The initial response of females towards congeneric males matches the propensity to hybridize
 in *Ophthalmotilapia*.

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26 Abstract:

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Cichlid radiations often harbour closely related species with overlapping niches and 28 29 distribution ranges. Such species sometimes hybridize in nature, which raises the question 30 how can they coexist. This also holds for the Tanganyika mouthbrooders Ophthalmotilapia 31 ventralis and O. nasuta. Earlier studies found indications of asymmetrical hybridisation with 32 females of O. ventralis accepting males of O. nasuta, but not the other way around. We 33 hypothesised that this was due to differences in the capacity for species recognition. Given the 34 higher propensity of O. ventralis females towards hybridization, we expect a reduced ability 35 for species recognition in O. ventralis females, compared to O. nasuta females. We staged 36 two experiments, one focusing on 22 female O. nasuta and one on 21 female O. ventralis. 37 These fish were placed in one half of a tank and briefly exposed to a conspecific or a 38 heterospecific male, a conspecific female, or nothing (control). Female response was 39 evaluated by scoring six tracking parameters and by noting the occurrence of ten discrete 40 behaviours before and during the first encounter. Females always responded to the presence 41 of another fish by approaching it. Remarkably, for both O. nasuta and O. ventralis, we did not 42 find a different response between encounters with conspecific males and females. However, 43 in agreement with our hypothesis, O. nasuta females behaved differently towards conspecific 44 or heterospecific males, whereas O. ventralis females did not. When presented with a 45 heterospecific male, O. nasuta females performed a lower number of 'ram' behaviours. Additionally, they never displayed the 'flee' behaviour, a component of the species' mating 46 47 repertoire that was seen in all but one of the presentations with a conspecific male. Our findings show that differences in species recognition at first encounter predict to a large 48 49 degree the outcome of the mating process, even in the absence of mating behaviour.

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## 51 Introduction

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Speciation is traditionally seen as a gradual build-up of reproductive isolation between 53 54 diverging populations (Mayr 1982; Coyne and Orr 2004). The classical view that this is a 55 slow process that occurs between allopatric populations has recently been challenged by 56 genomic findings (Margues et al. 2019) that showed how hybridization can drive rapid 57 speciation (Seehausen 2004; Jiggins et al. 2008). However, as unrestrained gene flow 58 inevitably homogenizes the genomes of diverging lineages (Roux et al. 2016), the question 59 remains what mechanisms keep incipient species separated. In scenarios of sympatric, closely 60 related species, the ability to correctly distinguish between conspecific and heterospecific 61 mates is probably crucial (Sullivan 2009).

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63 Mating is the end point of a complex decision-making process in which several potential 64 mates are evaluated (Luttbeg et al. 2001). An encounter with a potential mate can be seen as 65 the first step in this process. The outcome of this initial contact can have profound 66 implications on fitness, either through the refusal of suitable mates or through the acceptance 67 of suboptimal partners. Therefore, we expect intra- and interspecific differences in individual responses when confronted with a choice of partners. Additionally, the preference for a given 68 69 mate can depend on environmental, social and intrinsic parameters, explaining variation in 70 preference both between and within species, and between and within individuals (Pfennig 71 2008; Sommer-Trembo et al. 2017). Assortative mate selection was traditionally seen as a 72 sequential process in which individuals first assess whether 'the other' is a conspecific and 73 then assess its quality as a mate (Mayr 1982). However, empirical data and theory suggest 74 that assessment of species and quality are not independent steps (Sullivan 2009; Mendelson 75 and Shaw 2012) as the specific status of an individual could be judged using the same cues as

its quality. Additionally, adaptive hybridization has been observed in several taxa (overview in Mendelson and Shaw 2012), indicating that preferred mates are not necessarily always conspecific. Given the importance of the initial contact, we may expect that the early response to conspecific and heterospecific mates will predict the outcome of the mating process to a substantial degree. We test this hypothesis using a cichlid model.

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82 Cichlids of the large East African Lakes form endemic, species-rich radiations (Salzburger 83 2018). Several suggested "key adaptations" of cichlids, such as their pharyngeal jaws, help to 84 explain their evolution into numerous trophic niches (Kocher 2004). However, a large 85 proportion of these closely related species coexist without apparent eco-morphological 86 differences (Van Oppen et al. 1998). Since most cichlid assemblages are relatively young, 87 several taxa may be classified as incipient species and they retain the potential to hybridize. 88 The oldest East African lake, Lake Tanganyika, however, contains a mature cichlid radiation 89 (Salzburger 2018) in which most species are well-delineated (Ronco et al. 2020). However, 90 even between well-delineated biological species, boundaries can be permeable as molecular 91 studies identified several instances of inter-specific hybridization (Rüber et al. 2001; 92 Koblmüller et al. 2007; Nevado et al. 2011). Such examples allowed us to select a case to 93 study the importance of prezygotic, behavioural isolation after, or at the last stages, of the 94 speciation process.

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96 *Ophthalmotilapia* Pellegrin, 1904 species are maternal mouth brooders that occur on the 97 rocky and intermediate (rocky patches separated by sand) shores of Lake Tanganyika. The 98 genus contains four currently accepted valid species: *O. ventralis* (Boulenger 1898), *O. boops* 99 (Boulenger 1901), *O. heterodonta* (Poll and Matthes 1962) and *O. nasuta* (Poll and Matthes 100 1962) (Hanssens et al. 1999). They are sexually dimorphic maternal mouthbrooders with 101 territorial males that protect a spawning site, and females that aggregate in feeding swarms 102 when they are not breeding. Ophthalmotilapia males possess egg-shaped lappets at the distal ends of their elongated pelvic fins that are unique among Great Lake cichlids (Poll 1986). 103 104 These lappets function as egg dummies during the species' mating behaviour in a similar way 105 as the egg spots on the anal fins of the so-called 'modern' haplochromines (sensu Salzburger 106 et al. 2007; Theis et al. 2012). During the mating process, the female deposits the eggs and 107 almost immediately takes them into her mouth. By snapping at the egg dummies, which are 108 situated close to the genital opening of the male, the intake of sperm is facilitated, increasing 109 the fertilisation rate of the eggs within the female's mouth (Salzburger et al. 2007).

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111 The four species of *Ophthalmotilapia* have different but partially overlapping, distribution 112 ranges. *Ophthalmotilapia nasuta* is the sole species in the genus with a patchy but lake-wide 113 distribution. The sister species O. heterodonta and O. ventralis have non-overlapping ranges 114 with the former occurring in the northern half and the latter in the southern third of the Lake. 115 The fourth species, O. boops only occurs along a rather limited stretch of Lake Tanganyika's 116 south-eastern shoreline. There, it prefers sites where large stones are available (Konings 2019). 117 This is the only part of the lake where up to three species of Ophthalmotilapia occur in 118 sympatry (Hanssens et al. 1999).

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Although specimens of *Ophthalmotilapia* can be easily assigned to one of the valid species (with the possible exception of *O. heterodonta* and *O. ventralis*, see Hanssens et al. 1999), a phylogeographic study discovered gene flow among these species. Nevado et al. (2011) observed that specimens of *O. nasuta* often carried mitochondrial DNA of the other species, whereas the opposite was much less often the case. They suggested that this pattern either has a postzygotic, (*e.g.* by cyto-nuclear incompatibilities that affects mutual crossbreedings 126 differently) or a prezygotic (e.g. by an asymmetry in reproductive behaviour that results in a 127 different resistance towards hybridization) cause. The latter scenario implies that females of 128 all species would occasionally mate with O. nasuta males, while O. nasuta females would be 129 much less inclined to mate with heterospecific males. It also implies that the female hybrid 130 offspring would backcross into O. nasuta. This scenario agrees with the recent description of 131 a successful mating between a female *O. ventralis* and a male *O. nasuta* (Kéver et al. 2018). 132 Reproductive isolation in closely related species of East African cichlids is mostly maintained 133 through prezygotic isolation (Turner et al. 2001). Hence, models that describe speciation in 134 cichlids emphasize the importance of female mate choice in the initial stages of the speciation 135 process (Danley and Kocher 2001).

