Linking stream ecology with morphological variability in a native freshwater fish from semi-arid Australia

Samantha Lostrom¹,², Jonathan P. Evans¹, Pauline F. Grierson², Shaun P. Collin¹,³, Peter M. Davies⁴ & Jennifer L. Kelley¹,²

¹School of Animal Biology (M092), The University of Western Australia, 35 Stirling Highway, Crawley, Western Australia 6009, Australia
²Ecosystems Research Group and West Australian Biogeochemistry Centre, School of Plant Biology, The University of Western Australia, 35 Stirling Highway, Crawley, Western Australia 6009, Australia
³UWA Oceans Institute (M470), The University of Western Australia, 35 Stirling Highway, Crawley, Western Australia 6009, Australia
⁴Centre of Excellence in Natural Resource Management, The University of Western Australia, Albany, Western Australia, Australia

Keywords
Local adaptation, phenotype–environment correlation, phenotypic plasticity, Pilbara, polymorphism, population differentiation.

Correspondence
Jennifer L. Kelley, School of Animal Biology (M092), The University of Western Australia, 35 Stirling Highway, Crawley, WA 6009, Australia.
Tel: +61 (8) 6488 2239;
Fax: +61 (8) 6488 1029;
E-mail: Jennifer.kelley@uwa.edu.au

Funding Information
Funded by an Australian Research Council Linkage Grant LP120200002 with industry partners Rio Tinto and BHP Billiton.

Received: 18 December 2014; Revised: 21 May 2015; Accepted: 10 June 2015

Ecology and Evolution 2015; 5(16): 3272–3287
doi: 10.1002/ece3.1590

Abstract

Environmental variation is a potent force affecting phenotypic expression. While freshwater fishes have provided a compelling example of the link between the environment and phenotypic diversity, few studies have been conducted with arid-zone fishes, particularly those that occur in geographically isolated regions where species typically inhabit intermittent and ephemeral creeks. We investigated morphological variation of a freshwater fish (the western rainbowfish, Melanotaenia australis) inhabiting creeks in the Pilbara region of northwest Australia to determine whether body shape variation correlated with local environmental characteristics, including water velocity, habitat complexity, predator presence, and food availability. We expected that the geographic isolation of creeks within this arid region would result in habitat-specific morphological specializations. We used landmark-based geometric morphometrics to quantify the level of morphological variability in fish captured from 14 locations within three distinct subcatchments of a major river system. Western rainbowfish exhibited a range of morphologies, with variation in body depth accounting for a significant proportion (>42%) of the total variance in shape. Sexual dimorphism was also apparent, with males displaying deeper bodies than females. While the measured local habitat characteristics explained little of the observed morphological variation, fish displayed significant morphological differentiation at the level of the subcatchment. Local adaptation may partly explain the geographic patterns of body shape variation, but fine-scale genetic studies are required to disentangle the effects of genetic differentiation from environmentally determined phenotypic plasticity in body shape. Developing a better understanding of environment–phenotype relationships in species from arid regions will provide important insights into ecological and evolutionary processes in these unique and understudied habitats.

Introduction

Species typically exhibit considerable phenotypic variation across their geographic range (Endler 1986; Wade and Kalisz 1990; Foster and Endler 1999), where much variation can be explained by climate, habitat type, and predation pressure (Losos et al. 1998; Nagel and Schluter 1998; Schluter 2000; Langerhans and DeWitt 2004). Phenotypic variation can be attributable to numerous factors, including (1) individual genotypes producing different phenotypes in response to changing environments (i.e., phenotypic plasticity), (2) populations exhibiting fixed differences in phenotypic traits in response to selection (adaptive differentiation), and (3) processes such as genetic drift and developmental constraints (Price et al. 2003). Importantly, revealing the influence of environmental and ecological factors on phenotypic variation can provide valuable insights into evolutionary processes such as population differentiation and ecological speciation (Maynard Smith 1966; Schluter 1996; Rundle and Nosil 2005).
Freshwater fishes have provided some of the most compelling examples of the effect of environmental variation on phenotypic expression. In the guppy (*Poecilia reticulata*), for example, divergent patterns of selection associated with the risk of predation have generated populations that vary in life history traits (Reznick 1982; Reznick and Endler 1982), morphology (Alexander et al. 2006), coloration (Endler 1983), and behavior (Seghers 1974). Other selective agents may also play a role in contributing to population variation in body size and shape in guppies, including the level of canopy cover and water-flow rate (Hendry et al. 2006). Furthermore, in three-spine sticklebacks, variation in body armor (lateral plate and spine expression) has been attributed to the effects of predation, nutrient availability, and parasite abundance (Reimchen 1994; Marchinko 2009). Together, this evidence suggests that numerous and potentially interacting factors can drive patterns of phenotypic variation, thus cautioning against focusing on a single selective axis of ecological variance (e.g., high/low predation risk, benthic/limnetic habitat) when attempting to explain phenotypic variation among natural populations (Langerhans and Makowicz 2009).

Among the environmental factors influencing phenotypic variation in fishes, water flow can have an important affect on fish body shape due to the hydrodynamic effects of drag and turbulence on swimming efficiency (Enders et al. 2003; Langerhans 2008). Fishes inhabiting fast-flow habitats tend to have shallow, elongated (streamlined) bodies compared with those living in slow-flow regions, with the former becoming fusiform (spindle-shaped) through increased anterior body depth (Brinsmead and Fox 2002; Langerhans et al. 2003; Aguirre 2009; Franssen 2011; Drinan et al. 2012). Fishes with a streamlined morphology experience reduced drag and can cope with prolonged, steady swimming in moderate water flows (Schaef er et al. 1999; Wolfgang et al. 1999; Blake et al. 2005, 2009; Langerhans 2008, 2009; Blob et al. 2010). Water flow can also induce developmental shifts in morphology, as juvenile fish may develop a fusiform body shape when reared in fast flows and display a deep body form when reared in slow flows (Pakkasmaa and Piironen 2000; Paez et al. 2008). However, the reverse has also been observed, whereby fish develop deeper bodies in fast flows (Pakkasmaa and Piironen 2000; Peres-Neto and Magnan 2004; Kristjansson et al. 2012). It is likely that the complex and often unpredictable effects of water flow on body shape are in part attributable to interactions with other environmental and ecological factors.

Predation risk also has a profound effect on the phenotypic traits of many fishes, including body shape. Populations that coexist with predators typically possess deeper bodies than their low predation counterparts, which reduces their overall risk of predation relative to streamlined fishes (Andersson et al. 2006; Sass et al. 2006; Eklov and Jonsson 2007; Chivers et al. 2008; Blob et al. 2010). In crucian carp (*Carassius carassius L*.), for example, the exposure of individuals to chemical cues from predators induces a change in body shape, such that individuals became deeper-bodied when exposed to predatory cues from the piscivorous pike *Esox lucius* (Bronmark and Miner 1992; Bronmark and Pettersson 1994). Deep-bodied fish, in turn, are able to initiate enhanced “fast-start” responses during escape from predators due to the large surface area of the body that is available to produce thrust (Law and Blake 1996; Royle et al. 2006; Domenici et al. 2008). Deep-bodied prey are also less vulnerable to gape-limited piscivores (Magnhagen and Heibo 2001; Zimmerman 2007) and require a longer handling time than slim-bodied prey (Nilsson et al. 1995).

