Rare morph Lake Malawi mbuna cichlids benefit from reduced aggression from con- and hetero-specifics

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INTRODUCTION

Permanent polymorphism, the presence of multiple genetically determined morphological or behavioural phenotypes within a population, is common in nature and indicates some type of selective balance between morphs. Balancing selection is important for the maintenance of polymorphism as it can prevent either fixation of one morph through directional selection or genetic drift, or speciation by disruptive selection. Polychromatism, the presence of multiple genetically determined colour phenotypes, can be maintained if the fitness of alternative morphs depends on the relative frequency in a population. In aggressive species, negative frequency-dependent antagonism can prevent an increase in the frequency of rare morphs as they would only benefit from increased fitness while they are rare. Heterospecific aggression is common in nature and has the potential to contribute to rare morph advantage. Here we carry out field observations and laboratory aggression experiments with mbuna cichlids from Lake Malawi, to investigate the role of con- and heterospecific aggression in the maintenance of polychromatism and identify benefits to rare morphs which are likely to result from reduced aggression. We hypothesize that rare morph individuals receive less aggression than common morph individuals and therefore have an ecological advantage. Within species we found that males and females bias aggression towards their own morph, adding to the evidence that inherent own-morph aggression biases can contribute to balancing selection. Over-representation of rare morph territory owners may be influenced by two factors; higher tolerance of different morph individuals as neighbours, and the ability of rare morphs to spend more time feeding. Reduced aggression to rare morph individuals by heterospecifics may also contribute to rare morph advantage.

KEYWORDS

aggression, blotch polymorphism, cichlid, Malawi, rare morph advantage

Abstract

Balancing selection is important for the maintenance of polymorphism as it can prevent either fixation of one morph through directional selection or genetic drift, or speciation by disruptive selection. Polychromatism, the presence of multiple genetically determined colour phenotypes, can be maintained if the fitness of alternative morphs depends on the relative frequency in a population. In aggressive species, negative frequency-dependent antagonism can prevent an increase in the frequency of rare morphs as they would only benefit from increased fitness while they are rare. Heterospecific aggression is common in nature and has the potential to contribute to rare morph advantage. Here we carry out field observations and laboratory aggression experiments with mbuna cichlids from Lake Malawi, to investigate the role of con- and heterospecific aggression in the maintenance of polychromatism and identify benefits to rare morphs which are likely to result from reduced aggression. We hypothesize that rare morph individuals receive less aggression than common morph individuals and therefore have an ecological advantage. Within species we found that males and females bias aggression towards their own morph, adding to the evidence that inherent own-morph aggression biases can contribute to balancing selection. Over-representation of rare morph territory owners may be influenced by two factors; higher tolerance of different morph individuals as neighbours, and the ability of rare morphs to spend more time feeding. Reduced aggression to rare morph individuals by heterospecifics may also contribute to rare morph advantage.

1 | INTRODUCTION

Permanent polymorphism, the presence of multiple genetically determined morphological or behavioural phenotypes within a population, is common in nature and indicates some type of selective balance between morphs. Balancing selection is important for the
if the fitness of a phenotype depends on its relative frequency in a population (Henze et al., 2018; Hughes et al., 2013; Pérez i de Lanuza et al., 2017; Surmacki et al., 2013; Svensson, 2017).

In many taxa, species-recognition cues have diverged through reproductive or antagonistic character displacement to reduce hybridization or unnecessary exertion and risk of injury among conspecífics which are not in direct competition for mates or resources (Grether et al., 2009; Seehausen & Schluter, 2004). Rare colour morphs can benefit from lack of recognition by receiving less mating-related harassment (Takahashi et al., 2010) or less intrasexual aggression from conspecifics (Dijkstra et al., 2008; Lehtonen, 2014; Pérez i de Lanuza et al., 2017; Scali et al., 2020). In aggressive species, negative frequency-dependent antagonism, generated through either evolution of an own-morph bias (Dijkstra et al., 2008; Lehtonen, 2014; Scali et al., 2020) or by a dynamic common morph bias based on experience (Bolnick et al., 2016), can prevent an increase in the frequency of rare morphs as they would only benefit from increased fitness (due to reduced aggression) while they are rare (Bolnick et al., 2016; Dijkstra et al., 2007; Seehausen & Schluter, 2004).

The existence of conspecific aggression biases does not preclude heterospecific aggression completely. Indeed, recent studies suggest that heterospecific aggression as a result of resource competition and reproductive interference may be more common than previously assumed (Drury et al., 2020; Grether et al., 2009). Regardless of whether heterospecific aggression is due to convergence in territorial signals among species competing for resources or due to misdirection of aggression because closely related species still share similar signals (Losin et al., 2016), in a variety of taxa aggression is often higher among more similar coloured than more differently coloured species (Anderson & Grether, 2010; Genner et al., 1999; Losin et al., 2016; Pauers et al., 2008). In taxa, where multiple ecologically and phenotypically similar species co-exist in the same habitat there is therefore potential for rare morphs to benefit not only from reduced conspecific aggression, but also from reduced heterospecific aggression. A recent study of Midas cichlids, however, demonstrated increased aggression towards rare heterospecific morphs and suggested that this disadvantage may help to explain their lower frequency in natural populations (Lehtonen et al., 2015). The role of heterospecific aggression in relation to polychromatism requires further exploration to improve our understanding of how this may contribute to its evolution and maintenance.

