

RESEARCH ARTICLE

Sexual differences in exploration behavior in *Xenopus tropicalis*?Mathieu Videlier¹, Raphaël Cornette², Camille Bonneaud³ and Anthony Herrel^{1,4,*}

ABSTRACT

The two sexes of a species often differ in many ways. How sexes differ depends on the selective context, with females often investing more in reproductive output and males in territory defense and resource acquisition. This also implies that behavioral strategies may differ between the two sexes, allowing them to optimize their fitness in a given ecological context. Here, we investigated whether males and females differ in their exploration behavior in an aquatic frog (*Xenopus tropicalis*). Moreover, we explored whether females show different behavioral strategies in the exploration of a novel environment as has been demonstrated previously for males of the same species. Our results show significant sex differences, with males exploring their environment more than females. Yet, similar to males, female exploratory behavior varied significantly among individuals and broadly fell into three categories: shy, intermediate and bold. Moreover, like in males, behavioral strategies are decoupled from morphology and performance. Our results suggest that females are more sedentary than males, with males engaging in greater risk taking by exploring novel environments more. Male and female behaviors could, however, be classified into similar groups, with some individuals being bolder than others and displaying more exploration behavior. The decoupling of morphology and performance from behavior appears to be a general feature in the species and may allow selection to act on both types of traits independently.

KEY WORDS: Performance, Morphology, Frog, Locomotion, Sexual dimorphism

INTRODUCTION

Exploration behavior was defined by Scott as a sensory inspection of the environment (Scott, 1956). Exploration behavior is crucial as it allows animals to find food, reproductive partners and new territories. Moreover, longer distance dispersal is also driven to a large extent by individual exploration behavior. Different dispersal strategies have been observed in animals, with some being regular such as migration (Pulido et al., 2001) and others more episodic. The latter dispersal strategy is typically associated with escape from habitats that are being modified (Berg et al., 2010; Wittern and Berggren, 2007), and may be important to population survival in the context of resource limitation or catastrophic events. Whether an animal has a tendency to disperse is largely determined by its personality. Two contrasting behavioral syndromes are typically

identified in this context: bold and shy (Sih et al., 2004). These two syndromes are typically fixed throughout the life of an individual and have been demonstrated in a wide variety of animals including invertebrates such as crabs (Decker and Griffen, 2012) and crickets (Niemelä et al., 2012), but also in many vertebrates including birds (Carere et al., 2005), fish (Dziewieczynski and Crovo, 2011; Brown et al., 2007), turtles (Mafli et al., 2011) and primates (Uher et al., 2008). Bold individuals show a tendency to explore their environment more, thus increasing the risk of predation but also increasing the probability of finding mates, food or new territories. Shy individuals typically explore less, thus decreasing risk taking but also reducing opportunities. On average, these two strategies show equal fitness in neutral conditions, resulting in the maintenance of both (Smith and Blumstein, 2008).

Interestingly, not all individuals behave similarly and several factors are known to affect an animal's behavior including age and sex (Carere et al., 2005; Dingemanse and Réale, 2005). Differences between the sexes may arise as a result of sexual selection (Hedrick and Temeles, 1989) and result in differences in body size (Woolbright, 1983) and other morphological traits (Desjardins and Fernald, 2009). As the energetic cost of gamete production strongly differs between the sexes, this often results in differences in body size and life-history traits (Monroe and Alonzo, 2014). Moreover, exploration behavior may also be different in males and females because of the association between male exploration and territorial aggressiveness, as has been shown in birds and fish (Carere et al., 2005; Brown et al., 2007; Dziewieczynski and Crovo, 2011). In some birds, females also prefer bolder males and consequently sexual selection may also impact the evolution of exploration behavior (Dingemanse and Réale, 2005). Moreover, hormonal differences between the sexes are likely to impact their exploration behavior (Brandner, 2007). For example, in humans, men explore over greater distances than women do. This is caused by differences in orientation ability as well as constraints due to parental care that are not the same in the two sexes (Brandner, 2007).

