Life History Traits, Elasticity Analyses, and Phenotypic Plasticity of *Squaliobarbus curriculus* in the Pearl River Estuary, China

Teng Wang¹,²,³,⁴, Lin Lin¹,³,⁴, Yong Liu¹,²,³,⁴*, Ivan Jakovlić⁵, Chun-hou Li¹,²,³,⁴*, Ya-yuan Xiao¹,³,⁴ and Peng Wu¹,³,⁴

¹Key Laboratory of South China Sea Fishery Resources Exploitation & Utilization, Ministry of Agriculture and Rural Affairs, South China Sea Fisheries Research Institute, Chinese Academy of Fishery Sciences, Guangzhou, China, ²Southern Marine Science and Engineering Guangdong Laboratory (Guangzhou), Guangzhou, China, ³Scientific Observation and Research Field Station of Pearl River Estuary Ecosystem, Guangzhou, China, ⁴Guangdong Provincial Key Laboratory of Fishery Ecology and Environment, Guangzhou, China, ⁵Bio-Transduction Lab, Wuhan, China

*Squaliobarbus curriculus*, commonly known as red-eye fish, is widely distributed in East Asia. It is one of the important aquatic germplasm resource and economic species in the Pearl River. To give suggestions for better protection and management, we investigated its life history and conducted elasticity analyses. Samples (*n* = 451) were collected between 2016 and 2017 from the western Pearl River estuary. There were no significant differences between the length-weight relationships of females and males (*W* = 0.00001SL^3.121). The von Bertalanffy growth function was *L*ₚ = 553.2 [1 − e^−0.111(t+1.009)]. The estimated length at 50% sexual maturity for females was 209.6 and 200.0 mm for males, both at 3 years of age. Oocyte size-frequency distribution suggested batch spawning. Fecundity ranged between 9,407 and 175,086 eggs per fish (mean = 51,040, or 143.9 eggs/g of fish weight).

To better understand the ecological phenotypic plasticity of *S. curriculus* we conducted meta-analyses on all available life history data for this species. Our results showed that the standard lengths at ages 2 and 3 in the estuary were significantly smaller than in the upper reaches of the Pearl River basin, and there were also obvious differences in fecundity and oocyte size. For more, the standard lengths at ages 2 and 3 were correlated negatively with latitude. Elasticity analysis showed that juveniles’ (aged 1–3) survival had the largest contribution to the population growth rate, which suggests that management efforts should focus on the early life stages.

Keywords: growth, reproductive biology, salinity, ectotherms, latitude

INTRODUCTION

Due to its popularity with consumers and resilience to unfavorable environmental conditions, the importance of barbel chub *Squaliobarbus curriculus* (Richardson, 1846) for freshwater aquaculture and capture fisheries in southern China has been steadily growing during the last few decades (Lei et al., 2012). Belonging to the family Cyprinidae (Leuciscinae subfamily), and commonly known as red-eye fish for its red spots on the superior border of eyes, *S. curriculus* is mainly distributed in Asia, including China, western Korea, Vietnam, and Amur River drainage in Russia. As it is one of the dominant species in the western Pearl River Estuary (according to our unpublished 2016–2017...
survey), it is also an important commercial fishing species in this area. Importantly, the National Aquatic germplasm resources protection area for *S. curriculus* (and *Erythroculter pseudobreviceaula*) is also located in the Xijiang River, only 130 km upstream from our studied area.

Life history characteristics such as age, growth, and reproduction of fish populations are crucial for determining the fish recruitment and understanding the fish population dynamics (Quinn and Deriso, 1999; Lowerre-Barbieri et al., 2017). Therefore, life history studies are the most important prerequisite required for the assessment of exploited fish populations (Campana, 2001), and essential for fish population management and conservation (Coppel et al., 2017). By analyzing the life history of *S. curriculus*, we aim to improve our understanding of its population dynamics and produce data necessary for designing conservation measures to prevent the decline of its aquatic germplasm resources.

