No Evidence for Effects of Ecological and Behavioral Factors on Eye Size Evolution in Anurans

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Eye size varies markedly among taxonomic levels, and this variation is often related to the patterns shaped by phylogeny and ecological and behavioral factors. The selective pressures underlying eye size evolution are especially studied in fishes, anurans, birds, and mammals. However, selective pressures underlying the eye size evolution in anurans have inconsistent scaling rules. Here, we investigated the links between eye size and both ecological (e.g., light availability and habitat type) and behavioral factors (e.g., activity time, foraging mobility, defensive strategy, and mating system) among 252 species of anurans by using phylogenetically controlled generalized least-squared (PGLS) regression. Results show that anuran eye size scales hypo-allometrically with body size. However, eye size was not significantly influenced by ecological and behavioral factors, including habitat type, activity time, light availability, foraging mobility, defensive strategy, and mating system. Therefore, neither ecology nor behavior plays a key role in promoting eye size evolution in frogs.

Keywords: anurans, behavioral factors, body size, eye size, hypo-allometrical relationship

INTRODUCTION

The eyes can extract and exploit the information transmitted by light in animals at taxonomic groups (Land and Nilsson, 2012). Eye size varies extensively from tiny to giant among different animals (Martin, 1993; Land, 2009; Land and Nilsson, 2012). The size and dimension of an eye heavily affect the visual system and the abundance and quality of visual information, where animals extract from different environments (Walls, 1942). Although eye acuity is directly associated with eye size and lens diameter, it is also associated with other factors such as quality of the optic component, the angular spacing of the receptors, and the diameter of the photoreceptor (Veilleux and Kirk, 2014). Larger eyes possess larger abundance of photoreceptors and size of images, resulting in longer focal length for determining the retinal area size, where the images of the objects spread (Martin, 2007). Hence, selection for the increased resolving power leads to the enlarged eye size through the longer focal length (Martin, 1983).

While larger eyes can obtain information more effectively from changing environments compared with normal ones, eye size is physically and developmentally constrained (Huber et al., 1997; Moran et al., 2015). Considering the cost associated with benefiting from better vision by providing increased image resolution or enhance sensitivity to light, the eyes may reduce or even
lose during over evolutionary periods in species that move into dark or murky ecosystems (Caves et al., 2017; Porter and Summer-Rooney, 2018). In elasmobranchs, relative and absolute eye size varies considerably and species with smaller eyes tend to be costal, while species with larger eyes are oceanic species living in deep and dark environments (Lisney and Collin, 2007). Moreover, eye size is affected by the activity time, where nocturnal species have larger eye size associated with higher acuity than diurnal species (e.g., mammals: Veilleux and Kirk, 2014; birds: Garamszegi et al., 2002; Hall and Ross, 2007; geckos: Werner and Seifan, 2006).

Eye size is associated with behavioral factors, such as predator avoidance and prey tracking (Garamszegi et al., 2002; Møller and Erritzoe, 2010; Cronin et al., 2014; Thomas et al., 2020). For instance, predator pressure promotes eye size evolution in prey, because the better processing of visual information in species with large eyes enhances the survivability of birds and fishes (Douglas and Hawryshyn, 1990; Land and Nilsson, 2012; Starunov et al., 2017). Moreover, the cutaneous secretion of toxin is one of the main defensive mechanisms against predator risk, and species with granular glands may experience lower predator risk in anurans (Toledo and Jared, 1995; Prates et al., 2012; Huang et al., 2019). As a result, species with granular glands are associated with smaller eye size compared with species with non-granular glands.

For anurans, investigating the relationships of eye size and both ecological and behavioral factors among 44 species have revealed that eye size is not associated with activity time, foraging mobility, habitat type, defensive strategy, mating system, and water turbidity (Huang et al., 2019). However, a recent study has evaluated the eye size and six traits of natural history hypothesized to be related to eye size evolution among 220 species of anurans, and the results indicate that eye size is correlated with adult mating habitat and activity patterns (Thomas et al., 2020). Although anuran eye size was studied outside of a few families, potential relationships between eye size and ecological and behavioral factors are unclear.