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137 Although the Lake Tanganyika cichlids assemblage contains species with profoundly 138 different mating strategies (Ronco et al. 2020), Ophthalmotilapia stands out by its extreme 139 sexual dimorphism and female-biased reproductive investment. Although the correlation 140 between reproductive investment, sexual selection and choosiness is well-established, it still 141 remains debated whether choosiness is an evolutionary outcome (sensu Trivers 1972), or 142 rather a determinant of differences between the sexes in parental investment (Thomas and 143 Szekely 2005). Using Lake Tanganyika cichlids, Gonzales-Voyer et al. (2008) showed 144 support for the latter hypothesis. Regardless of the evolutionary mechanism, females should 145 be considered the choosy sex in Ophthalmotilapia (sensu Wirtz, 1999). Therefore, if a 146 prezygotic mechanism explains the asymmetric pattern observed in nature (Nevado et al. 147 2011), it would be caused by differences between females of the different species in accepting 148 matings with heterospecific males. As increased capacity for species recognition leads to 149 increased preference in the choosier sex (Kozak & Boughman, 2009), we predict to see an 150 interspecific difference in female response to conspecific and heterospecific males. As males 151 of the different *Ophthalmotilapia* species have very similar courtship behaviours, in which the 152 few species-specific elements are insufficient to prevent hybridization (Kéver et al. 2018), 153 females would mainly rely on other cues like colour patterns, body size and pheromones. 154 Although the reproductive behaviour of *Ophthalmotilapia* species is well documented 155 (Haesler et al. 2011; Immler and Taborsky 2009; Kéver et al. 2018), little is known on how 156 *Ophthalmotilapia* species recognize conspecifics and select potential mates.

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158 This study described and compared the behavioural response of O. ventralis and O. nasuta 159 females towards males of both species in an aquarium setting. We studied species recognition, 160 which, in spite of its shortcomings (Mendelson and Shaw 2012), was defined as "a 161 measurable difference in behavioural response toward conspecifics as compared to 162 heterospecifics". We expected a difference in species recognition between females of O. 163 ventralis and O. nasuta. Specifically, we hypothesized that O. nasuta females would be able 164 to differentiate between conspecific and heterospecific males at the initial stages of an 165 encounter. For O. ventralis females, however, we expected that this capacity would be less 166 pronounced or absent.

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168 Materials and Methods

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170 Experimental setting

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We performed two independent experiments using females and males of *O. ventralis* and *O. nasuta*. The first focused on the behaviour of focal *O. nasuta* females (ON experiment), the second on that of focal *O. ventralis* females (OV experiment). All individuals were wild caught off the coast of Ulwile Island or nearby Kala, on the mainland (Tanzania). Specimens

176 were acquired as juveniles and hence had no prior mating experience. They were kept in 177 aquaria per species until they reached maturity. Hence, all females used had similar ages. 178 Males were older and most had prior mating experience (Kéver et al. 2018). We verified the 179 origin of a random selection of these fishes by sequencing the mitochondrial control region, 180 and by comparing these sequences with the data collected by Nevado et al. (2011). The 181 experiments were performed at the aquarium facilities of the University of Liège. Prior to the 182 onset of the experimental trials, the sex of the specimens was checked by visually inspecting 183 their genital papillae. Female specimens were kept jointly but isolated from males and 184 heterospecific specimens in a separate tank for at least two weeks. This tank was devoid of 185 hiding places, in order to prevent the development of territoriality. During that period, males 186 were kept in monospecific tanks in which they were visually isolated from each other using 187 opaque partitions. We kept all specimens in the same condition for at least two weeks with 188 photoperiod: 12:12 h L:D, water temperature: 26±1°C, carbonate hardness: >8 dKH. Fishes 189 were fed daily ad libitum with 'Tropical Spirulina forte' mini-granules.

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We used three identical experimental aquaria (88cm\*50cm\*40cm with water level ca. 40cm), which we divided into two equal parts by a perforated transparent partition (separation wall), through which fishes could not pass, and by an opaque wall (visual barrier) that could be removed (Fig. 1A). A flower pot was placed on each side of the separation to allow the fish to take refuge. We kept the fishes in these aquaria for at least twelve hours before they were used in the experimental trials. During the trial, the visual barrier was removed.





198 Figure 1. Experimental setup. A: aquarium setup: A focal female of either O. nasuta or O. ventralis (here O. 199 nasuta) (A) was placed in one half of the experimental tank whereas no fish, a conspecific female or a hetero- or 200 conspecific male was placed in the other half (here an O. ventralis male) (B). The tank was divided in two by a 201 transparent wall (D) and a visual barrier (C), a microphone (E) was placed on the side of the non-focal specimen 202 and an empty flowering pot was placed in both halves of the tank, allowing fishes to take refuge. Video and 203 audio recordings were made 15 minutes prior and 45 minutes after the visual barriers were removed. B: 204 Contrasts tested using permanova: A: focal females presented with another fish vs. with no fish, B: focal females 205 presented with a conspecific female vs. a conspecific male, C: focal females presented with a conspecific vs. a 206 heterospecific male, D: conspecific females and males presented to a focal female and E: conspecific and 207 heterospecific presented to a focal female. Black and grey fishes represent different species, the white fish 208 represents all possible non-focal specimens used.

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We recorded the behaviour of focal specimens (O. nasuta females or O. ventralis females) in 210 211 four different experimental conditions. They were either exposed to (i) no other specimen 212 (Co), (ii) a conspecific female (CF), (iii) a conspecific male (CM) or (iv) a heterospecific 213 male (HM) (Supplement 1). For each experiment (ON and OV) and for each experimental 214 condition (i to iv), we conducted a minimum of five replicates. We filmed (using a CANON 215 Legria HF R606) the entire aquarium (i.e. focal and non-focal fishes) during one hour: from 216 15 minutes before to 45 minutes after the visual barrier was removed. Experimenters were 217 only briefly present in the room to remove the visual barrier. As O. ventralis males are known 218 to produce weak-pulsed sounds during the inviting behaviour (Kéver et al. 2018), we used a

219 HTI Min-96 hydrophone (-164.4 dB re. 1 V µPa-1; bandwidth 2 Hz and 30 kHz, MS, USA), 220 connected to a Tascam DR-05 recording (TEAC, Wiesbaden, Germany) at a 44.1 kHz 221 sampling rate to record sounds during the whole experiment. The hydrophone was positioned 222 near the separation wall, at half the height of the water column, on the side of the non-focal 223 specimen. At the start of each trial, we switched off the aeration of the tank so that sounds 224 could be recorded. However, these recordings were not analyzed, as we detected no 225 communication sounds. After each trial, the focal female was euthanized. Both the focal and 226 the non-focal specimens were weighed. Focal specimens were measured, dissected and the 227 stage of gonad development was scored following Panfili et al. (2006).

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229 We performed a total of 28 ON and 21 OV experimental trials, with a maximum of three trials 230 per day (Supplement 1). However, after the dissections (see below), we observed that six 231 focal O. nasuta females from the first set of trials possessed male or ambiguous gonads 232 (Supplement 2). These specimens were referred to as floater males and the recordings for 233 these trials were not analyzed. As we suspected that these specimens had changed sex, we 234 photographed the genital papillae of the focal females that were to be used subsequently, two 235 weeks before the onset of the experimental trials. A comparison between papillae of the same 236 individuals after two weeks confirmed that a sex change did indeed took place in several 237 specimens. These specimens were not studied. After each trial, the aquarium was cleaned and 238 the water fully renewed.

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240 Collection of tracking and qualitative behavioural data

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Video files were converted into JPG images using Adapter v2.1.6 (available at
 <a href="https://www.macroplant.com">https://www.macroplant.com</a>), capturing one frame per second and saving it as an 8-bit, gray-

scale JPG file. Images taken within ten seconds before or after an experimenter was performing an action (i.e. removing the wall) were discarded from analyses. We chose to analyse the same number of frames for all trials within the ON and the OV experiment, respectively. For the ON experiment, this resulted in a minimum of 721 and 2186 frames collected before and after the removal of the separation wall, respectively. For the OV experiment, 871 and 2685 frames were available for analyses. Both focal and non-focal specimens were tracked using the ImageJ v1.49 (Schneider et al. 2012) software package.