Amphibians, and especially frogs, are of interest in this context for two reasons: (1) the majority of species are dimorphic, with females typically being larger than males (Monnet and Cherry, 2002) and (2) amphibians are characterized by a low overall mobility, making them especially vulnerable to local habitat destruction and habitat fragmentation (Hillers et al., 2008). Thus, dispersal is likely under strong selection in fragmented habitats. Given known sexual dimorphism in morphology in many amphibian species (Monnet and Cherry, 2002; Shine, 1979), the two sexes may not be impacted to the same degree, however. Surprisingly, amphibian exploratory behavior remains poorly investigated. Here, we focused on *Xenopus (Silurana) tropicalis* (Gray 1864) as a model because this species is sexually dimorphic in body size, limb dimensions and locomotor performance (Herrel et al., 2012). The natural habitat of this species (the coastal rain forest belt of West Africa) is increasingly being fragmented as a result of direct human impacts and climate change (Hillers et al., 2008). This likely imposes selective pressures on locomotion and

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may affect the survival of animals with different exploration behaviors. An investigation of exploration behavior in male *X. tropicalis* showed a dissociation between performance and morphology on the one hand and exploration behavior on the other hand, suggesting that selection could act on behavior without affecting locomotor performance (Videliere et al., 2014). This circumvents the trade-off between endurance, important in the exploration of novel environments, and burst performance capacity, which is important in prey capture and predator escape.

We concentrated on the exploration behavior in females given the known differences in morphology and performance between the sexes. We specifically tested (1) whether females show behavioral syndromes related to the exploration of a novel environment, (2) whether behavioral strategies are similar in the two sexes and (3) whether and how the sexes differ in their overall exploration behavior. To complete, we tested whether exploration behavior is indeed decoupled from morphology and performance in females, as well as at the population level.

RESULTS

Female exploration behavior

Female *X. tropicalis* began to move 426 s after the onset of the experiment on average (range: 5.49–1639 s) and stopped moving 2348 s after the start of the trial on average (range: 145–3601 s). They covered a mean distance of 1.5 m in 1 h (range: 23–6393 cm). The mean duration of a roundtrip was 206 s (range: 29–1357 s) when including pauses and 13 s (range: 2.81–44 s) when excluding pauses. The pause is an important part of the exploration behavior as it allows an individual to screen its environment. Animals stopped moving 1.97 times on average (range: 0.5–12 times). Females moved with a mean overall speed of 5.36 cm s⁻¹ (range: 0.06–22.10 cm s⁻¹) and a mean movement speed of 15.24 cm s⁻¹ (range: 1.39–42.75 cm s⁻¹; see Table 1).

Three different groups were identified (35 individuals in the first, 21 in the second and four in the last; Fig. 1) in the Gaussian mixture model analysis using the 13 repeatable variables without imposing the number of groups. The three clusters showed differences in exploration behavior (Wilk's $\lambda=0.02$, $F_{26,90}=19.74$, $P<0.001$). Subsequent ANOVA showed significant differences for all variables ($P<0.05$), except the average number of pauses, which did not differ between the three groups ($F_{2,57}=1.25$, $P=0.30$). Group 1 showed numerous movements compared with the two others. Group 2 showed the opposite behavioral pattern, characterized by a reduced exploration of the tank. Group 3 showed an intermediate level of exploration as indicated by the number of movements, the distance traveled and the duration of exploration (Table 2). When testing whether the groups identified differed in morphology and performance, no significant differences were detected in body size, head, pelvic girdle or limb dimensions (see Table 3). Finally, no significant differences in locomotion performance were detected between the three groups (Wilk's $\lambda=0.82$, $F_{10,78}=0.79$, $P=0.64$).