Elasticity analysis quantifies the relative importance of vital life history traits (fecundity, growth, and survival) for the overall population growth rate. This, in turn, provides guidance for management efforts, which should generally focus on demographic parameters with the highest elasticity (Caswell, 2000). Elasticity analysis is one of the most widely used tools in demographic studies, evolutionary and population ecology studies, and particularly in studies aimed at the conservation of exploited and endangered species (Heppell et al., 2000; Manlik et al., 2018). It has been used to guide wildlife management (de Kroon et al., 2000; Manlik et al., 2018), not only in vertebrates (Sæther and Bakke, 2000; Heppell et al., 2000; Gerber and Heppell, 2004, Wang et al., 2017), but also in invertebrates (Sommerville et al., 2014) and plants (Gron, 2016).

In order to better protect and manage *S. curriculus* populations, it is also important to understand their phenotypic plasticity in different environments (Noss 2001). Salinity and temperature (latitude) are two important environmental factors that will affect the variation of the life history characteristics of *S. curriculus*. In estuaries, freshwater meets seawater, so the salinity in estuary habitats is affected by a number of factors, most notably sea tides and freshwater influx, and therefore it can vary notably over relatively short time periods (Bricheno et al., 2021). Due to the demands of maintaining osmotic balance, this variation causes metabolic stress in fish (Nordlie, 2006). Therefore, these environmental factors (salinity fluctuation) might be producing observable impacts on the life history traits of wild *S. curriculus* populations inhabiting the estuary of the Pearl River. The body sizes of organisms tend to be inversely correlated with latitude and temperature (Meiri, 2011; Rypel, 2014), but studies in ectotherms (including fish) often produce conflicting results (Bauer, 1992; Mousseau, 1997, Belk and Houston, 2002, Angilletta and Dunham, 2003, Ashton and Feldman, 2003, Heibo et al., 2005, Pincheira-Donoso et al., 2008, Chucholl, 2011, Rypel, 2014). Thus, a general explanation for the variation in size-latitude relationships of ectotherms has remained elusive (Angilletta and Dunham, 2003; Chucholl, 2011; Rubalcaba et al., 2019).

To propose appropriate fisheries management measures, we sampled the *S. curriculus* populations inhabiting the western part of the Pearl River estuary over a 1-year period, inferred their life history traits (age, growth, sex ratio, size at maturity, and fecundity), and then conducted elasticity analyses. Following this, the ecological phenotypic plasticity was studied to allow us to propose better protection and management measures for *S. curriculus*.

**MATERIALS AND METHODS**

**Study Site**

The Pearl River (length = 2,214 km, drainage basin = 452,000 km²) is the second largest river in China in terms of annual water discharge, with 3.26 × 10¹¹ m³ yr⁻¹. Although they merely share a common delta, Xijiang, Beijiang, and Dongjiang Rivers are considered tributaries of the Pearl River. The largest, Xijiang, accounts for ~70% of the total Pearl River freshwater discharge (China Bureau of Hydrology, Ministry of Water Resources, http://sqqx.hydroinfo.gov.cn/websq/). All samples for this study were collected in the Xijiang part (or the western part) of the Pearl River estuary, located in the vicinity of Jiangmen City, Guangdong Province, China (Figure 1).

**Sample Collection and Processing**

Sampling was conducted on 10 sites between December 2016 and September 2017. Each site was sampled thrice in each season: 22–24th December 2016, 24–26th March, 12–14th July, and 19–21st September 2017 (Figure 1). The sampling was conducted using 5 drift gillnets (100 m long × 1.5–2.3 m high with mesh sizes ranging from 4 to 7 cm) and 2 trap nets (20 m long × 0.6 m high, with mesh size 0.2 cm) set in the evening for 5 h. In total, 451 specimens were collected. The standard length (SL 1 mm) and weight (W 0.01 g) were recorded. The sexual characteristics were macroscopically analysed on 358 specimens (after discarding too small and damaged samples). Gonads were surgically removed, weighed (precision = 0.01 g), and preserved in 10% formalin for subsequent determination of the fecundity and oocyte size-frequency distributions. Gonad maturity stages, determined using a dissection microscope, were identified as stage I - inactive stage, stage II - recovering stage, stage III - early maturing stage, stage IV - late maturing stage, stage V - mature stage, and stage VI - spent stage (Bancroft and Stevens, 1996). Oocyte diameter was calculated as the average of the major and minor axes. Oocytes that did not contain yolk were not measured.