The mbuna cichlids of Lake Malawi (and the closely related ecologically similar Mbipi of Lake Victoria) provide an excellent system for the investigation of colour polymorphism. Mbuña inhabit densely packed multi-species communities in the shallow-waters and identify conspecific mates and rivals predominantly by their species-specific colour and pattern (e.g. Couldridge & Alexander, 2002; Seehausen & van Alphen, 1998; Jordan, 2008; Pauers et al., 2008). Several species display a polychromatism characterized by the presence of rare ‘blotched’ morph individuals, which occur at different frequencies in different species and populations (Konings, 2007; Lande et al., 2001; Ribbink et al., 1983). While it is likely that differences in conspicuousness among morphs (resulting in differences in the likelihood predation) has played some role in the evolution of this polychromatism (Maan et al., 2008; Roberts et al., 2009; Seehausen et al., 1999; Streelman et al., 2003), and mate choice may have been involved in the evolution of (partial) sex-linkage (Lande et al., 2001; Roberts et al., 2009; Seehausen et al., 1999), it is thought that intrasexual competition plays a large role in its maintenance (Dijkstra et al., 2008; Dijkstra, van Dijk, et al., 2009). Although in most species the frequency of rare morphs remains relatively low in all populations, in some, for example, Maylandia callainos at Thumbi West Island in Lake Malawi, rare morphs can occur with higher frequency, which allows greater ease of observation and collection. Here we used this population, which consists of a common blue and a rare white morph, to conduct field observations and laboratory behavioural experiments to test alternative hypotheses regarding aggression biases: Do both morphs preferentially direct aggression towards the common (presumably ancestral) blue morph, or is there an own-morph bias? An own-morph bias could be sufficient to maintain polymorphism through negative frequency-dependent selection because the advantage to the rare morph would be reduced as it becomes more common. A common/blue-morph bias, however, would result in continuous advantage to the rare/white-morph through reduced aggression even as it became more common. In the case of common-morph aggression bias an additional frequency-dependent process, or another factor resulting in lower survival/reproduction of the rare morph, would therefore be necessary to limit an increase in the number of rare morph individuals. We also test for aggression biases towards the common and rare morph from a closely related heterospecific (Maylandia zebra) to assess whether this may contribute to balancing selection. We aim to identify potential benefits to rare morphs, which may occur as a result of receiving less aggression, in the natural environment. Additionally, as differences in selection pressures on each sex, due to differences in the type of competition they experience (competition for mates among males and competition for non-mating resources among females) can result in sex differences in the types of aggressive behaviour used during contests (Arnott & Elwood, 2009), we also test for sex-differences in aggressive behaviour and aggression biases.

2 | METHODS

2.1 | Study system

Maylandia callainos (=Pseudotropheus callainos or Metriaclima callainos) is a member of the ‘mbuna’ complex of rocky shore cichlid fishes endemic to Lake Malawi. Populations of M. callainos are found in shallow water, with peak population density between 3 and 10 m (full range 0–25 m). Their natural range is confined to the northern end of Lake Malawi, where they are often found in sympathy with the more widely distributed ecologically similar congeneric M. zebra. However, due to human-mediated translocations, they are also found in some southern areas. Phenotypes of common and rare morph mbuna differ between species, but within populations,
the common morph is often BB (black vertical melanin bars on a blue/dark background) or solid blue/dark body colour, while rarer morphs have a disrupted melanin pattern of many or few blotches/spots on a light (orange/pink/white) body. Blotch polychromatism is not present in all *M. callainos* and *M. zebra* populations; at some localities only the plain blue (B) and BB morph are found, whereas at others, these common morphs may occur along side rare white (W) and orange-blotch (OB) and very rare white-blotch and orange (O) morphs. In this study we focus on a well-established translocated population of *M. callainos* at Thumbi West Island in the Lake Malawi National Park in the southwest arm of the lake which has both B and W morph fish (Figure 1a). The likely source population of the *M. callainos* at Thumbi West is Nkhata Bay, where they co-occur with a population of *M. zebra* comprised BB, OB and O morph individuals.

All fish were wild caught: *M. callainos* and *M. zebra* from Thumbi West Island (TW) in July 2010, *M. zebra* from Nkhata Bay (NB) and Chiofu Bay (CB—naïve to *M. callainos* in the wild and lab) in 2009. Males and females were used in this study (Figure 1c), partially because of the lower number of rare males, but also because both male and female aggression biases have previously been suggested to be important in colour polymorphism maintenance in cichlids (e.g. Dijkstra et al., 2008). Furthermore, unlike many species with blotch polychromatism, this one is less strongly female limited, as numerous white *M. callainos* males were found at the study/collection site.

### 2.2 Field observations (excluding aggression)

#### 2.2.1 Frequency of blue and white morph *M. callainos* at Thumbi West Island

Snorkel observations were used to estimate the ratio of B to W *M. callainos* morphs in the general population. Dominant mature adult males can be easily recognized by their behaviour and colour, but females and immature males are indistinguishable and are referred to as ‘apparent females’. Hence, the number of males and apparent females of each morph was counted along three 30 m transects covering an area half a meter each side of the line (n = 74 fish). The numbers of territory-holding males of each morph were counted in nine 5 m² quadrats (n = 142 fish). Although ideally, the comparison should be made between non-territorial males and territorial males, in practice this was not possible due to the difficulty in sexing fish without catching them. However, it is likely that in the whole population, rare morph males occur at a lower frequency than rare morph females (as found in other closely related species with blotch polychromatism, Lande et al., 2001; Maan & Sefc, 2013), which would make estimates of the ratio of rare to common morph males among non-territorial fish a conservative estimate; territorial W males would be present at a much lower frequency than predicted from the ratio of W morph in the general population.

#### 2.2.2 Territory distances between morphs

Territory maps were constructed by drawing the rocky substrate, within 5 × 5 m string quadrats (n = 9), on dive slates while snorkelling. Males frequently return to their spawning cave and focal observations allowed for accurate determination of the position of this territorial focal point for each male within the quadrats. The distance between the territory focal point of each male within the centre 3 m² (n = 27 B & 25 W) and closest white and blue neighbour (including fish nearer the edge of the quadrat) was then measured.