Sexual dimorphism in exploration behavior

When comparing the two sexes, males were found to move more quickly (average speed: 12.25 cm s⁻¹ for males versus 5.36 cm s⁻¹ for females) and showed fewer pauses (average number of pauses in males: 1.31 versus 1.97 in females), resulting in a shorter average roundtrip duration (average duration: 99.07 s in males versus 205.86 s in females). Moreover, the time spent hidden showed significant differences, with males spending more time hiding than females (average time spent hidden: 299.05 s in males versus 150.78 s in females). A MANOVA performed on the behavioral

Table 1. Exploration behavior in female *Xenopus tropicalis*

	Average	Maximal	Minimal
Number of roundtrips	10.04	40.00	0.03
Number of complete roundtrips	7.58	37.33	0.00
Number of movements	21.78	79.67	0.67
Total distance moved (cm)	1496.35	6393.16	23.08
Average speed (cm s ⁻¹)	5.36	22.10	0.06
Maximal speed (cm s ⁻¹)	12.27	49.72	0.06
Minimal speed (cm s ⁻¹)	1.67	18.38	0.03
Average speed without pauses (cm s ⁻¹)	15.24	42.75	1.39
Maximal speed without pauses (cm s ⁻¹)	23.00	112.44	1.39
Minimal speed without pauses (cm s ⁻¹)	9.18	37.24	1.39
Average duration of a roundtrip (s)	205.86	1356.89	28.76
Maximal duration of a roundtrip (s)	570.02	2023.38	64.27
Minimal duration of a roundtrip (s)	93.37	1093.96	0.71
Average duration of a roundtrip without pauses (s)	12.64	44.12	2.81
Maximal duration of a roundtrip without pauses (s)	28.71	121.71	2.96
Minimal duration of a roundtrip without pauses (s)	5.84	28.47	0.71
Latency of the first movement (s)	426.32	1639.22	5.49
Latency of the second movement (s)	560.67	1752.56	0.00
Latency of the last movement (s)	2347.91	3601.84	144.53
Duration of all movements with pauses (s)	1191.86	3032.24	100.24
Duration of exploration without pauses (s)	124.14	411.18	4.87
Total time spent hidden (s)	878.76	2539.66	0.00
Average time spent hidden (s)	150.78	837.92	0.00
Maximal time spent hidden (s)	393.26	1224.47	0.00
Minimal time spent hidden (s)	57.05	806.62	0.00
Average number of pauses	1.97	10.22	0.50
Maximal number of pauses	3.73	12.00	0.67
Minimal number of pauses	0.79	9.00	0.00
Number of movements away from the wall	2.54	8.67	0.00

Highlighted variables were not repeatable across trials.

variables showed significant differences between the sexes (Wilk's $\lambda=0.31$, $F_{15,80}=11.79$, $P<0.001$). Subsequent ANOVA showed that speed (average, minimal and maximal), the duration of a roundtrip, the number of pauses and the time spent hidden showed significant differences between the sexes (Fig. 2, Table 4).

The combined data set for 37 males and 61 females included 15 behavioral variables that were found to be repeatable. Our

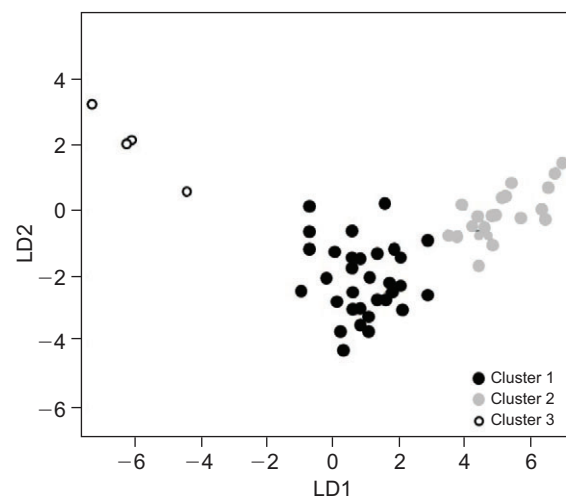


Fig. 1. Discriminate function based on the repeatable exploration behavior variables in females. The three (bold, shy and intermediary) behavioral syndromes are indicated (clusters 1–3, respectively).

