You can't always get what you want: size assortative mating by mutual mate choice as a resolution of sexual conflict

Sebastian A Baldauf*, Harald Kullmann†, Stefanie H Schroth†, Timo Thünken† and Theo CM Bakker†

Address: Institute for Evolutionary Biology and Ecology, University of Bonn, An der Immenburg 1, D-53121 Bonn, Germany

Email: Sebastian A Baldauf* - sbaldauf@evolution.uni-bonn.de; Harald Kullmann - h.kullmann@uni-muenster.de; Stefanie H Schroth - st.schroth@gmx.de; Timo Thünken - tthuenken@evolution.uni-bonn.de; Theo CM Bakker - t.bakker@uni-bonn.de

* Corresponding author †Equal contributors

Abstract

Background: Assortative mating patterns for mate quality traits like body size are often observed in nature. However, the underlying mechanisms that cause assortative mating patterns are less well known. Sexual selection is one important explanation for assortment, suggesting that i) one (usually the female) or both sexes could show preferences for mates of similar size or ii) mutual mate choice could resolve sexual conflict over quality traits into assortment. We tested these hypotheses experimentally in the socially monogamous cichlid fish Pelvicachromis taeniatus, in which mate choice is mutual.

Results: In mate choice experiments, both sexes preferred large mates irrespective of own body size suggesting mating preferences are not size-assortative. Especially males were highly selective for large females, probably because female body size signals direct fitness benefits. However, when potential mates were able to interact and assess each other mutually they showed size-assortative mating patterns, i.e. the likelihood to mate was higher in pairs with low size differences between mates.

Conclusion: Due to variation in body size, general preferences for large mating partners result in a sexual conflict: small, lower quality individuals who prefer themselves large partners are unacceptable for larger individuals. Relative size mismatches between mates translate into a lower likelihood to mate, suggesting that the threshold to accept mates depends on own body size. These results suggest that the underlying mechanism of assortment in P. taeniatus is mutual mate choice resolving the sexual conflict over mates, rather than preference for mates of similar size.

Background

Sexual selection is a general selective force that has been intensively investigated in the past decades [see [1]]. The form and intensity of sexual selection are highly variable among species. In species in which mating is non-random, females are often expected to be choosy, especially when male parental investment is low. However, choosiness may also occur in males and should be promoted in both sexes when variances in quality of potential mates and parental investment are high [2,3]. Hence, mating preferences in each sex could generate a zone for sexual conflict, e.g. over mate quality [3,4].
Body size plays an important role in sexual selection of many species [1,5,6], including humans [7,8]. Usually, larger individuals are favored. For example, larger males may be preferred by females because of direct benefits, for instance better territorial defence [e.g. [9]], thereby providing better environmental conditions for raising offspring [10]. However, in the field and laboratory size-assortative mating is often reported and therefore much research has concentrated on its occurrence and the causal mechanisms [e.g. [11-14]]. Nevertheless, knowledge about assortment and its underlying mechanisms in mating systems with mutual mate choice is scarce.

Many mechanisms may lead to assortative mating patterns, for example spatial or temporal distribution. However, mate choice is one important explanation for assortment [see [15] and citations therein], and provides several different mechanisms that may result in assortative mating patterns. First, one sex (usually the female) or both sexes may show assortative mating preferences, e.g. with respect to body size [16,17]. Second, assortment may be the consequence of intra-sexual competition [18], so that only the larger winning individual is able to choose its mating partner. For example, in the curculionid beetle Diaprepes abbreviatus only competitive large males chose large, highly fecund females, whereas the losers had to accept the females of lower quality [19,20]. Third, mutual mate choice could explain assortment by resolving sexual conflict over mate quality traits [3,4]. When both sexes show preferences [21] for high quality mates, irrespective of own quality, low quality mates may be unacceptable. As a consequence, only high quality individuals would find partners which meet their preference. In contrast, low quality individuals would fail to mate at all [4], unless they show a choice threshold that accepts potential mates correlating to their own quality. In this case, resulting mating patterns would be assortative.

To test this hypothesis, it is necessary to show 1) directional mating preferences of both sexes, i.e. to exclude assortative preferences, and 2) size-assortative mating patterns independent of intra-sexual competition.

