palumbi 1994, Bellwood et al. 2006) might strongly alter the expected dynamics of linked species and phenotypic diversification following evolutionary innovation by decreasing the probability of localized ecological speciation (e.g., Rocha and Bowen 2008; Budd and Pandolfi 2010). Instead, a constant rate of allopatric or peripatric speciation that is governed by physical processes and therefore unaffected by functional innovation would provide the dominant processes generating new lineages (see also, Budd and Pandolfi 2010). After the rise of each new lineage, secondary contact and character displacement (rather than ecological speciation) could then act as the primary pump driving phenotypic diversification into newly available niches (e.g., Price 2010). This model would yield an initial steady increase in disparity at a rate governed by the background rate of speciation. As the accumulating lineages filled niches over time, the rate of character displacement following instances of secondary sympatry in new lineages would slow; until all niches opened by the original innovation were filled. Thus, the eventual braking of the morphological diversification rate remains the same as in the classic adaptive radiation model.

We suggest that triggerfish may have evolved under this alternative model of diversification, with balistiform locomotion serving as the functional innovation catalyzing subsequent morphological diversification. As in the classic model of adaptive radiation, the origin of balistiform locomotion likely presented the potential for ancestral triggerfish to evolve novel fin and body shape combinations, partly as a consequence of the locomotor innovation providing access to novel ecological opportunities. However, as described above, this innovation in and of itself would not alter the background rate of species formation, as any potential for sympatric ecological diversification would be impeded by the presence of long-distance dispersal. Species formation would have instead been dominated by allopatry/peripatry and the primary control on morphological diversification would have been the background rate of isolation and secondary contact between lineages. Once the novel niches were filled, new species would still arise at the unchanged background rate of allopatric or parapatric speciation, but the rate of morphological diversification would decline and daughter species, having no open paths across the adaptive landscape, would tend to resemble their ancestors.

Although speculative, this model would explain the apparent constant rate of species diversification throughout the history of triggerfish and the inferred burst of morphological diversification at the base of their phylogeny. Such a model might also explain why many freshwater fish species radiations appear to be consistent with the traditional model of ecological radiation (e.g., Barbour 1973; Witte 1984; Hunt et al. 1997; Alesandrini and Bernardi 1999; Seehausen 2002; Day and Wilkinson 2006; Seehausen 2006), although species rich tropical marine fish families often are not (Santini et al. 2009; Alfaro et al. 2009a). Continued investigations of widespread tropical marine radiations will help reveal whether the decoupling of cladogenesis and morphological change observed in the triggerfish reflects the typical condition in coral reef fish groups. Filefish (Monacanthidae) that are balistiform swimmers with long-range dispersal that also share a close affinity with triggerfish, offer one excellent opportunity to test that hypothesis, but a full exploration should target groups with varied styles of locomotion and dispersal. Such future studies will aid in the conceptual development of adaptive radiation theory and determine if the lack of tropical coral reef fish species flocks reflects a lack of looking for them or fundamental differences in the evolutionary phenomena that shape biodiversity in tropical marine versus freshwater habitats.

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LITERATURE CITED


Supporting Information

The following supporting information is available for this article:

**Figure S1.** Landmarks and sliding semi-landmarks used for fin shape analysis.

**Figure S2.** Landmarks and sliding semi-landmarks used for body shape analysis.

**Figure S3.** Calculation of aspect ratio (Taken from Wright 2000).

**Figure S4.** Calculation of the null gamma statistic based on Pybus and Harvey’s (2000) Monte Carlo constant rates test.

**Figure S5.** First and second partial least squares axes summarizing major axes of covariance between anal and dorsal fin shapes.

**Figure S6.** First and second partial least squares axes summarizing major axes of covariance between dorsal fin and body shapes.

**Figure S7.** Comparison between the first dorsal fin and the first anal fin RWs.

**Table S1.** Sequences and Genbank numbers used in this study.

**Table S2.** AIC scores for phenotypic models of evolution in PGLS.

**Table S3.** Distribution of node ages.

**Table S4.** AICc scores for models of morphological evolution.

**Appendix S1.** Location of specimens examined for this study.

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

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