Trophic Plasticity of Bombay Duck (Harpadon nehereus) in the South-Central East China Sea Based on Stable Isotope Evidence

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High trophic plasticity is often invoked to explain the successful expansion of many aquatic species. Bombay duck (BD) outcompete other traditionally commercial fishes in the East China Sea (ECS) in recent years, displaying a continuous and sharp increase of catches, however, little is known whether the competition advantage of BD was related to its trophic plasticity. Using stable isotope analysis (SIA), we investigated the trophic ecology of BD in the south-central ECS in two periods. Significant differences in δ¹³C and δ¹⁵N values were found between sampling months and among body sizes. Modal (95% credibility limits) estimation showed that BD’s trophic position (TP) gradually increased from 3.56 (3.31–3.82) in G1 (120–150 mm for body size) to 4.09 (3.69–4.50) in G5 (241–270 mm) in November, and from 3.37 (2.92–3.88) in G2 (151–180 mm) to 3.66 (3.07–4.23) in G4 (211–240 mm) in April, respectively, suggesting obvious ontogenetic variation of δ¹⁵N and TP. More importantly, the TP of BD within the same body size showed wide amplitude, indicating BD possess high trophic plasticity, and can capture prey from low to high trophic levels. Further quantitative analysis based on the MixSIAR model showed that BD mainly fed on zooplankton and fish, but its feeding habit was characterized by the consumption of gradually reduced zooplankton and increased fish associated with increased body size. Interestingly, the fish’s contribution to BD showed obvious seasonal variation. It can be explained by varied food availability in ECS. Relatively low fish prey in April forced BD to adjust its diet, adding the ingestion of shrimps and cephalopods corresponding to temporal variations in food availability. The present study provides the evidence that high trophic plasticity exists in the feeding strategy of BD, which is reflected in the ability to adjust the feeding preference according to nutritional needs and food environmental changes; thus, it is competitively advantageous and may partly explain why BD is becoming the predominant species and displays an extensive distribution in the ECS.

Keywords: Bombay duck, stable isotope, East China Sea, trophic level, feeding habits
INTRODUCTION

Trophic plasticity is the adaptation of organisms to minimize potential competition for food by (partly) shifting their diets (MacNeil et al., 1997; Lefebvre et al., 2009; Mavraki et al., 2020). It is a form of trophic generalism, implying that a species is capable of feeding on a broad food spectrum, but only exploit a part depending on the competitive environment in which it occurs (Gutt, 2006; Riera, 2009, 2010). Species could adapt their foraging strategy in relation to variation of the availability of their trophic subsidies (Kopp et al., 2015; Timmerman et al., 2021) and the spatial-temporal variation in the food availability may greatly affect the food ingested and diet breadth, making species exhibit a trophic plasticity (Adite et al., 2013). Such plasticity may originate from changes either in direct predation in relation to food environmental variation such as changes in prey abundance or species competition (Zheng et al., 2014), or in indirect predation such as changes in the diet or trophic level of their prey (Timmerman et al., 2021). On the other hand, fish can adjust their diets with size to optimize their energetic return (Scharf et al., 2000). Ontogenetic shifts can be considered a mechanism to avoid, or at least minimize, intraspecific competition by allowing exploitation of different food resources at each developmental stage (Garica-Rodriguez et al., 2021). Species may shift their diet during ontogenesis and/or there may be changes in resource availability that affect the competitive interactions between them (Winemiller, 1989; Wood et al., 2016). The broader diet spectrum coupled with the trophic plasticity behavior is probably a competitive advantage because it enables species not only to colonize and to adapt to unstable and changing aquatic habitats, but also to invade and to well-establish in various ecosystems (Adite et al., 2013). Evidence suggests that introduced species with generalist diets have the potential for greatest impacts through competition and predation (Cathcart et al., 2019). It is important to well document the diet spectrum variation and the trophic plasticity to explore the species feeding strategies, would give a better insight into the trophic structure dynamic, an important tool and baseline information to conserve, manage and restore fish resources in their habitat (Adite et al., 2013; Garcia-Rodriguez et al., 2021).