The study organism, Pelvicachromis taeniatus, is a cave-breeding cichlid that is characterized by a high level of cooperative biparental care [22]. Parental care often has high costs [2,3,23], e.g. due to predation risk, and time and energy loss. Thus, the probability of future mating may be negatively influenced by parental effort. When the costs of biparental care are high, males and females should be choosy [2-4]. Both sexes of P. taeniatus make a large investment in brood care and were choosy in prior experiments [22,24]. Pairs are socially monogamous and alternative mating tactics have not been observed yet. The males' testis size is very small, even suggesting genetic monogamy; however, the sperm length is extraordinarily long [25]. Males and females show sexual dimorphism in body size. Males are usually larger than females. Furthermore, both sexes are conspicuously colored. Males show, among others, a yellow nuptial coloration of their ventral body, while females develop a violet ventral coloration. During courtship, both sexes present their ventral region by arching it towards the partner while intensely quivering the whole body. After mutual mate choice and spawning the female cares for the eggs in the cave while the male defends the territory against intruders. The fry swims free after about one week and is then guarded by both parents.

The aim of our study was to test (1) whether mating preferences for body size of males and females are present in P. taeniatus and (2) the impact of size differences between potential mates on the likelihood to mate when partners are able to interact and assess each other mutually. In order to measure mating preferences for body size, we conducted a series of choice experiments using computer animations of digital images of different sizes of the opposite sex as stimuli. Test fish that greatly varied in body size were chosen for the experiments, thus preferences for assortative mating could be tested. A general problem when investigating preferences for body size is that size is often correlated with other factors. For example, large individuals might differ in behavior from smaller ones. A striking advantage of computer-manipulated stimuli is a high degree of standardization between the stimuli [26,27], thus minimizing the effects of confounding variables like rapid changes in coloration or different responses in stimuli fish.

In a second experiment, the consequences of size difference between the sexes on the likelihood to mate as well as the brood care intensity after successful mating were tested. Furthermore, we investigated whether body size could signal potential mate quality by analyzing variables such as the amount of parental investment, egg number of females and the number of surviving offspring.

**Results**

**Size preferences in computer animations**

**Male mate choice**

Males showed a highly significant preference for the larger female stimulus in each treatment (small vs. medium: n = 19, z = 3.601, p < 0.001; small vs. large: z = -4.087, n = 21, p < 0.001; medium vs. large: n = 24, z = -2.754, p = 0.006; Fig. 1). The order of the treatments had no significant effect (LRT: all p > 0.078).

With the exception of the "medium vs. large" treatment, the males' standard length did not significantly correlate with the time spent in front of the stimuli (Pearson and Spearman correlations: all p > 0.177). The males' standard
length correlated significantly negative with the time spent in front of the large female stimulus in the "medium vs. large" treatment (Spearman correlation: \( n = 24, r = -0.536, p = 0.007 \)).

The preference for the larger female tended to be stronger in the "small vs. large" treatment compared to the "small vs. medium" treatment (Wilcoxon test: \( n = 19, z = -1.913, p < 0.056 \)), as well as in the "small vs. large" compared to the "medium vs. large" treatment (Wilcoxon test: \( n = 21, z = -1.899, p = 0.058 \)). No significant difference was found between the "small vs. medium" and "medium vs. large" treatment (paired t-test: \( n = 19, t = 0.894, p = 0.383 \)).

**Female mate choice**

Females associated high significantly longer with the larger male in the "small vs. large" treatment (paired t-test: \( n = 23, t = -5.042, p = 0.001 \); Fig. 2) but showed no significant preferences for the larger one in the two remaining treatments (paired t-tests: small vs. medium: \( n = 21, t = 0.812, p = 0.427 \); medium vs. large: \( n = 24, t = -0.944, p = 0.355 \); Fig. 2). However, a repeated measures analysis of

---

**Figure 1**

Male preference. Male preference expressed as percent of time spent in the association zone in front of female stimuli of different body size. Plotted are median, quartiles, percentiles. *** \( p < 0.001 \), ** \( p < 0.01 \), * \( p < 0.05 \), n.s. = not significant.

---

**Figure 2**

Female preference. Female preference expressed as percent of time spent in the association zone in front of male stimuli of different body size. Plotted are means + SD. *** \( p < 0.001 \), ** \( p < 0.01 \), * \( p < 0.05 \), n.s. = not significant.
variance revealed that females high significantly discrimi-
nated between the smaller or larger male stimulus
(ANOVA: F1,38 = 13.910, p = 0.001). The order of the treat-
ments had no significant effect (LRT: all p > 0.148). The females' standard length did not significantly correlate with the time spent in front of the stimuli (Pearson and Spearman correlations: all p > 0.138).

The preference for the larger male was significantly
stronger in the "small vs. large" than in both the "small vs.
medium" treatment (Wilcoxon test: n = 21, z = -2.950, p
< 0.003) and the "medium vs. large" treatment (Wilcoxon
test: n = 23, z = -2.516, p < 0.012). No significant differ-
ence was found between the "small vs. medium" and
"medium vs. large" treatments (paired t-test: n = 21, t = -
0.980, p = 0.339).