Sexual dimorphism, which is commonly observed in freshwater fishes (Proulx and Magnan 2004; McCairns and Bernatchez 2012; Naspleda et al. 2012), is another important source of morphological variation. Divergence between the sexes can be attributable to sexual selection, for example, where certain morphologies confer mating advantages through competition or mate choice (Forsgren, 1992; Quinn and Foote 1994). In the three-spine stickleback (*Gasterosteus aculeatus*), females exhibit mating preferences for males on the basis of body morphology and size, although these preferences depend on the females’ origin; benthic females prefer large males (irrespective of shape) while limnetic females prefer slender males (Head et al. 2013). Sexual dimorphism can also arise as environmental pressures and/or constraints impact the sexes differentially. For example, female guppies can utilize high-velocity areas of streams to avoid being harassed by males (Magellan and Magurran 2006), which may result in sex-specific changes in body shape due to selection imposed by water flow on female (but not male) body shape. Accounting for factors such as sex and body size is therefore an important prerequisite for understanding the relationship between body shape variation and environmental variability.

Most studies of morphological variation in freshwater fishes have focused on species in the northern hemisphere that inhabit semi-isolated environments and display discrete ecotypes (reviewed by Skulason and Smith 1995; Robinson et al. 1996; Schluter 1996). However, the process of population divergence can be viewed as a continuum from panmixia to total reproductive isolation, and understanding the factors that constrain or promote movement along this continuum can yield significant insights into evolutionary processes (Hendry 2009). Furthermore, few studies of morphological variation have been conducted in arid or semi-arid regions, where...
streams are often ephemeral or intermittent and where the hydrodynamics are highly unpredictable both spatially and temporally (Dogramaci et al. 2015). Such dynamic and often disconnected conditions, at both subcatchment and basin scales, have the potential to promote the development of habitat-associated morphological specializations (Tobler et al. 2008).

Here, we investigate body shape variation in relation to habitat heterogeneity in a native Australian freshwater fish, the western rainbowfish, *Melanotaenia australis*. This species is an ideal study model for exploring the links between ecological and environmental factors and body shape. Populations occupy a wide variety of habitats in arid northwest Australia, where fish are commonly found in ephemeral pools (Morgan and Gill 2004; Beesley and Prince 2010). Males tend to be deeper-bodied and have more pointed dorsal and ventral fins than females (Allen et al. 2002), while larger males gain an advantage during mating (Young et al. 2010). Previous research on *M. australis* has revealed that variation in body shape cannot be explained by predation, but there is a tendency for morphological variation to correlate with environmental factors, such as water flow and the availability of aquatic vegetation (Young et al. 2011). In two other species of rainbowfishes (*M. eucharis* and *M. dóboulayi*), variation in body shape (including fin position) is specific and has been attributed to a number of interacting factors, including habitat, sex, and water velocity (McGuigan et al. 2003). In this study, we characterize the level of variation in adult body shape across several natural populations of *M. australis* and determine whether this variation can be attributed to specific environmental and ecological characteristics of the local habitat. In accordance with previous studies (e.g., Langerhans et al. 2003, 2007; Aguirre 2009; Harrod et al. 2010), we expected fish morphology would reflect interpopulation differences in water velocity, predation, diet, and habitat complexity. Specifically, we anticipated that deep-bodied morphs would occur in habitats with a high availability of benthic prey, high risk of predation, high habitat complexity, and slow water velocity, while streamlined morphs would be found in sites with low habitat complexity, fast water flows, and low risk of predation.

**Materials and Methods**

**Study area**

The study was conducted within the Fortescue River catchment in the Pilbara region of northwest Australia. The Pilbara has a semi-arid to subtropical climate, characterized by hot summers (24–40°C) and mild winters (11–26°C) (www.bom.gov.au). Rainfall mainly occurs in the summer, and high evaporation relative to precipitation restricts drainages to contiguous or discrete pools or reaches maintained by groundwater. Pools can vary in depth (<1.5 m to >3 m) and serve as refuges for fishes until reconnection during flooding events (Beesley and Prince 2010). Large rainfall events, primarily associated with summer cyclones, recharge catchments and contribute to sustaining pools in creeks and rivers, such that the hydrology, biogeochemistry, and the ecology of pools are intimately linked (Fellman et al. 2011). At the time of sampling, relatively little rain (<50 mm) had fallen across the catchment in the preceding 4 months. Consequently, surface water in many tributaries and along the main channel had either dried or been reduced to a series of pools maintained largely by groundwater (Fellman et al. 2011).

**Field sampling**

A total of 14 sites encompassing the upper, mid, and lower subcatchments of the Fortescue River were sampled between May and November 2013. The Fortescue River is approximately 760 km in length, drains a 30,000 km² catchment of the Hamersley Basin, and only flows contiguously following exceptionally large flood events (Barnett and Commander 1985). The Fortescue River is divided into upper and lower sections that are separated by the Goodiadarrie Hills (Skrzypek et al. 2013). The lower Fortescue River drains in a westerly direction from the Hamersley Ranges toward the coast, whereas east of the hills the Fortescue Marsh receives drainage from the upper catchment. The upper and lower parts of the catchment are considered hydrologically disjunct (Skrzypek et al. 2013). We have designated pools on the lower Fortescue River but still within the Hamersley Ranges as “mid-Fortescue.” Overall, four pools were sampled in the upper Fortescue, two in the mid-Fortescue and eight in the lower Fortescue. The numbers of pools sampled varied among subcatchments owing to both the number of pools that occurred in each and the local abundance of western rainbowfish.

**Habitat characterization**

We collected physical environmental data (summarized in Table 1) at each site prior to capturing the fish. Five replicates of surface water velocity were measured at each pool using a water-flow probe (FP111; Global Water™, College Station, TX), which was placed 10 cm below the water’s surface. At each site, the proportion of open water, fine and coarse gravel substrates, bark/wood, rocks, and aquatic vegetation were assessed to generate a habitat.
Table 1. Location of sample sites, number of fish sampled, and a summary of the habitat characteristics. Complexity ranks ranged from 0 to 5, where 0 represented the lowest habitat complexity and five described the most complex sites.

