Color and behavior differently predict competitive outcomes for divergent stickleback color morphs

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Abstract

Our knowledge of how male competition contributes to speciation is dominated by investigations of competition between within-species morphs or closely related species that differ in conspicuous traits expressed during the breeding season (e.g. color, song). In such studies, it is important to consider the manner in which putatively sexually selected traits influence the outcome of competitive interactions within and between types because these traits can communicate information about competitor quality and may not be utilized by homotypic and heterotypic receivers in the same way. We studied the roles of breeding color and aggressive behaviors in competition within and between two divergent threespine stickleback Gasterosteus aculeatus color types. Our previous work in this system showed that the switch from red to black breeding coloration is associated with changes in male competition biases. Here, we find that red and black males also use different currencies in competition. Winners of both color types performed more aggressive behaviors than losers, regardless of whether the competitor was of the same or opposite color type. But breeding color differently predicted competitive outcomes for red and black males. Males who were redder at the start of competition were more likely to win when paired with homotypic competitors and less likely to win when paired with heterotypic competitors. In contrast, black color, though expressed in the breeding season and condition dependent, was unrelated to competitive outcomes. Placing questions about the role of male competition in speciation in a sexual signal evolution framework may provide insight into the “why and how” of aggression biases and asymmetries in competitive ability between closely related morphs and species.

Key words: male competition, speciation, sexual signal, color, threespine stickleback, Gasterosteus aculeatus

Darwin (1859, 1871) reasoned that intrasexual selection would lead to differential fitness outcomes when males varied in traits that enhance success in competition for mates or mating resources (Darwin 1871; Simmons 2001). Indeed, male competition is an important determinant of reproductive success. This is particularly true in systems in which males compete for access to pools of mating resources such as harems of females or territories (e.g., LeBouf 1974; Alatalo et al. 1986), and when male competition and female choice jointly determine mating outcomes (as when the two act simultaneously or sequentially, and when mate choice is based on male dominance; Wong and Candolin 2005; Hunt et al. 2009). Despite this, there has been a strong and persistent bias in the sexual selection literature toward investigating the role of female choice, rather than male competition, in driving diversification of sexually selected traits (McCallough et al. 2016) and generating reproductive isolation (Panhuis et al. 2001; Seehausen and Schluter 2004; Qvarnstrom et al. 2012; Tinghitella et al. in press). This is likely because early sexual selection work focused on mate choice (the more controversial of Darwin’s mechanisms of sexual selection; Anderson 1994), male ornaments are commonly correlated with female preferences and female sensory perception, implicating mate choice in ornament evolution (e.g. Boughman 2001; Mendelson and Shaw 2002; Rodriguez et al. 2006), and because there is a direct relationship between accepting or rejecting a mate and reproductive isolation (Qvarnstrom et al. 2012).
Evidence is accumulating in support of a role for male competition in speciation, however. For instance, recent work sheds light on how competitive phenotypes and competitor recognition coevolve (e.g., Martin and Mendelson 2016; Scordato 2017). Male competitive strategies (anything that leads to success in competition in a particular environment) can come to reflect variation in the ecological and social environments (predators and parasites, competitor densities, mating resources, habitat patchiness, and transmission properties of the environment, mate availability) with the result that well-adapted males can be socially dominant and gain access to mates. When divergent types come into contact, male competition researchers often find evidence of aggression biases (e.g., Dijkstra et al. 2006, 2007; Tyers and Turner 2013; Lackey and Boughman 2013; Lehtonen 2014; Tinghitella et al. 2015) and asymmetries in competitive ability (e.g., Owen-ASHLEY and Butler 2004; Dijkstra et al. 2005; Pryke 2009; Winkelmann et al. 2014; Martin et al. 2017). Each could contribute to reproductive isolation, though the consequences for hybridization and speciation are not well characterized. For instance, aggression biases may 1) facilitate co-existence of multiple morphs/species in sympathy through negative frequency dependent selection if aggression is directed toward homotypic males (Seehausen and Schluter 2004) or 2) lead to competitive exclusion (Hardin 1960) and habitat isolation if aggression is biased toward heterotypics (e.g., Peiman and Robinson 2007). Asymmetries in dominance can also facilitate habitat and reproductive isolation if one competitor is able to displace the other, compelling the latter to use less ideal habitat (Winkelmann et al. 2014) or drive local extinction of the lesser competitor. Previous work has nicely identified these patterns and their implications for speciation, but less attention has been paid to why those patterns occur from a sexual signal evolution (signalers/receivers) perspective. Here, we illuminate the “how and why” of aggression biases and asymmetries.