#### 2.2.3 Grazing differences between morphs

Each grazing action performed by focal individuals was recorded during 10-min observations of territorial males and non-territorial fish (n = 9 individuals of each morph for each social status).
2.3 | Field observations of aggressive interactions

During focal observation lasting 10 min per fish (n = 9 territorial males of each morph) all aggressive behaviours directed towards the two conspecific morphs were recorded. The vast majority of all aggressive acts recorded were ‘chases’, lateral displays were observed but rare, counts of each type of behaviour were summed for analysis. While collecting data on conspecific aggression biases, aggression towards each focal fish from heterospecifics was also recorded.

2.4 | Laboratory aggression trials

To test whether there are differences in the level aggression received by blue and white morph M. callainos from conspecifics and heterospecifics, three experiments were carried out using the same methods. Five minute pairwise aggression trials were conducted in two replicate tanks measuring 0.9 x 0.3 x 0.3 m. Each tank contained a central brick refuge to act as a territory focal point, two transparent (perforated) plastic jars to hold the stimulus fish, an air-driven box filter and an internal heater to maintain the water temperature at ca. 22–24°C (Figure 1b). Lights were kept on a 12:12 light:dark cycle. All fish were fed flake food once a day. Females and males were used, but stimulus fish were always the same sex as focal fish. Focal fish were allowed at least 24 h to acclimatize before the introduction of the stimulus and recording of focal fish behaviour began after emergence from the central refuge. Individual aggressive behaviours (frontal/lateral display, quiver, lunge/butt and bite) were recorded and combined to give an overall aggression count for each individual.

To control for potential tank side bias, two separate trials were carried out with each focal fish, each with a different stimulus fish pair and with morphs swapped between sides. To avoid pseudoreplication from the re-use of focal males, before analysis an average was taken of the aggressive behaviour observed in the two trials by each individual.

2.4.1 | Exp. 1: Interspecific aggression biases between species

Firstly, conspecific aggression bias was confirmed by presenting BB M. zebra males from CB (n = 10) with pairs of conspecific BB and heterospecific B stimulus fish.

2.4.2 | Exp. 2: Intraspecific aggression biases between morphs

For this experiment all available M. callainos were used as focal and stimulus fish (n = 10 B male, six B female, three W male and nine W female) to test for morph-specific aggression biases among conspecifics.

2.4.3 | Exp. 3: Interspecific aggression to different morphs

BB M. zebra focal fish from different populations (n = 12 male/12 female ‘TW’, 12 male/5 female ‘NB’, 12 male/12 female ‘CB’) were used to test for heterospecific aggression biases to B and W M. callainos stimulus pairs. Stimulus pairs consisted of the same M. callainos used in Exp. 2.

2.5 | Data analysis

Statistical analysis was carried out using RStudio (v. 1.2.5033; RStudio Team, 2019) with additional packages: MASS (for negative binomial models), lme4 (for mixed effects models), performance (for checking overdispersion of generalized linear models) and ggplot2 (for plotting). General and generalized linear models were used with different distributions (and canonical link functions) depending on the type of data: Gaussian distribution for analysis of continuous data (territory distances), Poisson or negative binomial models for data where the response variable is a single count (grazing frequency, total aggression counts and aggression received from heterospecifics in the field), binomial or quasibinomial distributions for count data resulting from a binary choice (aggression towards different morphs in the field and in pairwise laboratory aggression trials, and also the ratio of attacks:displays)—further details of model specifications for each analysis are given below. Summary statistics and significance values are reported from t-test for general linear models and z-test for generalized linear models. One M. zebra female in Exp. 3 (Interspecific aggression to different morphs) only left the central refuge briefly at the start of the trial, this fish was excluded from the analysis due to lack of territoriality/interaction with the stimuli fish.

2.5.1 | Frequency of blue and white morphs at Thumbi West Island

A G-test was used to compare the actual number of territory holding white (W) males observed with what would be expected given the proportions of blue (B) and W morphs in the general population.

2.5.2 | Territory distances between morphs

General linear models (Table 1a) were used to test for the effect of neighbour morph type on the average distance (in meters) between the territory focal points of: (i) each territory holding male and the nearest same and different morph neighbour. (ii) each B male and the nearest B and W neighbour, (iii) each W male and the nearest B and W neighbour.
### TABLE 1  Model construction for analysis of: (a) Differences in territory distances among *Maylandia callainos* morphs (results Table 2). (b) Differences in grazing frequency between morphs (results Table 3). (c) Aggression biases by and towards *M. callainos* morphs in the field (results Table 4). (d) Differences in total aggression displayed by the different species, populations and sexes in laboratory aggression trials (results Table 5). (e) Aggression biases in pairwise intruder choice tests in the laboratory (results Table 6)