| Site                        | Subcatchment | Latitude   | Longitude | Fish sampled | Complexity rank | Mean surface water velocity (ms⁻¹) | Predator presence | Proportion of surface invertebrates (%) | Invertebrate abundance | Green filamentous macroalgae (% cover) | Mean turbidity (NTU) |
|-----------------------------|--------------|------------|-----------|--------------|-----------------|-------------------------------------|-------------------|----------------------------------------|------------------------|----------------------------------------|---------------------|
| Outflow Creek (OC)          | Lower Fortescue | −21.5762  | 117.0860  | 25           | 4               | 0.20 ± 0.03                        | N                 | 88.89                                  | 9                      | 4                                      | 1.07 ± 0.03         |
| Deep Reach (DR)             | Lower Fortescue | −21.6104  | 117.1074  | 17           | 1               | 0.00 ± 0.00                        | Y                 | NA                                     | NA                     | 0                                      | 0.84 ± 0.01         |
| Jindawurranha Channel (JC)  | Lower Fortescue | −21.5904  | 117.0698  | 27           | 4               | 0.50 ± 0.00                        | N                 | 33.33                                  | 3                      | 0                                      | 0.15 ± 0.00         |
| Palm Pool (PP)              | Lower Fortescue | −21.5702  | 117.0536  | 30           | 3               | 0.33 ± 0.11                         | N                 | 0.00                                   | 10                     | 0                                      | 0.49 ± 0.00         |
| GB Creek (GB)               | Lower Fortescue | −21.5813  | 117.5813  | 29           | 2               | 0.00 ± 0.00                         | Y                 | 66.67                                  | 3                      | 0                                      | 0.24 ± 0.01         |
| Angular Pool (AP)           | Mid-Fortescue  | −22.4772  | 118.5631  | 20           | 0               | 0.02 ± 0.02                         | N                 | 42.86                                  | 7                      | 0                                      | 0.15 ± 0.00         |
| Flat Pool (FP)              | Mid-Fortescue  | −22.4776  | 118.5567  | 31           | 3               | 0.14 ± 0.05                         | Y                 | 33.33                                  | 3                      | 0                                      | 0.51 ± 0.01         |
| HD2.5                       | Upper Fortescue | −23.0043  | 119.6213  | 20           | 4               | 0.00 ± 0.00                         | N                 | 25.00                                  | 17                     | 20                                     | 0.39 ± 0.01         |
| HD2                         | Upper Fortescue | −23.0098  | 119.6199  | 16           | 5               | 0.00 ± 0.00                         | N                 | 5.51                                   | 151                    | 10                                     | 0.76 ± 0.01         |
| HD1.5                       | Upper Fortescue | −22.9897  | 119.6218  | 29           | 5               | 0.00 ± 0.00                         | N                 | 22.12                                  | 35                     | 3                                      | 0.28 ± 0.01         |
| Kalgan (K)                  | Upper Fortescue | −23.1873  | 119.6967  | 4            | 1               | 0.04 ± 0.02                         | N                 | 0.00                                   | 22                     | 0                                      | 2.54 ± 0.02         |
| Weeli Wolli 1 (WW1)         | Upper Fortescue | −22.9235  | 119.1953  | 24           | 3               | 0.033 ± 0.00                        | N                 | N/A                                    | N/A                    | 10                                     | 0.26 ± 0.02         |
| Weeli Wolli 2 (WW2)         | Upper Fortescue | −22.9136  | 119.2127  | 20           | 3               | 0.14 ± 0.01                         | N                 | N/A                                    | N/A                    | 2                                      | 0.41 ± 0.01         |
| Weeli Wolli 4 (WW4)         | Upper Fortescue | −22.8827  | 119.2357  | 20           | 3               | 0.33 ± 0.02                         | N                 | N/A                                    | N/A                    | 0                                      | 0.27 ± 0.02         |
Fish capture and photography

Adult western rainbowfish were captured using a 10 m or 4 m seine net (both with a 6 mm mesh), depending on pool size. Juvenile rainbowfish (identified by their small size and absence of body coloration) were not used in the analysis as ontogenetic effects may contribute to variation in body shape (Paez et al. 2008). Following capture, adult fish were placed on their right side and photographed on a perspex slate with a scale bar and a mini Munsell™ Colorchecker (Grand Rapids, MI) photography standard. Photographs were taken using an Olympus™ E-PL3 (Olympus Corporation, Tokyo, Japan) camera placed on a tripod at a fixed focal length to ensure photo uniformity. Photography was performed in the shade, under natural sunlight. Fish sex was determined by placing each fish in a transparent container and examining dorsal and anal fin morphology (Pusey et al. 2004).

Morphometric analyses of body shape

Body shape was quantified using geometric morphometric analyses (Zelditch et al. 2012) and TPS software (available at http://life.bio.sunysb.edu/morph/). Twenty landmarks were assigned to the outline of each individual using TPSDIG v. 2.17 (Rohlf 2006) (Fig. 1). Five of these landmarks were fixed, representing homologous points on the body, and were assigned to the mouth tip, center of the eye, center of the caudal peduncle edge, and on the dorsal and ventral body edges in line with the center of the eye (Fig. 1). A further 15 sliding semilandmarks were assigned to the edge of the body: seven dorsally and eight ventrally (see Fig. 1). These sliding semilandmarks were used to incorporate information about curvature in the subsequent geometric shape analysis. Together, the fixed and sliding semilandmarks produced a comprehensive shape outline that aided both the visual and analytical representation of morphological variation. The relative shape and size of fish in our sample were quantified using relative warps and centroid scores.

Figure 1. Landmark placement used in the morphological analysis of the western rainbowfish, Melanotaenia australis. The black markers represent the fixed landmarks while the white marks are sliding semilandmarks.
respectively, both of which were generated using the program TPSRELW v. 1.45 (Rohlf 2005). Relative warps are principle components that quantify a change in a group of body shape attributes (e.g., streamlined to deep body shape) (Zelditch et al. 2012). Centroid size was used as a measure of body size and was calculated as the square root of the summed squared distances of each landmark from the centroid position (i.e., the average position of the 20 landmarks) (Zelditch et al. 2012). In the absence of allometry, centroid size is uncorrelated with shape; it is therefore possible to test for allometry by assessing the significance of correlations between relative warp scores and centroid size (positive or negative correlations indicate positive and negative allometry, respectively, for any given axis of shape variance). In our analysis, we focused on the first five relative warp scores (i.e., RW1–5), which explained >87% of the total variance in body shape (Fig. 2).

Statistical modeling

A series of linear mixed-effects models combined with model selection procedures were used to evaluate the relative importance of the environmental (predictor) variables on fish morphology (relative warp scores), using the software program R (R Development Core Team 2012). We first used multivariate analysis of covariance (MANCOVA) to examine the effects of the environmental variables on total shape variation, while controlling for centroid size, which was fitted as the covariate. To visualize total variation in shape along the most important environmental axes, we used canonical variate analysis (CVA) in the software program MorphoJ (Klingenberg 2011). After evaluating total shape variation, we conducted a series of univariate tests to examine the effects of the predictor variables and their interactions on each of the relative warp scores separately. We used the lme4 package (Bates and Maechler 2009) to construct multiple models and determine the AIC (Akaike’s information criterion) values of the fit of each model. The AIC values were converted into AICc values to account for the effects of sample size (Symonds and Moussalli 2011). These were used to guide model selection (models with lower AICc values were considered more parsimonious) in combination with model weights \((w_i)\), which can be summed for each variable \((\sum w_i)\), to estimate their relative importance, ranging from 0 to 1 (Symonds and Moussalli 2011). Models with a change in AICc \((\Delta\text{AIC}c) > 10\) relative to the best model were excluded, while those with \(6 < \Delta\text{AIC}c < 10\) were considered unlikely and those with \(\Delta\text{AIC}c < 2\) were considered equal best models (Symonds and Moussalli 2011). A weight of >0.9 and a low \(\Delta\text{AIC}c\) value indicates that other candidate models can be excluded (Symonds and Moussalli 2011).