Table 2. Results of MANOVA testing for differences between behavioral groups in female *X. tropicalis*

	<i>F</i>	<i>P</i>	Group difference
Total distance moved	98.08	<0.001	3<1<2
Number of roundtrips	68.75	<0.001	3<1<2
Number of complete roundtrips	92.94	<0.001	3<1<2
Number of movements	87.28	<0.001	3<1<2
Maximal speed	72.73	<0.001	3<1<2
Number of movements away from the wall	19.10	<0.001	3<1<2
Average number of pauses	1.25	0.29	3=1=2
Maximal number of pauses	6.07	<0.001	3=1<2
Minimal number of pauses	10.40	<0.001	2=1<3
Latency of the last movement	9.46	<0.001	3=1<2
Minimal duration of a roundtrip without pauses	7.80	<0.001	2=1<3
Maximal duration of a roundtrip without pauses	9.42	<0.001	3=1<2
Duration of exploration without pauses	43.29	<0.001	3<1<2

Gaussian mixture analysis performed on the combined data set identified three groups (56, four and 36 individuals, respectively). A MANOVA performed on the behavioral variables indicated significant differences between groups (Wilk's $\lambda=0.02$, $F_{30,158}=31.90$, $P<0.001$). ANOVA showed significant differences in all variables ($P<0.05$) with the exception of the latency to the first movement, the maximal duration of a roundtrip and the average of number of pauses (Table 4). One group showed fewer and slower movements compared with the two others. The other two groups were generally similar, with the exception of the duration of a roundtrip, the number of pauses and the speed-related variables (Table 4). MANOVA testing for differences between the three groups showed significant differences in morphology ($P<0.05$), but not in performance (Wilk's $\lambda=0.85$, $F_{10,150}=1.24$, $P=0.27$). Subsequent ANOVA and *post hoc* tests showed that the differences in morphology were significant when comparing groups 1 and 3 (Table 5).

DISCUSSION

Female exploratory behavior

Across all females, three distinct and robust behaviors were detected using clustering analyses without *a priori* group definition. Two major and opposing behavioral syndromes, bold and shy, are typically identified in many species of animals including invertebrates such as spiders (Kralj-Fišer and Schneider, 2012) and crabs (Watanabe et al., 2012), but also a variety of vertebrates including rodents (Shillito, 2013), birds (Dingemanse et al., 2006) and fish (Wilson and Godin, 2009). Bold individuals typically explore more and make fewer stops or pauses. Yet, this pattern of exploration exposes an individual to risks such as

Table 3. Results of MANOVA testing for differences in locomotor performance and limb morphology across behavioral groups for females and the entire data set combining males and females

	Females			Males and females		
	Wilk's λ	<i>F</i>	<i>P</i>	Wilk's λ	<i>F</i>	<i>P</i>
Performance	0.82	0.79	0.64	0.85	1.25	0.27
Body size	0.93	0.95	0.44	0.67	9.64	<0.001
Head measures	0.86	0.98	0.46	0.81	2.33	0.02
Pelvic girdle dimensions	0.97	0.38	0.82	0.71	8.03	<0.001
Forelimb measures	0.75	1.49	0.15	0.80	1.99	0.04
Hindlimb measures	0.83	0.95	0.49	0.73	2.91	<0.01

Bold indicates significant *P*-values.

predation (Smith and Blumstein, 2008). At the opposite end of the spectrum, shy individuals explore less and spend more time hidden, but take fewer risks. However, these two strategies are likely to have similar overall fitness in the animal's natural unmodified environment. Whereas bold individuals may encounter more resources and sexual partners that allow them to increase the number of offspring, their overall survival may be decreased as a result of the greater risks taken (van Oers et al., 2004; Wolf et al., 2007). In female *X. tropicalis* the two major variables discriminating between bold and shy individuals are the speed of movement and the latency of the last movement. Bold females move fast, yet stop moving earlier.

The data sets for both females (this study) and males (Videliier et al., 2014) suggest the presence of a third pattern of exploration. This pattern has characteristics that are intermediate between the extremes of bold and shy. The individuals showing an intermediate pattern maintained some specific traits of both bold and shy individuals. Intermediate females moved faster and explored longer than shy ones during longer exploration events. However, during short exploration events they were similar to shy individuals, taking many pauses and stopping movements earlier. Both the number of pauses and the duration of a roundtrip are likely strongly linked to the overall risks encountered during exploration; while pauses allow an individual to screen the environment and thus decrease risks, short roundtrips limit the overall exposure to risks. Interestingly, females showing this intermediate strategy modulate their behavior depending on the exploratory phase, taking less risk in short bouts of exploration, yet greater risk during long ones. As movement speed is an important parameter in the context of predator escape (Husak et al., 2008), intermediate females may benefit from a faster exploration when engaging in longer exploration bouts. These intermediate individuals will likely have similar fitness to bold and shy ones because of their ability to balance risk taking and survival (van Oers et al., 2004).

