- Red operculum spots, body size, maturation and evidence for a satellite 1
- male phenotype in non-native European populations of pumpkinseed 2
- Lepomis gibbosus 3

- Grzegorz Zieba^{1,2} | Carl Smith^{2,3,4} | Michael G. Fox⁵ | Stan Yavno^{5,6} | Eva Záhorská⁷ | 5
- Mirosław Przybylski² | Gérard Masson⁸ | Julien Cucherousset⁹ | Hugo Verreycken¹⁰ | 6
- Hein H. van Kleef¹¹ | Gordon H. Copp^{1,5,12} 7

8

- 9 ¹Salmon and Freshwater Team, Centre for Environment, Fisheries and Aquaculture Science,
- 10 Pakefield Road, Lowestoft, NR33 0HT, UK
- ²Department of Ecology and Vertebrate Zoology, University of Łódź, Poland 11
- 12 ³Institute of Vertebrate Biology, Academy of Sciences of the Czech Republic, Brno, Czech
- 13
- ⁴School of Biology and Bell-Pettigrew Museum of Natural History, University of St Andrews, UK 14
- ⁵Environmental and Life Sciences Graduate Program, Trent University, Peterborough, Ontario, 15
- 16
- 17 ⁶Department of Zoology, Tel Aviv University and Inter-University Institute for Marine Sciences,
- 18 Eilat. Israel
- ⁷Comenius University, Faculty of Natural Sciences, Department of Ecology, Bratislava, Slovakia 19
- 20 ⁸Laboratoire Interdisciplinaire des Environnements Continentaux, Université de Lorraine, Metz,
- 21 France
- ⁹Laboratoire Evolution and Diversité Biologique (EDB UMR 5174), Université Paul Sabatier, 22
- CNRS, ENFA, UPS, Toulouse, France 23
- 24 ¹⁰Research Institute for Nature and Forest, Brussels, Belgium
- 25 ¹¹Bargerveen Foundation, Department of Animal Ecology and Ecophysiology & Department of
- Environmental Science, Faculty of Science, Radboud University Nijmegen, Nijmegen, The 26
- 27 Netherlands
- 28 ¹²Centre for Conservation and Environmental Change, School of Conservation Sciences,
- 29 Bournemouth University, Dorset, UK

30

- 31 Correspondence
- 32 Gordon H. Copp
- gordon.copp@cefas.co.uk; +44 01502 52 7751 33

34

Running title: Operculum spot of non-native pumpkinseed

3637

38

39

40

41

42

43

44

45

46

47

48

49

50

51

52

53

54

55

Abstract

Carotenoid-based pigmentation is a striking feature of many taxa, yet the function, if any, of colour traits is often unclear. Pumpkinseed Lepomis gibbosus, a widely introduced freshwater sunfish that exhibits alternative male mating strategies, express a striking, red operculum spot. To investigate the potential function of this red spot as a signal in this species' mating system, we determined the presence and measured the size of red operculum spots in fish collected from 12 populations in five European countries in which pumpkinseed is an established non-native species. We subsequently related the presence and size of the red spot to body size and mating strategy, based on an analysis of relative gonad size (gonado-somatic index, GSI), using a mixed modelling approach. The study demonstrated that the presence of a red operculum spot in pumpkinseed is associated with sexual maturation, with GSI frequency distributions suggesting that cuckolders in some non-native populations comprised both sneaker and satellite males, the latter not having previously been reported for this species. Further, the size of red spot correlated strongly with body size in parental and cuckolder males, though there was no difference in the presence or size of the red operculum spot between male mating strategies. The function of a red operculum spot in females is not clear but may be partly mediated by pleiotropic genetic mechanisms. Red operculum spots appear to function as signals of male maturation and body size in pumpkinseed, irrespective of mating strategy.

5657

58

59

Keywords alternative mating strategy · carotenoid · GLMM · sexual selection · visual signal · Centrarchidae

1 | INTRODUCTION

Visual signals play a key role in the environmental biology of teleost fishes (Marshall, 2000; Wootton, 1998). Pigmentation is an important component of visual signals in fishes, both within and among species. The expression of colour, which often comprises multicomponent signals, involves pigment-based and structural colours (Price, Weadick, Shim, & Rodd, 2008; Wootton & Smith, 2015). Carotenoid-based colouration, which typically appears as yellow, orange or red, has been of particular interest because these are costly to produce and thereby serve as a reliable indication of individual condition and foraging ability (Olson & Owens, 1998), with a potential role in mating system evolution (Wootton & Smith, 2015). Different carotenoid pigments are responsible for conveying different information to potential mates or rivals (McLennan, 2007). Colour traits can function in sexual selection; selection that acts on differences in fitness among individuals that arise due to the number and identity of their mates. Because sexual selection tends to act more strongly on males than females (Andersson, 1994), it is males that usually express the brightest and most elaborate visual traits, including nuptial colouration. Though a broad simplification, there is a tendency in many taxa for a preference to mate with more colourful and ornamented individuals (Andersson, 1994; Price et al., 2008). Understanding the adaptive value of mate choice (sensu Kokko, Brooks, Jennions & Morley, 2003) based on visual signals is a controversial field of research, but may arise through fitness benefits associated with mate choice, or an arbitrary mechanism (Andersson, 1994; Wootton & Smith, 2015).