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252 Given the presence of both light- and dark- coloured backgrounds in the aquarium setting, the 253 set of frames was studied twice. Specimens that were present before a light coloured 254 background were tracked by inverting black and white values whereas specimens present 255 before a dark-coloured background were tracked using non-inverted images. For 256 computational reasons, analyses were performed on subsets of the data containing a 257 maximum of 1,000 frames. For each set of images, a subset of 30 frames was used to create a 258 background using the plug-in 'Image stack merger plus'. Backgrounds were removed using 259 the image calculator and the resulting frames were transformed into black-and-white images 260 using the threshold function with 'MaxEntropy' as the methodology. Images were adjusted 261 using the 'erode' and 'dilate' functions to remove noise and to obtain a better representation 262 of the fishes. The resulting image series was then used for tracking using the plug-in 263 'MTrack2', in which tracks were summarized as the x- and y- coordinates of the centroids of 264 the tracked object. The quality of the automated tracking was checked by visually inspecting 265 each of the frames. When the software failed to track a specimen that was clearly present in 266 the final images, coordinates were added manually. Finally, tracks obtained from both 267 datasets, inverted and non-inverted, were combined. When a specimen was recognized by 268 both methods, e.g. when the fish was partially before a light- and partially before a dark-

269 coloured background, the average of the coordinates was used. When tracking data was 270 missing, the average value of the coordinates of the previous and the next positions were used. 271 This is justified, as missing data either corresponded to fish that remained stationary for many 272 frames, and could hence not be distinguished from the background, or to fish that hid behind 273 the flowering pots (Fig. 1). Frames collected before and after the removal of the visual barrier 274 were analysed separately. Coordinates were shifted using the lower- and anterior-most point 275 of the separation wall as the origin, and rotated by setting the anterior water level as a 276 reference for the horizontal plane. Finally, all coordinates were transformed from pixels to 277 centimetres using the dimensions of the aquaria. Tracks were visualized by plotting all 278 individual positions as well as the shift in average position of a specimen before and after the 279 removal of the barrier (Supplement 3).

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281 For each specimen, six tracking parameters were calculated from the coordinates (Table 1). 282 Each parameter was calculated three times: once using coordinates obtained for 721/871 283 seconds before (before) the removal of the visual barrier, once using coordinates obtained 284 during 721/871 seconds (after1) after the visual barrier was removed and finally using 285 coordinates obtained during 2186/2685 seconds (after2) after the visual barrier was removed, 286 hence including the after1 period (OV/ON experiment respectively). Additionally, ten specific 287 behaviours were defined based on Baerends and Baerends-Van Roon (1950). These were 288 encoded and recorded as point events in Boris v. 2.72 open source software (Friard and 289 Gamba 2016) (Table 1). This data was collected during the same three periods: before, after1 290 and after2. Behaviours displayed within ten seconds before or after an experimenter was 291 performing an action (removing the wall) were discarded from the analyses.

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## Table 1. Tracking parameters and point events recorded for the focal and non-focal individuals during

#### the experimental trials. For point events, interpretation of the behaviour was added.

Tracking parameters	Description	
Dist.wall	Distance to the visual barrier (% of length of	
	compartment).	
Dist.fish	Distance to the fish on the other side of the	
	transparent wall (cm).	
Sp	Average speed (cm/s).	
SpX	Horizontal speed (cm/s).	
SpY	Vertical speed (cm/s).	
Height	Mean height (% of height of water column).	
Point	Description	Interpretation
events	1	
Chase	The fish suddenly swims very fast towards the other fish and rams (or almost rams) the separation wall.	Contextual: agonistic and male courtship behaviour (Kéver et al. 2018)
Flee	The fish suddenly swims away from the other fish.	Contextual: agonistic and female courtship behaviour (Kéver et al. 2018)
Lateral	Lateral display: The fish positions itself perpendicular to the other fish, keeping its head slightly downwards, erects its fins and bends its body.	Signal movement (Baerends and Baerends-Van Roon 1950), agonistic behaviour (Kéver et al. 2017)
Frontal	Frontal display: The fish faces the other fish head up and erects its fins.	Signal movement (Baerends and Baerends-Van Roon 1950), agonisti behaviour (Kéver et al. 2018)
Bite	Biting the wall: The fish bites the separation wall (possibly trying to bite the other fish).	Signal movement (Baerends and Baerends-Van Roon 1950), agonisti behaviour (Kéver et al. 2018)
Ram	Ram into the wall: The fish tries to enter the other part of the aquarium and rams (not very fast) the separation wall.	
Sand	Sand picking: The fish takes sand in its mouth.	Courtship behaviour when linked to construction of bower (Kéver et al. 2018) signal movement when nipping off a substrate (Baerends and Baerends-Van Roon 1950).
Spasm	A quick, strong, and unilateral contraction of the trunk musculature that results in a displacement of the head and the caudal fin in the same direction.	Contextual: comfort behaviour or signa movement. Observed in courtship behaviour or inter-territorial fight depending on the genus (Baerends and Baerends-Van Roon 1950).
Tail	Tail-wagging: Exaggerated movements of the caudal fin (+ caudal part of the dorsal fin). At its zenith, the movement of the caudal fin is completely counterbalanced by backpedalling.	Signal movement (Baerends and Baerends-Van Roon 1950), courtship behaviour (Kéver et al. 2018).
Flicker	Pelvics flickering: The fish quickly and alternatively moves its right and left pelvic fins.	Comfort behaviour (Baerends and Baerends-Van Roon 1950)

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## 300 Statistical analyses

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302 Prior to testing differences in species recognition, we visually explored the combined datasets 303 of tracking parameters and point events. We did this by performing principal component 304 analysis (PCA) and canonical variate analyses (CVA) in Past 3.14 (Hammer et al. 2001). The 305 former allowed for an unbiased visualization of the variation in the data and was performed 306 on the correlation matrices. The later was conducted to maximize the differentiation between 307 the different groups. Separate analyses were performed for each period of time recorded 308 (before, after1, after2) and for each experiment (ON or OV). Point events that were not 309 recorded during one of these periods were disregarded and missing values (i.e. for Dist.fish in 310 the control condition Co) were treated using mean value imputation. Prior to the analyses, 311 each of the tracking parameters and point events was normalised. This was done for each 312 experiment (OV and ON), and for each of the time periods ('before', 'after1', 'after2' and 313 'shift') separately.

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315 We used permanova to compare the behaviour of both focal and non-focal specimens in five 316 different comparisons (Fig. 1B). We choose this approach since we wanted to measure 317 species recognition via differences in behaviour (sensu Mendelson and Shaw 2012) without 318 defining *a priori* in what variable specimens would differ. In order to reduce the number of 319 comparisons, we restricted ourselves to only biologically relevant contrasts. For focal 320 specimens, we compared the behaviour between (A) females that were presented with another 321 fish vs. with no fish, (B) focal females that were presented with a conspecific female vs. a 322 conspecific male, and (C) females that were presented with a conspecific vs. a heterospecific 323 male. Two additional comparisons were tested for the non-focal individuals. We tested (D) 324 whether conspecific females and males respond differently to a focal female and (E) whether

325 conspecific and heterospecific males respond differently to a focal female. Even though our 326 main goal was to test whether females of O. nasuta and O. ventralis differed in behavioural 327 response towards conspecific and heterospecific males (C), we tested the four other 328 comparisons as well, following the recommendations of Moran (2003) and Nakagawa (2004). 329 Tests were performed using non-parametric permanova, using the *pairwise.adonis* function, 330 of the R package *vegan* on the combined data of tracking parameters and point events. This 331 approach was chosen as the conditions for multivariate normality were violated. When 332 permanova revealed significant differences, we verified whether this could be due to 333 dispersion effects (Anderson 2006). For this, in view on the size of the dataset, non-334 parametric Mann-Whitney U tests were performed on the within group dispersions from the 335 mean, calculated using the betadisper function implemented in the R package vegan 336 (Oksanen et al. 2017). For each comparison revealed significant by permanova, Mann 337 Whitney U non-parametrical tests were performed on each of the variables separately in order 338 to detect which of these caused the difference between the treatments. We choose this non-339 parametric approach as the assumptions of normality were often not met. When significant, 340 the effect sizes of these variables were estimated using Hedge's g, which was calculated using 341 the estimation stats.com web application (Ho et al. 2019).

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In order to test whether the observed differences in behaviour depended on the visual presence of another specimen, and whether these differences were already visible at the first stages of the encounter, permanova tests were performed on data collected before the removal of the visual separation (before) as well as on data collected over a short (after1) and a long (after2) period of time after this separation was removed. Finally, an additional test was conducted which removed individual variation between the different treatments. For this, behavioural shifts were calculated for each tracking parameter and point event by subtracting

the values of the 'before' period before from those of the 'after1' period (shift). All tests were performed separately for the ON and the OV experiment. As behaviour can be influenced by gonad development and weight of the focal and non-focal specimens, Mann Whitney U tests were performed to check whether these differed between the treatments. Such tests were also performed on the amount of frames in which fishes could not be tracked. All statistical analyses were performed using Past 3.14 (Hammer et al. 2001) and R (R core team 2017).