**Length-Weight Relationships**

We examined the difference in SL distributions between sexes using the Kolmogorov–Smirnov test (K–S test). The standard length and weight relationship (W = aSLb) was converted into the logarithmic form lnW = ln(a + bLN). (Ricker, 1975), where a and b parameters were calculated using least-squares regression. SL–W relationships between sexes were compared using the analysis of covariance (ANCOVA). To test whether the growth of fish was isometric, the student’s T-test was applied to assess whether the b parameter significantly differed from the expected value of 3 (Pauly, 1984).
Age Estimates and Growth
The annual growth of scales was used to estimate the age of fish, needed for subsequent growth estimates. Scales were taken from the end of the pectoral fin and the beginning of the dorsal fin of 437 specimens and analysed under a dissecting microscope as described before (Wang et al., 2015). Each scale was interpreted twice by one reader, with an interval of at least 2 weeks between the two counts. The reader did not have any prior information on the sex, length, or capture time of specimens. Counts were accepted as correct if they were in agreement. If the two counts differed, a recount was conducted and accepted as correct only if it matched any of the previous two counts. If the count did not match any of the previous two, the specimen was not used for downstream analyses. Specimens in different age groups were all assigned a birth date of the 1st of May. The von Bertalanffy growth function (VBGF) based on length-at-age from all age readings were fitted by non-linear regression: $L_t = L_{\infty} \left[ 1 - e^{-k(t-t_0)} \right]$. $L_t$ is the length at age $t$, $L_{\infty}$ is the asymptotic length, $k$ is the growth coefficient, and $t_0$ is the age at length 0.

Reproduction
Chi-squared ($\chi^2$) test was used to determine whether the sex ratio deviated from 1:1 (Zar, 1999). Sexually mature specimens for both sexes were defined as possessing stage III to stage VI gonads during the spawning season. The SL at 50% maturity ($L_{50}$) was calculated by fitting the logistic curves to the proportion of fish designated as mature in each 10 mm length class using a non-linear least-squares procedure (the Marquardt method). The logistic equation used is $P = 100/[1 + \exp[-a \times (SL-L_{50})]]$, where $P$ is the percentage of mature specimens at SL, SL is the standard length (mm), and $a$ is the slope of the curve.

Phenotypic Plasticity
To better understand the ecological phenotypic plasticity of $S. curriculus$, we tested the impacts of varying salinity on life history traits and correlation between size and latitude by conducting a meta-analysis using all available life history data for this species, across a broad range of latitudes and habitat types (Table 1). We obtained all published literature on the life history of $S. curriculus$, and calculated the length at the ages of 2 and 3 according to VBGF. If there was data available for the length at the ages of 2 and 3 in the literature, we directly used the data from the literature. We obtained latitude data from Google Maps. In the analysis of length and latitude, we excluded the relevant data for the estuary.

Survivorship, Mortality Rates, and Elasticity Analysis
The population catch curve was established from the age-frequency distribution to estimate the total mortality rate ($Z$), corresponding to the sum of natural mortality ($M$) and fishing mortality ($F$). The natural log of the number of specimens ($\ln N$) was plotted against the age. The slope ($b$) of the descending right limb of the age-frequency distribution curve was found through
TABLE 1 | Length at age and sampling site for S. curriculus populations from China analysed in this study. L2 and L3 indicate length at the ages of 2 and 3 respectively.