Pigmentation in North American sunfishes (Centrarchidae), in common with most other teleosts, is more extensive in males than in females (Scott & Crossman, 1973). In the pumpkinseed *Lepomis gibbosus* males have a striking, red operculum spot that is associated with male aggression (Stacey & Chiszar, 1978). In aquarium studies, breeding males were shown to attack fish models that intruded into their nesting area, and models with red colour on the operculum spot or iris received more attacks and aggressive displays than those without colour (Stacey & Chiszar, 1978). It is recognised that dominance relationships among sunfishes are related to colouration, with dominant, brightly-coloured parental (territorial) pumpkinseed males building and defending nests, and performing courtship behaviour to attract females to their nest for spawning. Male sunfishes also adopt alternative mating strategies (*sensu* Wootton, 1984), whereby subdominant 'cuckolder' males attempt to intrude into the nest of a parental male that is in the act of spawning (Gross, 1979, 1982). Cuckolder males are early-maturing, do not

perform courtship and avoid fitness costs associated with parental care (Smith & Wootton, 1995). Small cuckolders, typically possessing disproportionately large testes, are referred to as 'sneakers' due to their rapid entry into the nest of a spawning territorial male to release sperm at the moment of spawning. Large cuckolders, which have testes of intermediate size relative to parental and sneaker males, are termed 'satellites' and participate in spawning by mimicking female characteristics and entering the nest of a territorial male as a potential mate (Gross, 1982; Garner & Neff, 2013). Despite aggressive defence of their nest by parental males, cuckolder males frequently intrude successfully during spawning events (Rios-Cardenas, 2003). In the close congener, bluegill *Lepomis* macrochirus, a model taxon for research on alternative reproductive strategies (Taborsky, 2008), Gross (1982) characterised two cuckolder phenotypes, based on a population in Lake Opinicon, Canada. Sneaker bluegill are small and have relatively large testes relative to parental males, with a mean gonado-somatic index (GSI) of 4.6 (contrasting with parental males with a mean GSI of 1.1). In contrast, satellite male bluegill are larger than sneakers, with GSI values (mean 3.3) that are intermediate between those of parental males and sneakers. In applying these criteria to pumpkinseed from the same lake, Gross (1982) identified the sneaker phenotype, but not that of the satellite. In a subsequent paternity study. Neff and Clare (2008) noted that the satellite strategy had yet to be reported for pumpkinseed. In a subsequent genetic study of bluegill-pumpkinseed hybridisation, Garner and Neff (2013) did not distinguish between satellites and sneakers, referring to them collectively as cuckolders. Thus, there are no reports that the satellite phenotype (sensu Gross, 1982) is expressed in pumpkinseed and hereafter we use the term 'cuckolder' to refer to the alternative male mating strategy in pumpkinseed to the parental strategy.

94

95

96

97

98

99

100

101

102

103

104

105

106

107

108

109

110

111

112

113

114

115

116

117

118

119

120

121

122

123

124

125

126

127

The approach, onset and end of the reproductive period in fishes can often be identified by variation in GSI, the relationship between gonad weight and somatic weight (Wootton, 1998). Male mating polymorphisms, particularly the cuckolder strategy, are associated with sperm competition, which occurs when the spermatozoa of different males compete for the same eggs (Parker, 1970). Males expressing alternative mating strategies tend to experience a higher "risk" of sperm competition; i.e. an elevated probability that their sperm will compete with the sperm of other males, because they usually mate in the presence of at least one competitor, typically the territorial male and often other cuckolder males (Wedell, Gage, & Parker, 2002; Wootton & Smith, 2015). An outcome is that cuckolders are predicted to show adaptations for sperm competition, including relatively

(or even absolutely) larger testis size, larger ejaculates, and faster swimming spermatozoa (Montgomerie & Fitzpatrick, 2009). Consequently, individual GSI can be used to identify male mating strategy.

Pumpkinseed have been widely established outside their native North American range. Most introductions have been in Europe, where the species was introduced in the late 19th and early 20th centuries, establishing populations in at least 28 countries (Copp & Fox, 2007). Most of the research undertaken on pumpkinseed in Europe has focused on female growth and life-history traits (reviewed in Fox & Copp, 2014), with relatively few studies on males (e.g. Almeida, Tobes, Miranda, & Copp, 2012; Valente et al., 2016). In its native range, Neff & Clare (2008) observed temporal variability in the patterns of pumpkinseed paternity and suggested that these reflect temporal variation in mating opportunities, parental male defence ability, or cuckolder densities. Given this temporal variability, spatial variations are also feasible, possibly extending to visual signals, such as the operculum spot, which is assumed to be important for males for attracting mates (Stacey & Chiszar, 1978).

The aim of the present study was to determine whether the presence and size of the red operculum spot serves as a signal of sex and/or mating strategy in pumpkinseed. To do this we used estimates of GSI to categorise males as nesters or cuckolders and fitted models to test whether spot presence and size varied between the sexes and between males adopting different reproductive strategies. We predicted that parental males, which experience intrasexual contests and intersexual mate choice, would be more likely than cuckolder males and females to express a red operculum spot and that these males would also express larger red spots.