Many of the study systems in which the role of male competition in speciation has been addressed are systems in which alternate color morphs are maintained within species (e.g., Kissenda Island cichlids (Dijkstra et al. 2005; Dijkstra et al. 2009), sticklebacks (Tinghitella et al. 2015; Bolnick et al. 2016), European wall-lizards (While et al. 2015), or in which color is a primary phenotype that differs between closely related species [e.g., lake Victoria cichlids (Seehausen and Schluter 2004); Nicaraguan cichlids (Lehtonen et al. 2015); darters (Martin and Mendelson 2016; Roberts and Mendelson 2017); ruby-spot damselflies (Anderson and Grether 2010a, 2010b); limnetic-benthic stickleback species pairs (Lackey and Boughman 2013; Keagy et al. 2016)]. In a handful of these systems, variation in the sensory environment is implicated in color shifts (Reimchen 1989; Boughman 2001; Scott 2001; Maan et al. 2006; Seehausen et al. 2008). Environment-associated differences in color can influence male aggressive responses to heterotypic competitors (e.g., Macedonia et al. 1994; Macedonia and Stamps 2010; Macedonia et al. 2013, 2015; Tinghitella et al. 2015; Bolnick et al. 2016), as identifying appropriate competitors is a first step in competitive interactions. If agonistic color signal evolution is hypothesized to contribute to reproductive isolation, it is important to understand how divergent colors function in male competition (the extent to which and which components of putative signals dictate competitive outcomes) and whether the relationship between signals and competitive outcomes depends on the competitor (receiver) type.

We tested whether two threespine stickleback *Gasterosteus aculeatus* color types have diverged in the manner in which male-limited breeding color is associated with male competition outcomes. The study system is one in which male competition interacts with female choice to influence reproductive success and male color functions in both male competition and female choice contexts (e.g., Candolin 1999). During the breeding season, most male threespine sticklebacks *Gasterosteus aculeatus*, including the marine ancestors of the freshwater fish studied here, express a carotenoid-based red throat that can extend from the mouth to the pelvic spines. The red throat is strongly preferred by females in mate choice because it is correlated with health, condition, and parental care ability (e.g., Milinski and Bakker 1990; Frischknecht 1993; Candolin 2000; Boughman 2007; Kurtz et al. 2007) and is also used in male competition (reviewed in Bakker and Sevenster 1983; McLennan and McPhail 1989; Rowland 1994; Rowland et al. 1995; Bolyard and Rowland 1996; McKinnon 1996; Baube 1997; Rick and Bakker 2008; but see McKinnon and McPhail 1996). Male threespine sticklebacks compete with one another for breeding territories and males must have a territory and a nest built before they can court potential mates (Tinbergen and van Iersel 1947; van Iersel 1953), so male competition is an important determinant of reproductive success. Females then search and choose amongst nesting males, and males subsequently provide all parental care. In some studies, brighter red males are more aggressive (Rowland 1984; McLennan and McPhail 1989) and more likely to win in dominance tests (Bakker and Sevenster 1983; McKinnon 1996; Baube 1997). The dominance status of a male may also reflect his parental ability (McKinnon 1996; Candolin 2000). Males frequently destroy one another’s nests, and conspecifics eat and steal eggs regularly (Wootton 1976), so dominance likely influences success in parenting.