Bombay duck (BD, Harpadon nehereus) is a small, inshore, bottom-dwelling predatory fish species, which is widely distributed in the waters of the Indian Ocean and the Northwest Pacific Ocean (Khan, 1989; Ghosh et al., 2009; Lin, 2009; Luo et al., 2012; Firdaus et al., 2017). In China, BD has been regarded as a secondary commercial fish species in the past and is usually caught by coastal nets, so there were no relevant production records available (Lin, 2009). Since the 1980s, the rapid increase in the number of motorized fishing boats has led to an increase in catches year by year, modifies the structure and function of marine ecosystems. Most of the fish species caught in the East China Sea (ECS) have shown characteristics of miniaturization and early maturation, and the resources of some traditional commercial fish species, such as Larimichthys crocea, have declined drastically (Mi, 1997; Lin et al., 2004, 2006; Xu and Liu, 2007; Ma et al., 2019). Under the background of the serious decline in the catch yield in the ECS, BD, as a supplementary fishery resource, is becoming more abundant and gradually becoming one of the main commercial fishes in the ECS (Song et al., 2012; Zhong et al., 2016; Du, 2018).

BD possesses some ecological traits that make it a successful abundant species and ferocious predator in China’s coastal waters. The spawning period of BD is from May to September (Luo et al., 2012), which corresponds to summer fishing closure and ensures the safe development of juvenile individuals. At the beginning of the period, BD moves to inshore waters for spawning and then moves to open sea areas for feeding (Luo et al., 2012). A large mouth split and short intestine allow them to have many choices in the size and species of its prey, ranging from tiny zooplankton to small fish species of the genera Engraulis and Sardinella (Chen et al., 2004; Du, 2018), also suggesting great potential in trophic plasticity for the species. Although limited evidence from stomach content analyses has indicated that BD can switch its diet composition with increased size class, but to date there is little information on BD trophic plasticity (Zhang, 2005; Lin et al., 2010; Pan, 2011; Ghosh, 2014; Du, 2018), some outstanding issues remain. Previous studies have shown that trophic plasticity is to be one of the reasons for the continued high yield of Trichiurus lepturus and the gradual recovery of Larimichthys polyactis in the East China Sea (Yan et al., 2011; Lin and Gao, 2013). BD outcompete other traditionally commercial fishes in the ECS in recent years, displaying a continuous and sharp increase of catches, however, little is known whether the competition advantage of BD was related to its trophic plasticity. Therefore, the trophic plasticity of BD and how the trophic plasticity is reflected in the process of environmental adaptation and growth need to be further determined.

Stomach content analysis (SCA) is a commonly used method in fish biology. However, SCA only provides information on prey that have been ingested most recently. The contribution of digested prey items is neglected even though their assimilation has contributed to the energy budget of an individual (Wolf et al., 2009). Comparing to SCA, Stable isotopic analysis (SIA) can avoid these deficiencies by indicating the relative contributions of prey items or prey characteristics assimilated over a longer time period, and provide a wider understanding of fish trophic ecology (Zorica et al., 2021). SIA has been widely used in understanding the trophic ecology of marine organisms, such as trophic position, biological migration, and food web dynamics (Zanden et al., 1999; Cherel et al., 2008; Match et al., 2010; Nilsson et al., 2011; Rolla et al., 2020). Here, we employed an SIA of carbon and nitrogen to investigate variations in the feeding behaviors and trophic ecology of BD in the south-central East China Sea (scECS). We aimed to test whether BD possesses large trophic plasticity and to preliminarily analyze the possible reasons for the resource gradually increasing of BD in the scECS from the perspective of trophic ecology.