2 | Methods

Previously unpublished data for 1118 pumpkinseed were obtained from established non-native populations in Belgium, England, France, the Netherlands and Poland (Table 1), with all data elements being available for 924 specimens of which 465 were males, 387 were females, and 266 were juveniles (Table 2). The fish were collected by a variety of sampling gear (Table 1), depending upon location characteristics and gear availability. Note that sampling of the two Netherlands populations was incorrectly reported as by funnel trap in Cucherousset et al. (2009). Where possible, the sampling aimed to obtain ≈ 100 specimens for growth and life-history trait analysis (e.g. Villeneuve, Copp, Fox, & Stakėnas, 2005; Cucherousset et al., 2009) and so was not quantitative. In the three

English sites, sampling was by catch-per-unit-effort (CPUE) with minnow traps (see Fox, Villeneuve & Copp, 2011). The data on female growth and life-history traits, but not operculum spot occurrence or size, have been previously published for all these populations (Villeneuve et al., 2005; Cucherousset et al., 2009; Fox et al., 2011; Valente et al., 2016), except the two populations from northwest Poland (Table 1): i) a former clay pit (Brodowski Pond) of ≈ 0.9 ha (maximum depth of 7 m) situated in Szczecin with a water temperature that ranges from 4 °C under thick ice cover in winter to 25 °C in summer; and ii) an artificial channel (4 km long, mean depth of 4.5 m), situated on the lower River Oder near the city of Gryfino that receives heated water effluent from a power plant, resulting in daily water temperature fluctuations of 5–15 °C, with maximum temperatures of 30 °C in summer and 15 °C in winter. There is currently no published study on the North American source populations of pumpkinseed in Europe.

For all populations, pumpkinseed were collected at the beginning of the spawning season (between May and June), immediately killed with a lethal dose of anaesthetic following licenced procedures in the country of collection and cooled to freezing for subsequent processing in the laboratory. After thawing, specimens were photographed (left side only) using a Nikon CoolPix 5000 camera for morphological analysis. The presence of operculum spots (denoted as present or absent) was recorded, including the rare occurrence of deviations from red (i.e. yellow or orange, or reddish shades thereof). Note that the reflectance spectra, hue or intensity of operculum spots were not measured, nor was the spectral sensitivity of pumpkinseed in the red region of the light spectrum known. The presence of an operculum spot was recorded for both sides of each specimen assessed, but only the left side of the fish (the standard for fish measurements) was used for quantifying the size of the red operculum spot, which was estimated as the surface area of the spot (ImageJ, 1.48v). Fish were measured for standard length (SL, nearest 1 mm) and mass (nearest 1 g) and dissected to determine sex and gonad mass (in mg) when present. GSI was calculated as: 100 × gonad mass ÷ total body mass) (Wootton & Smith, 2015).

2.1 | Statistical analysis

To categorise males as parentals or cuckolders, a frequency distributions of male GSI was plotted by country. The resulting multimodal GSI frequency distributions were decomposed to normal distributions using the Bhattacharya method (Bhattacharya, 1967), with a meaningful separation of normal distributions identified with a separation index (SI)

> 2 (Gayanilo, Soriano, & Pauly, 1989). GSI values below the threshold value were categorised as parental and those equal to or above the threshold value were categorised as cuckolders. Data were pooled by country, rather than population, to maximise the precision with which threshold values could be identified.

To test whether parental males were more likely to express a red operculum spot than other males and females, the presence of a red spot in pumpkinseed was modelled using a Bernoulli Generalised Linear Mixed Model (GLMM). Fish SL and mass were collinear, and mass was subsequently dropped from the analysis. Only a single fish that was classified as a juvenile expressed a red spot, and juveniles were subsequently dropped from the analysis. The distribution for the model response variable, link function and predictor function were specified as:

```
207 Spot_{ij} \sim Binomial(\pi_{ij}, n_{ij})

208 E(Spot_{ij}) \sim n_{ij} \times \pi_{ij} and var(Spot_{ij}) = n_{ij} \times \pi_{ij} \times (1 - \pi_{ij})

209 \eta_{ij} = \beta_1 + \beta_2 \times SL_{ij} + \beta_3 \times strategy_{ij} + population_i

210 logit(\pi_{ij}) = \eta_{ij}

211 population_j \sim N(0, \sigma^2_{population})
```

Where $Spot_{ij}$ is the probability of fish i in population j having a red operculum spot in n_{ij} independent individuals, which was assumed to follow a binomial distribution with an expected probability (E) of expressing a spot of mean $\pi_i N_i$ and variance $\pi_i \times (1-\pi_i)$, with a logit link function. The logit function ensures the fitted probability of a red spot falls between 0 and 1. The variable strategy $_{ij}$ is a categorical covariate with three levels, corresponding with fish mating strategy; female, territorial or cuckolder. The model also contained a linear effect for fish SL (SL_{ij}). Exploratory analyses demonstrated differences among populations in the size and expression of red operculum spots. To accommodate this effect in the model, the random intercept $population_j$ was included to introduce a correlation structure between observations for different fish from the same population, with variance $\sigma_{population}$ distributed normally and equal to 0.