Comparison of preferences between the sexes
In all treatments, males spent significantly relative more
time than females with the larger stimulus of the opposite
sex, whereas females spent significantly more absolute
time in association zones than males (Table 1).

Control experiment
Both sexes significantly preferred the large stimulus
(males: mean ± SD = 78.41% ± 21.39%; females: mean ±
SD = 79.57% ± 26.54%) over the small stimulus (males:
mean ± SD = 21.59% ± 21.39%; females: mean ± SD =
20.43% ± 26.54%) when the extent of nuptial coloration
was kept constant between both stimuli (males: paired t-
test: n = 11, t = 4.404, p = 0.001; females: paired t-test: n
= 13, t = 4.018, p = 0.002).

Mating and brood care experiment
Overall, 15 pairs mated and nine pairs did not mate. The
likelihood to mate was significantly explained by the dis-
tance of pairs (LRT: χ2 = -4.508, df = 1, p = 0.034; Fig. 3)
and the relative size difference of pairs (LRT: χ2 = -4.344,
df = 1, p = 0.037). Within the group of unmated pairs,
females smaller than the mean female standard length of the population showed significantly more courtship
behavior towards the males than females larger than the
mean female standard length (Mann-Whitney U test: Nmale smaller = 4, Nmale larger = 5, z = -2.205, p = 0.032). The
number of days until spawning was not significantly
explained by the females’ standard length (Cox regression:
LRT1,24 = 0.83, beta = 0.545 ± 0.571, p = 0.358). Body size
did not significantly correlate with different aspects of
parental investment during brood care (e.g. the time
males guarded caves, the female cared for eggs, the fre-
quency of warning signals, time near the young; all p > 0.313). The standard length of the females tended to
explain the number of eggs (LRT: F1,12 = 4.403, p = 0.057)
and significantly explained the number of surviving off-
spring (LRT: F1,12 = 6.810, p = 0.023).

Discussion and conclusion
Males, as well as females of P. taeniatus showed preference
for larger mates. However, the likelihood to mate in P. tae-
niatus was higher, when the size difference between poten-
tial mates was low, thus resulting in size-assortative
mating. This corresponds to previous findings in cichlids
concerning assortative mating patterns [21,28-31] and
preference for larger mates [9,32-34].

The results allow the discussion of the underlying mecha-
nism of the assortment because both individual prefer-
ces for body size and their consequences on the
likelihood to mate within the same species were investi-
gated. The hypothesis of assortment as a consequence of
individual preference for size-assortative mating can be
ruled out because no significant positive relationships and

| Treatment            | mean ± SD or median (quartiles) t or z n p |
|----------------------|------------------------------------------|----------------|
| a)                   |                                          |                |
| small vs. medium     | Males: 78.27% ± 19.50%                   | 4.568          |
|                      | Females: 45.59% ± 24.91%                 |                |
| small vs. large      | Males: 100% (96.38%; 100%)               | -2.752         |
|                      | Females: 86.59% (50.00%; 100%)           |                |
| medium vs. large     | Males: 84.56% (51.35%; 100%)             | -2.478         |
|                      | Females: 50.00% (37.95%; 84.05%)         |                |
| b)                   |                                          |                |
| small vs. medium     | Males: 60.00 s ± 25.00 s                 | 4.427          |
|                      | Females: 93.52 s ± 22.89 s               |                |
| small vs. large      | Males: 77.48 s ± 34.79 s                 | -2.377         |
|                      | Females: 97.41 s ± 17.10 s               |                |
| medium vs. large     | Males: 66.75 s (51.00 s; 58.63 s)        | -3.177         |
|                      | Females:104.25 s (88.50 s; 113.38 s)     |                |

Differences between males and females concerning a) the relative amount of time at the larger stimulus side and b) the absolute time in both association zones. Differences between the sexes were tested with Wilcoxon or t-tests. t or z = test statistics; n = sample size; p = probability.
one negative correlation between individual body size and preference for the larger stimulus were observed in both sexes. Thus, when given free choice, both sexes of *P. taeniatus* show preference for large potential mates. Furthermore, intra-sexual competition as a mechanism was excluded by the experimental design, although under natural conditions it might also play a role. Due to the mutual character of courtship in the mating system, both males and females are able to accept or reject mates, depending on their individual size compared to that of the mating partner. Consequently, large males could reject small females and *vice versa*. The rejection of relatively small mates is supported by our data since large males did not mate with small females, although these females showed intense courtship behavior. Rejected individuals may fail to mate at all, or have a lower threshold to accept mates that are almost as small as themselves [4]. Thus, the underlying mechanism of size-assortment in *P. taeniatus* is the resolution of sexual conflict over large mates by mutual mate choice [3,4].