MATERIALS AND METHODS

Description of Study Area

The scECS is one of the most productive sea areas for fishery resources in China. The shallow water depth enables sufficient
light intensity for photosynthesis in continental shelf waters. The front that is formed by the intersection of the Taiwan warm current and the coastal current brings abundant prey and provides favorable hydrological conditions for the fishing grounds in this area (Zheng et al., 2003). These factors make the scECS a suitable ground for breeding, feeding, and overwintering for some commercial fish species (Mi, 1997; Lin et al., 2004). Corresponding to the improvement of fishing equipment and changes in fishing methods, the catch in the ECS experienced three periods: slow growth (1951 to the 1990s), rapid growth (1991 to the 2000s), and a high yield period after the 2000s (Chen, 2004). High-intensity fishing pressure has brought immense economic benefits, but it has also driven a great change in the catch composition in the ECS that is characterized by a decreasing proportion of traditional high-quality catches (such as L. crocea, L. polyactis, and T. lepturus) and an increase in the catch of secondary commercial species (such as Scomber japonicus and H. nehereus) (Zheng et al., 2003; Chen, 2004).

**Sampling Collection and Processing for Stable Isotopic Determination**

The study was conducted in November 2018 and April 2019 in the scECS, and a total of 49 sites were chosen within 7 transects, within a water depth range of 30–200 m (Figure 1). The survey vessel was a local bottom trawler with a total power of 258 KW that was assembled with a fishery cod-end net with a width of 27 m and a mesh size of 20 mm. Trawling at each site lasted approximately 1 h, and the ship’s speeds were maintained at approximately 3 knots. All specimens, including BD and potential prey nekton (i.e., fishes, shrimps, and cephalopods), were collected and frozen (−20°C). In the laboratory, the biomass (g/hour) of nekton samples was calculated for analysis, and the body size (mm) and weight (g) of BD were recorded for all individuals. A total of 42 individuals (in 14 stations) of BD in November was used for SIA and 58 individuals (in 22 stations) in April (Supplementary Table S1). Generally, stable isotope (SI) analysis studies rely on knowledge of isotopic turnover rates (Winter et al., 2019). The metabolic processes in organisms determine the rate at which the isotopic values of a food item are assimilated into body tissue. For fish, muscle tissue was obtained from the dorsal part of fish species. Although there are no studies on the turnover rate of BD, a variable turnover rate might be less than the maximum estimate (85 days). BD dorsal muscle in this study was used to analyze the feeding information in two periods (November 2018 vs. April 2019). For shrimps and cephalopods, the soft body parts (flesh) was collected. Additionally, zooplankton was reported to be an important food for BD (Lin et al., 2010; Pan, 2011); therefore, zooplankton were collected using a plankton net (mesh size of 505 µm) using a vertical tow and held in 0.22-µm filtered seawater for 24 h to evacuate the gut contents. Whole body of zooplankton was used for stable isotope analyses. All samples were freeze-dried at −50°C for 48 h, ground to a fine powder and stored dry until subsequent isotope analysis.

Measurements of stable isotope ratios were performed via continuous flow - elemental analysis - isotope ratio mass spectrometry (CF-EA-IRMS) at the Instrument sharing platform of the Third Oceanographic Institute, Ministry of Natural Resources, CHN, using a vario ISOTOPE cube Elemental analyzer (Elementar Analysensysteme GMBH, Hanau, Germany) coupled to an IsoPrime100 isotope ratio mass spectrometer (Isoprime, Cheadle, United Kingdom). The samples were combusted into CO₂ and N₂ under a high temperature, and the values of ¹³C/¹²C and ¹⁵N/¹⁴N were compared to international standards (Pee Dee Belemnite and atmospheric N₂, respectively).

In this study, IAEA-600 was used as working standard material. ¹³C and ¹⁵N were expressed using the following equation:

\[
\delta^{13}C \text{ or } \delta^{15}N = \left( \frac{R_{\text{Sample}}}{R_{\text{Standard}}} - 1 \right) \times 10^3
\]

*R correspondingly represents ¹³C/¹²C and ¹⁵N/¹⁴N, and the standard deviations (SD) of the ¹³C and ¹⁵N values were generally <0.1 and <0.2‰, respectively.