In the present study, we investigated the relationships between eye size and both ecological (e.g., light availability and habitat type) and behavioral factors (e.g., defensive strategy, activity time, foraging mobility, and mating system) in 252 anuran species within seven families in China. First, we predicted that species living in aquatic habitats would display increased eye size, because they inhabit light environments. Second, we predicted that nocturnal species would have large eyes to maximize sensitivity in low-light conditions. Third, we predicted that species approaching slowly and capturing prey would have smaller eyes than species approaching quickly and often chasing prey. Finally, we predicted that species with granular glands would possess small eyes, because they experience low predator risk.

**MATERIALS AND METHODS**

**Data Collection**

The data of eye diameter and snout-vent length (SVL) were collected from “Fauna Sinica,” which includes 3,171 males and 1,820 females in 252 anuran species (Fei et al., 2009). We calculated the average eye size and SVL based on males and females for all analyses (Supplementary Table 1). All species in this study have available phylogenetic information.

**Categorical Variables**

Following the method of Thomas et al. (2020) and our filed observation, adult habitat type for each species on a six-point scale were classified into scansorial [primarily associated with plants; up off the ground (arboreal/shrubs/reeds)], ground-dwelling (primarily active on the ground), subfossorial (primarily active under leaf litter; shallow burrowers), fossorial (primarily active in deeper burrows (not burrowing simply for aestivation/long periods of inactivity)], aquatic (primarily found in water; rarely/never leaves water), and semi-aquatic (strongly associated with/commonly found in water but also frequently uses land habitats). Activity time was classified on a two-point scale, namely, nocturnal (primarily active at night), and both diurnal and nocturnal (active during both day and at night) (Huang et al., 2019). Foraging mobility was categorized based on behavioral phenotype, including species approaching slowly and capturing prey and species approaching quickly and often actively chasing prey (Huang et al., 2019). Light availability of species was classified into strong light (i.e., forest is an open environment) and weak light (i.e., light does not penetrate much) (Huang et al., 2019). Defensive strategy was also classified on a two-point scale, namely, species with granular glands in dorsal and ventral skin and species with non-granular glands in skin (Huang et al., 2019). Mating system for each species was classified as polyandry and monandry (Zeng et al., 2016).

**Phylogeny**

The molecular phylogeny was constructed in the 252 species based on three mitochondrial genes and three nuclear genes (Figure 1). The mitochondrial genes included the large and small subunits of the mitochondrial ribosome genes (12S/16S) and cytochrome b (CYTB). The nuclear genes included the rhodopsin (RHOD), the tyrosinase (TYR), and the recombination-activating gene 1 (RAG1). All sequences were aligned using the MUSCLE function in MEGA v.6.0.6 (Tamura et al., 2013), and we determined the best nucleotide substitution model of each gene by using the Akaike Information Criterion in jModelTest v.2.1.2 (Darriba et al., 2012). The best substitution model was TYR, HKY + G for RAG1 and RHOD, GTR + G for 12S and GTR + I + G for CYTB and 16S. Considering the lack of fossil dates, we used these models to construct the phylogenies based on BEAUTi and BEAST v.1.8.3 (Drummond et al., 2012; Mai et al., 2019; Mai et al., 2020), with a relaxed uncorrelated lognormal clock, unlinked substitution models, a Yule speciation process, and no calibration points. The effective sample size values for each tree statistics showed the satisfying convergence of the Bayesian chain and adequate model mixing in Tracer v.1.6.0 (Rambaut and Drummond, 2014). Then, we generated a maximum clade credibility tree with a 20% burn-in and mean node heights before ending the analysis by using TreeAnnotator v.1.8.3 (Drummond et al., 2012; Chen et al., 2021).
Data Analysis
All data analysis were performed in the statistical software R v. 4.1.0 (R Development Core Team, 2021). We analyzed the evolutionary link between eye and body size among species by using phylogenetically controlled generalized least-squared (PGLS) regression with log_{10}-transformed data in the APE-package in R software (v. 5.4-1) (Paradis, 2012; Cai et al., 2020; Huang et al., 2020). In PGLS analysis, we estimated the phylogenetic scaling parameter ($\lambda$) by using the maximum-likelihood method. In the model residuals, $\lambda$ represented the phylogenetic signal. The parameter $\lambda$ estimated the effect of phylogenetic signal on the relationship between eye and body size ($\lambda = 1$ indicates strongly phylogenetic signal, and $\lambda = 0$ indicates no phylogenetic signal). For comparison with allometric studies, where the phylogenetical relationships were corrected, we used the ordinary least-squares (OLS) regressions in stats v. 4.1.0 (R Development Core Team, 2021) and the standardized major axis (SMA) regressions in smatr v. 3.4.8 (Warton et al., 2012) with the same log_{10}-transformed data.