In several locations across the west coast of North America, male sticklebacks no longer display red nuptial coloration and instead have full-body black coloration (McPhail 1969; Moodie 1972; Reimchen 1989; von Hippel 1999; Boughman 2001; Catchen et al. 2013; Bolnick et al. 2016). Black color is expressed during the breeding season and is condition-dependent (Scott 2011), suggesting a sexual selection function. The best-supported explanation for the switch from red to black coloration is sensory drive, which posits that signals and sensory systems coevolve to maximize signal detectability in a given habitat (Endler 1992; Boughman 2002; Maan and Seehausen 2011). Black populations are typically found in locations with red-shifted water color, whereas red populations are found in clear water (Reimchen 1989; Boughman 2001; Scott 2001; Tinghitella, unpublished data). Habitats with red-shifted and full-spectrum light within each drainage are not separated by physical barriers and there are documented polymorphic regions where red and black fish interact (Hagen and Moodie 1979). The difference in coloration between red and black males affects male competitive behavior in both regions where it has been investigated (Tinghitella et al. 2015; Bolnick et al. 2016). In simulated mixed color assemblages in the lab, black males from Washington state (WA) rivers biased their aggression toward males with red nuptial coloration, whereas red males show no bias in aggressive behavior (Tinghitella et al. 2015). This pattern of aggression (bias toward heterotypic males) suggests that red males are the recipients of more aggression overall which could allow black males to exclude red males from preferred breeding sites (e.g., Adams 2004; Peiman and Robinson 2007; Vallin and Qvarnström 2011; Winkelmann et al. 2014; Lehtonen et al. 2015), enhancing habitat use differences. However, our understanding of how male competition might contribute to genetic divergence (through male competition outcomes)
system has been limited by not knowing whether black nuptial coloration itself is related to male competition outcomes.

We conducted dyadic within and between color morph male competition trials to determine whether and how red and black breeding color contribute to competition outcomes. Previous work also supports a possible relationship between measurements of male aggressiveness (performing more aggressive behaviors) and territory establishment in threespine stickleback males with red coloration (Bakker and Sevenster 1983), so we also tested for a role of aggressive behaviors. We predicted that red throat coloration would predict competitive outcomes, with males who have a larger proportion body coverage of red winning more territories (Bakker and Sevenster 1983; Baube 1997). Given that black body coloration likely evolved as a consequence of sensory drive, is conspicuous to the stickleback visual system (Boughman 2001), and is condition-dependent (Scott 2011), we also predicted that melanin-based black body coloration would predict competitive outcomes, with more melanin males winning territories. There is reason to believe this may be the case. Melanin-based coloration is correlated with increased aggression and sexual activity in a wide variety of vertebrates through pleiotropic effects of the melanocortin system (reviewed in Ducrest et al. 2008). However, if signals and competitor recognition have co-evolved within color types, we predict that body color (red or black) and/or the number of aggressive competitive behaviors will be differentially associated with competitive outcomes in homotypic versus heterotypic interactions and that more homotypic than heterotypic trials will end with territory establishment (assuming homotypic competitors have mutually “understood” signals).