| Site                                      | L2 (mm) | L3 (mm) | Latitude (N) | Sample size | Sampling year | References     |
|-------------------------------------------|---------|---------|--------------|-------------|---------------|----------------|
| Jialing river, Chongqing                  | 238.43  | 297.50  | 31.16        | 122         | 1976-1977     | He and Yang (1997) |
| Fuhe river, Hebei                        | 186.00  | 236.12  | 30.86        | 267         | 2003-2005     | Yang et al. (2006) |
| River in Guangxi                          | 258.29  | 311.37  | 23.53        | 892         | 1981-1984     | GXRIF 1984      |
| Huijiang river, Guangxi                  | 259.00  | 309.00  | 22.77        | 1981-1984   | GXRIF 1984    |                |
| Xijiang river, Guangxi                   | 254.00  | 294.00  | 23.44        | 1981-1984   | GXRIF 1984    |                |
| Lijiang river, Guangxi                   | 258.00  | 309.00  | 24.26        | 1981-1984   | GXRIF 1984    |                |
| Youjiang river, Guangxi                  | 231.00  | 281.00  | 23.75        | 1981-1984   | GXRIF 1984    |                |
| Lake Baiyangding, Hebei                  | 165.12  | 217.95  | 38.87        | 30          | 1978          | XDRF 1984      |
| Lake Suoyu, Henan                        | 229.00  | 270.00  | 33.04        |             |               |                |
| Songhua river, Heilongjiang              | 118.00  | 154.00  | 46.18        |             |               |                |
| Lake Dongting, Hunan                     | 208.00  | 271.00  | 28.89        | 179         | 1958-1959     | Li (2009)       |
| Lake Nandagang, Hebei                    | 147.50  | 250.60  | 38.50        |             | 1981-1983     | GXRIF 1984      |
| Haife river, Tianjin                     | 167.80  | 220.76  | 39.20        |             |               | Nicolsky (1963) |
| Hanjing river, Hebei                     | 186.00  | 253.00  | 32.05        | 166         | 1977          | Deng et al. (1981) |
| Changjiang river, Anhui                  | 191.54  | 218.31  | 31.29        | 338         | 2006-2007     | Guo et al. (2009) |
| Beijing river, Guangdong                 | 218.00  | 279.00  | 31.29        | 70          | 1981-1983     | Pan (1989)      |
| Dongjiang river, Guangdong               | 185.00  | 245.20  | 24.25        | 304         | 1981-1983     | Lu (1990)       |
| Xijiang river, Guangxi                   | 197.00  | 258.00  | 23.42        | 892         | 1981-1983     | Lu (1990)       |
| Xijiang river/estuary, Guangdong         | 172.00  | 225.00  | 23.05        | 307         | 1981-1983     |                |
| *Xijiang river, Guangdong                | 171.30  | 227.85  | 23.04        | 296         | 2006-2008     | Zhu et al. (2013) |

*GXRIF: Guangzhou Zhuang Autonomous Region Fisheries Research Institute.
*XDF: Xinxiang Normal University, Department of Biology, Fish Writing Team.
*HFRI: Hunan Fisheries Research Institute.
*Estuary area.

Least-squares linear regression, with b estimating the Z (Ricker, 1975).

The methodology described by Chen and Watanabe (1989) was used to estimate the M (natural mortality) for each age-class t:

$$M(t) = \frac{k}{1 - e^{-kt}}$$

and

$$M(t, t_{max}) = \left[\frac{1}{t_{max} - t}\right] \times \ln \left[\frac{e^{kt_{max}} - e^{kt}}{e^{kt_{max}} - e^{kt_{0}}}\right]$$

for t ≤ t_{mat},

in which t_{mat} is the age at 50% sexual maturity, t_{max} is the maximum age in a population (life span), k and t_{0} are the von Bertalanffy growth curve parameters.

Survival rates of the age zero (S_{0}) can be calculated according to the population growth rate (\lambda).

Survival values (S) for each mortality rate were estimated using the formula described by Ricker (1975): S = e^{-\lambda t}.

For the cases when t ≤ t_{mat}, age-specific survivorship and mortality rates were estimated using the methodology described by Chen and Watanabe (1989); for t_{mat} < t, age-specific survivorship and mortality rates were estimated using the population catch curve.

The mean number of female offspring per female was multiplied by 1/2 of the fecundity. Fecundity was then converted to age-specific fecundity, according to the relationship between fecundity and the age of the fish species.