**Data Analysis**

Bayesian model using the R package tRophicPosition (Quezada-Romegialli et al., 2018) were applied to estimated trophic position (TP). 95% credibility interval of BD was calculated. The Bayesian model includes isotopic variation in the baseline indicator, the consumer and in the trophic discrimination in order to provide a robust estimation of consumer TP at the population level. Bivalves with a relatively long lifespan can integrate the isotopic signature at a time scale and from different basal carbon sources, therefore it was commonly used as baseline organisms for estimating trophic level in coastal systems (Zanden and Rasmussen, 2001; Gong et al., 2017; Lin et al., 2021). However, bivalves were found at only a few sites in the scECS. Thus, zooplankton collected at each site were more suitable to be selected as the baseline, as used in the Baltic Sea and the northern Okhotsk Sea (Gorbatenko et al., 2013; Lesutienė et al., 2014). We used the ¹³N values of zooplankton with particle sizes greater than 505 µm from the study area as the trophic baseline (λ = 2), and assumed a trophic discrimination factor (δ¹³N) of 2.9 ± 0.3‰ (McCutchan et al., 2003). We set the model parameters as follows: baseline = onebaseline, iterations = 20,000, number of chains (n.chains) = 5.

Bayesian mixing models were applied to estimate the proportion of prey source contributions to the consumer based on stable isotope analyses using MixSIAR R package (Stock et al., 2018). MixSIAR uses Markov Chain Monte Carlo method to identify the probability of contributions of food sources to the consumer, taking account of the variances in isotopic composition of food sources, consumers and trophic enrichment factor (TEF). Since there are no available data in literature on TEF for the BD and the available data for other carnivorous teleost varied widely among species and tissues.
We estimated and compared the proportion of prey sources contributions to each group.

One-way analysis of variance (one-way ANOVA) followed by Tukey’s HSD post hoc test was applied to examine the differences in stable isotope composition or TP in different sampling periods or size classes. Normality and homogeneity of variance were examined before data analysis, and the K-W test (Kruskal-Wallis test) was carried out if the above data did not conform to the assumptions of ANOVA. Pearson correlation analysis was applied to confirm the correlation between stable isotope values and body size of BD. An independent sample T test was used to examine the differences in the stable isotope composition of the prey among seasons.

RESULTS

Spatial and Temporal Variation in the Stable Isotope Composition and Biomass of Bombay Duck Prey

In the scECS, the prey was extremely high in biomass, in which planktivorous fishes were predominantly composed of *Bethosema pterotum*, and piscivorous fishes of *H. nehereus* (Table 1). Additionally, the biomass of the prey displayed interspecies and temporal differences. The biomass was significantly higher in November than in April for total nektonic prey and almost all the fishes, but shrimp showed the opposite trend, especially *Solenocera crassicornis*. *Euprymna berryi* and *Abralia*...
multihamata were two abundant cephalopods, and there was no obviously difference between the two sampling periods.

The $\delta^{13}C$ mean values of the prey ranged from $-21.1$ to $-16.2\%$, while $\delta^{15}N$ ranged from 6.6 to 13.0\% (Table 1). Temporal differences in $\delta^{13}C$ and $\delta^{15}N$ values of the prey were detected between sampling periods (Table 2 and Figure 2). There was no significant temporal difference in $\delta^{13}C$ and $\delta^{15}N$ values for zooplankton ($P = 0.663$ and 0.398, respectively), while a significant difference in $\delta^{15}N$ was detected for most of the nektonic prey (Table 2). Although 7 out of 13 species (i.e., B. pterotum, T. lepturus, H. nehereus, Johnius belangerii, Acetes chinensis, E. berryi, and A. multihamata) had more $^{13}C$-enriched values in November than in April, the $\delta^{13}C$ difference between the two sampling periods was minor, most of which were less

### TABLE 1 | Stable isotope signatures (mean ± SD) and biomass (B) of Bombay duck’s prey; $n$ represents the replicates of samples.