To test whether parental males expressed larger red operculum spots than other males and females, the size of red spots was modelled on a zero-truncated subset of the data using a Gaussian GLMM, specified as:

```
226 log_{10}Spotsize_{ij} \sim N(\mu_{ij}, \sigma^2)

227 E(log_{10}Spotsize_{ij}) = \mu_{ij} and var(log_{10}Spotsize_{ij}) = \sigma^2

228 \mu_{ij} = \beta_1 + \beta_2 \times log_{10}SL_{ij} \times \beta_3 \times strategy_{ij} + population_i

229 population_j \sim N(0, \sigma^2_{population})
```

Where $log_{10}Spotsize_{ij}$ is the log_{10} size of red spot on fish i in population j and $log_{10}SL_{ij}$ is the log_{10} standard length of fish i in population j. Expected $log_{10}Spotsize_{ij}$ is μ_{ij} with variance σ^2 . Analysis was conducted using the lme4 library (Bates, Maechler, Bolker, & Walker, 2014) in R (R Development Core Team, 2017). Model residuals were examined to ensure assumptions were met and that effects were adequately accounted for by the model.

3 | RESULTS

Male GSI distributions were multimodal within countries (Figure 1). With the exception of data from the Netherlands, the overlap of normally-distributed GSI groups was low; SI was always >2 and thereby provided meaningful separation of modal groups, even for the Netherlands specimens (separation index = 2.68). Decomposition of GSI distributions showed the threshold GSI value that distinguished parental and cuckolder males varied among countries, from 2.3 (Poland) to 4.2 (England) (Table 2, Figure 1). In Belgian, French and Polish populations, separation of male strategies was ambiguous, with Bhattacharya decomposition revealing two thresholds within these data (Figure 1). In each case, the higher threshold was used as the cut-off between male strategies.

The probability of expressing a red operculum spot was predicted by male SL, with a significantly greater likelihood of a red spot in larger males (Table 3; Figure 2). The probability of a red operculum spot was also predicted by sex, with both cuckolder and territorial males significantly more likely to express a red spot than females (Table 3; Figure 2). Among pumpkinseed that expressed a red spot, there was a significantly stronger interaction between SL and spot size in males than in females (Table 4; Figure 3), though there was no difference between male strategies classified by GSI (Figure 3). Two specimens were observed to possess a yellow operculum spot, a mature female from Schoapedobbe Pond and a male from Brière Marsh (Table 1).

4 | DISCUSSION

Our analysis of introduced pumpkinseed populations across Europe demonstrated that the expression of a red operculum spot was more likely in mature males than females, (Table 3), whereas only a single sexually immature juvenile expressed a red spot. Further, a significantly stronger interaction was observed (Table 4) between red spot size and body size in parental and cuckolder males than in females. Taken together, these findings indicate that in pumpkinseed, the presence and size of a red operculum spot signals

sexual maturity and correlates strongly with body size in males. This signal, however, does not distinguish between the male mating strategies of parental and cuckolder.

Colour signals can have multiple functions, potentially conveying information to rivals, mates or both. In pumpkinseed, the function of the red operculum spot is equivocal. The present findings are consistent with previous studies in demonstrating that a red spot has a potential role in male-male interactions by eliciting an aggressive response by males (Stacey & Chiszar, 1978). Notably, during aggressive intra-sexual contests, male sunfishes flare their opercula (Colgan & Gross, 1977), displaying the red spot to a rival facing them head-on. Our results support a role for the red operculum spot in male contests, with the expression of a red spot primarily associated with sexually mature males and correlating with body size. However, this finding contradicts our prediction that parental males, which experience intrasexual contests and intersexual mate choice, should be more likely than cuckolder males to express a red operculum spot.

The expression of a red carotenoid-based colour signal may convey information about male condition, which could be functional in the context of male-male contests. Carotenoids are acquired solely in the diet by vertebrates and these compounds have a number of critical physiological functions, making carotenoid-based signals a potentially reliable indication of individual condition and fighting ability (Olson & Owens, 1998). Whether additional information is conveyed by the size, reflectance spectra and intensity of the red colour spot, or whether the red operculum spot comprises one component of a multimodal signal in pumpkinseed, remains to be investigated. Experimental studies would be particularly effective in differentiating these aspects of signal evolution in pumpkinseed.

Surprisingly, we detected no significant difference between parental and cuckolder males, either in the probability of expressing a red spot, or in the strength of the relationship between red spot size and male size. Alternative mating strategies in fish, which are exclusively shown by males in teleost fishes, are reflected most strikingly in reproductive behaviour, but also in external appearance and physiology. In contrast to Gross' (1982) interpretation of his sunfish model to a single native population of pumpkinseed, our analysis of non-native populations in Europe suggests that, while a clear distinction can be seen between the parental and cuckolder male reproductive strategies in relative testes size, the GSI value threshold that distinguishes parental and cuckolder males varies among populations, ranging from 2.3 (Poland) to 4.2 (England).

A possible alternative explanation for the observed range of GSI values may relate to variation in pumpkinseed population density, which has consequences for the reproductive success of cuckolder males. In water bodies where the populations occurred at high density, traits associated with sperm competition, including large testes size, are favoured, thereby potentially selecting for relatively larger gonads (high GSI) in both parental and cuckolder males, with a consequent upward shift in the threshold between them. In the three ponds in England for which quantitative density data were available (Fox et al., 2011), pumpkinseed was virtually the only species present in the two ponds where the species have been observed in high density and males in these ponds had higher mean and maximum male GSI values than observed in the low-density pond. In water bodies where populations occur at low densities, where the risk of sperm competition for parental males is low, relative testes size is predicted to be smaller and the GSI threshold between parental and cuckolder males lower (Wootton & Smith 2015).