Whereas males preferred the larger female in all treatments, females selected the larger male only when the size difference between both stimuli was maximal. Females may not have been able to perceive small differences between stimuli due to the set-up, given that 2D-stimuli may look similar in size when the female is not in the middle of the tank. However, reference objects had been installed. Moreover, males were able to discriminate between the smaller and the larger female stimulus in all treatments, although female stimuli were on average smaller than male models.

Females had a less strong preference for larger mates than males, but spent more time in the association zones. On the one hand this could mean, that females may be less selective for the trait and must present themselves to the choosier sex [1]. On the other hand, each cave-occupying male may initially be attractive for females [35], but an increase of size difference among males may enhance the importance of male body size for female mate choice. A greater difference between stimuli led to a significant preference for the larger male stimulus, indeed. Thus, females may be choosy, but a smaller difference in male size may not be important. As paternal investment of *P. taeniatus* in brood care seems to be independent of males' body size in our study, the offspring may gain the same investment from a medium-sized or large male. However, other possible advantages of larger males, e.g. protection against predators, were not investigated in this study.

The adaptive significance of sexual selection on body size has been discussed in many studies [14,36]. In the present study, female body-size in *P. taeniatus* indicated the number of surviving offspring, thus supporting previous findings that females' body size reveals fecundity [e.g. [21] and citations therein, [37]]. Thus, preference for large females could increase male fitness. The impact of large male body size was not investigated in the present study. Larger males may be more successful in intra-sexual competition, offering better territories or protection against predators [9,38,39]. Furthermore, if larger males in *P. taeniatus* have advantages in intra-sexual competition (TT, unpublished data) like larger males in other species [e.g. [40-42]] females may prevent the loss of a brood when preferring the larger male.

The parental investment of males – as well as females – seemed to be independent of body size during brood care in *P. taeniatus*. On the one hand this could mean, that body size does not signal the quality of a potential mating partner concerning brood care. On the other hand, brood care may be performed by each sex on a constant high level independent of its own or the partner's body size. However, in our study parental investment of *P. taeniatus* was measured in size-assortative mated pairs which do

---

**Figure 3**

**Likelihood to mate.** The likelihood to mate in *P. taeniatus* was significantly explained by the relative body size distance of pairs (see equation 1 and Results section for statistics). An equal distance was measured in four pairs (filled gender symbols), thus 20 instead of 24 data points are shown. Gender symbols reveal whether the male (male symbol) or the female (female symbol) was relatively larger in a pair. In one case an equal distance between pairs with a relatively larger male and a relatively larger female was measured (male and female symbol combined).
not represent the largest individuals of the population used. Of course, parental investment was assessed under optimal laboratory conditions leaving an influence of body size under more stressful conditions open.

There are two general problems when investigating the effects of body size on mating decisions. First, in many species body size may be correlated to other traits. Thus, it is impossible to disentangle which trait is ultimately selected for. Second, the relative proportion of other traits with respect to body size may be more important, making body size relevant as a combined trait [e.g. [32]]. The control experiment in which the area of nuptial coloration was kept constant independent of body size revealed a strong preference for the larger stimulus in both sexes, underlining that body size is decisive for each sex independent of the extent of the nuptially colored area. We cannot exclude that there are unknown sexual traits that shift with the downscaling of the stimulus' body size. Nonetheless, this method of investigating body size provides the highest possible degree of standardization.

In general, the sex with the higher reproductive costs is expected to be more selective which is usually assumed to be the female [see [43] for a recent model explaining assortment by female mate choice]. Our results show that *P. taeniatus* may be a good model species for research questions concerning mutual mate choice. A recent review revealed that mutual mate choice plays a role in many species [see [44] for an overview of species showing mutual mate choice or positive assortment]. Thus, assortment by mutual mate choice may be an important mechanism for the resolution of sexual conflict over quality traits in many species.

**Methods**

**Experimental animals**

All individuals of *P. taeniatus* were bred and maintained under standardized laboratory conditions [see [22]]. The parents of the test subjects originated from the river Moliwe in Cameroon (04°04’N/09°16’E), West Africa. Individuals were F1- and F2-offspring from 23 outbreeding pairs and were raised in mixed-sex family tanks (80 × 30 × 30 cm). The tanks were surrounded with opaque plastic sheets to avoid visual contact to other aquaria. Test individuals were F1- and F2-offspring from 23 outbreeding pairs and were raised in mixed-sex family tanks (80 × 30 × 30 cm). The tanks were surrounded with opaque plastic sheets to avoid visual contact to other aquaria. Test fish were 1–2 years old and reproducitively active. The water temperature was kept at 25 ± 1°C and natural light conditions were given (L/D 12/12). Nutrition was provided once a day with a mixture of frozen chironomid larvae and *Artemia* ssp.