| Model | Distribution | Link | Response variable (and sample size) | Fixed factors | Random effect |
|-------|--------------|------|------------------------------------|---------------|---------------|
| (a) Territory distances between morphs | | | | | |
| i) | Gaussian | Identity | Distance (m) from all fish \( (n = 52) \) | Morph of neighbour (same or different) | |
| ii) | Gaussian | Identity | Distance (m) from B fish \( (n = 27) \) | Morph of neighbour (B or W) | |
| iii) | Gaussian | Identity | Distance (m) from W fish \( (n = 25) \) | Morph of neighbour (B or W) | |
| (b) Grazing differences between morphs | | | | | |
| i) | Negative binomial | Logit | Grazing count \( (n = 36) \) | Status (territorial or not) + Morph (B or W) | |
| (c) Field observations of aggressive behaviour | | | | | |
| i) | Binomial | Logit | Aggression count: All focals \( (n = 18) \) | Morph of aggressed intruder (B or W) * Focal Morph (B or W) | Focal ID |
| ii) | Binomial | Logit | Aggression count: All focals \( (n = 18) \) | Morph of aggressed intruder (B or W) | Focal ID |
| iii) | Binomial | Logit | Aggression count: B focals \( (n = 9) \) | Morph of aggressed intruder (B or W) | Focal ID |
| iv) | Binomial | Logit | Aggression count: W focals \( (n = 9) \) | Morph of aggressed intruder (B or W) | Focal ID |
| v) | Poisson | Log | Aggression received from heterospecifics \( (n = 18) \) | Morph of focal fish (B or W) | |
| (d) Laboratory aggression trials: Aggression levels | | | | | |
| i) | Negative binomial | Logit | Aggression count: all focal fish \( (n = 103) \) | Species \( (M. cal \) or \( M. zeb \)) + Sex (female or male) | |
| ii) | Negative binomial | Logit | Aggression count: male subset \( (n = 35) \) | Exp (1, 2, 3) | |
| iii) | Quasibinomial | Logit | Attack:Display: Exp 2&3 focals \( (n = 92) \) | Species \( (M. cal \) or \( M. zeb \)) + Sex (female or male) | |
| iv) | Negative binomial | Logit | Total aggression count: \( M. zeb \) \( (n = 65) \) | Population (CB, NB, TW) + Sex (female or male) | |
| (e) Laboratory aggression trials: Aggression biases | | | | | |
| Exp 1; BB focal fish, conspecific/heterospecific stimulus fish | | | | | |
| i) | Quasibinomial | Logit | Aggression count: BB males \( (n = 10) \) | Stimulus morph (BB or B) | |
| Exp 2; B&W focal fish, conspecific stimulus fish | | | | | |
| ii) | Quasibinomial | Logit | Aggression count: B&W focals \( (n = 28) \) | Focal morph (B or W) * Stimulus morph (B or W) + Sex (female or male) | |
| iii) | Quasibinomial | Logit | Aggression count: B&W focals \( (n = 28) \) | Stimulus morph (own or other) + Stimulus morph (common or rare) | |
| Exp 3; BB focal fish, heterospecific stimulus fish | | | | | |
| iv) | Quasibinomial | Logit | Aggression count: BB focals \( (n = 64) \) | Stimulus morph (B or W) + Sex (female or male) | |
| v) | Quasibinomial | Logit | Aggression count to B: BB focals \( (n = 64) \) | Population (CB, NB, TW) + Sex (female or male) | |

Abbreviations: CB, Chiofu Bay; *M. cal*, *M. callainos* (B, blue morph; *M. zeb*, *M. zebra* (BB, blue/black/barred morph); NB, Nkhata Bay; TW, Thumbsi West Island; W, white morph).
2.5.3 | Grazing differences between morphs

A generalized linear model was used to test for the effect of dominance status and morph/colour on grazing frequency in the field. Due to the overdispersion of the initial Poisson regression model, a negative-binomial model was chosen (Table 1b).

2.5.4 | Field observations of aggressive behaviour

Generalized linear models (Table 1c) were used to test for aggression biases by and towards the *Maylandia callainos* morphs in the field. Aggression directed towards territorial intruders of each of the two colour morphs was recorded and mixed effect binomial models (with focal fish ID included as a random factor) were used to test: (i) the effect of an interaction between focal and intruder colour (morph) on the aggression count, (ii) the effect of intruder colour on the amount of aggression received (regardless of focal fish morph), (iii) the effect of intruder colour on the amount of aggression received from territorial B males, and (iv) the effect of intruder colour on the amount of aggression received from territorial W males. Poisson regression was used to test for (v) the effect of *Maylandia callainos* territorial male colour on the amount of aggression received from heterospecific fish.

2.5.5 | Laboratory aggression trials

Generalized linear models (Table 1d) were used to test for the effects of species, population and sex on the level of aggression (total aggression count to both stimulus fish) recorded in pairwise intruder choice experiments. As initial Poisson regression models were all overdispersed, other models suitable for analysis of overdispersed count data were compared: Negative binomial models resulted in lower dispersion ratio, larger standard error and more conservative p-values than quasipoisson models and were therefore chosen to test: (i) all data for the effect of species and sex on level of aggression, (ii) a subset of males for the effect of species on the level of aggression. A quasibinomial model was used to test for (iii) the effect of species and sex on the ratio of attacks:displays. And finally, a negative binomial model was used to test for (iv) the effect of population and sex on the level of aggression displayed by *M. zebra* focal fish when presented with pairs of B and W *M. callainos* stimulus fish.

Generalized linear models were also used to test for aggression biases in the pairwise intruder choice experiments. Two statistical approaches were compared: Binomial mixed effect models with ‘focal ID’ as a random effect (data included aggression count from each focal fish to each of the two stimulus types) were overdispersed and the random effect variance was always zero (see Table S1 for the results of these models). The random effect was therefore omitted in order to use quasibinomial models (which are suitable for analysis of overdispersed count data and gave larger standard errors and more conservative p-values—Table 1e) to (i) confirm that conspecific aggression biases could be detected in this experimental set-up, (ii) test for an interaction between focal and stimulus morph on the aggression from *M. callainos* focal fish to different morph conspecifics, (iii) check for common-morph or own-morph aggression bias in *M. callainos*, (iv) test whether *M. callainos* stimulus fish colour has an effect on the amount of aggression received from *M. zebra* and (v) test whether *M. zebra* population has an effect on the aggression bias (depending on whether they co-occur with *M. callainos* or not).

3 | RESULTS

3.1 | Field observations (excluding aggression)

3.1.1 | Frequency of the blue and white morphs at Thumbi West Island

There were significantly more territory holding rare W morph males than would be expected given the proportion of W and B morph fish in the general population (Territorial B = 83, W = 59; General population B = 51, W = 23. G-test: $G_1 = 6.91, p = 0.009$; Figure 2a).