We considered predictor variables that previous studies suggested were likely to affect morphology (Garamszegi 2011; Richards et al. 2011). Fixed effects included invertebrate abundance, the proportion of surface invertebrates, predator presence, habitat complexity rank, turbidity, surface water velocity, filamentous macroalgae cover, centroid

![Figure 2](image-url)
size (a covariate), and sex. Site nested within subcatchment was included in every model as a random effect to account for sites that were located in the same subcatchment. A number of a priori interactions were also tested, including the interaction between size and sex (morphological differences between the sexes may depend on body size); turbidity and predation (turbidity may reduce predation risk; Snickars et al. 2004); sex and habitat complexity (due to differential habitat use by the sexes; Magellan and Magurran 2006); predation and habitat complexity (complex habitats provide refuges for prey; Walker 1997; Eklov and Svanback 2006; Sass et al. 2006); predation and sex (predators may preferentially target one of the sexes; Godin and McDonough 2003; Moyaho et al. 2004); and water flow and habitat complexity (water flow is affected by habitat; Kilsby and Walker 2012). All models were compared to a null model, which contained only the random effect. We tested the assumptions of the models by inspecting plots of the fitted models against the residual values.

Results

Body shape morphology

There was considerable variation in body shape among fish sampled in this study, which was predominantly attributable to differences in body depth (Table 2). Specifically, RW1, which accounted for >42% of the variance in body shape, described variation in both body depth and the length of the caudal peduncle. RW2 accounted for >19% of variation in body shape and described increased body curvature with a slightly downward-facing head to a relatively cylindrical body and an upward-facing head. In subsequent warps (RW3–RW5), changes in morphology and the proportion of variation explained were relatively minor: RW3 described reduced head length combined with increased peduncle length; RW4 described a range between downward and upward body bending; and RW5 described decreased posterior body depth. Cumulatively, RW3–RW5 explained ~25% of the overall variation in body shape.

Effect of sex and body size

The MANCOVA models revealed that sex was the most important determinant of total shape variation (estimate of effect of sex: $-3.25e-3$, SE = 6.36e-4, $t = -5.12$); models containing all the other predictor variables had AIC values >10 and were therefore considered less plausible. Extraction of the canonical variates along the major environmental axes revealed that for the first canonical variate (CV1), 38.1% of the total variation in shape was associated with water velocity and 48.0% with macroalgae.
cover. Positive CV1 scores were associated with slow water flows and high macroalgae cover and were linked with body deepening, while negative CV1 scores were associated with body narrowing, particularly at the anterior end (Fig. 3). Fish habitats where predators were present displayed anterior narrowing and deeper caudal peduncles (negative CV1) relative to those in habitats without predators (positive CV1) (Fig. 3).

When the sexes were analyzed separately, the top three MANCOVA models (ΔAICc < 10) describing total variation in shape contained the fixed effects of water velocity (males: estimate ± SE = 0.004 ± 0.007, t = 0.53; females: = 0.005 ± 0.006, t = 0.86), macroalgae cover (males: −0.004 ± 0.0001, t = −2.84; females: = −0.0003 ± 0.0001, t = −2.55), and predation risk (males: 0.81e-3 ± 0.003, t = −0.32; females: = −0.0008 ± 0.002, t = −0.39). For males, models containing the parameters water velocity and macroalgae were equally likely (ΔAICc < 2), while for females, water velocity was also included in the best-fitting model (ΔAICc = 0), but models containing predation risk and macroalgae also had some support and both were equally likely (ΔAICc < 2).

In the univariate tests, all of the best-fitting models (ΔAICc < 10) for RW1 included the effect of sex (Σw1 = 1; Table 3). Sex was also an important predictor of variation in RW2, with an overall summed weight (Σw2) of 0.62. The predictor “sex” also appeared in the top models for RW4 and RW5; however, these models were poor candidates due to low weights (RW4: w3 = 0.17; RW5: w3 = 0.02) and relatively large AICcs (ΔAICc RW4: 3.32; RW5: 8.32). In fact, models for RW2, RW4, and RW5 did not significantly differ (ΔAICc < 2) from the null model, which contained only the random effects, suggesting that none of our predictor variables accounted for variation in these body shape attributes. The RW1 scores revealed that males displayed deep bodies, short caudal peduncles, and a curved body, while females tended to have slimmer, bullet-shaped bodies with upward-facing heads. Centroid size was included in all of the top RW1 and RW3 models (ΔAICc < 10), indicating allometry in body depth, head, and peduncle length.

### Environmental predictors

None of the measured environmental fixed effects (predator presence, habitat complexity rank, turbidity, surface water velocity, filamentous macroalgae cover, sex) was included in the top model set for any of the relative warp scores. There was a common reoccurrence of surface water velocity in the suboptimal models for RW1, RW2, RW3, and RW5; however, these models were poor candidates owing to their AICc values (ΔAICc 3.48–9.78) and low weights (0.01–0.15) (Table 3). Predation was present in one of the RW1 models but had low weight (0.01) and an AICc value that was considerably higher (indicating a poorer fit) than the top model (ΔAICc = 9.96). Conducting partial correlations (controlling for centroid size) on the mean relative warp scores with the site means for the environmental parameters revealed a significant negative correlation between both RW2 and RW5 and the mean percentage cover of macroalgae (RW2: r13 = -0.68, P = 0.008; RW5: r13 = -0.69, P = 0.006), and a negative correlation between RW3 and mean water velocity (r13 = 0.53, P = 0.049). This result suggests that increasing macroalgae cover is associated with a downward-facing head and increased body curvature while increased water velocity is linked with changes in head and caudal peduncle length.

### Geographical effects of subcatchment and site

Subcatchment and site were included in all optimal models, and the relative importance of each varied depending on the relative warp (Table 3). Subcatchment explained a
higher proportion of the random variation than site in the optimal models for RW1 (50.6% compared to 19.4% for site) and RW5 (20.9% compared to 15.4% for site). Subcatchment variance was lower (0.0%–12.6%) than (nested) site variance in all other relative warps (14.3%–23.6%) in the optimal models.

RW1 and RW2 formed three distinct morphological groups when plotted according to subcatchment, particularly when considering variation along the RW1 axis (Fig. 4). The mid-Fortescue populations had slim bodies while the lower Fortescue populations tended to have deep, cylindrical bodies with shorter caudal peduncles and upward-facing heads. The upper Fortescue populations tended to have downward-facing heads and increased body curvature relative to the lower and mid subcatchment sites. However, not all populations fell into this distinct pattern. In particular, fish from the Weeli Wolli Creek sites (which have received continuous artificial discharge from mine de-watering since 2007; Dogramaci et al. 2015) tended to be morphologically more similar to those in the mid-Fortescue populations (i.e., slim-bodied) than those from other sites in the upper Fortescue (Fig. 5).

**Discussion**

We found that western rainbowfish exhibited considerable variation in body shape and marked sexual dimorphism, with males displaying deeper bodies and shorter caudal peduncles than females. Geographic patterns of morphological differentiation were largely consistent with the species’ distribution among discrete subcatchments. Nevertheless, some populations did not conform to this geographic pattern and levels of morphological similarity among subcatchments were not consistent with the hierarchical stream order (i.e., pool – creek – subcatchment – catchment). The environmental factors measured in this study, including water velocity, habitat complexity, the presence of predators, and food availability, played only a minor role in explaining overall variation in body shape. Consequently, our findings suggest that, at the population
level, variation in habitat characteristics may not be the main driver of phenotypic diversity; instead, the morphological variability of the western rainbowfish may reflect underlying patterns of genetic differentiation and sex-specific selective pressures.