A caveat to these conclusions was the finding that many females also expressed red spots in their opercula, albeit with lower probability than males (Figure 2). The size of red operculum spots in females also conveyed less information about size than in males (Figure 3). If red operculum spots function primarily in male-male contests, then it is unclear why females would display this trait. For example, the red operculum spot may have multiple functions, potentially signalling male size and fighting ability, but also undergoing intersexual selection in the context of mate choice. If the red operculum spot is selected through intersexual selection, then it is possible that females express the trait for the same reason as males. Female ornamentation is surprisingly common in teleosts, and in the context of the pumpkinseed mating system, in which males are the sole providers of parental care, the evolution of male mate choice and female ornamentation is possible (Wootton & Smith 2015).

The expression of red operculum spots in males is adaptive solely through intra-sexual selection, whereas in females the expression of red spots may be through a genetic correlated response. In threespine stickleback (*Gasterosteus aculeatus*), red throat colouration is known to function in intra- and intersexual selection on males and is a reliable indicator of body condition and parasite resistance (Milinski & Bakker, 1990; Barber, Arnott, Braithwaite, Andrew, & Huntingford, 2001). The red ornament, which has a strong genetic component, is also found in females and its expression in females is consistent with pleiotropy (Yong, Peichel & McKinnon, 2016). Pleiotropy may also mediate the expression of a red operculum spot in female pumpkinseed, and possibly also in

cuckolder males, though this conclusion relies on there being a shared genetic basis to red spots in males and females and no adaptive value in expressing a red spot in females and cuckolders, and these assumptions remain to be tested.

The presence of two male GSI thresholds in pumpkinseed populations in Belgium, France and Poland provides evidence that, contrary to Gross, (1982), both the satellite and sneaker strategies may be present in some non-native European pumpkinseed populations (Figure 1). Alternatively, these findings may simply reflect the greater heterogeneity in age at maturity within some European populations (Copp & Fox, 2007; Fox & Copp, 2014), which is a possible outcome of multiple sources of introduction or adaptation to conditions outside the natural range of the species. Behavioural studies, complemented with morphological analyses, are needed to clarify the existence of a satellite strategy in European pumpkinseed populations. Similarly, variation in predation among populations has the capacity to modify the selective landscape for red operculum spots, since sexual selection for visual signals can be limited through natural selection by predators in teleosts (Endler, 1980; Wootton & Smith, 2015). Although outside the scope of the present study, the effect of selective predation on pumpkinseed in response to the expression of the red operculum spot is not known and warrants further research.

In conclusion, the present study demonstrates that the presence of a red operculum spot in pumpkinseed is associated with sexual maturation, primarily in males. The size of red spot correlates with body size in parental and cuckolder males, although there was no difference in the presence or relative size of the red operculum spot between male mating strategies. The observed GSI distributions suggest the possible existence of the satellite mating strategy in some European populations, but alternatively this may be due to variations in pumpkinseed population density or to selective predation. The function, if any, of a red operculum spot in females is not clear but is consistent with pleiotropy.

ACKNOWLEDGEMENTS

This study, which derives from work initiated as part of a N.A.T.O. Collaborative Linkage Grant (LST-CLG No. 979499), supported by consecutive research contracts (SF0238 and SF0248) from the UK Department of Environment, Food and Rural Affairs (to GHC), which were complemented by a Marie Curie post-doctoral fellowship (PIEF-GA-2008-219707) and a subsequent research grant from the National Science Centre, Poland (decision No DEC-2011/01/D/NZ8/01807) to GZ. We thank two anonymous reviewers for their constructive and perceptive comments.

366

Author contribution

- The study was conceived by GHC and MGF. Specimens were collected and processed
- by GHC, GZ, MGF, SY, EZ, GM, JC, HV and HHvK. Data were analysed by GHC, SY,
- 369 GZ, MP and CS. The initial draft of the manuscript was prepared by SY, EZ, GZ and GHC,
- then extensively revised by GZ, CS and MP, with co-author contributions.