Sexual dimorphism in exploration behavior

When comparing our data for females with previously published data for males of the same species (Videliier et al., 2014), we were able to identify similarities among groups of behaviors. Indeed, three distinct exploration behaviors were identified in each case: the 'classic' bold and shy groups, as well as an intermediate group. However, the intermediate group was not strictly identical in males and females. While intermediate males were characterized by a long latency to initiate the exploration of their environment and are thus more similar to shy individuals, they displayed exploratory behavior that was more similar to that of bold individuals. In females, however, intermediate animals explored at the same speed as bold individuals did, but stopped moving earlier, similar to shy ones. Moreover, the intermediate females appeared to modulate their behavior depending on the duration of exploration. Another interesting similarity between males and females was the relatively low number of shy individuals detected (four females and five males). This low number might be explained by two different scenarios. Firstly, as one of the principal characteristics of shy individuals is to reduce risk taking by staying hidden, the probability of capture is likely lower. Thus, it is relatively common to find only a few shy individuals (Carter et al., 2012; Garamszegi et al., 2009). Secondly, a population often involves the presence of many individuals with one trait and few with an alternative trait. When the environmental context changes, however, the rare trait may become advantageous and invade the entire population at the expense of the initially common trait (Hedrick, 2007). As such, it may remain fixed in the population at low frequency.

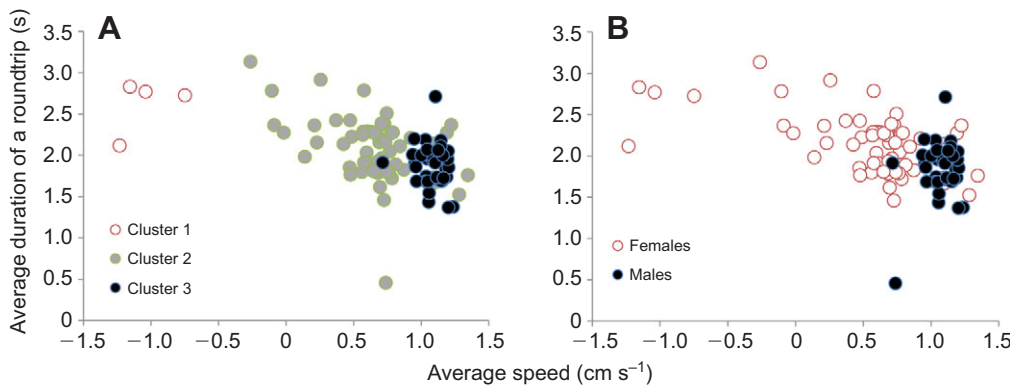


Fig. 2. Graphs of average duration of a roundtrip as a function of average speed. (A) The three behavioral groups identified with the Gaussian mixture analysis are indicated (clusters 1–3). (B) The sex of the individuals is indicated, with open symbols representing females and filled symbols representing males.

When analyzing the data for all individuals of both sexes combined, three groups could be identified as well. Among these three groups, one corresponded to shy individuals, whereas the other two appeared to correspond to bold individuals. Analyses of behavioral traits differing between groups indicated that these groups varied in speed of movement, average duration of a roundtrip and the number of pauses taken. Group 3 appeared to be the boldest and was composed entirely of males, suggesting that, on average, males were bolder than females in their exploration behavior (Fig. 2, Table 4). When comparing sex differences in exploration behavior, we found that males moved more and more quickly and took fewer pauses, resulting in shorter average roundtrip times. Dispersal in *X. tropicalis* is typically observed during the rainy season and is facilitated by both the establishment of new, but temporary, aquatic connections between ponds and overland dispersal during periods of heavy rain. Males are likely more mobile than females as male frogs in general need to find as many sexual partners as possible (Wells, 1977). Boldness is often connected to aggression (Mafli et al., 2011; Wilson and Godin, 2009) and *Xenopus* males are known to engage in male–male competitive fighting (Rabb and Rabb, 1965; Rabb, 1969; Wells, 1977). Moreover, females in some species of animals are known to have a preference for bolder males, which may impose selection on male behavior (Dingemanse and Réale, 2005; Godin and Dugatkin, 1996; Smith and Blumstein, 2008). The relationship between boldness and aggressive behavior is thought to be related to the levels of circulating hormones such as

cortisol, testosterone and melanin, which are typically higher in males (Dingemanse and Goede, 2004; Mafli et al., 2011; Thomson et al., 2011).