According to the above data, elasticity analyses were calculated using the projection matrix A, or Leslie matrix (Caswell 1989):

$$F_i = S_0 \times c \times \frac{P_i}{F_{i-1}}$$

$$A = \begin{bmatrix} F_1 & F_2 & \ldots & F_i \\ P_1 & 0 & 0 & 0 \end{bmatrix}$$

$$\lambda = e^{\sum \log a_{ij}}$$

$$e_{ij} = \frac{a_{ij}}{\lambda}$$

where a_{ij} is the (i, j) element of the matrix A, v and w are the left and right eigenvectors of the projection matrix A, and <w, v> is the scalar product of the two vectors \(|v_1 \times w_1 + v_2 \times w_2 + \ldots|\).

From there, fertility elasticity, juvenile survival elasticity and adult survival elasticity were calculated according to Heppell et al. (2000) and Wang et al. (2017). Elasticity analyses were performed using “Popbio” R package (http://cran.r-project.org/web/packages/popbio/).

In the present paper, we defined juvenile as age-1 to age at maturity, and adult as age after maturation.

RESULTS

Length-Weight Relationships

The standard length ranged from 40 to 350 mm (mean ± SD = 185.1 ± 53.4 mm), and weight from 1.00 to 986.21 g (mean ± SD = 147.3 ± 127.0 g). The K–S test found a significant difference between the SL distributions of females and males (H = 1.448,
p = 0.030), but the difference was not significant after removing 11 individuals larger than 290 mm (H = 1.121, p = 0.162; among these 11 individuals, there was only one male specimen, which skewed the data distribution).

Length–weight relationships were calculated separately for females and males. The regression equations were $W = 0.00001SL^{3.095}$ ($r^2 = 0.966$, $n = 174$) for females and $W = 0.00001SL^{3.215}$ ($r^2 = 0.974$, $n = 147$) for males. No statistically significant differences were detected for SL–W relationships between sexes (ANCOVA after log-transformation, $n = 321$, $F = 3.641$, $p = 0.057$). The regression equation derived from pooled data was $W = 0.00001SL^{3.121}$ ($r^2 = 0.986$, $n = 451$). The allometric index value ($b = 3.121$) obtained from the function was significantly larger than 3 ($t$-test, df = 450, $t = 6.94$, $p < 0.01$), which indicates a positive allometric growth (Figure 2).

**Age Structure and Growth**

The age varied from 1 to 7 years for females and 1–6 years for males (Figure 3). The VBGFs fitted to length-at-age data were described as $L_t = 552.2 \left[1 - e^{-0.111(t+1.123)}\right]$ for females ($n = 174$) and $L_t = 616.5 \left[1 - e^{-0.091(t+1.21)}\right]$ for males ($n = 147$). As there were no significant differences in SL–W relationships between sexes, the VBGFs fitted to all of the length-at-age data together was: $L_t = 553.2 \left[1 - e^{-0.111(t+1.099)}\right]$ (Figure 4).
Reproduction

Among the 321 successfully sexed specimens, 174 were females and 147 were males. This puts the overall sex ratio (F/M) at 1.18, not significantly different from the expected 1:1 ratio ($\chi^2 = 2.271$, df = 1, $p = 0.132$). Analysis of seasonal gonadal maturation stages variation showed that maturity stage IV of females and males was found between March and September, with the highest frequency in July (summer), whereas stage III was found throughout the year (Figure 5).

Size-at-maturity estimates were based on the examination of the 282 specimens collected during the spawning period (March to September). Logistic curves describing the proportion of
mature specimens in each 10 mm interval of SL were expressed as:

\[ P = 100/1 + e^{0.3949(\text{SL} - 20.9)} \]  
\[ (r^2 = 0.934, n = 160) \] for females and

\[ P = 100/1 + e^{0.3949(\text{SL} - 20.9)} \]  
\[ (r^2 = 0.841, n = 122) \] for males (Figure 6). The estimated length at 50% sexual maturity (SL50) was 209.6 mm for females and 200.0 mm for males, both at 3 years of age according to the VBGFs. The SLs of the smallest sexually mature female and male were 175 mm (aged 2) and 172 mm (aged 3), respectively.