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357 Results

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359 We separately analysed two experiments, one focusing on O. nasuta and one on O. ventralis 360 females (ON and OV experiment). In the ON experiment, two males of O. ventralis 361 performed advanced courtship behaviours. After the encounter, these males started to swim in 362 circles, in fast and erratic movements. This was accompanied by tail wagging, generally 363 displayed close to the partition wall. These males often bit the hydrophone and picked up and 364 moved around sand (49 and 45 times within 45 min vs. 0 for the other males). One of these 365 two males (O. ventralis male presented to ON38) also tried to chase the female (79 times) and 366 presented the egg dummies of its pelvic fins (5 times). This behaviour stopped immediately 367 when the experimenter removed the female fish. During the encounter, these males turned 368 dark grey, to almost black, which was swiftly reversed after the experimental trial. As we 369 designed our experiment to study behavioural response in the absence of courtship behaviour, 370 we removed these outliers from all analyses. All ten point events were observed in at least one 371 of the fishes in the ON experiment, whereas 'tail' (*i.e.* tail wagging) was never observed in 372 the OV experiment (Supplement 4).

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#### 375 Visualization of the behavioural data

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377 We visually explored the data using PCA and CVA to compare the behaviour of all 378 specimens used in each of the two experiments. In the PCAs conducted on the behavioural 379 data collected before the removal of the barrier, values of all females as well as of conspecific 380 males overlapped (Fig. 2A.B), suggesting a highly similar behaviour. However, heterospecific 381 males were (somewhat) separated from all other specimens by their higher values for PC1 382 (ON experiment) or PC2 (OV experiment). This difference was due to a more active 383 swimming behaviour (Sp, SpX, SpY) higher up in the water column (height) for O. ventralis 384 males (ON experiment) and a higher number of point events (ram, sand, bite) performed at 385 the floor of the aquarium (height) for the O. nasuta males (OV experiment), prior to their 386 presentation to a heterospecific female (Supplement 5.1). The CVAs also reflected the 387 behavioural differences of heterospecific males (Fig. 3A,B), as they had higher values for the 388 first CVs. The behaviours that contributed strongly to the separating PCs, also contributed to 389 the main CVs (Supplement 5.2).

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391 The PCAs performed on the data collected 15 minutes after the removal of the barrier 392 foremost showed how the focal females that were used as controls behaved differently than 393 the specimens that were presented with another specimen (Fig. 2C,D). For both experiments, 394 this can be explained by control females spending less time closer to the wall (Dist.wall), and 395 performing less agonistic behaviour (ram, lateral, flee). We did not observe any additional 396 separation in the PCA of the OV experiment (Fig. 2D). In the ON experiment, values for 397 (heterospecific) O. ventralis males stood out by their high values for PC1, whereas those of 398 (conspecific) O. nasuta males had mostly low values for this axis (Fig. 2C). The tracking 399 parameters Sp, SpX, SpY all had high, positive contributions to PC1 (Supplement 5),

400 reflecting that *O. ventralis* males kept swimming actively when the barrier was removed in401 the ON experiment.

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403 We carried out additional CVAs on the same datasets (Fig. 3C,D). For both experiments, the 404 control females stood out by their high values for CV1. This could again be explained by their 405 higher values for Dist. Wall. In the ON experiment, (heterospecific) O. ventralis males stood 406 out by their low values for CV1, which would be attributed to their more active swimming 407 behaviour (Sp, SpX, SpY). Conspecific O. nasuta males stood out by their low values for 408 CV2, which could be due to the higher occurrence of 'sand' and 'bite' behaviour. Values for 409 O. nasuta females that were presented with another fish had more intermediate values for 410 CV1 and CV2. However, values of O. nasuta females that were presented to a conspecific 411 male clustered between values of those males and of those of the other females. Similarly, 412 females that were presented to a heterospecific male had values that were intermediate 413 between those of these males and those of the other females (Fig. 3C). This suggests that the 414 behaviour of focal females shares characteristics with the behaviour of the non-focal fishes 415 presented to them. In the CVA of the OV experiment, (heterospecific) O. nasuta males stood 416 out by their low values for CV1 and high values for CV2. This was most influenced by the 417 higher occurrence of point behaviours (ram, spasm, sand, flicker). Values of O. ventralis 418 males overlapped with those of female specimens that were presented with another fish (Fig. 419 3D). These patterns remained present when performing similar analyses on the data collected 420 45 minutes after the removal of the visual barrier (Supplement 6). Plotting the shift in average 421 position before and after the removal of the barrier revealed how almost all specimens moved 422 towards the wall when presented with another specimen. Additionally, this showed that O. 423 nasuta specimens, on average, spent more time closer to the bottom whereas O. ventralis 424 specimens were more often found higher up in the water column (Supplement 7).



Figure 2. Principal component analyses performed on the behavioural data collected 15 min before (A, B) and 15 min after (C, D) the visual barrier was removed in the ON (*O. nasuta*, left) and the OV (*O. ventralis*, right) experiments. Symbols on the scatter plots for the ON and OV experiment as in E and F, respectively, with full circles denoting focal females presented with no fish (red), a conspecific female (blue) an *O. ventralis* male (purple), and an *O. nasuta* male (green), empty circles denote non-focal conspecific females and full squares *O. ventralis* (purple) and *O. nasuta* (green) males. Explained variances are added to the axes.



Figure 3. Canonical variate analyses on the behavioural data collected 15 min before and 15 min after the visual barrier was removed in the ON (*O. nasuta*, left) and the OV (*O. ventralis*, right) experiment.
Symbols on the scatter plots for the ON and OV experiments as in E and F, respectively, with full circles denoting focal females presented with no fish (red), a conspecific female (blue) an *O. ventralis* male (purple), and an *O. nasuta* male (green), empty circles denote non-focal conspecific females and full squares *O. ventralis*(purple) and *O. nasuta* (green) males. Explained variances are added to the axes.

440

433

#### 442 Behaviour of focal females

443

444 We tested whether the behavioural responses of females of O. nasuta and O. ventralis differed 445 in three different comparisons (Fig. 1B), i.e. when they were presented to (A) another fish vs. 446 no fish, (B) a conspecific female vs. a conspecific male, and (C) a conspecific vs. a heterospecific male. Within each of the comparisons in the ON experiment, we detected no 447 448 significant difference in the gonad development of the focal females, in their weights, in the 449 weights of the non-focal fishes, and in the percentage of missing frames. The same applies for 450 the OV experiment, although here, non-focal females and conspecific males differed in body 451 weight. Prior to the removal of the visual barrier, no significant difference in behaviour was 452 recorded for focal females from the different treatments in both experiments, and for all three 453 comparisons (Table 2).

454

455 For the three comparisons, and for both experiments (ON and OV), we performed permanova 456 on the behavioural data recorded during the 15 minutes after the removal of the visual barrier 457 (Table 2). For both the OV and the ON experiment, this revealed a significant difference in 458 the behaviour of focal females that were not presented to another fish (controls) and focal 459 females that were presented with another fish (comparison A). Although we could not exclude 460 that these differences stem from dispersion effects, these groups were also well-separated on 461 PCA (Fig. 2C,D). Mann-Whitney U tests revealed that controls differed from other focal 462 females by their higher values for the variable Dist.wall (ON: 46.3 +-18.2 vs. 20.1 +-8.8, 463 p=0.005, g=1.32; OV: 55.2 +-13.3 vs. 20.8 +-9.7, p=0.002, g=1.13) and their lower values for 'ram' (ON: 8.6+-12.1 vs. 74.5+-54.0, p=0.005, g=-2.14; OV: 8.0 +-11.9 vs. 48.8 +-38.4, 464 465 p=0.014, g=-3.12).