Our findings contrast with previous studies that have documented a strong effect of water velocity (Brinsmead and Fox 2002; Langerhans et al. 2003; Aguirre 2009; Franssen 2011; Drinan et al. 2012), predation pressure (Basolo and Wagner 2004; Langerhans and DeWitt 2004; Langerhans et al. 2007), and habitat type and diet (Hjelm et al. 2001; Berner et al. 2008; Svanback et al. 2008; Aguirre 2009) on the morphology of wild-caught fishes. A possible explanation for this point of difference is that we did not measure the morphology of the fins, which are important determinants of thrust, drag, and maneuverability while swimming (Plaut 2000; Drucker and Lauder 2001; Nauen and Lauder 2002). For example, caudal fins provide thrust and counteract drag while swimming. Thus, the height of the caudal fin is often increased in fish that are exposed to fast-flowing water (Pakkasmaa and Piironen 2000; Brinsmead and Fox 2002; Imre et al. 2002). We did not measure the fins because extending and positioning the fins for photography is difficult on live, nonanaesthetized fish.

Previous studies have revealed that freshwater fishes commonly exhibit sexual dimorphism for both body size and shape (e.g., Quinn and Foote 1994; Caldecutt et al. 2001; McGuigan et al. 2003; Hendry et al. 2006). Our analysis of body shape supports prior observations that male rainbowfish tend to be deeper-bodied than females (Allen et al. 2002). Sexual selection via female choice and male–male competition commonly explains sexual dimorphism, and both of these processes operate in *M. australis* (Young et al. 2010). Although female western rainbowfish prefer large males, it would be interesting to investigate whether there is also sexual selection on body shape and whether this varies among populations (Head et al. 2013). It would also be worthwhile investigating the functional significance of the observed sexual dimorphism; a previous study with other species of rainbowfishes found that variation in body shape was not attributed to hydrological habitat, but the sexes differed in critical swim speeds (McGuigan et al. 2003). These findings suggest that environmental factors such as water flow may lead to differential habitat use by the sexes and sexual segregation.

Sexual dimorphism may also arise due to natural selection operating differentially on the sexes. The sexes are often exposed to different levels of predation risk, for example, because bright colors that attract females also increase the males’ conspicuousness to visual predators (Godin and McDonough 2003; Moyaho et al. 2004). Male rainbowfish tend to be more brightly colored than females (Young et al. 2011b) and may therefore compensate for their increased risk of predation by developing deeper bodies that provide protection from gape-limited predators (Nilsson et al. 1995; Domenici et al. 2008). However, in this study, predator presence was associated with an overall narrowing of the anterior of the body while predator absence was associated with body deepening. This finding might be an adaptation for fast-start escape.
responses from predators because a narrow head should reduce drag while a deep caudal peduncle should increase thrust (Langerhans and DeWitt 2004). We also found some evidence for an effect of macroalgae cover on increased body deepening and a downward-facing head, which could be advantageous for prey capture efficiency and navigation in complex habitats (Blake et al. 2005; Eklov and Svanback 2006; Sass et al. 2006; Svanback and Eklov 2006). In short, differences between the sexes in morphological traits may reflect a complex interplay between natural and sexual selection that ultimately favor different body morphologies in males and females.

Most of the variation in our morphological data was explained by geographic (subcatchment) effects, which might be explained by underlying genetic differentiation or the distinctive habitat characteristics of each region, or an interaction between the two. A recent study of the western rainbowfish found that morphological variation across the species’ range is consistent with a hierarchical pattern of genetic divergence (Young et al. 2011a), lending support to the notion that, on a regional scale (among drainages), local adaptation is the main process driving phenotypic differentiation and may have contributed to the evolution of body shape polymorphism in
this species (Young et al. 2011a). Wet season dispersal is considered to be an important determinant of genetic structuring in M. australis, particularly over small spatial scales such as within creek lines (Phillips et al. 2009). In the present study, sites that are located within the same creek lines are in relatively close proximity (within 40 km) and likely to become connected during summer rainfall (Fellman et al. 2011). In contrast, subcatchments are isolated by large distances (~180 km), with the upper subcatchment being separated from the mid and lower subcatchments by the Goodiadarrie Hills (Barnett and Commander 1985; Skrzypek et al. 2013). Nonetheless, recent studies of the biogeography of the Pilbara, an area of some 500,000 km², have revealed that patterns of genetic diversification are often not consistent with biogeographical region, particularly at a fine spatial scale (Pepper et al. 2013). Indeed, there is increasing evidence that the Hamersley Ranges of the Pilbara (encompassing the upper subcatchment of our study) is characterized by high levels of endemism and “cryptic diversity” of both animal and plant species (Cracraft 1991; Unmack 2001).

Aspects of our study have highlighted some of the challenges of conducting research on fishes inhabiting arid and remote regions; sample sites are often sporadic and separated by hundreds of kilometers, which means that sample size is necessarily limited. In the current study, 14 sample sites were split across three distinct geographic subcatchments. We considered the geographical features of the landscape using subcatchment as a random effect; however, the effect of region may have masked any responses to the environmental parameters, particularly because not all habitat types were represented in each region. This may have limited our ability to detect any environmental effects or their interactions. Hydrological conditions in the study region are also typically highly variable among years, leading to unpredictable environmental conditions and strong temporal effects on pool connectivity. The observed morphological variation we observed may therefore reflect past ecological conditions, that is, hydrological events occurring during the fish’s development, which would encompass periods of several months, rather than necessarily the conditions at the time of sampling. Indeed, a lack of morphological differentiation in pumpkinseed fish (Lepomis gibbosus) has been attributed to the strong seasonal variation in water flows that occurs in the Mediterranean, suggesting that dynamic environments may confound morphological responses (Naspleda et al. 2012). Furthermore, flow at one of the creeks in this study (Weeli Wolli Creek; encompassing three sample sites) has been modified for ~7 years due to mining activities in the area. This previously ephemeral creek now has continuous surface flows for ~24 km due to the discharge of groundwater (Dogramaci et al. 2015). Interestingly, fish from these sites tended to possess slender bodies relative to other populations within their (upper) subcatchment, perhaps due to the occurrence of relatively fast-flowing water (>0.4 m·s⁻¹) at this site. This finding is consistent with models of fish swimming biomechanics; fish exposed to fast-flowing water are expected to develop fusiform body shapes for optimal steady swimming performance while those in low flows should develop deeper bodies that maximize thrust and stability for “burst” swimming (Langerhans 2008). Although previous fish studies have reported morphological responses to anthropogenic modifications such as impoundment (Haas et al. 2010; Franssen 2011; Franssen et al. 2012, 2013), further sampling of streams receiving high rates of discharge is required to determine whether the apparent morphological response observed here can be generalized.

In summary, we found limited evidence that body shape variation in M. australis corresponded with strong habitat differentiation or in response to differences in key environmental factors such as water flow. However, patterns of morphological differentiation were largely consistent within three geographically distinct subcatchments, suggesting that these regions may present distinct and isolated habitats that may promote differentiated patterns of morphology. Nonetheless, fine-scale genetic studies are required to determine whether patterns of genetic structuring are consistent with the geographic features of the Pilbara landscape and its hydrological connectivity. We found some evidence that fish have altered their morphology in a creek affected by mine dewatering; however, further study is required before we can establish whether this provides an example of morphological responses to anthropogenic habitat alteration (Franssen 2011; Franssen et al. 2012, 2013).