371

27

372

373

REFERENCES

- 374 Almeida, D., Tobes, I., Miranda R., & Copp G. H. (2012). Cuckoldry features of introduced
- pumpkinseed sunfish (Lepomis gibbosus) in contrasting environmental conditions in
- southern Europe. Canadian Journal of Zoology, 90, 1051–1057.
- 377 https://doi.org/10.1139/z2012-073
- 378 Andersson, M. (1994). Sexual Selection. Princeton University Press, Princeton.
- Barber, I., Arnott, S. A., Braithwaite, V. A., Andrew, J., & Huntingford, F. A. (2001). Indirect
- fitness consequences of mate choice in sticklebacks: offspring of brighter males grow
- 381 slowly but resist parasitic infections. *Proceedings of the Royal Society of London Series*
- 382 *B-Biological Sciences*, 268(1462), 71–76. https://doi.org/10.1098/rspb.2000.1331
- Bates, D., Maechler, M., Bolker, B., & Walker, S. (2014). Ime4: Linear mixed-effects
- models using Eigen and S4. R package version 1(7), 1–23.
- 385 Bhattacharya, C. G. (1967). A simple method of resolution of a distribution into Gaussian
- 386 components. *Biometrics*, 23, 115–135. https://doi.org/10.2307/2528285
- Colgan, P. W., & Gross M. R. (1977). Dynamics of aggression in male pumpkinseed
- 388 sunfish (Lepomis gibbosus) over the reproductive phase. Zeitschrift für
- 389 *Tierpsychologie*, 43, 139–151. https://doi.org/10.1111/j.1439-0310.1977.tb00064.x
- 390 Copp, G. H., & Fox, M. G. (2007). Growth and life history traits of introduced pumpkinseed
- 391 (Lepomis gibbosus) in Europe, and the relevance to its potential invasiveness. In F.
- 392 Gherardi (Ed.), Biological Invaders in Inland Waters: Profiles, Distribution, and Threats,
- 393 (pp. 289–306). Springer, Dordrecht. https://doi.org/10.1007/978-1-4020-6029-8_15
- Cucherousset, J., Copp, G. H., Fox, M. G., Sterud, E., Van Kleef, H. H., Verreycken, H.,
- 395 & Záhorská, E. (2009). Life-history traits and potential invasiveness of introduced
- pumpkinseed Lepomis gibbosus populations in northwestern Europe. Biological
- 397 *Invasions*, 11, 2171–2180. https://doi.org/10.1007/s10530-009-9493-5

- Endler, J. A. (1980). Natural selection on color pattern in *Poecilia reticulata*. *Evolution*, 34,
- 399 76–91.
- 400 Fox, M. G., & Copp, G. H. (2014). Old world versus new world: life-history alterations in a
- 401 successful invader introduced across Europe. Oecologia, 174, 435-446.
- 402 https://doi.org/10.1007/s00442-013-2776-7
- 403 Fox, M. G., Villeneuve, F., & Copp, G. H. (2011). Seasonal reproductive allocation, local-
- 404 scale variation and environmental influences on life history traits of introduced
- pumpkinseed (Lepomis gibbosus) in Southern England. Fundamental and Applied
- 406 Limnology, 178, 231–243. https://doi.org/10.1127/1863-9135/2011/0178-0231
- 407 Garner, S. R., & Neff, B. D. (2013). Alternative male reproductive tactics drive
- asymmetrical hybridization between sunfishes (Lepomis spp.). Biology Letters, 9(6),
- 409 20130658. https://doi.org/10.1098/rsbl.2013.0658
- Gayanilo, F. C., Soriano, M., & Pauly, D. (1989). A draft guide to the Complete ELEFAN.
- International Center for Living Aquatic Resources Management, Manila.
- 412 (www.worldfishcenter.org/content/draft-guide-compleat-elefan).
- 413 Gross, M. R. (1979). Cuckoldry in sunfishes (*Lepomis*: Centrarchidae). *Canadian Journal*
- 414 of Zoology, 57, 1507–1509. https://doi.org/10.1139/z79-197
- 415 Gross, M. R. (1982). Sneakers, satellites and parentals: polymorphic mating strategies in
- North American sunfishes. Zeitschrift für Tierpsychologie, 60, 1–26.
- 417 https://doi.org/10.1111/j.1439-0310.1982.tb01073.x
- 418 Kokko H., Brooks, R., Jennions, M. D., & Morley, J. (2003) The evolution of mate choices
- and mating biases. Proceedings of the Royal Society of London Series B-Biological
- 420 *Sciences*, 270(1515), 653–664. https://doi.org/10.1098/rspb.2002.2235
- 421 Marshall, N. J. (2000). The visual ecology of reef fish colours. In Y. Espmark, T. Amundsen
- 422 & G. Rosenqvist (Eds.), Signaling and Signal Design in Animal Communication, (pp.
- 423 83–120). Tapir Academic Press, Trondheim.
- 424 McLennan D. A. (2007) The umwelt of the three-spined stickleback. In S. Ostlund-Nilsson,
- I. Mayer & F. Huntingford F (Eds.), Biology of the Three-Spined Stickleback (pp. 179–
- 426 224). CRC Press, Boca Raton.
- 427 Milinski, M., & Bakker, T. C. (1990). Female sticklebacks use male coloration in mate
- 428 choice and hence avoid parasitized males. *Nature*, 344(6264), 330–333.
- 429 https://doi.org/10.1038/344330a0