Decomposition of locomotion patterns: the dissociation of exploratory behavior and performance/morphology

A previous study of male *X. tropicalis* revealed that individuals in the three behavioral groups did not significantly differ in morphology and performance (Videliér et al., 2014). The fact that a similar result was observed for females suggests that this disconnect may be a general feature of this species. However, when analyzing behavioral variation across both males and females, differences among behavioral groups in morphology, but not performance, were detected. Specifically, differences in morphology were significant between groups 1 and 3. However, an exploration of the data shows that this difference also corresponds to differences between males (group 3) and females (group 1). This result is consistent with previously reported sex differences in morphology (Herrel et al., 2012). Thus, mobility in *X. tropicalis* appears to be composed of two independent sets of traits: the exploration behavior on the one hand and performance/morphology on the other hand.

In the context of the ongoing habitat fragmentation, selection on overall mobility is likely great because of the increase in the distance between optimal habitat sites (Hillers et al., 2008). The maintenance of niche networks is, however, necessary to maintain gene flow within and between populations. Without gene flow, the

Table 4. Results of ANOVA testing for differences in exploration behavior across the behavioral groups identified in the data set consisting of data for males and females combined

	Behavior			Sex		
	<i>F</i>	<i>P</i>	Group	<i>F</i>	<i>P</i>	Group difference
Total distance	23.18	<0.001	2<3=1	0.85	0.36	Male=female
Number of complete roundtrips	8.85	<0.001	2<3=1	0.00	0.97	Male=female
Total number of movements	18.57	<0.001	2<3=1	1.14	0.29	Male=female
Number of movements away from wall	6.78	<0.01	2<3=1	0.62	0.43	Male=female
Time of all movements with pauses	12.01	<0.001	2<3=1	0.39	0.53	Male=female
Average time spent hidden	58.07	<0.001	2<3=1	9.02	<0.01	Male>female
Average speed	122.60	<0.001	2<3<1	35.91	<0.001	Male>female
Maximal speed	138.10	<0.001	2<3<1	11.64	<0.001	Male>female
Minimal speed	84.66	<0.001	2<3<1	132.30	<0.001	Male>female
Average duration of a roundtrip	9.82	<0.01	3<1<2	18.36	<0.001	Male<female
Maximal duration of a roundtrip	2.33	0.10	2=1=3	10.77	<0.01	Male<female
Average number of pauses	5.02	0.01	(1<3)=2	11.92	<0.001	Male<female
Latency of the first movement	2.23	0.11	2=1=3	0.16	0.69	Male=female
Latency of the second movement	93.10	<0.001	2<3=1	1.98	0.16	Male=female
Latency of the last movement	5.20	<0.01	(3<2)=1	3.11	0.08	Male=female

Bold values indicate significant *P*-values and highlighted cells indicate variables with significant differences between clusters 1 and 3.

Table 5. Results of ANOVA testing for differences in locomotor performance and limb morphology across the behavioral groups identified in the data set consisting of data for males and females combined