Data accessibility

Data will be made publicly available on the general-purpose repository Dryad (https://datadryad.org).

Acknowledgments

We are very grateful to S. Luccitti, S. Wild, and J. Iles for field assistance and logistical support. A. Storey, J. Delaney, A. Siebers, D. Ford, G. Skrzypek, E. Skrzypek, M. Donn, J. Hehre, and K. Bowler provided valuable help with the collection and processing of data, while M. Renton and G. Page gave statistical advice. We would also like to acknowledge the support provided by the rangers at Karijini and Millstream National Parks. This manuscript was greatly improved as a result of comments provided by an anonymous reviewer. The project was approved by the UWA...
Animal Ethics Committee, approval number RA/3/100/1176. This project was funded by an Australian Research Council Linkage Grant LP120200002 with industry partners Rio Tinto and BHP Billiton.

**Conflict of Interest**

None declared.

**References**

Aguirre, W. E. 2009. Microgeographical diversification of threespine stickleback: body shape-habitat correlations in a small, ecologically diverse Alaskan drainage. Biol. J. Linn. Soc. 98:139–151.

Alexander, H. J., J. S. Taylor, S. S. Å. Wu, and F. Breden. 2006. Parallel evolution and vicariance in the guppy (Poecilia reticulata) over multiple spatial and temporal scales. Evolution 60:2352–2369.

Allen, G. R., S. H. Midgley, and M. Allen. 2002. Field guide to the freshwater fishes of Australia. Western Australian Museum, Perth.

Andersson, J., F. Johansson, and T. Soderlund. 2006. Interactions between predator- and diet-induced phenotypic changes in body shape of crucian carp. Proc. Biol. Sci. 273:431–437.

Barnett, J. C., and D. P. Commander. 1985. Hydrogeology of the western Fortescue Valley, Pilbara region, Western Australia. Western Australia Geological Survey, Perth.

Basolo, A. L., and W. E. Wagner. 2004. Covariation between predation risk, body size and fin elaboration in the green swordtail, Xiphophorus helleri. Biol. J. Linn. Soc. 83:87–100.

Bates, D., M. Maechler and B. Bolker. 2009. lm4: Linear mixed-effects models using [S4] classes. [R] package version 0.999375-32, http://CRAN.R-Project.org/package.

Beesley, L. S., and J. Prince. 2010. Fish community structure in an intermittent river: the importance of environmental stability, landscape factors and within-pool habitat descriptors. Mar. Freshw. Res. 61:605–614.

Berner, D., D. C. Adams, A. C. Grandchamp, and A. P. Hendry. 2008. Natural selection drives patterns of lake-stream divergence in stickleback foraging morphology. J. Evol. Biol. 21:1653–1665.

Blake, R. W., T. C. Law, K. H. S. Chan, and J. F. Z. Li. 2005. Comparison of the prolonged swimming performances of closely related, morphologically distinct three-spined sticklebacks Gasterosteus spp. J. Fish Biol. 67:834–848.

Blake, R. W., J. Li, and K. H. S. Chan. 2009. Swimming in four goldfish Carassius auratus morphotypes: understanding functional design and performance employing artificially selected forms. J. Fish Biol. 75:591–617.

Blob, R. W., S. M. Kawano, K. N. Moody, W. C. Bridges, T. Maie, M. B. Ptacek, et al. 2010. Morphological selection and the evaluation of potential tradeoffs between escape from predators and the climbing of waterfalls in the Hawaiian stream goby Sicyopterus stimpsoni. Integr. Comp. Biol. 50:1185–1199.

Brinsmead, J., and M. G. Fox. 2002. Morphological variation between lake- and stream-dwelling rock bass and pumpkinseed populations. J. Fish Biol. 61:1619–1638.

Bronmark, C., and J. G. Miner. 1992. Predator-induced phenotypical change in body morphology in crucian carp. Science 258:1348–1350.

Bronmark, C., and L. B. Pettersson. 1994. Chemical cues from piscivores induce a change in morphology in crucian carp. Oikos 70:396–402.

Caldecutt, W. J., M. A. Bell, and J. A. Buckland-Nicks. 2001. Sexual dimorphism and geographic variation in dentition of threespine stickleback, Gasterosteus aculeatus. Copeia 2001:936–944.

Chivers, D., X. Zhao, G. Brown, T. Marchant, and M. O. Ferrari. 2008. Predator-induced changes in morphology of a prey fish: the effects of food level and temporal frequency of predation risk. Evol. Ecol. 22:561–574.

Cracraft, J. 1991. Patterns of diversification within continental biotas: hierarchical congruence among the areas of endemism of Australian vertebrates. Aust. Syst. Bot. 4:211–227.

Dogramaci, S., P. Hedley, G. Firmani, G. Skrzypek, and P. F. Grierson. 2015. Evaluating recharge to an ephemeral dryland stream using a hydraulic model and water isotope-chlorine mass balance. J. Hydrol. 521:520–532.

Domenici, P., H. Turesson, J. Brodersen, and C. Brönmark. 2008. Predator-induced morphology enhances escape locomotion in crucian carp. Proceed. Royal Soc. B: Biol. Sci. 275:195–201.

Drinan, T. J., P. McGinnity, J. P. Coughlan, T. F. Cross, and S. S. Harrison. 2012. Morphological variability of Atlantic salmon Salmo salar and brown trout Salmo trutta in different river environments. Ecol. Freshw. Fish 21:420–432.

Drucker, E. G., and G. V. Lauder. 2001. Locomotor function of the dorsal fin in teleost fishes: experimental analysis of wake forces in sunfish. J. Exp. Biol. 204:2943–2958.

Eklov, P., and P. Jonsson. 2007. Pike predators induce morphological changes in young perch and roach. J. Fish Biol. 70:155–164.

Eklov, P., and R. Svanback. 2006. Predation risk influences adaptive morphological variation in fish populations. Am. Nat. 167:440–452.

Enders, E. C., D. Boisclair, and A. G. Roy. 2003. The effect of turbulence on the cost of swimming for juvenile Atlantic salmon (Salmo salar). Can. J. Fish. Aquat. Sci. 60:1149–1160.

Endler, J. A. 1983. Natural and sexual selection on color patterns in Poeciliid fishes. Environment. Biol. Fish 9:173–190.

Endler, J. A. 1986. Natural selection in the wild. Princeton Univ. Press, Princeton, NJ.
Fellman, J. B., S. Dogramaci, G. Skrzypek, W. Dodson, and P. F. Grierson. 2011. Hydrologic control of dissolved organic matter biochemistry in pools of a subtropical dryland river. Water Resour. Res. 47:W06501.

Forsgren, E. 1992. Predation risk affects mate choice in a gobiid fish. Am. Nat. 140:1041–1049.

Foster, S. A., and J. A. Endler. 1999. Geographic variation in behavior: perspectives on evolutionary mechanisms. Oxford Univ. Press, New York.

Franssen, N. R. 2011. Anthropogenic habitat alteration induces rapid morphological divergence in a native stream fish. Evol. Appl. 4:791–804.