## 467 Table 2. PERMANOVA performed on the behavioural parameters of the ON (O. nasuta) and OV (O.

468 ventralis) experiment. Tests were performed on the data collected during 15 minutes before (B), 15 minutes 469 after (A1) and 45 minutes after (A2) the removal of the opaque wall as well as on behavioural shifts (S) 470 calculated by subtracting data recorded during 15 minutes before from that of 15 min after the removal of the 471 wall (A1-B). We tested five comparisons (Fig. 1B): by comparing the behaviour of focal females that were 472 presented with (A) another fish vs. with nothing, (B) with a conspecific female vs. a conspecific male, and (C) 473 with a conspecific vs. a heterospecific male. We further compared (D) the behaviour of non-focal conspecific 474 males vs. females, and (E) conspecific vs. heterospecific males. Behaviours that were neither recorded before or 475 after the removal of the wall, 'tail' and 'sand' for the ON, and 'flicker' and 'sand' for the OV experiment, were 476 excluded. For the first comparison, Dist.fish was excluded, as it could not be calculated. Values in bold are 477 significant at the 0.05 level. For these comparisons, ° and † denote that the assumption of equal dispersion was 478 violated at the 0.05 and 0.01 levels.

	С	А	В	С	D	Е
ON	В	F:0.31; p: 0.789	F:0.95; p: 0.423	F:0.91; p: 0.474	F:1.17; p: 0.339	F:1.15; p: 0.24
	A1	F:7.28; p: 0.012 †	F:0.42; p: 0.768	F:11.33; p: 0.016 °	F:0.45; p: 0.651	F:0.55; p: 0.73
	A2	F:4.71; p: 0.027 †	F:0.49; p: 0.553	F:8.13; p: 0.040	F:0.66; p: 0.545	F:0.85; p: 0.47
	S	F:8.40; p: 0.006 †	F:1.03; p: 0.416	F:8.75; p: 0.018 †	F:1.18; p: 0.332	F:0.25; p: 0.93
OV	В	F:0.23; p: 0.762	F:1.62; p: 0.203	F:0.85; p: 0.387	F:3.19; p: 0.061	F:5.13; p: 0.013
	A1	F:8.00; p: 0.005	F:0.91; p: 0.386	F:0.40; p: 0.546	F:0.28; p: 0.633	F:0.48; p: 0.524
	A2	F:4.28; p: 0.026	F:0.78; p: 0.438	F:0.48; p: 0.610	F:0.09; p: 0.837	F:0.49; p: 0.525
	S	F:7.61; p: 0.004	F:0.89; p: 0.418	F:0.87; p: 0.407	F:0.42; p: 0.679	F:0.73; p: 0.447

479

480 Unexpectedly, in both experiments, we did not observe a difference in behaviour between 481 focal females that were presented with a conspecific female or a conspecific male 482 (comparison B). However, when comparing the behaviour of focal females presented with 483 conspecific and heterospecific males (comparison C), a difference became evident between 484 the ON and the OV experiment. In support of our hypothesis, females of O. nasuta responded 485 differently towards conspecific and heterospecific males, whereas females of O. ventralis did 486 not (Table 2). Mann-Whitney U tests revealed that this was due to the lower number of 487 observed 'ram' behaviours (43.8+-17.9 vs. 138+-52.9; p=0.04, g=2.25) in O. nasuta females 488 that were presented to *O. nasuta* males compared to those presented to *O. ventralis* males. 489 Additionally, O. nasuta females never performed a 'flee' behaviour when being presented 490 with an O. ventralis male, whereas this was observed in all but one of the O. nasuta females

491 presented to a *O. nasuta* male (0 *vs.* 2.6+-2.3; p=0.04, g=-1.33). We obtained highly similar 492 results when analyzing the data collected 45 minutes after the visual barrier was removed, or 493 when analyzing the shift data (Table 2).

494

495 Behaviour of non-focal specimens

496

497 We tested whether the behavioural response of conspecific females and males differed when 498 presented to a focal female (D) and whether the behavioural response of conspecific and 499 heterospecific males differed when presented to a focal female (E). Unexpectedly, permanova 500 revealed a difference in the behaviour of O. nasuta and O. ventralis males in the OV 501 experiment, prior to the removal of the barrier. Mann-Whitney U tests revealed that this was 502 due to the higher average vertical swimming speed of O. ventralis males compared to O. 503 *nasuta* males (SpY 1.7+-0.5 vs. 0.7+-0.4, p=0.014, g=2.05). None of the other comparisons 504 was shown to be significantly different, neither before, nor after the removal of the barrier 505 (Table 2).

506

507 Discussion

508

510

We tested the initial response of *O. nasuta* and *O. ventralis* females towards conspecific and heterospecific males. In support of our hypothesis, *O. nasuta* females differentiated between conspecific and heterospecific males, whereas *O. ventralis* females did not. Visualisation of the data revealed that *O. nasuta* females mirrored the behaviour of the males to which they were presented. We also presented females of both species with a conspecific female or with

<sup>509</sup> Summary

516 nothing (control). Although females always responded to the presence of another fish, their 517 behaviour did not differ when presented with conspecific males and females. Comparisons of 518 non-focal specimens didn't reveal any differences in behaviour after presentation to a focal 519 female. However, before the removal of the wall, males of *O. ventralis* and *O. nasuta* behaved 520 differently in the OV experiment.

521

# 522 Responses of Ophthalmotilapia females

523

Nevado et al. (2011) discovered signatures of unidirectional hybridization in *Ophthalmotilapia*, which could either be explained by cyto-nuclear incompatibilities, or by asymmetric mate choice. The latter explanation implies that *O. nasuta* females would discriminate stronger against heterospecific males, than females of *O. ventralis*. This is supported by our experiments.

529

530 As we did not present focal females with heterospecific females, we cannot say that the 531 observed species recognition in *O. nasuta* females was due to a different response towards 532 males, or towards all specimens of the other species. However, as females in 533 *Ophthalmotilapia* are non-territorial and therefore often encounter heterospecific congeners, 534 we expect that the female response is specific to heterospecific males. Unexpectedly, females 535 of both species behaved similarly towards conspecific males and females. This suggests that 536 we observed the routine behaviour of a (isolated) female that encounters a conspecific 537 individual, rather than sexually motivated behaviour. In the wild, non-breeding females of 538 both species aggregate in large feeding groups (Konings 2019). Hence, being isolated for 12 539 hours, as was the case prior to the start of the experiment, represents an unnatural situation for 540 Ophthalmotilapia females. It would, therefore, not be unlikely if Ophthalmotilapia females

are behaviourally hardwired to reunite immediately with conspecifics, regardless of whetherthese are female or male.

543

544 Females of O. nasuta only performed the 'flee' behaviour towards conspecific males and 545 displayed the 'ram' behaviour less frequently. Although we found no evidence that this behaviour is sexually motivated, male chasing and female fleeing (i.e. 'flee') form the first 546 547 steps in the mating process of Ophthalmotilapia (Kéver et al. 2018). The ram behaviour, on 548 the other hand, was seen in all experimental trials in which a focal female was presented to 549 another fish. Indeed, the main difference in behaviour between focal females of both species 550 that were, or were not, presented with another fish was the amount of time spent close to the 551 wall, and in the display of the 'ram' behaviour.

552

We discovered that several of the females that we planned to use in the experiments changed into males. This was also observed in one of the non-focal females in the ON experiment, which was kept isolated from males after the experiment (Supplement 2). Although there are several reports of sex changes occurring in cichlids (Peters 1975; Naish and Ribbink 1990), evidence hereof remained, until now, limited.

558

### 559 Interpretation

560

Although we uncovered a significant difference in the behaviour of *O. nasuta* females that were presented with a conspecific and a heterospecific male, permanova did not reveal a significant interspecific difference in the behaviour of the males. This could indicate that *O. nasuta* females interpreted behaviour differently when displayed by *O. nasuta* or by *O. ventralis* males. Such species- or sex-dependent interpretation of behaviour is known for

566 several cichlid species, in which territorial males present themselves identically towards both 567 visiting females and intruding males (Baerends and Baerends-Van Roon 1950). Visual 568 exploration of the data, however (Fig. 3C, Supplement 6), revealed a potential difference in 569 male behaviour, which was mirrored by female response. As female O. ventralis did not 570 appear to differentiate between conspecific and heterospecific males, one could ask why 571 hybridization is not even more prevalent. However, we only examined the very first stage in a 572 potential mating process, so other differences that are present in the mating behaviour 573 between both species could be responsible for this. For example, O. ventralis males display a 574 specific late mating behaviour, called 'invite', which O. nasuta males never display (Kéver et 575 al. 2018). Additionally, hybrids might have a lower fitness. In order to reach mitochrondrial 576 introgression, female hybrids would also need to mate with O. nasuta males. This is not 577 unlikely given that, in cichlids, female mate choice is influenced, via imprinting, by the 578 maternal phenotype (Verzijden and ten Cate 2007; Verzijden et al. 2008).