RESULTS
Eye Size Scales Hypo-Allometrically With Snout-Vent Length
A hypo-allometric (slope < 1) interspecific scaling was observed between eye size and SVL in 252 species of anurans (PGLS:...
slope = 0.818, $R^2 = 0.821$, $P < 0.0001$, $\lambda = 0.415$; Figure 2). The OLS and SMA models produced similar fits to PGLS models, though the slopes of SMA models were the highest (slope = 0.889, $P < 0.001$), whereas those of PGLS were the lowest. SMA tests also indicated that the scaling of eye size with SVL was hypo-allometric.

### Eye Size Correlates With Ecological and Behavioral Traits

Analysis with phylogenetic ANCOVA in separate models showed that the relative eye size was not affected by habitat type (Supplementary Table 2; $F_1 < 0.001$, $p = 0.988$), activity time ($F_1 = 0.128$, $p = 0.721$), foraging mobility ($F_1 = 0.017$, $p = 0.897$), light availability ($F_1 = 0.996$, $p = 0.319$), defensive strategies ($F_1 = 0.632$, $p = 0.427$), and mating system ($F_1 < 0.001$, 0.992).

PGLS in a single model revealed that the relative eye size was not affected significantly by habitat type, activity time, light availability, foraging mobility, defensive strategy, and mating system (Table 1). Nocturnal species did not possess larger eyes than diurnal and nocturnal species (Figure 3A). Species approaching slowly and capturing prey did not have smaller eyes than species approaching quickly and often actively chasing prey (Figure 3B). Species living under strong light did not have smaller eyes than species living in weak light (Figure 3C). Species with granular glands in dorsal and ventral skin did not have smaller eyes.

**FIGURE 2** Hypo-allometrical relationships between eye diameter and snout-vent length across 252 anurans species.
eyes than species with non-granular glands in skin (Figure 3D). The effects of habitat type and mating system on eye size is shown in Supplementary Figures 1A,B.

**DISCUSSION**

Eye size scales with body size among 252 species of anurans in this study, in which species with larger bodies have larger eye size, indicating a hypo-allometric relationship between body and eye size. However, eye size in anurans is not affected by ecological and behavioral factors (e.g., adult mating habitats, activity time, foraging mobility, light availability, defensive strategy, and mating system). Scansorial species do not have larger eyes than ground-dwelling, fossorial, and aquatic species. Nocturnal species living under weak light do not possess larger eyes than both nocturnal and diurnal species living under strong light. Moreover, species quickly and often actively chasing preys do not display larger eyes than species approaching slowly preys. Moreover, species with granular glands did not have larger eyes than those without granular glands. Hence, these traits are not correlated with eye size when correcting for body size and phylogeny, suggesting that light availability and foraging behavior cannot play a key role in shaping eye size evolution in anurans.

Body size is the strongest predictor of eye size, which is expected as larger species usually possess larger eyes than smaller species in sharks, reptile, birds, and mammals (Brooke et al., 1999; Kiltie, 2000; Garamszegi et al., 2002; Ross et al., 2006; Werner and Seifan, 2007; Lisney and Collin, 2007; Liu et al., 2012). For anurans, eye size scales hypo-allometrically with SVL (Huang et al., 2019; Thomas et al., 2020) and we found the same patterns as Huang and Thomas that larger species had larger eyes, displaying hypo-allometrical relationship between eye size and body size.

Eye size varied among adult habitats in animal groups. For instance, evidence suggests that larger eye in anurans are beneficial to scansorial species by accommodating fast temporal resolutions during jumping process and high acuity in visually complex arboreal habitats (Thomas et al., 2020). Likewise, fossorial species reduced eye size as adaptations to dark and/or abrasive habitats in mammals (Borghi et al., 2002), fishes (Eageder and Adrians, 2010), caecilian amphibians (Mohun et al., 2010), and reptiles (Yovanovich et al., 2019). However, the variation in the eye size of the 44 species of anurans investigated seems largely independent of adult habitats, possibly because all specimens were obtained from one, albeit large, geographical region (Huang et al., 2019). In the present study, eye size did not increase from fully fossorial to subfossorial to non-fossorial (e.g., ground-dwelling, semiaquatic, scansorial) species, indicating that the fossoriality degree cannot determine eye size evolution across 252 species of anurans.