| Prey items | Species | November 2018 | April 2019 |
|------------|---------|---------------|------------|
|            |         | $\delta^{13}C$ (%) | $\delta^{15}N$ (%) | B (g/h) | $\delta^{13}C$ (%) | $\delta^{15}N$ (%) | B (g/h) |
| Zooplankton (>505 μm) | | | | | | | |
| Planktivorous fishes | B. pterotum | 20 | $-20.4 ± 0.9$ | 7.0 ± 1.1 | / | 9 | $-20.2 ± 1.1$ | 6.6 ± 1.4 | / |
| | B. maclellandii | 19 | $-18.3 ± 0.5$ | 11.5 ± 0.6 | 45601.4 | 13 | $-21.1 ± 0.8$ | 8.3 ± 1.4 | 12140.1 |
| | A. japonicum | 16 | $-18.3 ± 0.6$ | 11.9 ± 0.5 | 8724.2 | 5 | $-18.9 ± 0.4$ | 9.5 ± 0.3 | 524.5 |
| Piscivorous fishes | T. lepturus | 12 | $-18.5 ± 0.3$ | 12.0 ± 0.9 | 26248.8 | 8 | $-19.1 ± 1.1$ | 9.2 ± 1.1 | 24820.7 |
| | H. nehereus | 10 | $-17.6 ± 0.5$ | 11.5 ± 0.5 | 41040.1 | 21 | $-18.3 ± 0.8$ | 10.2 ± 0.8 | 7684.1 |
| Benthivorous fishes | J. distinctus | 15 | $-16.9 ± 0.7$ | 13.0 ± 0.6 | 1891.3 | 12 | $-17.0 ± 0.4$ | 12.0 ± 0.4 | 393.7 |
| | J. belangerii | 15 | $-16.2 ± 0.3$ | 13.0 ± 0.5 | 2192.6 | 7 | $-16.8 ± 0.5$ | 11.6 ± 0.4 | 269.3 |
| | A. hexanema | 12 | $-17.5 ± 0.6$ | 12.4 ± 0.4 | 1052.1 | / | / | / |
| | O. lacepedii | / | / | / | 8 | $-17.6 ± 0.3$ | 10.8 ± 0.5 | 543.8 |
| Omnivorous fishes | L. polyactis | 49 | $-16.8 ± 0.6$ | 12.5 ± 0.7 | 1448.5 | 16 | $-17.3 ± 0.4$ | 11.3 ± 0.4 | 313.9 |
| | C. capensis | 20 | $-18.9 ± 0.3$ | 11.6 ± 0.5 | 1201.3 | / | / | / |
| | E. cardinalis | / | / | / | 8 | $-18.8 ± 0.7$ | 11.0 ± 0.8 | 1193.2 |
| Shrimps | A. chinensis | 18 | $-17.8 ± 0.5$ | 9.7 ± 0.7 | 873.2 | 9 | $-19.6 ± 0.4$ | 7.2 ± 0.5 | 740.3 |
| | P. tenella | 3 | $-18.7 ± 0.4$ | 10.0 ± 0.1 | 87.6 | / | / | / |
| | P. izumiae | 3 | $-19.0 ± 0.2$ | 10.4 ± 0.3 | 264.1 | 4 | $-19.4 ± 0.6$ | 8.2 ± 1.3 | 478.6 |
| | M. barbata | 3 | $-18.3 ± 0.5$ | 11.1 ± 0.2 | 39.4 | / | / | / |
| | S. crassicornis | 13 | $-17.9 ± 0.9$ | 12.3 ± 0.4 | 651.9 | 4 | $-16.9 ± 0.1$ | 9.1 ± 0.5 | 2010.4 |
| Cephalopods | E. b erryi | 13 | $-18.9 ± 0.7$ | 10.7 ± 0.8 | 387.2 | 7 | $-19.6 ± 0.5$ | 8.3 ± 1.1 | 488.7 |
| | A. multihamata | 41 | $-18.5 ± 0.5$ | 11.8 ± 0.7 | 4216.6 | 6 | $-19.5 ± 0.5$ | 9.1 ± 1.1 | 5348.9 |