**Mate choice using computer animations**

**Preparation of artificial stimuli**

We took digital photographs (Olympus Camedia Widezoom 5060) of a nuptially colored male and female to obtain source data for two-dimensional fish models. 2D-models are a sufficient method to test mating preferences in *P. taeniatus* for certain traits like body size (SAB, HK, TT, TCMB unpublished data). The pictures were saved in RAW-format to avoid the loss of coloration data due to algorithmic compression. They were white-balanced during import to Adobe Photoshop CS2. The size dimensions of the pictures were manipulated for the three treatments (Table 2).

To achieve moving animations of the models, we used "The GIMP 2.20 with animation package". A grey background image (1024 × 400 px) was created (RGB: 238,238,238) including a reference object for each sex (plant for test males/breeding cave for test females, in the middle of the image). Each animation consisted of 30 frames per second which is an established method to present artificial stimuli to test fish [26,27,45]. Each stimulus moved a horizontal pathway from one side of the monitor to the other for a period of 15 seconds, including a second stop in the middle. After that, it recurred horizontally and moved back in the same time frame.

For each sex we created three different experimental treatments (Table 2) with different body size of the stimuli. The body size of each stimulus was adapted for a monitor and video resolution of 1024 pixel width.

**Experimental design**

Experiments were conducted between 01 June 2007 and 02 August 2007. Before the start of the trials 30 males and 24 females, that varied greatly in body size (Table 3), were randomly chosen and individually isolated in separate tanks (25 × 15 × 15 cm) for a minimum period of two days. The mating readiness of each test fish was determined visually on the basis of the ventral coloration, as well as the display of courtship behavior in the family tanks [22,24]. The isolation tanks were surrounded at the broad sides by print-outs of the animation's background image and opaque, grey partitions at the longer sides, thus ensuring that fish did not interact with other isolated individuals and were able to habituate to the background and reference objects also used in the trials. Java moss, *Vesicularia dubynana*, was added to provide shelter for the females, whereas the male tanks were equipped with a breeding cave. All other conditions were similar to those of the mixed-sex tanks. Each individual conducted the three experimental treatments in random order. Test fish were transported to the experimental set-up in their isolation tank, thus reducing stress by leaving the fish in its familiar habitat.

The set-up was illuminated by a fluorescent tube (37W) installed one meter above the middle of the tank. Additionally, white Styrofoam surrounded the tank. Tanks
containing the test fish were placed between two CRT monitors of the same model (EIZO FlexScan F520, 85 Hz, connected to a Matrox G550 PCIe graphic board). The level of the bottom of the tank was justified so that it coincided with the lower margins of the monitor screens. An association zone of 5 cm in front of each monitor was marked on the white Styrofoam under the tank creating a 15 cm neutral zone in between.

It was determined randomly which stimulus was presented on which monitor. During an acclimatization period of 15 minutes both screens showed the grey background. After acclimatization, the stimuli appeared simultaneously on both monitors [46]. In which direction the stimuli were moving (animations) was randomly determined. We recorded two minutes after the fish had visited the first association zone, which is a time frame established in recent studies [47-49]. After recording, the empty background was shown again for five minutes. The trial was then repeated with switched monitor sides of the stimuli.

After the three experimental treatments, the standard length and mass of the test fish was measured. A novice observer analyzed the video recordings. Mating preferences were measured as association time near a stimulus of the opposite sex, which reliably predicts mating decisions in both male and female P. taeniatus ([22] and references therein, [24], TT, TCMB, N. Henning, HK unpublished data). The time spent in each association zone was calculated over a period of two minutes after the fish had visited an association zone. In the analyses, relative proportion of time spent in one of the association zones was used. For each test fish, we averaged the time spent in front of each stimulus in the first and the second trial, thus excluding possible side biases. In each treatment a different number of test fish completed both trials, thus sample sizes varied among treatments. To compare the preferences for stimuli between the three treatments, we subtracted the relative time spent in front of the smaller stimulus from the larger stimulus for each sex.