Franssen, N. R., J. Harris, S. R. Clark, J. F. Schaefer, and L. K. Stewart. 2012. Shared and unique morphological responses of stream fishes to anthropogenic habitat alteration. Proceed. Royal Soc. B: Biol. Sci. 280:21022715.

Franssen, N. R., L. K. Stewart, and J. F. Schaefer. 2013. Morphological divergence and flow-induced phenotypic plasticity in a native fish from anthropogenically altered stream habitats. Ecol. Evol. 3:4648–4657.

Garamszegi, L. 2011. Information-theoretic approaches to statistical analysis in behavioural ecology: an introduction. Behav. Ecol. Sociobiol. 65:1–11.

Godin, J.-G. J., and H. E. McDonough. 2003. Predator preference for brightly colored males in the guppy: a viability cost for a sexually selected trait. Behav. Ecol. 14:194–200.

Haas, T. C., M. J. Blum, and D. C. Heins. 2010. Morphological responses of a stream fish to water impoundment. Biol. Lett. 6:803–806.

Harrod, C., J. Mallela, and K. K. Kahlilainen. 2010. Phenotype-environment correlations in a putative whitefish adaptive radiation. J. Anim. Ecol. 79:1057–1068.

Head, M. L., G. M. Kozak, and J. W. Boughman. 2013. Female mate preferences for male body size and shape promote sexual isolation in threespine sticklebacks. Ecol. Evol. 3:2183–2196.

Hendry, A. P. 2009. Ecological speciation! Or the lack thereof? This perspective is based on the author’s J.C. Stevenson Memorial Lecture delivered at the Canadian Conference for Fisheries Research in Halifax, Nova Scotia, January 2008. Can. J. Fish. Aquat. Sci. 66:1383–1398.

Hendry, A. P., M. L. Kelly, M. T. Kinnison, and D. N. Reznick. 2006. Parallel evolution of the sexes? Effects of predation and habitat features on the size and shape of wild guppies. J. Evol. Biol. 19:741–754.

Hjelm, J., R. Svanback, P. Bystrom, L. Persson, and E. Wahlstrom. 2001. Diet-dependent body morphology and ontogenetic reaction norms in Eurasian perch. Oikos 95:311–323.

Imre, I., R. L. McLaughlin, and D. L. G. Noakes. 2002. Phenotypic plasticity in brook char: changes in caudal fin induced by water flow. J. Fish Biol. 61:1171–1181.

Kilsby, N. N., and K. F. Walker. 2012. Behaviour of two small pelagic and demersal fish species in diverse hydraulic environments. River Res. Appl. 28:543–553.

Klingenberg, C. P. 2011. MorphoJ: An integrated software package for geometric morphometrics. Mol. Ecol. Resour. 11:353–357.

Kristjansson, B. K., S. Skulason, S. S. Snorraslon, and D. L. Noakes. 2012. Fine-scale parallel patterns in diversity of small benthic Arctic char (Salvelinus alpinus) in relation to the ecology of lava/groundwater habitats. Ecol. Evol. 2:1099–1112.

Langerhans, R. B. 2008. Predictability of phenotypic differentiation across flow regimes in fishes. Integr. Comp. Biol. 48:750–768.

Langerhans, R. B. 2009. Trade-off between steady and unsteady swimming underlies predator-driven divergence in Gambusia affinis. J. Evol. Biol. 22:1057–1075.

Langerhans, R. B., and T. J. DeWitt. 2004. Shared and unique features of evolutionary diversification. Am. Nat. 164:335–349.

Langerhans, R. B., and A. M. Makowicz. 2009. Shared and unique features of morphological differentiation between predator regimes in Gambusia caymanensis. J. Evol. Biol. 22:2231–2242.

Langerhans, R. B., C. A. Layman, A. K. Langerhans, and T. J. Dewitt. 2003. Habitat-associated morphological divergence in two neotropical fish species. Biol. J. Linn. Soc. 80:689–698.

Langerhans, R. B., L. J. Chapman, and T. J. Dewitt. 2007. Complex phenotype-environment associations revealed in an east African cyprinid. J. Evol. Biol. 20:1171–1181.

Law, T., and R. Blake. 1996. Comparison of the fast-start performances of closely related, morphologically distinct threespine sticklebacks (Gasterosteus spp.). J. Exp. Biol. 199:2595–2604.

Losos, J. B., T. R. Jackman, A. Larson, K. de Queiroz, and L. Rodriguez-Schettino. 1998. Contingency and determinism in replicated adaptive radiations of island lizards. Science 279:2115–2118.

Magellan, K., and A. E. Magurran. 2006. Habitat use mediates the conflict of interest between the sexes. Anim. Behav. 72:75–81.

Magnhagen, C., and E. Heibo. 2001. Gape size allometry in pike reflects variation between Lakes in prey availability and relative body depth. Funct. Ecol. 15:754–762.

Marchinko, K. B. 2009. Predation’s role in repeated phenotypic and genetic divergence of armor in threespine stickleback. Evolution 63:127–138.

Maynard Smith, J. 1966. Sympatric speciation. Am. Nat. 100:637–650.

McCa irns, R. J., and L. Bernatchez. 2012. Plasticity and heritability of morphological variation within and between parapatric stickleback demes. J. Evol. Biol. 25:1097–1112.
McGuigan, K., D. Zhu, G. R. Allen, and C. Moritz. 2000. Phylogenetic relationships and historical biogeography of melanotaenid fishes in Australia and New Guinea. Mar. Freshw. Res. 51:713–723.

McGuigan, K., C. E. Franklin, C. Moritz, and M. W. Blows. 2003. Adaptation of rainbow fish to lake and stream habitats. Evolution 57:104–118.

Morgan, D. L., and H. S. Gill. 2004. Fish fauna in inland waters of the Pilbara (Indian Ocean) Drainage Division of Western Australia — evidence for three subprovinces. Zootaxa 636:1–43.

Moyaho, A., C. M. Garcia, and J. Manjarrez. 2004. Predation risk is associated with the geographic variation of a sexually selected trait in a viviparous fish (Xenotoca variata). J. Zool. 262:265–270.

Nagel, L., and D. Schluter. 1998. Body size, natural selection, and speciation in sticklebacks. Evolution 52:209–218.

Naspleda, J., A. Vila-Gispert, M. G. Fox, L. Zamora, and A. Ruiz-Navarro. 2012. Morphological variation between non-native lake- and stream-dwelling pumpkinseed Lepomis gibbosus in the Iberian Peninsula. J. Fish Biol. 81:1915–1935.

Nauern, J. C., and G. V. Lauder. 2002. Hydrodynamics of caudal fin locomotion by chub mackerel, Scomber japonicus (Scombridae). J. Exp. Biol. 205:1709–1724.

Nilsson, P. A., C. Bronmark, and L. B. Pettersson. 1995. Benefits of a predator-induced morphology in crucian carp. Oecologia 104:291–296.

Paez, D. J., R. Hedger, L. Bernatchez, and J. J. Dodson. 2008. The morphological plastic response to water current velocity varies with age and sexual state in juvenile Atlantic salmon, Salmo salar. Freshw. Biol. 53:1544–1554.

Pakkasmaa, S., and J. Piironen. 2000. Water velocity shapes juvenile salmonids. Evol. Ecol. 14:721–730.