Materials and Methods

Competition trials

We collected sexually mature sticklebacks in regions of WA where red and black fish are allopatric and transferred them to the University of Denver by air in summers of 2014–2017. Fish with red nuptial coloration were collected from the Chehalis River (46°56′22″N, 123°18′46″W) and Campbell Slough (47°2′40″N, 124°3′33″W), and fish with black nuptial coloration were collected from Vance Creek (46°59′48″N, 123°24′43″W), Scatter Creek (46°49′20″N, 123°3′11″W), Conner Creek (47°4′10″N, 124°9′58″W), and Black River (46°49′45″N, 123°8′1″W). We housed fish in visually isolated 110-L tanks (77 cm × 32 cm × 48 cm) separated by collection location and sex at densities of no more than 10% at the start of the trial. Males were given two days from introduction into the tank to interact and to establish a territory and/or build a nest. This time period was chosen because male sticklebacks begin to build nests within two days of introduction to a new nesting area when alone, and must have an established territory in order to build a nest (Timbergen and van Iersel 1947; van Iersel 1953). Twenty-four hours and 48 h after introduction to the competition tanks, we presented competing males with a gravid female in a clear glass jar for 10 min near the nesting site (following Lackey and Boughman 2013). This reinforced that sexually mature females were present and motivated males to compete and build a nest. We also recorded whether males had established a territory, and inspected the sand substrate for signs of nest-building. When a territory has been established, the losing male becomes restricted to a small area of the tank, and upon moving toward the territory edge is attacked by the dominant fish (Rowland 1989). Forty-eight hours after introduction to the competition tank and immediately after experience with the gravid female, we observed the competing males for 20 min and recorded all aggressive behaviors (approaches, bites, and chases) using the event recorder JWatcher (Blumstein et al. 2006). During approaches a male swims, sometimes rapidly, towards the other, often at territorial boundaries, coming within one body-length. These have variously been described as “charges” in the literature as well. Bites can range in intensity from nips to sustained biting. Chases occur when one male swims vigorously behind the other (within one body-length), and can involve movement across the length of the tank multiple times.

Color quantification

To measure nuptial color, we photographed fish immediately before placing them into competition tanks. For 187 of the 195 competition trials, we also photographed males immediately after the 20-min focal observation period. Red breeding color is quite plastic, including during competitive interactions between males (Candolin 1999, personal observation). Photographing males before and after competition trials allowed us to capture the starting conditions and changes in body color that might occur during the course of the trial. All photographs were taken with a digital camera (Canon PowerShot G15) under standardized lighting (four evenly spaced xenon 20 W bulbs) inside a photobox that held the camera and blocked ambient light. In each photo the fish was on its right side, unanesthetized, against a neutral background with a millimeter ruler in view for scale. The process was brief, minimizing the potential for color change and stress.

Using FIJI (Schindelin et al. 2012), we quantified red and black coloration as a proportion of total body area from each photo (Figure 1). To obtain red color area, we selected red color using the Threshold Color plugin (Y = 32–255, U = 0–143, V = 141–255; following Wong et al. 2007). To obtain black area, we selected black coloration using the Threshold Color plugin (Y = 0–20; U = 0–255; V = 0–255). We determined total body area using the SIOX: Simple Interactive Object Extraction and scaled all measured areas using the millimeter ruler in each photo.

Statistical analysis

We assessed the effects of proportion red body color and aggressive behaviors of the focal red male on competitive outcomes in dyadic
trials between two red competitors and between one red and one black male. A focal red male was chosen randomly within each red × red trial for analyses. To examine territory establishment by red focal males, we ran two sets of models: one using “before-competition” red area and the other using difference in red area, calculated from before and after competition photos, as measures of male color. “Before-competition” red area captures color at the start of competition trials and the difference in red area (after competition – before competition) captures color change. We also ran a model using “after” red area as a predictor variable, which we provide in the Appendix. To examine the effect of aggressive behaviors on competitive outcomes, we combined approaches, bites, and chases using PCA as a variable reduction technique. All factors loaded evenly onto PC1, which explained 70.2% of the variation. We tested the effects of red color and behavior on competition outcomes using logistic regression via a generalized linear mixed model (family = binomial) in the lme4 package (version 1.2–12) in R (version 3.3.1; RStudio version 1.0.143; Bates et al. 2017). The “before-competition” model included before red color area, aggressive behaviors (PC1-behaviors), trial type (homotypic or heterotypic), and the interactions between color and trial type and between aggressive behaviors (PC1-behaviors) and trial type as independent variables. Trial outcome [whether or not the focal (red) male established a territory] was the dependent variable. The model also included the difference in mass between the two males in a trial as a covariate and the source populations of the red and black fish as random effects. The difference in red area model had the same structure. “Before-competition” color measurements were square-root-transformed prior to analysis to meet assumptions of normality.