	<i>F</i>	<i>P</i>	Group difference
Body size			
Snout–vent length (mm)	9.19	<0.001	(1=2)≠3
Mass (g)	20.11	<0.001	(1=2)≠3
Head dimensions			
Length (mm)	3.45	0.04	2=(1≠3)
Width (mm)	4.27	0.02	2=(1≠3)
Height (mm)	8.41	<0.001	2=(1≠3)
Lower jaw length (mm)	0.98	0.38	1=2=3
Pelvic girdle			
Ilium length (mm)	16.16	<0.001	2=(1≠3)
Ilium width (mm)	7.50	<0.001	(1=2)≠3
Forelimb segments			
Humerus (mm)	0.90	0.41	1=2=3
Radius (mm)	1.07	0.35	1=2=3
Hand (mm)	2.33	0.10	1=2=3
Finger (mm)	0.24	0.78	1=2=3
Forelimb length (mm)	0.38	0.69	1=2=3
Hindlimb segments			
Femur (mm)	2.71	0.07	1=2=3
Tibia (mm)	10.92	<0.001	2=(1≠3)
Foot (mm)	9.30	<0.001	2=(1≠3)
Toe (mm)	1.76	0.18	1=2=3
Hindlimb length (mm)	1.36	0.26	1=2=3
Performance			
Average swimming speed (cm s ⁻¹)	0.43	0.65	1=2=3
Peak swimming speed (cm s ⁻¹)	0.04	0.96	1=2=3
Peak swimming acceleration (cm s ⁻²)	2.45	0.09	1=2=3
Maximal time jumped (s)	1.91	0.15	1=2=3
Maximal distance jumped (cm)	0.25	0.78	1=2=3

Bold indicates significant *P*-values.

inbreeding risk increases and genetic variability decreases, thus increasing the fixation of deleterious mutations (Dixo et al., 2009). Given that we expect bold and intermediate individuals to be positively selected in such a context, given their greater tendency to explore novel environments, this may lead to a reduced genetic diversity within fragmented populations, with the rare shy phenotypes being lost. Moreover, previous studies have shown that strong selection on locomotor capacity and mobility may have a significant impact on life-history traits and reproductive output (Phillips et al., 2007; Seebacher and Franklin, 2011). However, to evolve, traits have to be heritable. Previous studies have demonstrated the heritability of morphology and performance (e.g. Le Galliard and Ferrière, 2008; Garland et al., 1990) and exploration behavior has also been found to be heritable in some vertebrate species (Dingemans et al., 2006; Drent et al., 2003; Pulido et al., 2001). Whether the heritability of these behavioral traits is as strong as that of morphological and performance traits remains, however, unknown. Future studies exploring the genetic basis of these behaviors, as well as the underlying genetic basis of variation in mobility would be especially insightful.

In summary, our data demonstrate the presence of three stable behavioral patterns of exploration in female *X. tropicalis*. Although males also show three behavioral groups, the behaviors are different between the two sexes, with males being bolder and exploring more than females. Finally, in both males and females, behavior appears to be decoupled from morphology and performance, suggesting that selection can act on both sets of traits independently.

MATERIALS AND METHODS

Animals

Individuals of *X. tropicalis* were caught in the wild in Cameroon in 2009. An additional 10 individuals that were bred in captivity were added to the data set. Animals were housed at the Muséum National d'Histoire Naturelle (MNHN) in Paris and maintained in 21 l tanks mounted on three-shelf stand-alone *Xenopus* frog racks (Aquanearing, Inc., San Diego, CA, USA) with the water temperature set at 24°C. This temperature is close to the optimal performance temperature of *Xenopus* (Herrel and Bonneaud, 2012) and similar to temperatures measured under field conditions for ponds in the forest (Careau et al., 2014). Animals were fed with beef heart and mosquito larvae twice weekly. All individuals were pit-tagged (Nonatec, Rodange, Luxembourg), allowing unique identification of each individual. Data for morphology and performance of these same individuals were published previously (Herrel et al., 2012). Moreover, data on the exploration behavior in males from the same populations have been published recently (Videliere et al., 2014) and are used here for comparison with the data obtained for females. A total of 61 females were used in the current analysis.

Morphology and performance

Body measures included mass, snout–vent length, ilium length and width, head length, width and height as well as lower jaw length, forelimb segments (humerus, radius, hand and the length of the longest finger) and hindlimb segments (femur, tibia, foot, longest toe length). These measures are important in locomotion and have been described in a previous study (Herrel et al., 2012). Performance measures included swimming velocity and acceleration, as well as terrestrial endurance capacity (time and distance jumped until exhaustion; see Herrel et al., 2012, 2014).