Pepper, M., P. Doughty, and J. S. Keogh. 2013. Geodiversity and endemism in the iconic Australian Pilbara region: a review of landscape evolution and biotic response in an ancient refugium. J. Biogeogr. 40:1225–1239.

Peres-Neto, P. R., and P. Magnan. 2004. The influence of swimming demand on phenotypic plasticity and morphological integration: a comparison of two polymorphic char species. Oecologia 140:36–45.

Phillips, R. D., A. W. Storey, and M. S. Johnson. 2009. Genetic structure of Melanotaenia australis at local and regional scales in the east Kimberley, Western Australia. J. Fish Biol. 74:437–451.

Plaut, I. 2000. Effects of fin size on swimming performance, swimming behaviour and routine activity of zebrafish Danio rerio. J. Exp. Biol. 203:813–820.

Price, T. D., A. Qvarnstrom, and D. E. Irwin. 2003. The role of phenotypic plasticity in driving genetic evolution. Proc. Biol. Sci. 270:1433–1440.

Proulx, R., and P. Magnan. 2004. Contribution of phenotypic plasticity and heredity to the trophic polymorphism of lacustrine brook charr (Salvelinus fontinalis M.). Evol. Ecol. Res. 6:503–522.

Pusey, B. J., M. J. Kennard, and A. H. Arthington. 2004. Freshwater fishes of north-eastern Australia. CSIRO Publishing, Collingwood, Victoria.

Quinn, T. P., and C. J. Foote. 1994. The effects of body size and sexual dimorphism on the reproductive behaviour of sockeye salmon, Oncorhynchus nerka. Anim. Behav. 48:751–761.

R Development Core Team. 2012. A language and environment for statistical computing. R Foundation for Statistical Computing, Vienna, Austria.

Reimchen, T. E. 1994. Predators and morphological evolution in threespine stickleback. Pp. 240–276 in M.A. Bell and S. A. Foster, eds. The evolutionary biology of the threespine stickleback. Oxford University Press, Oxford.

Reznick, D. 1982. The impact of predation on life history evolution in Trinidadian guppies: genetic basis of observed life history patterns. Evolution, 36:1236–1250.

Reznick, D., and J. A. Endler. 1982. The impact of predation on life history evolution in Trinidadian guppies (Poecilia reticulata). Evolution 36:160–177.

Richards, S. A., M. J. Whittingham, and P. A. Stephens. 2011. Model selection and model averaging in behavioural ecology: the utility of the IT-AIC framework. Behav. Ecol. Sociobiol. 65:77–89.

Robinson, B. W., D. S. Wilson, and G. O. Shea. 1996. Trade-offs of ecological specialization: an intraspecific comparison of pumpkinseed sunfish phenotypes. Ecology 77:170–178.

Rohlf, F. J. 2005. tpsRelw, relative warp analysis, version 1.45. Stony Brook University, New York.

Rohlf, F. J. 2006. TpsDig. Department of Ecology and Evolution, State University, New York.

Royle, N. J., N. B. Metcalfe, and J. Lindström. 2006. Sexual selection, growth compensation and fast-start swimming performance in Green Swordtails, Xiphophorus helleri. Funct. Ecol. 20:662–669.

Rundle, H. D., and P. Nosil. 2005. Ecological speciation. Ecol. Lett. 8:336–352.

Sass, G. G., C. M. Gille, J. T. Hinke, and J. F. Kitchell. 2006. Whole-lake influences of littoral structural complexity and prey body morphology on fish predator–prey interactions. Ecol. Freshw. Fish 15:301–308.

Schafer, J., W. Lutterschmidt, and L. Hill. 1999. Physiological performance and stream microhabitat use by the Centrarchids Lepomis megalotis and Lepomis macrochirus. Environ. Biol. Fishes 54:303–312.

Schluter, D. 1996. Ecological speciation in postglacial fishes. Philos. Trans. Royal Soc. London Series B-Biol. Sci. 351:807–814.

Schluter, D. 2000. The ecology of adaptive radiation. Oxford Univ. Press, Oxford, U.K.

Seghers, B. H. 1974. Schooling behaviour in the guppy (Poecilia reticulata): an evolutionary response to predation. Evolution 28:486–489.
Skrzypek, G., S. Dogramaci, and P. F. Grierson. 2013. Geochemical and hydrological processes controlling groundwater salinity of a large inland wetland of northwest Australia. Chem. Geol. 357:164–177.

Skulason, S., and T. B. Smith. 1995. Resource polymorphisms in vertebrates. Trends Ecol. Evol. 10:366–370.

Snickars, M., A. Sandström, and J. Mattila. 2004. Antipredator behaviour of 0+ year Perca fluviatilis: effect of vegetation density and turbidity. J. Fish Biol. 65:1604–1613.

Svanback, R., and P. Eklov. 2006. Genetic variation and phenotypic plasticity: causes of morphological and dietary variation in Eurasian perch. Evol. Ecol. Res. 8:37–49.

Svanback, R., P. Eklov, R. Fransson, and K. Holmgren. 2008. Intraspecific competition drives multiple species resource polymorphism in fish communities. Oikos 117:114–124.

Symonds, M. R. E., and A. Moussalli. 2011. A brief guide to model selection, multimodel inference and model averaging in behavioural ecology using Akaike’s information criteria. Behav. Ecol. Sociobiol. 65:13–21.

Tobler, M., T. J. DeWitt, I. Schlupp, F. J. García de León, R. Herrmann, P. G. D. Feulner, et al. 2008. Toxic hydrogen sulfide and dark caves: phenotypic and genetic divergence across two abiotic environmental gradients in Poecilia mexicana. Evolution 62:2643–2659.

Unmack, P. J. 2001. Biogeography of Australian freshwater fishes. J. Biogeogr. 28:1053–1089.

Wade, M. J., and S. Kalisz. 1990. The causes of natural selection. Evolution 44:1947–1955.

Walker, J. A. 1997. Ecological morphology of lacustrine threespine stickleback Gasterosteus aculeatus L. (Gasterosteidae) body shape. Biol. J. Linn. Soc. 61:3–50.

Wolfgang, M. J., J. M. Anderson, M. A. Grosenbaugh, D. K. Yue, and M. S. Triantafyllou. 1999. Near-body flow dynamics in swimming fish. J. Exp. Biol. 202:2303–2327.

Young, M., L. Simmons, and J. Evans. 2010. Pre- and post-mating sexual selection both favor large males in a rainbowfish. Behav. Ecol. Sociobiol. 64:915–925.

Young, M. J., J. P. Evans, and L. W. Simmons. 2011a. Population genetic structure and a possible role for selection in driving phenotypic divergence in a rainbowfish (Melanotaeniidae). Biol. J. Linn. Soc. 102:144–160.

Young, M. J., L. W. Simmons, and J. P. Evans. 2011b. Predation is associated with variation in colour pattern, but not body shape or colour reflectance, in a rainbowfish (Melanotaenia australis). J. Anim. Ecol. 80:183–191.

Zelditch, M. L., D. L. Swiderski, and H. D. Sheets. 2012. Geometric morphometrics for biologists: a primer, 2nd ed. Academic Press, London.

Zimmerman, M. S. 2007. A field study of brook stickleback morphology: multiple predators and multiple traits. Can. J. Zool. 85:250–260.