We assessed the effects of proportion black body color and aggressive behaviors of the focal black male on competitive outcomes in dyadic trials between two black competitors and between one red and one black male. Again, a focal black male was chosen at random within each black × black trial for analyses. We performed a parallel analysis for focal black males using “before-competition” black area and difference in black area (after competition – before competition) as our measures of male color in separate models. Again, we also ran a model using only “after” black color area which we discuss in the Appendix. The measure of focal male aggressive behavior was again PC1-behaviors from the PCA described above and the model structure was as described above.

For all models, we followed a model comparison method in which the full models were compared to reduced models using χ² tests (performed using the anova function in lme4 in R) to assess whether the fit of the model decreased significantly when the effects of interest were systematically removed (Winter 2013). For interaction effects, we plotted logistic curves using geom_smooth (method = “glm”, family = “binomial”) in the ggplot 2 package (version 2.2.1) in R (Wickham 2017) and for fixed effects, we plotted logistic curves using sjp.glmer in the sjPlot package (version 2.3.3) in R (Lüdecke and Schwemmer 2017).

**Results**

When competing with homotypic competitors (other red males), males with greater “before” red area were more likely to establish a territory, but when competing with heterotypics (black males) red males with less “before” red body area were more successful in establishing a territory (A in Table 1 and Figure 2A). Red males were also more likely to establish a territory, regardless of trial type, when they performed more aggressive behaviors. We found no interaction between behaviors and trial type and no effect of trial type alone or “before” red area alone on whether red focal males established a territory (A in Table 1).

The difference in red area (after competition – before competition) was also associated with territory establishment (B in Table 1 and Figure 3A). Males who established a territory gained more red body area during competition, regardless of trial type. As above, red males that successfully established a territory performed more aggressive behaviors (Figure 3B). We found no interaction between difference in red and trial type, no interaction between behaviors and trial type, and no effect of trial type alone on whether red focal males established a territory (B in Table 1). Because the relationships between “before” red color and territory establishment and difference in red color and territory establishment were different (see A versus B in Table 1), we also investigated the relationship between...
“before” red color and difference in red color. Males that entered a competition trial with less red body area (before) gained more red coloration than males who entered a competition trial with more red body area ($\chi^2_{1}=65.7$, $P<0.0001$; mixed effects linear model with red focal male population = random).

In contrast to red body area, and contrary to our hypotheses, we found no effect of “before” black body area or the difference in black body area (after competition — before competition) on territory establishment by black males, regardless of competitor type (C and D in Table 1; Figure 3C). The only measured variable significantly related to the likelihood of a black male establishing a territory was aggressive behaviors; black males performed more aggressive behaviors when successfully establishing a territory (Figure 3D). The interactions between “before” black body area and trial type and difference in black area and trial type on territory establishment were non-significant. Trial type alone did not affect territory establishment in models using either black measurement (C and D in Table 1).

Of 195 male competition trials, 148 (~76% overall) ended with a clear winner who had established a territory. A total of 20 of 23 red × red trials, 51 of 74 black × black trials, and 77 of 98 red × black trials ended with territory establishment. The proportion of trials ending with a territory did not differ among the three trial types ($\chi^2=3.89$, $P=0.14$). There was a trend toward a greater proportion of red × red trials than black × black trials ending with territory establishment, but this was not significant ($\chi^2=2.91$, $P=0.09$). Males performed 135% more aggressive behaviors (PC1-behaviors) in trials in which a territory was established (LS means ± SE: territory = 0.19 ± 0.20, no territory = −0.54 ± 0.30; $\chi^2=5.13$, $P=0.02$), but the number of aggressive behaviors performed during competition did not differ across trial types (LS means ± SE: red × red = −0.33 ± 0.48, red × black = 0.25 ± 0.23, black × black = −0.09 ± 0.26; $\chi^2=2.18$, $P=0.34$). Neither color type was more likely to win in heterotypic competition (red wins = 42, black wins = 35, $\chi^2=0.46$, $P=0.50$), and both red and black focal males performed similar numbers of aggressive behaviors (LS means ± S.E.; red males = −0.33 ± 0.49, black males = 0.10 ± 0.21; $\chi^2=0.861$, $P=0.35$).