Behavioral analysis

Frogs were filmed for 60 min with a Quickcam Pro 500 (Logitech, Inc., Romanel-sur-Morges, Switzerland) set at 15 frames s⁻¹. Animals were released into a rectangular tank (height 0.98 m, length 0.40 m, width 0.20 m) with a water level of 0.20 m maintained at 24±2°C (see Videliere et al., 2014) and left quietly for 5 min before the onset of the recording. Shelters were placed at the two extremities to provide a hiding place. Each individual was tested three times at different times of the day (morning: 09:00 h–12:00 h; early afternoon: 12:00 h–16:00 h; late afternoon: 16:00 h–20:00 h) in a randomized way. This allowed us to test the repeatability of behavior across different activity periods. Videos were analyzed using ProAnalyst software (Xcitex, Inc., Cambridge, MA, USA) by tracking all movements of frogs during their exploration of the environment for 1 h. Coordinates of the snout tip were extracted and used to quantify exploration behavior.

The following variables were extracted for each video: the total distance (cm) moved in 1 h; the number of all movements and the number of roundtrips (complete or not); the average, minimal and maximal speed of movement (cm s⁻¹) extracted from the videos; the speed of movement without pauses (cm s⁻¹); the latency to the first, second and last movement (s); the average, minimal and maximal duration of a roundtrip with or without pauses (s); the total duration of exploration with or without pauses (s); the total, average, minimal and maximal time spent hidden between two roundtrips (s); the average, minimal and maximal number of pauses; and the number of roundtrips away from the wall of the aquarium. In total, 29 variables were extracted for each 1 h video.

All individuals were in good health and were still alive at the time of the submission of this paper and showed no signs of weight loss. Experiments were approved by the institutional ethics committee at the MNHN.

Statistical analyses

To confirm the assumptions of normality and homoscedasticity, all the data were log₁₀ transformed and tested using Shapiro tests. One outlier was detected in the initial exploration of the female data and showed extreme movements in one recording and no movement at all in the two others. This extreme variability was not present in other individuals and thus this female was removed from the data set. The repeatability of each variable was tested using Pearson correlations (Pearson, 1909). Parameters without correlation

between at least two of the recordings were excluded from the analysis. Thirteen parameters were repeatable: the total distance moved (cm) in 1 h; the number of all movements and roundtrips (complete or not); the maximal speed of movement (cm s^{-1}); the latency of the last movement (s); the minimal and maximal duration of a roundtrip without pauses; the total duration of all movements without pauses; the average, minimal, maximal number of pauses; and the number of roundtrips away from the wall of the aquarium.

To describe variation in the female exploration behavior, a Gaussian mixture model (Fraley and Raftery, 2007) was used without imposed group number. Three groups were detected, and group membership was saved for each individual. To identify the variables that differed between clusters, a MANOVA coupled to subsequent ANOVA and *post hoc* tests was run on the three clusters. Next, we tested whether behavioral groups differed in performance and morphology using MANOVA. Twelve individuals were excluded because of missing performance data. Finally, we combined previously published data for males ($N=37$; Videlier et al., 2014) with our data on females to explore differences between sexes in exploration behavior. When combining the two data sets, 15 variables were found to be repeatable across all individuals: the total distance moved in 1 h (cm); the number of all movements and complete roundtrips; the maximal, minimal and average speed of movement (cm s^{-1}); the latency of the first, the second and the last movement (s); the average and maximal duration of a roundtrip with pauses (s); the duration of exploration with pauses (s); the average duration spent hidden between two roundtrips (s); the average number of pauses; and the number of roundtrips away from the wall of the aquarium. A Gaussian mixture analysis was run and extracted three groups without *a priori* group definition. MANOVA and ANOVA were then run to test for differences between sexes and clusters in exploration behavior, and to test for differences in morphology and performance between the groups identified in the overall data set.

All analyses were performed in R (R Development Core Team, 2010) using the Mclust, Class and conventional packages.

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Competing interests

The authors declare no competing or financial interests.

Author contributions

C.B. and A.H. conceived the study; M.V. did the experiments; M.V., R.C. and A.H. analyzed the data; all authors drafted and revised the paper.

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