Many nocturnal vertebrates display enlarged eye size to maximize visual sensitivity (e.g., primates: Kirk, 2006; birds: Hall and Ross, 2007; reef fish: Schmitz and Wainwright, 2011). For instance, nocturnal birds evolve larger eyes to provide a wide pupil and improve light sensitivity during night compared with diurnal birds (Martin, 1985; Brooke et al., 1999). Significant influences were observed in the activity pattern of relative eye size among the 220 species of anurans, in which nocturnal species evolved larger eyes compared with both diurnal species and nocturnal and diurnal species (Thomas et al., 2020). However, previous studies have indicated that activity pattern cannot affect variation in relative eye size among species because most of the 44 species of anurans can be seen foraging and mating also at night (Huang et al., 2019; Mai et al., 2020). Similar to the findings of Huang et al. (2019), we found that activity pattern did not affect relative eye size among the 252 species of anurans, possibly because the common anuran ancestor is likely to be nocturnal, and the majority of the species of extant anurans retain this active pattern (Anderson and Wiens, 2017). Large eyes are associated with larger abundance of photoreceptors and produce larger image sizes, thereby collecting more light per solid angle of image than small eyes (Martin, 2007). For fishes, the larger eyes increase the chance of photon capture when detecting small bioluminescent flashes at low levels of sunlight (Warrant, 2000; Warrant and Locket, 2004). Moreover, to improve light sensitivity at night, it may be beneficial to enlarge not only retinal area, but also pupil diameter in dim light conditions (Martin, 1985). However, light availability did not affect the eye size evolution in anurans, where species living under weak light did not have larger eyes than those living under strong light. Hence, nocturnal species with higher light availability did not evolve larger eyes than diurnal species, suggesting that light availability was not correlated with detection of predator risks.

Vision plays an important role in searching for foods for birds, as indicated by marked differences in eye size across the types of foraging mobility, where species approaching quickly and actively capturing preys displayed an increase in eye size, whereas those approaching slowly preys displayed a decrease in eye size (Garamszegi et al., 2002). In a previous study on 44 species of anurans, foraging mobility did not affect eye size evolution, suggesting that anurans approaching quickly preys may live in low-light conditions, which cannot lead to increased eye size compared with anurans approaching quickly preys (Huang et al., 2019). All the 44 used species are possibly at least partially active at night. However, after sampling a larger number of species in our study, the results did not confirm the positive effect of capture behavior on eye size evolution.
Predator risk is the strongest selective pressure in shaping eye size evolution (Garamszegi et al., 2002). Considering species with larger eyes can easily spot predators early (Striedter, 2005; Kotrschal et al., 2015), stronger predator risk coincides with the larger-eyed birds (Møller and Erritzøe, 2010). The relationship between eye size and predator risk may be as prominent as it is for instance in birds with eye size that is positively correlated with fight distance (Blumstein et al., 2004). Consistent with a previous study (Huang et al., 2019), we found that species with granular glands in skin did not exhibit smaller eyes than species with non-granular glands in skin, suggesting that species depending on chemical defense cannot shape eye size evolution in anurans. Moreover, relative eye size is neither correlated with sexual size dimorphism nor mating system among the 44 species of anurans (Huang et al., 2019). In the present study, we found that eye size variation in 252 species of anuran cannot be explained by...
mating system contrary to the prominent role of sexual selection in eye size evolution.

In conclusion, eye size variation cannot be explained by ecological and behavioral factors including activity time, light availability, foraging mobility, and defensive strategy in our sample of 252 Chinese frogs. Eye size is not correlated with activity pattern and light availability, showing that a potential nocturnal bottleneck is not observed in anuran eyes evolution. Moreover, foraging mobility is not associated with the eye size in anurans, highlighting that foraging behavior cannot shape eye size evolution. Considering that defensive strategy (i.e., poison glands) does not affect eye size, the anti-predator ability in anurans cannot promote the enlarged eye size.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding author/s.

ETHICS STATEMENT

The animal study was reviewed and approved by China West Normal University.

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AUTHOR CONTRIBUTIONS

YJ and CC participated in laboratory work, data analysis, and manuscript drafting. LJ and WBL conducted data analysis and visual representation of the data. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

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