- 430 Montgomerie, R., & Fitzpatrick, J. L. (2009). Testes, sperm, and sperm competition. In B.
- 431 G. M. Jamieson (Ed.), Reproductive Biology and Phylogeny of Fishes (Agnathans and
- 432 Bony Fishes) (pp. 1–53). Science Publishers, Enfield (NH).
- 433 Morrell, L. J., Backwell, P. R., & Metcalfe, N. B. (2005). Fighting in fiddler crabs *Uca*
- 434 *mjoebergi*: what determines duration? *Animal Behaviour*, 70, 653–662.
- 435 https://doi.org/10.1016/j.anbehav.2004.11.014
- Neff, B. D., & Clare, E. L. (2008). Temporal variation in cuckoldry and paternity in two
- sunfish species (*Lepomis* spp.) with alternative reproductive tactics. *Canadian Journal*
- 438 of Zoology, 86, 92–98. https://doi.org/10.1139/Z07-121
- Olson, V. A., & Owens, I. P. (1998). Costly sexual signals: are carotenoids rare, risky or
- required? *Trends in Ecology & Evolution*, 13, 510–514. https://doi.org/10.1016/S0169-
- 441 5347(98)01484-0
- Parker, G. A. (1970). Sperm competition and its evolutionary consequences in insects.
- 443 Biological Reviews, 45, 525–567. https://doi.org/10.1111/j.1469-185X.1970.tb01176.x
- 444 Price, A. C., Weadick, C. J., Shim, J., & Rodd, F. H. (2008). Pigments, patterns, and fish
- behavior. Zebrafish, 5, 297–307. https://doi.org/10.1089/zeb.2008.0551
- 446 Rios-Cardenas, O. (2003). The use of Molecular Genetic Analyses to Examine Mating
- Patterns a Paternal Care in the Pumpkinseed Sunfish (PhD dissertation). State
- 448 University of New York at Buffalo.
- Scott, W. B., & Crossman, E. J. (1973). Freshwater Fishes of Canada. Fisheries Research
- 450 Board of Canada, Ottawa.
- Smith, C., & Wootton, R. J. (1995). The costs of parental care in teleost fishes. *Reviews*
- in Fish Biology and Fisheries, 5, 7–22.
- Stacey, P. B., & Chiszar, D. (1978). Body color pattern and the aggressive behavior of
- 454 male pumpkinseed sunfish (Lepomis gibbosus) during the reproductive season.
- 455 Behaviour, 64, 271–297. https://doi.org/10.1163/156853978X00062
- Taborsky, M. (2008). Alternative reproductive tactics in fish. In R. F. Oliveira, M. Taborsky
- 457 & H. J. Brockmann (Eds.), Alternative Reproductive Tactics: An Integrative Approach
- 458 (pp. 251–299). Cambridge University Press, Cambridge.
- Valente, E., Masson, G., Maul, A., Fox, M. G., Meyer, A., & Pihan J. C. (2016). Seasonal
- gonadal development and age-related maturity patterns of introduced pumpkinseed
- (Lepomis gibbosus Linnaeus, 1758) in a heated thermal reservoir and an adjacent river
- 462 reach. Journal of Thermal Biology, 58, 60–71.
- 463 https://doi.org/10.1016/j.jtherbio.2016.03.004

- Villeneuve, F., Copp, G. H., Fox, M. G., & Stakenas, S. (2005). Interpopulation variation
- in the growth and life history traits of the introduced sunfish, pumpkinseed *Lepomis*
- 466 gibbosus, in Southern England. Journal of Applied Ichthyology, 21, 275–281.
- 467 https://doi: 10111/j.1439-0426.2005.00690.x
- Wedell, N., Gage, M. J. G., & Parker, G. A. (2002). Sperm competition, male prudence
- and sperm-limited females. Trends in Ecology and Evolution, 17, 313-320.
- 470 https://doi.org/10.1016/S0169-5347(02)02533-8
- Wootton R. J. (1984) Introduction: tactics and strategies in fish reproduction. In G. W.
- Potts & R. J. Wootton (Eds.), Fish Reproduction: Strategies and Tactics (pp. 1–12).
- 473 Academic Press, London.
- Wootton, R. J. (1998). Ecology of Teleost Fishes, 2nd edn. Elsevier, Dordrecht.
- Wootton, R. J., & Smith, C. (2015). Reproductive Biology of Teleost Fishes. Wiley-
- 476 Blackwell, Oxford.
- 477 Yong, L., Peichel, C. L., & McKinnon, J. S. (2016). Genetic architecture of conspicuous
- red ornaments in female threespine stickleback. G3-Genes Genomes Genetics, 6(3),
- 479 579–588. https://doi.org/1534/g3.115.024505

TABLE 1. Country, population, location (WGS84 latitude and longitude coordinates) and capture method for male pumpkinseed *Lepomis gibbosus*, including number of specimens (*n*), mean and SD of standard length (mm) and the mean, minimum (min.) and maximum (max.) of the gonado-somatic index (GSI) for each population by water body.

							GSI		
Country	Population	Latitude	Longitude	Method [†]	n	SL (SD)	mean	min.	max.
Belgium	Webbekomsbroek Pond	50.975833N	5.076111E	ELF	76	61.2 (15.7)	2.4	0.2	5.3
France	Briere Marsh	47.366111N	2.312500W	ELF	63	90.9 (16.4)	1.9	0.5	4.6
	Grand-Lieu Lake	47.083333N	1.650000W	ELF	51	69.4 (21.2)	3.1	0.2	8.3
Netherlands	Schoapedobbe Pond	52.951944N	6.258333E	NET	30	124.2 (16.6)	2.0	0.7	5.6
	Meeuwven Pond	51.391111N	5.499722E	NET	58	67.3 (3.8)	3.5	1.3	10.7
Poland	Brodowski Pond	53.450342N	14.565856E	ELF	24	77.1 (11.1)	2.0	0.2	5.2
	Power Plant discharge channel	el53.213983N	14.468975E	ELF	25	77.1 (18.4)	1.8	0.1	5.7
England	Cottesmore Pond (Lower)	51.089167N	0.219444W	TRC	31	64.5 (12.5)	2.3	0.4	9.5
	Cottesmore Pond (Upper)	51.088056N	0.219444W	TRC	82	64.9 (12.3)	2.6	8.0	8.3
	Cottesmore Pond (School)	51.087222N	0.219444W	TRC	56	64.4 (16.2)	1.8	0.2	2.9
	Batts Stream at Cackle Street	51.018611N	0.074444W	ELF	28	75.2 (12.4)	1.7	0.7	6.4
	Tanyard Specimen Lake 3	51.019843N	0.012377W	ANG	39	92.7 (16.4)	2.4	0.9	12.5