Control experiment

Small stimuli in the computer animations were down-sized versions of the larger stimuli. Thus, other sexual traits possibly playing a role in sexual selection were also down-scaled like the extent of nuptial coloration of each sex. We therefore carried out a control experiment to test whether preference for body size in each sex was independent from the extent of nuptial coloration. Experiments were conducted between 03 December 2007 and 14 December 2007. The methods complied with the set-up described above. We used the small and the large stimulus for each sex from the former experiments as stimuli, but kept the area of nuptial coloration constant in both stimuli. Both stimuli showed the extent of nuptial coloration of the small stimulus.

Mating and brood care experiment

Observations were made in winter 2007. We formed 24 outbreeding pairs differing in relative body size distance by taking nuptially colored individuals of each sex from family tanks, measuring their standard length, and sorting males (range: 4.9–7.5 cm) in descending and females (range: 3.4–5.0 cm) in ascending order of body size. Then the largest male was combined to the smallest female and vice versa (i.e. largest relative size distance, see equation 1 and 2 below) with a continuous shift of relative size distance of pairs in between. The pairs were allocated randomly to aquaria (30 × 40 × 43 cm), which were filled with 2/3 parts of tap water and 1/3 parts of osmotic water and tempered at 25 ± 1°C. Two observers (SHS, SAB) recorded behavioral patterns of each pair for 10 min daily over a period of eight weeks. The observation order was random and determined daily. Unmated pairs were stimulated weekly by renewing 1/3 of the water volume with the 2:1 tap/osmotic water mix, but observations were stopped for pairs that were still unmated after three weeks.

After spawning, the cave was shortly removed and the number of eggs was counted. The egg number of one female could not be determined. We recorded several behavioral patterns during courtship and parental care [see [22] for details]. We recorded every 30 seconds whether the male guarded the cave or not, or whether the female cared for the eggs. Mating was counted as successful when fry left the breeding cave and the parents showed brood care for the free swimming offspring. The number of surviving young was counted four weeks after the fry had left the breeding cave. The sample size here was reduced by one, as one pair cannibalized their fry after three weeks. Fry were fed with living Artemia nauplii, and

Table 2: Total lengths of the manipulated images

| Treatment       | Female stimuli size (cm) | Male stimuli size (cm) |
|-----------------|--------------------------|------------------------|
| small vs. medium| 3.5 vs. 4.5               | 5.0 vs. 6.5             |
| small vs. large | 3.5 vs. 5.5               | 5.0 vs. 8.0             |
| medium vs. large| 4.5 vs. 5.5               | 6.5 vs. 8.0             |

Overview of pairs of artificial stimuli used in measuring mating preferences for body size. Each test fish conducted all three treatments in random order.
when they grew older with a mix of frozen Artemia, chironomid, and mosquito larvae.

In order to analyze the likelihood to mate we calculated the body size distance of each pair relative to the mean population size difference, called "distance of pairs" from now on (Equation 1). However, males and females achieve different maximum body sizes: a large female and a large male have different absolute body sizes, but may be the same in relative size compared to other individuals of their own sex. Thus, we additionally calculated the relative size difference between a male and a female in relation to the mean body size of each sex in the population, so called "relative size difference of pairs" from now on (Equation 2).

\[
\text{Distance of pair}_{(i)} = \frac{((\text{male}_{(i)} \text{ standard length} - \text{female}_{(i)} \text{ standard length}) - \text{mean (male standard length}_{(\text{population})} - \text{female standard length}_{(\text{population})})}{100} \\
\text{Relative size difference of pair}_{(i)} = \frac{((\text{male}_{(i)} \text{ standard length} / \text{mean male standard length}_{(\text{population})} \times 100) - ((\text{female}_{(i)} \text{ standard length} / \text{mean female standard length}_{(\text{population})} \times 100))}{100}
\]

**Statistics**

Parametric statistics were used when data did not significantly deviate from normal distribution according to Kolmogorov-Smirnov tests with Lilliefors correction. Given test probabilities are two-tailed throughout. p-values <0.05 were considered statistically significant. Male body preferences in the computer animation experiments were additionally analyzed by a repeated measures analysis of variance. Because male data were not normally distributed (not even after transformation), only non-parametric test statistics are shown. The likelihood to mate in relation to the distance of pairs (see equation 1) and relative size difference of pairs (see equation 2) were analyzed with generalised linear models ("glm"), with binomial error distribution and logit link function. The impact of body size on the number of eggs or surviving young was analyzed with linear models ("lm"). Likelihood-ratio tests (LRT) assessed whether the removal of a variable caused significant decrease in model fit. Reported p-values of models refer to the increase in deviance when the respective variable was removed ("lm": F-statistics; "glm": Pearson's chi-square). Analyses were performed using SPSS 12.0, and LRTs were calculated using R 2.6.1 statistical package.