579

#### 580 The role of males

581

582 Asymmetric propensities towards hybridization are known for a variety of animal taxa 583 including lungless salamanders (Verrell 1990), spadefoot toads (Pfennig 2007), swordtails 584 (Crapon de Caprona and Ryan 1990), pupfishes (Strecker and Kodric-Brown 1999), and 585 several cichlids (Egger et al. 2008; Nevado et al. 2011). Although most examples are related 586 to female mate choice, these patterns can also be caused by asymmetries in male choosiness 587 (Svensson et al. 2007). Although male mate choice is common in fishes (Wong and Jennions 588 2003; Werner and Lotem 2003), we choose to focus on the role of females (see introduction, 589 Seehausen et al. 2008; Sefc et al. 2017). However, the mode of fertilization in 590 Ophthalmotilapia could also have an influence on male choosiness. Haesler et al. (2011) 591 studied the reproductive behaviour of O. ventralis, but it can be assumed that the behaviour of 592 its congeners is highly similar. In O. ventralis, a ripe female will visit the territories of several 593 males, either to spawn, or just to collect additional ejaculates. Subsequently, sperm 594 competition will take place within her mouth, resulting in clutches with multiple sires 595 (Haesler et al. 2011). Given that this dilutes the effect of a 'wrong' choice, a female can 596 afford to be less choosy. Differences in both male and female courtship effort towards 597 genetically distant or similar mates have been documented in another mouth brooding cichlid: 598 Tropheus Boulenger, 1898 (Zoppoth et al. 2013). However, Tropheus species are sexually 599 monomorphic and both sexes are territorial. Additionally, *Tropheus* males invest significantly 600 more in raising the clutch, by providing the female access to their feeding territories. As 601 males of Ophthalmotilapia do not share their resources, we can expect these males to be less 602 choosy than those of *Tropheus*. Additionally, a substantial role of male mate choice is not 603 supported by our data, as we did not observe a difference in behaviour between non-focal 604 males of O. nasuta and O. ventralis when presented with females of the two species. It should 605 be noted, however, that two O. ventralis males that displayed mating behaviour towards O. 606 nasuta females were excluded from the analyses.

607

608 Whereas our experiments only revealed the capacity for species recognition in females of O. 609 *nasuta*, we cannot conclude that males cannot distinguish between females of the two species. 610 Whereas the males of O. nasuta and O. ventralis behaved differently when a visual barrier 611 was present, no significant difference was found after its removal. This could imply that 612 males of the two species behave in a similar way when presented with a conspecific or a 613 heterospecific female. However, an alternative explanation would be that males recognise 614 conspecific and heterospecifics, and use this knowledge to court females using a repertoire 615 appropriate to the species. Although this was found in sister species pairs of freshwater

sticklebacks (*Gasterosteus* spp. L. 1758) (Kozak et al. 2009), our experimental design did not
allow us to test this in *Ophthalmotilapia*.

618

619 We cannot exclude that morphological, physiological and behavioural features that 620 distinguish O. nasuta males from males of congeners could have caused the asymmetric 621 pattern of introgression. Foremost, O. nasuta males become larger and possess longer pelvic 622 fins. This feature could render them more attractive as O. ventralis females have a preference 623 towards males with strongly elongated pelvic fins (Haesler et al. 2011). As a change in the 624 feature associated with attractiveness can alter species recognition in the mating process 625 (Phelps et al. 2006), the extra-long pelvic fins of O. nasuta males could serve as a super-626 natural stimulus (sensu Tinbergen 1948). Additionally, even though Haesler et al. (2011) 627 found no correlation between female choice and male body length in O. ventralis, they did 628 observe that larger males outcompeted their rivals in sperm competition within the females' 629 mouth. Additionally, sperm of O. nasuta remains viable for a significantly longer amount of 630 time than that of O. ventralis (Morita et al. 2014). Lastly, O. nasuta males construct true 631 bowers (elaborate, crater-shaped sand mounts), whereas the nests of males of the other 632 species of *Ophthalmotilapia* only consist of a small area of cleaned rock, or of a small pit in 633 the sand (Konings 2019).

634

635 *The importance of visual cues* 

636

Although animals can use multiple kinds of cues to assess the quality of a potential mate, their
final assessment depends on the overall information available. This is exemplified by female
mate choice in the allopatric swordtail species *Xiphophorus nigrensis* Rosen 1960 and *X. pygmaeus* Hubbs and Gordon 1943. Here, mating preferences differed depending on whether

641 visual, olfactory or a combination of both cues were available (Crapon de Caprona and Ryan 642 1990). Different responses to visual and olfactory cues were also shown for females of sympatric Cyprinodon Lacipède, 1803 pupfish species from Lake Chichancanab (Mexico). 643 644 Here, different degrees of asymmetric discrimination of males were observed depending on 645 whether females had access to visual or olfactory information (Strecker and Kodric-Brown 646 1999). In species-rich systems and in species that form leks, such as *Ophthalmotilapia* spp., 647 females must be able to rapidly assess the quality of a potential mate (Barlow 2002). Males 648 therefore evolved morphological characteristics, build conspicuous bowers and/or perform 649 stereotyped displays to distinguish them from sympatric congeners. However, even though 650 multiple cues can be involved, mate choice decisions in radiations are often based on just a 651 small amount of (combinations of) these traits (Hohenlohe and Arnold 2010).

652

653 The separation wall used in our experiments contained holes that allowed for the exchange of 654 water between both compartments. Hence, besides visual clues, the fishes most likely also 655 received olfactory and acoustic information. Although visual cues were suggested to be the 656 primary factor in species-isolating, female mate choice in other cichlids (Jordan et al. 2003; 657 Kidd et al. 2006), we cannot rule out the importance of other types of information. Studies 658 have shown that olfactory (Blais et al. 2009; Plenderleith et al. 2005), acoustic (Nelissen 659 1978; Amorim et al. 2004; Kéver et al. 2018) and behavioural (Barlow 2002) information can 660 also influence the mating process. Although Seehausen and van Alphen (1998) showed a 661 certain hierarchy of information, where other cues are taken into account when visual 662 information is absent or masked, other experiments showed that female cichlids are more 663 likely to select the right male when both olfactory and visual cues are available (Plenderleith 664 et al. 2005; Blais et al. 2009). When visual information suffices for mate recognition, the 665 behaviour throughout the mating process, i.e. potentially leading toward spawning, doesn't need to diverge between closely-related species (Barlow 2002). This may explain why
spawning behaviour of *Ophthalmotilapia* is remarkably similar across the genus (Kéver et al.
2018) and why differently-coloured, sympatric mbuna cichlids from Lake Malawi have
identical courtship behaviours (McElroy and Kornfeld 1990).

670

## 671 Ecological reasons for asymmetric hybridization

672

673 Although they can be found in sympatry, O. ventralis is more associated with the rocky 674 shores of Lake Tanganyika, whereas O. nasuta has a wider ecological tolerance. At rocky 675 shores, *O. ventralis* can be one of the most abundant cichlid species (Sturmbauer et al. 2008). 676 Hence, for an O. ventralis female, a random encounter with another Ophthalmotilapia male is 677 much more likely to result in a conspecific than a heterospecific encounter. In contrast, for an 678 O. nasuta female venturing into the preferred O. ventralis habitat, a conspecific encounter 679 would be less often the case. Therefore, the ability to discriminate between conspecific and heterospecifics would be less important for females of O. ventralis than for those of O. nasuta. 680 681 A similar interpretation was given to explain asymmetries in female discrimination of 682 sympatric *Cyprinodon* species, where the species with the highest abundance had the lowest choosiness (Strecker and Kodric-Brown 1999). Although a species' propensity for 683 684 discrimination could be a consequence of its distribution range, the opposite could also hold. 685 Species that are better in recognizing conspecifics are more likely to maintain the integrity of their gene pool. Hence, they could be better in colonising habitats that have already been 686 687 occupied by related species. Finally, we showed that substantial behavioural differences can 688 be observed between closely-related species. This should be a warning to be cautious when 689 assuming similarities in the behaviour of certain (model) organisms and related taxa.