[†]ANG = angling; ELF = continuous electrofishing; NET = small seine netting; TRC = minnow traps, catch-per-unit-effort (see Fox et al. 2011)

TABLE 2. Mean (SD) pumpkinseed body mass, standard length (SL) and gonado-somatic index (GSI; n/a = not available) among countries of origin and reproductive status. Male reproductive strategy (either cuckolder or parental) was identified by separation of GSI frequency distributions by Bhattacharya's (1967) method.

Country of origin	Reproductive status	n	Mass ((g)	SL (mm)	GSI
Belgium	Female	61	8.33	(9.89)	61.3 (16.0)	n/a
	Juvenile	10	3.85	(1.63)	52.5 (6.5)	n/a
	Cuckolder	21	4.81	(2.59)	55.9 (7.3)	3.60 (0.86)
	Parental	24	14.86	(10.22)	75.2 (16.4)	1.26 (0.59)
France	Female	66	17.81	(11.68)	81.1 (18.5)	n/a
	Juvenile	16	1.28	(0.75)	38.9 (6.1)	n/a
	Cuckolder	15	5.03	(2.95)	55.6 (8.9)	4.90 (1.11)
	Parental	91	22.58	(14.21)	88.2 (17.1)	2.00 (0.83)
Netherlands	Female	62	23.77	(23.99)	86.7 (30.2)	n/a
	Juvenile	1	7.00		66.6	n/a
	Cuckolder	33	7.25	(1.16)	66.3 (4.5)	4.25 (1.32)
	Parental	51	36.35	(26.58)	101.3 (30.4)	2.10 (0.63)
Poland	Female	18	20.25	(11.76)	76.4 (14.5)	n/a
	Juvenile	0	n/a		n/a	n/a
	Cuckolder	10	11.01	(7.24)	64.2 (10.3)	4.19 (1.05)
	Parental	39	23.79	(12.42)	80.4 (14.4)	1.26 (0.62)
England	Female	180	14.46	(14.64)	76.5 (19.9)	n/a
	Juvenile	45	2.56	(3.14)	46.2 (13.4)	n/a
	Cuckolder	16	5.69	(7.08)	59.4 (14.4)	6.20 (2.23)
	Parental	165	12.29	(10.04)	76.1 (16.2)	n/a

TABLE 3. Summary of Bernoulli GLMM to model the probability of pumpkinseed expressing a red operculum spot. Fish from different populations were fitted as random intercepts, with standard deviation of 1.47. $N_{\text{obs}} = 897$.

Model parameter	Estimate	SE	Р	
Intercept _(female)	-8.02	0.80	<0.001	
SL [†]	80.0	0.01	<0.001	
Strategy _(cuckolder)	2.30	0.34	<0.001	
Strategy _(territorial)	2.35	0.23	<0.001	

† SL = standard length

TABLE 4. Summary of a Gaussian GLMM to model the size of pumpkinseed red operculum spots. Fish from different populations were fitted as random intercepts, with standard deviation of 0.24. $N_{\text{obs}} = 344$.

Model parameter	Estimate	SE	Р	
Intercept _(female)	-7.43	0.99	<0.001	
Log ₁₀ SL [†]	1.82	0.22	<0.001	
Strategy _(cuckolder)	-5.31	2.77	0.055	
Strategy _(territorial)	-4.38	1.06	<0.001	
Log ₁₀ SL x Strategy _(cuckolder)	1.40	0.66	0.033	
Log ₁₀ SL x Strategy _(territorial)	1.18	0.24	<0.001	

[†] SL = standard length

518 Figure legends

- 519 **FIGURE 1.** Observed gonado-somatic index (GSI): frequency plots for pumpkinseed
- among country of origin with Gaussian distribution curves and separation indices (SI),
- 521 generated using Bhattacharyya's (1967) method. Threshold GSI values for separating
- 522 frequency distributions for male mating strategies are indicated with arrows.
- 523 **FIGURE 2.** Mean fitted probability (solid line) of pumpkinseed expressing a red operculum
- spot as a function of standard length (SL in mm) with 95% confidence intervals (shaded
- 525 area) for females, cuckolder males and parental males. Data were modelled with a
- 526 Bernoulli GLMM. Fish from different populations were fitted as random intercepts in the
- 527 model.
- 528 **FIGURE 3.** Mean fitted Log₁₀ area (mm²) of red operculum spot (solid line) of
- 529 pumpkinseed as a function of log₁₀ standard length (SL in mm) with 95% confidence
- 530 intervals (shaded area) for females, cuckolder males and parental males fitted with a
- 531 Gaussian GLMM. Fish from different populations were fitted as random intercepts in the
- 532 model.