**Authors' contributions**

SAB participated in the design of the study, collected data, performed statistical analyses and wrote the paper. HK participated in the design of the study, the statistical analyses and the writing of the paper. SHS collected data and performed statistical analyses. TT participated in the design of the study, statistical analysis and the writing of the paper. TCMB participated in the design of the study and the writing of the paper. All authors read and approved the final manuscript.

**Acknowledgements**

We thank A. Hornberg and S. Brungs for assistance during the control experiments, L. Engevist and T. Schmoll for statistical support and R. Modarressie, J. G. Frommen and I. P. Rick for discussion. M. Hiermes is acknowledged for improving our English. Four anonymous referees gave valuable comments on an earlier version of the manuscript.

**References**

1. Andersson M: Sexual selection. Princeton: Princeton University Press; 1994.
2. Kokko H, Johnstone RA: Why is mutual mate choice not the norm? Operational sex ratios, sex roles and the evolution of sexually dimorphic and monomorphic signalling. Philosophical Transactions of the Royal Society of London B 2002, 357(1419):319-330.
3. Parker GA: Sexual conflict over mating and fertilization: an overview. Philosophical Transactions of the Royal Society of London B 2006, 361:235-259.
4. Parker GA: Mate quality and mating decisions. In Mate Choice Edited by: Bateson P. Cambridge: Cambridge University Press; 1983:141-166.
5. Darwin C: On the descent of man and selection in relation to sex. London: Murray; 1871.
6. Arnqvist G, Rowe L, Krupa JJ, Sih A: Assortative mating by size: patterns in water striders a meta-analysis of mating. Evolutionary Ecology 1996, 10:265-284.
7. Ginsburg E, Livshits G, Yakovenko K, Kobyliansky E: Major gene control of human body height, weight and BMI in five ethnically different populations. Annals of Human Genetics 1998, 62:307-322.
8. Russell RJH, Wells PA, Rushton JP: Evidence for genetic similarity detection in human marriage. Ethology and Sociobiology 1985, 6:183-187.
9. Keenleyside MHA, Rangeley RW, Kuppers BU: Female mate choice and male parental defence behaviour in the cichlid fish Cichlasoma nigrofasciatum. Canadian Journal of Zoology 1985, 63(11):2489-2493.
10. Eberhard JR, Ewald PW: Food availability, intraspecific competition and territory size – an experimental study of annas hummingbirds (Calypte anna). Behavioral Ecology and Sociobiology 1994, 34:11-18.
11. Wright S: Systems of mating. III. Assortative mating based on somatic resemblance. Genetics 1921, 6:144-166.
12. Cooke F, Davies JC: Assortative mating, mate choice, and reproductive fitness in snow geese. In Mate Choice Edited by: Bateson P. Cambridge: Cambridge University Press; 1983:279-296.
13. Burley N: The meaning of assortative mating. Ethology and Sociobiology 1983, 4:191-203.
14. Ridley M: The explanation of organic diversity: the comparative method and adaptations for mating. Oxford: Clarendon Press; 1983.

15. Crespi BJ: Causes of assortative mating in arthropods. Animal Behaviour 1989, 38(6):980-1000.

16. Beeching SC, Wack CL, Ruffner GL: Female convict cichlids (Archocentrus nigrofuscatus) prefer to consort with same-sized males. Ethology Ecology and Evolution 2004, 16:209-216.

17. Kraak SBM, Bakker TCM: Mutual mate choice in sticklebacks: attractive males choose big females, which lay big eggs. Animal Behaviour 1998, 56:859-866.

18. Hudman SP, Gotelli NJ: Intra- and intersexual selection on male body size are complementary in the fathead minnow (Pimephales promelas). Behaviour 2007, 144:1065-1086.

19. Harari AR, Handler AM, Landolt PJ: Size-assortative mating, male choice and female choice in the cichlid fish Diaprepes abbreviatus. Animal Behaviour 1999, 58(1):1191-1200.

20. Noonan KC: Sex and Evolution. New Jersey: Princeton University Press; 1975.

21. McKay KR: Mate choice and size assortative pairing by the cichlid fishes of Lake Jiloa, Nicaragua. Journal of Fish Biology 1986, 29:135-150.

22. Thünken T, Bakker TCM, Baldauf SA, Kullmann H: Active inbreeding in a cichlid fish and its adaptive significance. Current Biology 2007, 17:225-229.

23. Trivers RL: Parental investment and sexual selection. In Sexual Selection and the Descent of Man Edited by: Campbell B. Chicago: Aldine; 1972:139-179.