Oocyte size-frequency distribution in stage IV consisted of up to two cohorts of oocytes, which indicates two batches in the spawning period (Figure 7). Fecundity was estimated from 30 mature females that ranged from 213 to 350 mm SL. The estimated values of fecundity ranged between 9,407 and 172,086 eggs per fish, and the mean was 51,040 eggs (SD 37,945). The relative fecundity was between 25.4 and 37,945. The fecundity of S. curriculus increased linearly with increasing standard length and total weight (Figure 8), and the fitted regression equation was:

\[ F = 592.89SL - 100561, r^2 = 0.2573 \] and

\[ F = 138.17W + 1622.7, r^2 = 0.4198 \] respectively. The fecundity also increased linearly with increasing age (Figure 8), and the best function was described as:

\[ F = 21527A - 42780, r^2 = 0.3949 \] .

### Phenotypic Plasticity

The standard lengths at ages 2 (157.00 mm) and 3 (199.00 mm) determined in the present study are significantly smaller than in the upper reaches of the Pearl River basin (Table 1): 254.00 and 294.00 mm in the Xijiang River, 258.00 and 309.00 mm in the Liujiang River, 231.00 and 281.00 mm in the Youjiang River (all three in the Guangxi province, China, and all are tributaries of the Pearl River). The corresponding two values were also somewhat larger in a study of this species conducted in the years 1981–1983 in the Xijiang River/estuary (Guangdong province): 172.00 and 225.00 mm. The fecundity of 3- (21,801 eggs) and 4-year-olds (43,328 eggs) is also significantly lower than in the upper reaches of the river (54,238 and 126,639 eggs, respectively) (Pearl River Fishery Resources Survey Editorial Committee, 1985). As opposed to this, the oocyte size in stage IV (0.70–1.20 mm Figure 7) is larger than in the upper reaches of the river (0.37–1.09 mm) (Pearl River Fishery Resources Survey Editorial Committee, 1985).

Our meta-analysis of S. curriculus populations in China indicates that standard lengths at ages 2 (L2 = -4.7809N + 348.93, r^2 = 0.6643, p < 0.01) and 3 (L3 = -4.6586N + 399.43, r^2 = 0.6422, p < 0.01) were correlated negatively with latitude (the latitude ranged from 23.04 to 46.18; Table 1).

### Elasticity Analyses

According to the relationship between fecundity and age of S. curriculus in this study, and the fact that it breeds twice a year, the age-specific fecundities of 3–7 year-old females were 21,801, 43,328, 64,855, 86,382, and 107,909 eggs per fish, respectively. We divided the number of specimens aged 2 and 3 in September 2017 by the number of specimens aged 2 and 3 in December 2016, thereby calculating the finite rate of population increase:

\[ \lambda = 1.069 \] . Using this number, the survival rate of the age zero (S0) was 0.000035 year^-1. Thus, F0 values of 3–7 year-old females were 0.766, 1.521, 2.277, 3.033, and 3.789 offsprings per fish, respectively. Age-specific survival rates of 1–6 year-olds were 0.499, 0.838, 0.944, 0.290, 0.290, 0.290 year^-1, respectively. Based on the above results, the inferred Leslie matrix (age-based projection matrix model) is:

| Age | 0 | 0.766 | 1.521 | 2.277 | 3.033 | 3.789 |
|-----|---|-------|-------|-------|-------|-------|
| 0   | 0 | 0     | 0     | 0     | 0     | 0     |
| 0.499 | 0 | 0     | 0     | 0     | 0     | 0     |
| 0.838 | 0 | 0     | 0     | 0     | 0     | 0     |
| 0   | 0.944 | 0 | 0     | 0     | 0     | 0     |
| 0   | 0 | 0.290 | 0     | 0     | 0     | 0     |
| 0.290 | 0 | 0     | 0     | 0     | 0     | 0     |

Elasticity analyses based on the age-based projection matrix model showed that: fertility elasticity (age 0–1 elasticity) of S. curriculus was 0.242, juvenile (specimens aged 1–3) survival elasticity was 0.485, and adult (specimens aged > 3) survival elasticity was 0.273.