3.1.2 | Territory distances between morphs

First, whether a neighbour is of the same or different morph has a significant effect on the territory distance between males...
Males and their nearest neighbor are found at a greater distance from each other than territory neighbors (from the distance between territorial males and their nearest W distance recorded between territorial males, and significantly different (distance between B and nearest B is on average the same as, and not should on average be the smallest distance. This is not the case: The males, if the distance between morphs was only based on frequency, based on the observation of a lower frequency of W morph territorial W and W are found furthest apart, as this would be the assumption trial males of different morphs. While it is not surprising to find that p focal fish (n significant effect on the amount of aggression received from territorial p fish (Table 3): Compared to territorial males, non territorial fish grazed significantly less (p < 0.001), but regardless of social status B morph fish grazed significantly less than W morph fish (p < 0.001; Figure 2c).

### 3.3.1 | Laboratory aggression trials

The analysis of all data from the laboratory aggression experiments indicated that species had a significant effect on the overall level of aggression displayed (i.e. total aggression count to both stimulus fish); M. callainos were more aggressive than M. zebra, but sex had no significant effect on the level of aggression (Species p = 0.002; Sex p = 0.963—Table 5 model i). To test whether this difference in aggressiveness was a real difference between species, or due to M. callainos being presented more often with the possibility of being aggressive towards conspecific fish in these experiments, a subset of M. zebra CB and M. callainos males from the three experiments was compared (Table 5 model ii); M. zebra displayed a significantly (p = 0.001) higher level of aggression in the experiment where the stimulus pair consisted of one conspecific and one heterospecific (Exp. 1) compared to more aggression than their W morph counterparts. However, there is also a significant interaction between focal and intruder morph type (p = 0.026—Table 4 model i) and aggression bias appears to differ between the morphs: B morph intruders receive significantly more aggression from B morph territory holders than W morph intruders do (p < 0.001—Table 4 model iii), whereas B and W morph intruders do not differ significantly in the amount of aggression received from territorial W morph males (p = 0.332—Table 4 model iv; Figure 3a). Although on average B morph territorial males received more aggression than W morph males from heterospecifics this difference is not significant (p = 0.068—Table 4, model v; Figure 3c).

### 3.3 | Laboratory aggression trials

#### 3.3.1 | Differences in focal fish behaviour between species/(experiment), populations and sexes

Overall, the morph of an M. callainos territorial intruder has a significant effect on the amount of aggression received from territorial focal fish (p < 0.001—Table 4 model ii); B morph intruders receiving more aggression than their W morph counterparts. However, there is also a significant interaction between focal and intruder morph type (p = 0.026—Table 4 model i) and aggression bias appears to differ between the morphs: B morph intruders receive significantly more aggression from B morph territory holders than W morph intruders do (p < 0.001—Table 4 model iii), whereas B and W morph intruders do not differ significantly in the amount of aggression received from territorial W morph males (p = 0.332—Table 4 model iv; Figure 3a). Although on average B morph territorial males received more aggression than W morph males from heterospecifics this difference is not significant (p = 0.068—Table 4, model v; Figure 3c).

### 3.2 | Field observations of aggressive interactions

Both dominance status and morph colour had a significant effect on grazing frequency (Table 3): Compared to territorial males, non territorial fish grazed significantly less (p < 0.001), but regardless of social status B morph fish grazed significantly less than W morph fish (p < 0.001; Figure 2c).

### 3.1.3 | Grazing differences between morphs

The analysis of all data from the laboratory aggression experiments indicated that species had a significant effect on the overall level of aggression displayed (i.e. total aggression count to both stimulus fish); M. callainos were more aggressive than M. zebra, but sex had no significant effect on the level of aggression (Species p = 0.002; Sex p = 0.963—Table 5 model i). To test whether this difference in aggressiveness was a real difference between species, or due to M. callainos being presented more often with the possibility of being aggressive towards conspecific fish in these experiments, a subset of M. zebra CB and M. callainos males from the three experiments was compared (Table 5 model ii); M. zebra displayed a significantly (p = 0.001) higher level of aggression in the experiment where the stimulus pair consisted of one conspecific and one heterospecific (Exp. 1) compared to...
the experiment where they were presented with two heterospecific stimulus fish (Exp. 3). There was no significant difference ($p = 0.782$), however, in the level of aggression between the species in the experiments in which the stimulus pairs contained one conspecific (Exp. 1: *M. zebra* focal fish) or two conspecifics (Exp. 2: *M. callainos* focal fish) stimulus fish. This suggests that the presence or absence of a conspecific stimulus fish contributed to the overall difference in the level of aggression observed between the species in these experiments. There is therefore no evidence here of differences in intrinsic level of aggression between these species, rather that aggression among heterospecifics is lower than among conspecifics.

Species did not have a significant effect on the proportion of attacks:displays, but sex did. On average females attack more than they display, whereas males perform roughly equal proportions of attacks and displays ($p < 0.001$—Table 5 model iii).

Maylandia zebra population had a significant effect on the level of aggression displayed: sympatric Thumbi West (TW) fish, were significantly more aggressive to the heterospecific stimuli fish overall than those from the allopatric population at Chiofu Bay (CB), sex did not have a significant effect ($p < 0.001$—Table 5 model iv).

### 3.3.2 Aggression biases in pairwise intruder choice tests

Experiment 1: As expected, *M. zebra* stimulus fish received more aggression than *M. callainos* stimulus fish from *M. zebra* focal fish—that is, *M. zebra* focal fish showed a significant conspecific aggression bias ($p < 0.001$—Table 6 model i).
Experiment 2: Within *M. callainos*, sex had no effect on aggression count, but there was a significant interaction between morph of focal and stimulus fish (Interaction $p = 0.002$; Sex $p = 1.000$—Table 6). To further clarify whether the difference in the proportion of aggression directed to stimulus fish of each colour morph is due to an own morph aggression bias (i.e. each colour morph is more...
aggressive to other fish of the same colour) or an overall common morph aggression bias (i.e. fish of both colour morphs direct more aggression towards common/blue morph fish), stimulus type was recoded from blue/white to other/own and common/rare: Own-morph stimulus fish received significantly more aggression than other-morph fish, whereas whether the stimulus was common or rare had no significant effect (own/other $p = 0.001$; common/rare $p = 0.667$—Table 6 model iii, Figure 3b).