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692

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- 715 Baerends GP, Baerends-Van Roon JM. 1950. An introduction to the study of the ethology of
- the cichlid fishes, Leiden, The Netherlands, E. J. Brill.
- 717 Blais J, Plenderleith M, Rico C, Taylor MI, Seehausen O, van Oosterhout C, Turner GF. 2009.
- 718 Assortative mating among Lake Malawi cichlid fish populations is not simply predictable
- from male nuptial colour. BMC Evol Biol. 9(53):53.
- 720 Coyne JA, Orr HA. 2004. Speciation. Sunderland, MA: Sinauer Associates.
- 721 Crapon de Caprona MD, Ryan MJ. 1990. Conspecific mate recognition in swordtails,
- 722 *Xiphophorus nigrensis* and *X. pygmaeus* (Poeciliidae): olfactory and visual cues. Anim Behav.
- 723 39(2):290-296.
- Danley PD, Kocher TD. 2001. Speciation in rapidly diverging systems: lessons from Lake
  Malawi. Mol Ecol. 10:1075-1086.
- 726 Egger B, Obermüller B, Eigner E, Sturmbauer C, Sefc KM. 2008. Assortative mating
- 727 preferences between colour morphs of the endemic Lake Tanganyika cichlid genus *Tropheus*.
- 728 Hydrobiologia. 615:37-48.
- 729 Friard O, Gamba M. 2016. BORIS: A Free, Versatile Open-Source Event-Logging Software
- for Video / Audio Coding and Live Observations. Methods Ecol Evol. 7(11):1–6.
- Gonzalez-Voyer A, Fitzpatrick JL, Kolm N. 2008. Sexual selection determines parental care
  patterns in cichlid fishes. Evolution 62(8): 2015-2026.
- Haesler MP, Lindeyer CM, Otti O, Bonfils D, Heg D, Taborsky M. 2011. Female
  mouthbrooders in control of pre- and postmating sexual selection. Behav Ecol. 22(5):1033–
  1041.
- Hammer Ø, Harper DA, Ryan PD. 2001. PAST: Paleontological statistics software package
  for education and data analysis. Palaeontologia Electronica. 4:9pp.

- 738 Hanssens M, Snoeks J, Verheyen E. 1999. A Morphometric Revision of the Genus
- 739 Ophthalmotilapia (Teleostei, Cichlidae) from Lake Tanganyika (East Africa). Zool J Linn
- 740 Soc-Lon. 125(4):487–512.
- 741 Ho J, Tumkaya T, Aryal S, Choi H, Claridge-Chang A. 2019. Moving beyond P values: data
- analysis with estimation graphics. Nat Met. 16(7):565-566.
- 743 Hohenlohe PA, Arnold SJ. 2010. Dimensionality of mate choice, sexual isolation, and
- 744 speciation. PNAS. 107(38):16583-16588.
- 745 Immler S, Taborsky M. 2009. Sequential polyandry affords post-mating sexual selection in
- the mouths of cichlid females. Behav Ecol Sociobiol. 63(8):1219–1230.
- 747 Jiggins CD, Salazar C, Linares M, Mavarez J. 2008. Hybrid trait speciation and Heliconius
- 748 butterflies. Philos T R Soc B. 363(1506):3047-3054.
- Jordan R, Kellogg K, Juanes F, Stauffer J. 2003. Evaluation of female mate choice cues in a
  group of Lake Malawi mbuna (Cichlidae). Copeia. 2003(1):181-186.
- 751 Kéver L, Parmentier E, Derycke S, Verheyen E, Snoeks J, Van Steenberge M, Poncin P. 2018
- 752 Limited possibilities for prezygotic barriers in the reproductive behaviour of sympatric
- 753 *Ophthalmotilapia* species (Teleostei, Cichlidae). Zoology. 126:71-81.
- Kidd MR, Danley PD, Kocher TD. 2006. A direct assay of female choice in cichlids: all the
- 755 eggs in one basket. J Fish Biol. 68(2):373–384.
- Kocher TD. 2004. Adaptive evolution and explosive speciation: the cichlid fish model. NatRev Genet. 5(4):288-298.
- 758 Koblmüller S, Duftner N, Sefc KM, Aibara M, Stipacek M, Blanc M, Egger B, Sturmbauer C.
- 759 2007. Reticulate phylogeny of gastropod-shell-breeding cichlids from Lake Tanganyika the
- result of repeated introgressive hybridisation. BMC Evol Biol. 7:7.
- 761 Konings A. 2019. Tanganyika cichlids in their natural habitat. 4<sup>th</sup> edition. El Paso, TX:
- 762 Cichlid press.

- 763 Kozak GM, Boughman JW. 2009. Learned conspecific mate preference in a species pair of
- 764 sticklebacks. Behav Ecol. 20: 1282–1288.
- 765 Kozak GM, Reisland M, Boughmann JW. 2009. Sex differences in mate recognition and
- conspecific preference in species with mutual mate choice. Evolution. 63: 353-365.
- 767 Luttbeg B, Towner MC, Wandesforde-Smith A, Mangel M, Foster SA. 2001. State-dependent
- 768 mate-assessment and mate-selection behavior in female threespine sticklebacks (Gasterosteus
- 769 *aculeatus*, Gasterosteiformes: Gasterosteidae). Ethology. 107(6):545-558.
- 770 Mayr E. 1982. The growth of biological thought: Diversity, evolution and inheritance.
- 771 Cambridge, MA: Harvard University Belknap Press.
- 772 Marques DA, Meier JI, Seehausen O. 2019. A combinatorial view on speciation and adaptive
- radiation. Trends Ecol Evol. 34(6):531-544.
- Mendelson TC, Shaw KL. 2012. The (mis)concept of species recognition. Trends Ecol Evol.
  27(8):421-427.
- McElroy DM, Kornfeld I. 1990. Sexual selection, reproductive behaviour, and speciation in
  the mbuna species fock of Lake Nyasa (Pisces: Cichlidae). Environ Biol Fish. 28:273-284.
- 778 Moran MD. 2003. Arguments for rejecting the sequentional Bonferroni in ecological studies.
- 779 Oicos. 100(2):403-405.
- 780 Morita M, Awata S, Yorifuji M, Ota K, Kohda M, Ochi H. 2014. Bower-building behaviour is
- associated with increased sperm longevity in Tanganyikan cichlids. J Evolution Biol.
  27(12):2629–2643.
- Naish KA, Ribbink AJ. 1990. A preliminary investigation of sex change in *Pseudotropheus lombardoi* (Pisces: Cichlidae). Environ Biol Fish. 28:285–294.
- 785 Nakagawa S. 2004. A farewell to Bonferroni: the problem if low statistical power and
- publication bias. Behav. Ecol. 15:1044-1045.

- 787 Nelissen MHJ. 1978. Sound production by some Tanganyikan cichlid fishes and a hypothesis
- for the evolution of their communication mechanisms. Behaviour. 64(1-2):137–147.
- 789 Nevado B, Fazalova V, Backeljau T, Hanssens M, Verheyen E. 2011. Repeated unidirectional
- 790 introgression of nuclear and mitochondrial DNA between four congeneric Tanganyikan
- 791 cichlids. Mol Biol Evol. 28(8):2253-2267.
- 792 Oksanen JF, Blanchet G, Friendly M, Kindt R, Legendre P, McGlinn D, Minchin PR, O'Hara
- RB, Simpson GL, Solymos PM, Stevens HH, Szoecs E, Wagner H. 2017. vegan: Community
- Ecology Package. R package version 2.4-5.
- Panfili J, Thior D, Ecoutin J-M, Ndiaye P, Albaret J-J. 2006. Influence of salinity on the size
- at maturity for fish species reproducing in contrasting West African estuaries. J Fish Biol.69(1):95-113.
- Peters HM. 1975. Hermaphroditism in cichlid fishes. pp. 228–235.In: R. Reinboth (ed.)
  Intersexuality in the Animal Kingdom. Springer-Verlag, Berlin.
- Pfennig KS. 2007. Facultative mate choice drives adaptive hybridization. Science, 318(5852):965.
- Phelps SM, Rand AS, Ryan MJ. 2006. A cognitive framework for mate choice and species
  recognition. Am Nat. 167(1):28–42.
- Plenderleith M, van Oosterhout C, Robinson RL, Turner GF. 2005. Female preference for
  conspecific males based on olfactory cues in a Lake Malawi cichlid fish. Biol Lett. 1(4):411414.
- Poll M. 1986. Classification des cichlidae du Lac Tanganyika: Tribus, genres et espèces. Bull
  Acad R Belg. 8:1-163.
- 809 R core team. 2017. R: A Language and environment for statistical computing. R foundation
- 810 for statistical computing, Vienna, Austria.