### DISCUSSION

S. curriculus is a medium-sized fish (its maximum total length is 48.8 cm according to the FishBase) with a short lifespan (the maximum age in this study was 7 years old). It has early sexual maturity (the earliest sexual maturity was 2 years old), high fecundity, long breeding period, and it can reproduce twice a year. This life history characteristics underlie its strong adaptability to environmental and human disturbances. This study showed through meta-analysis that the life history characteristics of S. curriculus were affected by fluctuations in environmental factors (such as salinity and temperature), and elasticity analysis proved that the early life history stage was the most critical life history stage to maintain its population growth.

### Estuarine Fish Life History Characteristics

The estuary is a region that is affected both by the sea tides and widely varying freshwater influx, which can result in a strong variation in salinity (Bricheno et al., 2021). In such an environment, maintaining homeostasis can be energetically very costly, and affect the amount of energy available for growth and reproduction (Schreck, 2010). Therefore, fluctuating salinity often has highly significant effects on life history traits, such as growth and reproduction (Stearns, 1992; Martin et al., 2009). The S. curriculus populations studied in this research also demonstrate this effect, for the individuals of the same age in the estuary were significantly smaller in size and fecundity than individuals in the upper reaches, but the oocyte size in stage IV was larger. Although there could be a large number of variables synergistically producing these observed phenomena, we hypothesise that they may be at least partially attributed to increased metabolic costs of osmoregulation in a fluctuating-salinity environment (Gan et al., 2016). For example, a similar decrease in growth and reproduction parameters with increasing salinity has been reported in Oreochromis niloticus and Colossoma macropomum (Lowe et al., 2012; Fiúza et al., 2015).
Intriguingly, the “environmental stability hypothesis” predicts that more environmentally variable habitats select for organisms that have higher reproductive allotment (per unit body mass), larger brood sizes, and smaller offspring size relative to organisms from more stable habitats (Stearns, 1983, 1992). Examples of species fitting this pattern are the sailfin molly Poecilia latipinna, the western mosquitofish Gambusia affinis, and the least killifish Heterandria formosa (Martin et al., 2009). The discrepancy between our results and this hypothesis may be caused by different adaptation characteristics of the fish to the environment. We should also mention that some of the differences in fecundity and oocyte size may be partially caused by the comparison of studies spanning several decades, so changes in climate and water environment may also have an impact.

### Size-Latitude Relationships

Variation in life-history traits with latitude (temperature) can be explained by phenotypic plasticity (Roff, 2002; Ren et al., 2020), and a ubiquitous rule is that body sizes of organisms tend to be smaller at high temperatures and low latitudes, and larger at low temperatures and high latitudes (Meiri, 2011; Rypel, 2014). Our meta-analysis of *S. curriculus* populations in China indicates exactly the opposite. This rule was also disproven in many other freshwater fishes (Belk and Houston, 2002; Heibo et al., 2005, Rypel, 2014), as well as other ectotherm animals, such as insects (Mousseau, 1997), snakes (Ashto and Feldman, 2003), and lizards (Pincheira-Donoso et al., 2008). However, many ectotherms are known to exhibit a larger size in colder environments (Angilletta and Dunham, 2003), such as the freshwater pearl mussel *Margaritifera* (Bauer, 1992), chelonian turtles (Ashto and Feldman, 2003), some crayfish (*Procambarus clarkii*) (Chucholl, 2011). A general explanation for the variation in size-latitude relationships of ectotherms remains elusive (Angilletta and Dunham, 2003; Chucholl, 2011; Rubalcaba et al., 2019).

Intriguingly, among the freshwater fishes on the North American continent, cool- or cold-water species were in agreement with the rule, whereas the opposite was found for warm-water species (Rypel, 2014). Temperature is perhaps the most important environmental factor for fish growth, as fish have a thermal optimum (T<sub>opt</sub>, Zarco-Perello et al., 2012); at lower temperatures, metabolic rate declines and growth rate decreases (Reynolds, 2002; Chung et al., 2021), but temperatures higher than T<sub>opt</sub> are likely to lead to declines in growth (Zarco-Perello et al., 2012). Since the environmental temperature decreases with latitude, our meta-analysis of *S. curriculus* populations in China shows that size is positively correlated with the average temperature. As our data comprised only localities north of the 23 degrees north latitude, they don’t comprise the T<sub>opt</sub> of *S. curriculus*; this explains the positive association between growth and temperature found in this study.