### Discussion

In this study, we first asked whether male nuptial coloration and aggressive behavior function differently in the competitive interactions of two diverging stickleback color types, one with red nuptial coloration (the ancestral type; McLennan 1996) and one with black nuptial coloration (the derived type). Overall, we found that different combinations of traits influence the outcome of dyadic competitive interactions involving red and black males. Males of both color types who performed more aggressive behaviors were more likely to establish a territory, regardless of whether the competitor was homotypic or heterotypic (Figure 3B, D; Table 1). But, whereas competition outcomes for red males varied with “before-competition” red body area and the degree to which red body area changed during competition (difference in red area), black body area was...
unrelated to competitive success (Table 1; Figure 3A, C). This is contrary to our original hypothesis that condition-dependent (Scott 2011) melanism expressed during the breeding season is sexually selected and that more melanic males would win territories, but supports the idea that competitive strategies (anything that leads to success in competition in a particular environment) of red and black males have diverged. Divergent competitive strategies may lead to aggression biases and asymmetries upon secondary contact.

What, then, is the function of black coloration, if any? Several authors have suggested that black coloration was favored through sensory drive and increases male conspicuousness in red-shifted light habitats (Boughman 2001; Scott 2001; Boughman 2002). Black coloration may function in female choice, as suggested in some work (Scott 2004; but see Tinghitella et al. 2015 and below). Other evidence suggests that black is maximally expressed during care for eggs and fry (Scott 2004) and thus may signal parental care ability to female conspecifics (Scott 2011) and/or act as a threat to egg predators or conspecific egg thieves (McPail 1969; Hagen et al. 1980; but see Scott and Foster 2000). Alternatively, black coloration may not be a signal at all, but instead have been “pulled along” by selection on another trait (Ducrest et al. 2008). Experiments in which black body coloration is manipulated directly to assess effects on sexual selection or success in parental care would reveal whether black coloration is a signal and, if so, of what.

We also asked whether color or aggressive behavior varied with competitive outcomes differently when the competitor is homotypic versus heterotypic, and whether competition between homotypic males was more likely to be resolved than competition between heterotypic males. First, we found no evidence that homotypic trials were resolved more often that heterotypic trials, though there was a trend toward more red–red than black–black trials ending with a territory holder. In Pacific field crickets that have lost the ability to produce an aggressive song that mitigates the costs of fighting, fights involving one or more silent males are more physically aggressive than those involving two calling males (Logue et al. 2010). Fights between two threespine stickleback males who do not have the ancestral red threat signal did not involve more physically aggressive behaviors. However, red males with greater “before-competition” red color were more likely to establish a territory in homotypic red x red trials (consistent with previous work in threespine sticklebacks; Bakker and Sevenster 1983; Baube 1997), and the same trait (extensive red color) led to reduced success in competition with heterotypic (black) males (A in Table 1; Figure 2A). The relationship between black color (either measure) and competitive outcomes did not differ depending on whether a male faced a homotypic or heterotypic competitor (C and D in Table 1; Figure 2B). We interpret this to mean that the ancestral color, which does function in competition among red males, does not influence competitive outcomes with the derived color type in the same way—competitive strategies have diverged and the relationship between color and competitive outcomes differs depending on the receiver. This supports the hypothesis that signals and competitor recognition have co-evolved within color types.
In a previous study, we found that in mixed sex assemblages in the lab, black males biased their aggressive behaviors toward red males, while red males had no aggression bias (Tinghitella et al. 2015). The aggression bias we observed previously and difference in competition currencies (which traits are correlated with success in competition) uncovered here are likely related. For instance, less red (dull) males may be more successful in competition with black males because black males bias aggression toward red competitors. Said another way, if black males bias aggression toward red males with greater red body area, it would be reasonable to expect males with greater red body area to lose in one on one competitive interactions with black males, relative to dull red males (as we find here).