- 811 Ronco F., Büscher H. H., Indermauer A., Salzburger W. 2020. The taxonomic diversity of the
- 812 cichlid fish fauna of ancient Lake Tanganyika, East Africa. J. Great Lakes Res. 46: 1067-1078.
- 813 Schneider CA, Rasband WS, Eliceiri KW. 2012. NIH Image to ImageJ: 25 years of image
- 814 analysis. Nat Met. 9(7):671-675.
- 815 Roux C, Fraïsse C, Romiguier J, Anciaux Y, Galtier N, Bierne N. 2016. Shedding light on the
- 816 grey zone of speciation along a continuum of genomic divergence. PLOS Biol. 817 14(12):e2000234.
- 818 Rüber L, Meyer A, Sturmbauer C, Verheyen E. 2001. Population structure in two sympatric 819 species of the Lake Tanganyika cichlid tribe Eretmodini: Evidence for introgression. Mol 820 Ecol. 10(5):1207-1225.
- 821 Salzburger W. 2018. Understanding explosive diversification through cichlid fish genomics. 822 Nat Rev Genet. 19:705-717.
- 823 Salzburger W, Braasch I, Meyer A. 2007. Adaptive sequence divergence in a colour gene 824 involved in the formation of the characteristic egg-dummies of male Haplochromis cichlid 825 fishes. BMC Biol. 5(51):13pp.
- 826 Seehausen O. 2004. Hybridisation and adaptive radiation. Trends Ecol Evol. 19(4):198-207.
- Seehausen O, van Alphen JJM. 1998. The effect of male coloration on female mate choice in 828 closely related Lake Victoria cichlids (Haplochromis nyererei complex). Behav Ecol 829 Sociobiol. 42:1–8.
- 830 Seehausen O, Terai Y, Magalhaes IS, Carleton KL, Mrosso HDJ, Miyagi R, van der Sluijs I,
- 831 Schneider MV, Maan ME, Tachida H, Imai H, Okada N. 2008. Speciation through sensory
- 832 drive in cichlid fish. Nature. 455:620-626.

- 833 Sefc KM, Mattersdorfer K, Ziegelbecker A, Neuhüttler N, Steiner O, Goessler W, Koblmüller 834 S. 2017. Shifting barriers and phenotypic diversification by hybridisation. Ecol Lett.
- 835 20(5):651-662.

- 836 Sommer-Trembo C, Plath M, Gismann J, Helfrich C, Bierbach D. 2017. Context-dependent
- female mate choice maintains variation in male sexual activity. R Soc Open Sci. 4: 170303.
- 838 Strecker U, Kodric-Brown A. 1999. Mate recognition systems in a species flock of Mexican
- 839 pupfish. J Evolution Biol. 12(5):927-935.
- 840 Sturmbauer C, Fuchs C, Harb G, Damm E, Duftner N, Maderbacher M, Koch M, Koblmüller
- 841 S. 2008. Abundance, distribution, and territory areas of rock-dwelling Lake Tanganyika
- cichlid fish species. Hydrobiologia. 615(1):57-68.
- Sullivan BK. 2009. Mate recognition, species boundaries and the fallacy of "species
  recognition". Open Zool. 2:86-90.
- 845 Svensson EI, Karlsson K, Friberg M, Eroukhmanoff F. 2007. Gender differences in species
- recognition and the evolution of asymmetric sexual isolation. Curr Biol. 17(22):1943–1947.
- 847 Tinbergen N. 1948. Social releasers and the experimental method required for their study.848 Wilson Bull. 60(1):6-51.
- 849 Turner GF, Seehausen O, Knight ME, Allender CJ, Robinson RL. 2001. How many species of
- cichlid fishes are there in African lakes. Mol Ecol. 10(3):793-806.
- Theis A, Salzburger W, Egger B, 2012. The function of anal fin egg-spots in the cichlid fish *Astatotilapia burtoni*. PLoS ONE 7(1):e29878.
- Thomas GH, Székely T. 2005. Evolutionary pathways in shorebird breeding systems: sexual conflict, parental care, and chick development. Evolution 59(10): 2222–2230.
- Trivers RL. 1972. Parental investment and sexual selection. in Sexual selection and the descent of man, EBG. Campbell, ed. (136-179). Aldine, Chicago.
- 857 Van Oppen MJH, Turner GF, Rico C, Robinson RL, Deutsch JC, Genner MJ, Hewitt GM.
- 858 1998. Assortative mating among rock-dwelling cichlid fishes supports high estimates of
- species richness from Lake Malawi. Mol Ecol. 7(8):991-1001.

- Verrell PA. 1990. Frequency of interspecific mating in salamanders of the plethodontid genus *Desmognathus*: different experimental designs may yield different results. J Zool. 221(3):441451.
- 863 Verzijden MN, Korthof REM, ten Cate C. 2008. Females learn from mothers and males learn
- 864 from others. The effect of mother and siblings on the development of female mate preferences
- and male aggression biases in Lake Victoria cichlids, genus *Mbipia*. Behav Ecol Sociobiol.
- 866 62: 1359–1368.
- 867 Verzijden MN, ten Cate, C. 2007. Early learning influences species assortative mating
  868 preferences in Lake Victoria cichlid fish. Biol lett 3(2): 134-136.
- 869 Werner NY, Lotem A. 2003. Choosy males in a haplochromine cichlid: first experimental
- evidence for male mate choice in a lekking species. Anim Behav. 66(2):293-298.
- 871 Wirtz, P. 1999. Mother species-father species: unidirectional hybridization in animals with
- 872 female choice. Animal Behaviour. 58:1–12.
- Wong BMB, Jennions MD. 2003. Costs influence male mate choice in a freshwater fish. P
  Roy Soc B-Biol Sci. 270(Suppl 1):S36-S38.
- Zoppoth P, Koblmüller S, Sefc KM. 2013. Male courtship preferences demonstrate
  discrimination against allopatric colour morphs in a cichlid fish. J Evolution Biol. 26(3): 577586.
- 878

878 Additional material

879

880 Supplement 1. Summary of the experiments.

881

Supplement 2. Sex change observed in *O. nasuta* females. Top row: gonads of some of the specimens that were female in external phenotype but had male or ambiguous gonads (ON33, 35, 26, 43). Vertical left: female gonads in developmental stage 3 (ON34), 4 (ON24) and 5 (ON37) respectively (Panfili et al. 2006). Bottom: two non-focal *O. nasuta* females used in the ON experiment, one of which underwent transition after the experiments, and horizontal right, the ventral area of the same specimens, with A: anal pore and UG: urogentital pore.

888

889 Supplement 3. Visualization of tracking data. The position of each fish is plotted for each 890 second in which specimens were tracked with the positions recorded before and after the 891 removal of the opaque wall (grey) coloured differently. Ellipses denote the area in which 90% 892 of tracks are situated, large dots denote the average positions before and after the removal of 893 the barrier and arrows shows the change in mean position. The separation wall is visualized as 894 a meshed partition. The average speed before (v0) and after (v1) the removal of the barrier is 895 plotted for each tracked specimen with data for focal specimens given in bold. Abbreviations 896 (ON: O. nasuta, OV, O. ventralis, F: female, M; male)

897

898 Supplement 4. Summary of the data.

899

Supplement 5. Loadings and variance of the main axes of the PCAs and CVAs conducted inthe study.

Supplement 6. Principal component analyses and Canonical variate analyses performed on the
behaviours recorded 45 min after the visual barrier was removed of the ON (*O. nasuta*, left)
and the OV (*O. ventralis*, right) experiment. Symbols on the scatter plots for the ON and OV
experiment as in E and F, respectively, with full circles denoting focal females presented with
no fish (red), a conspecific female (blue) an *O. ventralis* male (purple), and an *O. nasuta* male
(green), empty circles denote non-focal conspecific females and full squares *O. ventralis*(purple) and *O. nasuta* (green) males. Explained variances are added to the axes.

910

911 Supplement 7. Shift in average position of the specimens analysed 15min before and 15min 912 after the removal of the separation wall for the ON (above) and OV (below) experiment. 913 Focal specimens are all visualised on the left, and non-focal specimens on the right. 914 Dimensions in cm, with the vertical bar representing the separation wall. Dashed arrows 915 represent individual fishes, bold arrows the average per treatment. Colours, for focal females 916 (ON and OV) presented with no fish (red), a conspecific female (turquoise) an O. ventralis 917 male (pink), and an O. nasuta male (light green), and non focal specimens (right): conspecific 918 females (blue), O. ventralis males (purple) and O. nasuta males (green).