### Elasticity Analysis

Elasticity analysis quantifies the contributions of different variables to the population growth rate (Heppell et al., 2000), so it can help us predict the response of populations to disturbance and decide which life stages should be protected (Heppell et al., 2000; Manlik et al., 2018). Our results indicate that management efforts should focus on protecting the juvenile *S. curriculus*, as this life stage had the highest contribution to the population growth rate (0.485). In the context of mostly overfished fishery resources in China (Chen et al., 2009), these life history characteristics of *S. curriculus* might be advantageous. This may be the explanation for the increase in the relative abundance of this species in recent decades (Wang et al., 2017), with the finite rate of population increase of 1.069 inferred in our study. Although this indicates that *S. curriculus* should be able to withstand the current fishing pressure, excessive overfishing would still be likely to produce negative impacts on this population. At present, a yearly fishing ban from March to June is the main protection and management measure employed in the Pearl River Basin. This measure is primarily aimed at protecting the fish during the reproductive season and early life history stages. According to our results, the contribution of the 0–1-year-old life stage of *S. curriculus* to population growth is merely 0.242, whereas the contribution of the 1–3-year-old stage is much higher (0.485). This indicates that protection (or fishing intensity regulation) measures aimed at juveniles are more likely to produce positive outcomes on population growth and stability than the measures currently employed. Thus, implementing a minimum catch size and/or minimum mesh size is particularly important. In the study area, the minimum catch size should not be less than 200 mm (3 years old), and the minimum mesh size of the fishing net should also be adjusted accordingly.

According to our research, the spawning period of *S. curriculus* is from March to September, which may actually be longer than the fishing ban period mentioned above. Therefore, if this species is to be better protected and managed, the fishing ban period needs to be extended. The unique environment of the estuary has a significant impact on the growth of this species, which directly shows that changes in the aquatic habitat, such as environmental pollution, are likely to affect its growth. In summary, only by formulating reasonable management methods based on the life history characteristics of *S. curriculus* can we ensure the permanent sustainability of this resource.

### DATA AVAILABILITY STATEMENT

The raw data supporting the conclusions of this article will be made available by the authors, without undue reservation.

### ETHICS STATEMENT

The animal study was reviewed and approved by Key Laboratory of South China Sea Fishery Resources Exploitation & Utilization, Ministry of Agriculture and Rural Affairs, South China Sea Fisheries Research Institute, Chinese Academy of Fishery Sciences, Guangzhou, China.
AUTHOR CONTRIBUTIONS

Conceptualization, TW, YL, and C-HL; investigation, TW, Y-YX, LL; methodology, TW, Y-YX; formal analysis, TW; writing original draft, TW, IJ, and LI; writing-review & editing, TW, YL, IJ, C-HL, and PW; data curation, YL, IJ, and LI; funding acquisition, C-HL and PW.

FUNDING

This research was supported by the National Key R&D Program of China (2019YFD0901201, 2019YFD0901204), Guangdong Basic and Applied Basic Research Foundation (2019B1515120065), Science and Technology Planning Project of Guangdong Province (2019B121201001), Key Special Project for Introduced Talents Team of Southern Marine Science and Engineering Guangdong Laboratory (Guangzhou) (GML2019ZD0605), Central Public-interest Scientific Institution Basal Research Fund, South China Sea Fisheries Research Institute, CAFS (2016TS35), Central Public-interest Scientific Institution Basal Research Fund, CAFS (No. 2020TD16), Science and Technology Program of Guangzhou, China (202102080509).

ACKNOWLEDGMENTS

We would like to thank Sun T., Chen D. F., Wei F. S., and Huang H. Y. for their help in collecting the survey data.

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