Experiment 3: M. callainos B-morph stimulus fish received significantly more aggression than W-morph fish from heterospecific M. zebra focal fish, and sex had no effect on this bias (stimulus morph $p < 0.001$; sex $p = 1.000$—Table 6 model iv, Figure 3d). The strength of aggression bias (proportion of aggression to B morph) differs between M. zebra populations: TW focal fish displayed a significantly lower proportion of aggression towards the blue morph than those from CB (i.e. a weaker bias), there was no difference between NB and CB and sex had no significant effect (population CB–TW $p = 0.047$, CB–NB $p = 0.400$; sex $p = 0.200$—Table 6 model v; Figure 3e).

### 4 | DISCUSSION

Our field observations of $M$. callainos, a polychromatic mbuna cichlid from Lake Malawi, indicated that common (blue) morph territorial intruders received more aggression than rare (white) morph intruders. Pairwise intruder choice tests in a controlled laboratory setting demonstrated that males and females of each morph bias aggression towards their own morph. These results add to the evidence that inherent own-morph aggression biases, which result in negative frequency-dependent selection on rare colour morphs, can contribute to balancing selection and thereby promote the maintenance of polychromatism (Dijkstra et al., 2008; Lehtonen, 2014; Scali et al., 2020). As there is little possibility for imprinting in these fish, and it seems unlikely that they can see their own colour, such an own-morph bias is likely the result of an inherent (genetically determined) behavioural/preferece polymorphism.

While this and previous studies (e.g. Dijkstra et al., 2008; Dijkstra, Hemelrijk, et al., 2009; Lehtonen, 2014; Scali et al., 2020; Seehausen & Schluter, 2004) have shown that aggression biases can
be involved in stabilizing polychromatism, to our knowledge this is the first study to identify benefits to rare morph cichlids which may result from receiving less aggression in the natural environment. We found there to be significantly more territory holding rare morph males than would be expected given the proportions of the two colours in the general population. Our observations suggest that the over-representation of rare morph territory owners may be influenced by two factors. First, different morph males appear to have higher tolerance of each other as neighbours, being found on average significantly closer to each other than same morph males. Additionally, the smallest average distance was found to be between white and blue males. Second, both territorial and non-territorial white morph individuals spend more time feeding, which suggests that the rare colour morph may benefit from lack of recognition during competition for non-mating related resources (Dijkstra et al., 2008; Lehtonen, 2014; Pérez i de Lanuza et al., 2017; Scali et al., 2020).

Further to showing that rare morph individuals can benefit from reduced intraspecific aggression, we found that a closely related ecologically similar heterospecific (M. zebra) also biases aggression towards the M. callainos blue morph. While these results are in conflict with those from another cichlid fish system, which suggest that rare morphs may be disadvantaged by greater heterospecific aggression (Lehtonen et al., 2015), given that aggression among heterospecifics is often higher among more similar coloured than more differently coloured species (Anderson & Grether, 2010; Genner et al., 1999; Losin et al., 2016; Pauers et al., 2008), it is not surprising to find that in some cases rare morph individuals may receives less aggression from a heterospecific which is more similar in colour to the common morph.

There was no difference in the total aggression displayed by males and females, nor in the aggression biases observed. There was however a difference in the number of attacks and displays used. We speculate that the greater use of direct attacks compared to display behaviours by females (this study and Arnott & Elwood, 2009) may be due to the difference in competition among females and males (i.e. greater use of attack behaviours may be more beneficial than displays when trying to win or defend caves among the rocks which are desirable shelters for mouthbrooding females).

In cichlids and other taxa, laboratory studies have shown that in species which differ in colour among allopatric populations, males tend to bias aggression towards males from their own population (Bolnick et al., 2016; Cooke & Turner, 2018; Tyers & Turner, 2013; Yang et al., 2018). In this study, we found that heterospecific aggression also varies depending on whether a pair of species occurs in sympatry or allopatry. The level of aggression (total aggression count to both M. callainos morphs) differs between M. zebra populations: M. zebra from Thumbi West (TW), which co-occur with high numbers of M. callainos, were significantly more aggressive to M. callainos than those from Chiofu Bay (CB), which are naïve to M. callainos. M. zebra from Nkhata Bay (NB), which co-occur with lower numbers of M. callainos showed intermediate level of aggression. These findings support the hypothesis that aggression among heterospecifics may often not simply be due to misdirected aggression among species (Peiman & Robinson, 2010), which would be indicated by higher levels of aggression from the allopatic M. zebra population (CB). The persistence of heterospecific aggression at NB support the idea that it has an adaptive function in long-term co-existing multi-species communities (Losin et al., 2016; Peiman & Robinson, 2010). Although there are no M. callainos at Chiofu Bay, this location is home to another closely-related species (M. esterae) which has blue males, and brown, orange and orange blotch females. M. zebra at Chiofu Bay therefore do co-occur with a similar blue morph fish, but no white morph fish and we found that the M. zebra from this location has a stronger blue-morph aggression bias than the other M. zebra populations which co-occur with blue and white M. callainos. A previous study of a polymorphic frog species found stronger aggression biases among morphs when they occur in allopatry compared to when they are found in sympatry (Yang et al., 2018).

Our results show a similar pattern in heterospecific aggression; a weaker blue morph aggression bias in M. zebra populations which coexist with both colour morphs.