In the mixed sex assemblage experiment described above, we found no evidence that the preferences of females from red and black populations had diverged; all females, regardless of color type preferentially directed courtship interest behaviors toward red males (Tinghitella et al. 2015; but see Scott 2004). Ongoing no-choice courtship trials in the lab support this pattern (R.T., unpublished data). It is relatively straightforward to imagine how male competition and female choice might act in concert to drive speciation, for instance, if both favor the same alternative male trait combinations or females choose mates on the basis of competition outcomes. If females retain a preference for ancestral type males, however, there must be some other mechanism contributing to reproductive isolation.

To illuminate whether male competition contributes to reproductive isolation in this system, we suggest several lines of inquiry. It will be critical to know, for instance, whether the patterns we have uncovered in the lab also exist in the field. Are red males the recipients of more aggression in the field, and does this pattern lead red males to nest in less preferred habitat, reinforcing isolation? Are competitive outcomes and female preferences environment-dependent, varying, for instance, with light environment? If competitor types do differ in habitat use, natural selection can further differentiate competitive phenotypes and selection against migrants and hybrids might facilitate ecological specialization and speciation. Where the color types co-occur (in regions of polymorphism or upon secondary contact), black males biasing aggression toward red males should lead to a pattern whereby black males’ nearest neighbors are other black males or “dull” red males, rather than red males with extensive red coloration. This could be easily observed and tested in situ. However, while black males that nest adjacent to homotypics may enjoy a fitness advantage if they lose fewer mating opportunities to red males who are preferred by female conspecifics, it is unclear whether nesting near “dull” red males would reinforce divergence between the color types. Further, comparing and contrasting competitor recognition and competitive outcomes between red and black sticklebacks from regions of their distribution where the two color types are allopatric versus sympatric will be particularly revealing. Where the two color types commonly co-occur, for instance, we might expect to find greater divergence in aggressive signals and competitive responses relative to allopatric populations, such that heterotypic competition is rare (e.g., Anderson and Grether 2010a, 2010b). Finally, future experiments similar to those conducted here, but with fish from regions where red and black males interact frequently would reveal whether competitors who commonly interact easily establish dominance through mutually “understood” signals.

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Author contributions
RMT conceived of and designed the experiment. RMT and VFL conducted male competition trials. VFL quantified color from photos with undergraduate assistants, using methods developed by WRL. WRL analyzed the data and made the figures. All co-authors contributed to writing and editing the manuscript.

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APPENDIX

Table A1. Relationship between after color measures, aggressive behaviors, and trial type on territory establishment by focal red or black males

|          | \( \chi^2 \) | df | P      |
|----------|--------------|----|--------|
| Red focal males |               |    |        |
| After red area * trial type | 0.85 | 1 | 0.36 |
| Aggressive behaviors * trial type | 1.03 | 1 | 0.31 |
| After red area | 1.26 | 1 | 0.26 |
| Aggressive behaviors | 14.20 | 1 | <0.001 |
| Trial type | 2.38 | 1 | 0.12 |
| Black focal males |               |    |        |
| After black area * trial type | 0.35 | 1 | 0.55 |
| Aggressive behaviors * trial type | 1.25 | 1 | 0.26 |
| After black area | 0.56 | 1 | 0.45 |
| Aggressive behavior | 55.06 | 1 | <0.0001 |
| Trial type | 0.41 | 1 | 0.52 |