24. Thünken T, Bakker TCM, Baldauf SA, Kullmann H: Direct familiarity does not alter mating preferences for sisters in male Pelvicachromis taeniatus (Cichlidae). Ethology 2007, 113:1107-1112.

25. Thünken T, Bakker TCM, Kullmann H: Extraordinarily long sperm in the socially monogamous cichlid fish Pelvicachromis taeniatus. Naturwissenschaften 2007, 94:489-491.

26. Baldauf SA, Kullmann H, Bakker TCM: Technical restrictions of computer-manipulated visual stimuli and display units for studying animal behaviour. Ethology 2008, 114(8):737-751.

27. Kunzler R, Bakker TCM: Computer animations as a tool in the study of mating preferences. Behaviour 1998, 135:1137-1159.

28. Barlow GW, Rogers W, Cappeto RV: Incompatibility and assortative mating in the midas cichlid. Behavioral Ecology and Sociobiology 1977, 2:249-59.

29. Beeching SC, Hopp AB: Male mate preference and size-assortative pairing in the convict cichlid. Journal of Fish Biology 1999, 55:1001-1008.

30. Taborsky B, Guyer L, Taborsky M: Size-assortative mating in the absence of mate choice. Animal Behaviour 2009, 77(2):439-448.

31. Awata S, Kohda M: Parental roles and the amount of care in a bi-parental substrate brooding cichlid: The effect of size differences within pairs. Behaviour 2004, 141:1135-1149.

32. Nuttall DB, Keenleyside MHA: Mate choice by the male convict cichlid (Cichlasoma nigrofasciatum; Piscis, Cichlidae). Ethology 1993, 95:247-258.

33. Werner NY, Lotem A: Experimental evidence for male sequential mate preference in a lekking species. Ethology 2006, 112(7):657-663.

34. Noonan KC: Female mate choice in the cichlid fish Cichlasoma nigrofasciatum. Animal Behaviour 1983, 31(4):1005-1010.

35. Markert JA, Arnegard ME: Size-dependent use of territorial space by a rock-dwelling cichlid fish. Oecologia 2007, 154(3):611-621.

36. Monaghan P, Metcalfe N, Houston DC: Male finches selectively pair with fecund females. Proceedings of the Royal Society of London B 1996, 263:1183-1186.

37. Clutton-Brock TH: Reproductive success: studies of individual variation in contrasting breeding systems. Chicago: University of Chicago Press; 1988.

38. Barlow GW: Contrasts in social behavior between central american cichlid fishes and coral-reef surgeon fishes. American Zooologist 1974, 14:9-34.

39. McKay KR: Behavioural aspects of cichlid reproductive strategies: patterns of territoriality and brood defence in Central American substratum spawners versus African mouth brooders. In Fish Reproduction: Strategies and Tactics Edited by: Wootom RJ, Potts GW. New York: Academic Press; 1984:245-273.

40. Leiser JK, Gagliardi JL, Itzkowitz M: Does size matter? Assessment and fighting in small and large size-matched pairs of adult male convict cichlids. Journal of Fish Biology 2004, 64:1339-1350.

41. Turner GF: The fighting tactics of male mouth-brooding cichlids: the effects of size and residency. Animal Behaviour 1994, 47:655-662.

42. Lehtonen TK, Lindsrom K: Females decide whether size matters: plastic mate preferences tuned to the intensity of male-male competition. Behavioral Ecology 2008, 20(1):195-199.

43. Castro L, Toro MA: Assortative mating through a mechanism of sexual selection. Journal of Theoretical Biology 2006, 243:386-392.

44. Hooper PL, Miller GF: Mutual mate choice can drive costly signaling even under perfect monogamy. Adaptive Behavior 2008, 16(1):53-70.

45. Rowland WJ: Studying visual cues in fish behavior: a review of ethological techniques. Environmental Biology of Fishes 1999, 56(3):285-305.

46. MacLaren RD, Rowland WJ: Differences in female preference for male body size in Pocelia latipinna using simultaneous versus sequential stimulus presentation designs. Behaviour 2006, 143:273-292.

47. Mazzi D, Kunzler R, Bakker TCM: Female preference for symmetry in computer-animated three-spined sticklebacks, Gasterosteus aculeatus. Behavioral Ecology and Sociobiology 2003, 54(2):156-161.

48. Kunzler R, Bakker TCM: Female preference for single and combined traits in computer animated stickleback males. Behavioral Ecology 2001, 12(6):681-685.

49. Rowland WJ, Bolyard KJ, Jenkins JJ, Fowler J: Video playback experiments on stickleback mate choice: female motivation and attentiveness to male color cues. Animal Behaviour 1995, 49(6):1559-1567.