Our results indicate that a rare colour morph may benefit from lack of recognition as a resource competitor, by both conspecifics and heterospecifics. This results in rare morph individuals receiving less aggression and gaining improved access to territories and food. This can benefit rare morph individuals while they are rare, but then what prevents them from increasing in frequency until fixation? Firstly, we found that rare (white) morph individuals were more aggressive towards their own morph than they were to the common (blue) morph, which would result in white morph individuals experiencing increasing levels of aggression as they became more common. Secondly, the lower level of heterospecific aggression bias towards the common morph, in populations with blue and white morphs, suggests that heterospecifics learn or evolve the ability to recognize rare morph individuals as competitors. The ability to recognize rare morph individuals may increase as they become more common: TW has the highest frequency of white morph individuals and the weakest common morph aggression bias by heterospecifics. Finally, female preference for common-morph males may result in a disadvantage to rare morph males (Roberts et al., 2009). The genes responsible for the expression of the melanin-disrupted (‘blotched’) morphs are almost always closely linked to a dominant female determinant, and so are generally much more common in females. This suggests that these colour phenotypes are disadvantageous to males, although they may be advantageous to females by providing increased crypsis or reduced aggression from conspecifics and/or heterospecifics. The reason for the differences in the types of rare morphs and frequency of rare morph individuals in different populations remains unclear but is likely due to multiple factors. Further investigation of differences in aggression biases and rare morph advantages in different mbuna communities, along with a detailed study of differences in morph crypsis and predation pressure in different populations/environments would greatly increase our understanding of the variation in polychromatism in these fish.
5 | CONCLUSIONS

Our results support previous studies indicating that negative frequency-dependent antagonism can be generated by own-morph aggression biases among conspecifics in cichlids which display polymorphism. We find that heterospecifics show reduced aggression to rare morph individuals, suggesting that heterospecific aggression may also facilitate invasion of rare colour morphs into a population. We identify potential advantages to rare morph individuals in the field, in terms of territory and foraging.

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CONFLICT OF INTEREST

The authors have no conflict of interest to declare.

AUTHOR CONTRIBUTION

Alexandra M. Tyers carried out all data analysis and laboratory experiments. Alexandra M. Tyers/Gavan M. Cooke/George F. Turner contributed to collection of field data and fish. Alexandra M. Tyers/George F. Turner drafted the manuscript.

DATA AVAILABILITY STATEMENT

The data that support the findings of this study can be viewed at https://doi.org/10.5061/dryad.0gb5mkm21.

PEER REVIEW

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REFERENCES

Anderson, C. N., & Grether, G. F. (2010). Interspecific aggression and character displacement of competitor recognition in Heteraerina damselfishes. Proceedings of the Royal Society of London B: Biological Sciences, 277, 549–555. https://doi.org/10.1098/rspb.2009.1371

Arnott, G., & Elwood, R. W. (2009). Gender differences in aggressive behaviour in convict cichlids. Animal Behaviour, 78, 1221–1227. https://doi.org/10.1016/j.anbehav.2009.08.005

Bolnick, D. I., Hendrix, K., Jordan, L. A., Veen, T., & Brock, D. (2016). Intruder colour and light environment jointly determine how nesting male stickleback respond to simulated territorial intrusions. Biology Letters, 12, 20160467. https://doi.org/10.1098/rsbl.2016.0467

Cooke, G. M., & Turner, G. F. (2018). Team colours matter when playing away from home: Aggression biases in geographically isolated mbuna cichlid populations. Hydrobiologia, 809, 31–40. https://doi.org/10.1007/s10750-017-3442-6

Couldridge, V. C. K., & Alexander, G. J. (2002). Colour patterns and species recognition in four closely related species of Lake Malawi cichlid. Behavioral Ecology, 13, 59–64. https://doi.org/10.1093/beheco/arv13.1.59

Dijkstra, P. D., Hemelrijk, C. K., Seebohm, O., & Groothuis, T. G. G. (2009a). Colour polymorphism and intrasexual selection in assemblages of cichlid fish. Behavioural Ecology, 20, 138–144. https://doi.org/10.1093/beheco/arm125

Dijkstra, P. D., Seebohm, O., & Groothuis, T. G. G. (2008). Intrasexual competition among females and the stabilization of a conspicuous colour polymorphism in a Lake Victoria cichlid fish. Proceedings of the Royal Society of London B: Biological Sciences, 275, 519–526. https://doi.org/10.1098/rspb.2007.1441

Dijkstra, P. D., Seebohm, O., Pierrotti, M. E. R., & Groothuis, T. G. G. (2007). Male-male competition and speciation: Aggression bias towards differently coloured rivals varies between stages of speciation in a Lake Victoria cichlid species complex. Journal of Evolutionary Biology, 20, 496–502. https://doi.org/10.1111/j.1420-9101.2006.01266.x

Dijkstra, P. D., van Dijk, S., Groothuis, T. G. G., Pierrotti, M. E. R., & Seebohm, O. (2009). Behavioral dominance between female color morphs of a Lake Victoria cichlid fish. Behavioural Ecology, 20, 593–600. https://doi.org/10.1093/beheco/arp036

Drury, J. P., Cowen, M. C., & Grether, G. F. (2020). Competition and hybridization drive interspecific territoriality in birds. Proceedings of the National Academy of Sciences of the United States of America, 117, 12923–12930. https://doi.org/10.1073/pnas.1921380117

Genner, M. J., Turner, G. F., & Hawkins, S. J. (1999). Resource control by territorial male cichlids in Lake Malawi. Journal of Animal Ecology, 68, 522–529. https://doi.org/10.1111/j.1365-2656

Grether, G. F., Losin, N., Anderson, C. N., & Okamoto, K. (2009). The role of interspecific interference competition in character displacement and the evolution of competitor recognition. Biological Reviews, 84, 617–635. https://doi.org/10.1111/j.1469-185X.2009.00089.x

Henze, M. J., Lind, O., Mappes, J., Rojas, B., & Kelber, A. (2018). An aposematic colour-polyomorph moth seen through the eyes of conspecifics and predators - Sensitivity and colour discrimination in a tiger moth. Functional Ecology, 32, 1797–1890. https://doi. org/10.1111/1365-2435.13100

Hughes, K. A., Houde, A. E., Price, A. C., & Rodd, F. H. (2013). Mating advantage for rare males in wild guppy populations. Nature, 503, 108–110. https://doi.org/10.1038/nature12717

Huxley, J. (1955). Morphism and evolution. Heredity, 9, 1–52. https://doi. org/10.1038/hdy.1955.1

Jordan, R. C. (2008). Color-based association among heterospecics in Lake Malawi rock-dwelling cichlids. Ethology, 114, 272–278. https://doi.org/10.1111/j.1439-0310.2007.01460.x

Kim, K. W., Jackson, B. C., Zhang, H., Toews, D. P. L., Taylor, S. A., Greig, E. I., Lovette, I. J., Liu, M. M., Davison, A., Griffith, S. C., Zeng, K., & Burke, T. (2019). Genetics and evidence for balancing selection of a sex-linked colour polymorphism in a songbird. Nature Communications, 10, 1852–1861. https://doi.org/10.1038/s41467-019-09806-6

Konings, A. (2007). Malawi cichlids in their natural habitat (4th ed.). Cichlid Press.

Lande, R., Seebohm, O., & van Alphen, J. J. M. (2001). Mechanisms of rapid sympatric speciation by sex reversal and sexual selection in cichlid fish. Genetica, 112–113, 435–443.

Lehtonen, T. K. (2014). Colour biases in territorial aggression in a Neotropical cichlid. Oecologia, 175, 85–93. https://doi.org/10.1007/s00442-013-2879-1

Lehtonen, T. K., Sowersby, W., & Wong, B. B. M. (2015). Heterospecific aggression bias towards a rare colour morph. Proceedings of the
Pérez i de Lanuza, G., Carretero, M. A., & Font, E. (2017). Intensity of
Ribbink, A. J., Marsh, B. A., Marsh, A. C., Ribbink, A. C., & Sharp, B. J. (1983). A preliminary survey of the cichlid fishes of rocky habitats in Lake Malawi. South African Journal of Zoology, 18, 149–308. https://doi.org/10.1080/02541858.1983.11447831

Roberts, R. B., Ser, J. R., & Kocher, T. D. (2009). Sexual conflict resolved by invasion of a novel sex determination in Lake Malawi cichlid fishes. Science, 326, 998–1001. https://doi.org/10.1126/science.1174705

RStudio Team. (2019). RStudio: Integrated development for R. Rstudio. PBC. http://www.rstudio.com/

Scali, S., Mangiacotti, M., Sacchi, R., Coladonato, A. J., Falaschi, M., Saviano, L., Rampoldi, M. G., Crozi, M., Perotti, C., Zucca, F., Gozzo, E., & Zuffi, M. A. L. (2020). Close encounters of the three morphs: Does color affect aggression in a polymorphic lizard? bioRxiv. https://doi.org/10.1101/2020.05.02.074146

Seehausen, O., & Schluter, D. (2004). Male–male competition and nuptial-colour displacement as a diversifying force in Lake Victoria cichlid fishes. Proceedings of the Royal Society of London B: Biological Sciences, 271, 1345–1353. https://doi.org/10.1098/rspb.2004.2377

Seehausen, O., & van Alphen, J. J. M. (1998). The effect of male coloration on female mate choice in closely related Lake Victoria cichlids (Haplochromis nyererei complex). Behavioral Ecology and Sociobiology, 42, 1–8. https://doi.org/10.1007/s002650050405

Seehausen, O., van Alphen, J. J. M., & Lande, R. (1999). Color polymorphism and sex ratio distortion in a cichlid fish as an incipient stage in sympatric speciation by sexual selection. Ecology Letters, 2, 367–378. https://doi.org/10.1046/j.1461-0248.1999.00098.x

Streelman, J. T., Albertson, R. C., & Kocher, T. D. (2003). Genomemapping of the orange blotch colour pattern in cichlid fish. Molecular Ecology, 12, 2465–2471. https://doi.org/10.1046/j.1365-294x.2003.01920.x

Surmacki, A., Ożarowska-Nowicka, A., & Rosin, Z. M. (2013). Color polymorphism in a land snail Cepaea nemoralis (Pulmonata: Helicidae) as viewed by potential avian predators. Naturwissenschaften, 100, 533–540. https://doi.org/10.1007/s00114-013-1049-y

Svensson, E., & Sefc, K. M. (2013). Colour variation in cichlid fish: Developmental mechanisms, selective pressures and evolutionary consequences. Seminars in Cell & Developmental Biology, 24, 526–528. https://doi.org/10.1016/j.semcdb.2013.05.003

Tyers, A. M., & Turner, G. F. (2013). Signal and preference divergence among populations of the non-endemic basal Lake Malawi cichlid fish Astatotilapia calliptera (Perciformes: Cichlidae). Biological Journal of the Linnean Society, 110, 180–188. https://doi.org/10.1111/bij.12105

Wellenreuther, M., Svensson, E. I., & Hansson, B. (2014). Sexual selection and genetic colour polymorphisms in animals. Molecular Ecology, 23, 5398–5414. https://doi.org/10.1111/mec.12935

Yang, Y., Dugas, M. B., Sudekum, H. J., Murphy, S. N., & Richards-Zawacki, C. L. (2018). Male–male aggression is unlikely to stabilize a poison frog polymorphism. Journal of Evolutionary Biology, 31, 457–468. https://doi.org/10.1111/jeb.13243

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