Total biomass 151546.1 70016.6

### Figure 2 | Biplots of $\delta^{13}C$ and $\delta^{15}N$ values of Bombay duck and their prey items. Small color dots represent different size classes of Bombay duck and prey items are shown as mean (±SD).
The isotopic composition of BD ranged widely, from −18.6 to −16.0‰ for δ¹³C and 10.7 to 14.0‰ for δ¹⁵N in November and from −20.2 to −16.7‰ for δ¹³C and 8.5 to 13.2‰ for δ¹⁵N in April (Figure 2; Table 3; Supplementary Table S1). There were significant differences in δ¹³C and δ¹⁵N values between November and in April (both P < 0.001). The δ¹³C and δ¹⁵N values in April were much lower than those in November (Table 3). The δ¹³C isoscape showed a gradual decline gradient from the northern to southern in November, while in April, it was completely reversed. For δ¹⁵N, the spatial distribution trend is similar to that of δ¹³C, but not so obvious (Figure 3).

Linear regression analysis showed significant correlations between body size and δ¹⁵N values of BD in November and April (r = 0.584, P < 0.01 for November and r = 0.460, P < 0.01 for April). Similarly, there was a significant correlation between body size and the δ¹³C values of BD in November (r = 0.475, P < 0.01) but not so obvious (Figure 3).

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### Table 2: Temporal variation in stable isotope signatures (δ¹³C and δ¹⁵N) for Bombay duck’s prey in the south central East China Sea.

| Species          | df  | δ¹³C (%) | Comparison | t     | P    | Comparison | t     | P    |
|------------------|-----|----------|------------|-------|------|------------|-------|------|
| Zooplankton      | 27  | −0.441   | 0.663      | Nov. = Apr. | 0.859 | 0.398      | Nov. = Apr. | 0.315 |
| B. pterotum      | 33  | 9.493    | <0.001     | Nov. > Apr. | 8.791 | <0.001     | Nov. > Apr. | 5.776 |
| B. maclelandi    | 29  | 0.174    | 0.884      | Nov. = Apr. | 7.711 | <0.001     | Nov. > Apr. | 3.517 |
| A. japonicum     | 19  | 1.954    | 0.066      | Nov. = Apr. | 14.037| <0.001     | Nov. > Apr. | 4.767 |
| T. lepturus      | 18  | 2.537    | 0.021      | Nov. > Apr. | 6.755 | <0.001     | Nov. > Apr. | 2.026 |
| H. neheusen      | 29  | 4.983    | <0.001     | Nov. > Apr. | 9.094 | <0.001     | Nov. > Apr. | 3.863 |
| J. distinctus    | 25  | 1.026    | 0.315      | Nov. = Apr. | 3.863 | <0.001     | Nov. > Apr. | 1.021 |
| J. belangerii    | 20  | 2.35     | 0.029      | Nov. > Apr. | 5.776 | <0.001     | Nov. > Apr. | 0.859 |
| L. polyactis     | 63  | 1.617    | 0.111      | Nov. = Apr. | 6.028 | <0.001     | Nov. > Apr. | 0.376 |
| A. chinenis      | 25  | 9.315    | <0.001     | Nov. > Apr. | 7.697 | <0.001     | Nov. > Apr. | 0.301 |
| P. izumiae       | 5   | 0.953    | 0.385      | Nov. = Apr. | 2.377 | 0.063      | Nov. = Apr. | 0.026 |
| S. crassicornis  | 15  | −0.735   | 0.474      | Nov. = Apr. | 9.301 | <0.001     | Nov. > Apr. | 0.036 |
| E. berryi        | 18  | 3.615    | 0.004      | Nov. > Apr. | 6.431 | <0.001     | Nov. > Apr. | 0.036 |
| A. muthamata     | 45  | 5.154    | <0.001     | Nov. > Apr. | 7.011 | <0.001     | Nov. > Apr. | 0.036 |

Bold values indicate significant differences (P < 0.05).

### Table 3: Stable isotopic signatures of Bombay duck in different size groups during both sampling periods.

| Season | Group | Size classes (mm) | n | δ¹³C (%) | Range | Mean ± SD | δ¹⁵N (%) | Range | Mean ± SD |
|--------|-------|-------------------|---|----------|-------|-----------|----------|-------|-----------|
| November | G1    | 121–150           | 10 | −18.6−−−−16.7 | −18.6 ± 0.5<sup>a</sup> | 10.7−−−−12.3 | 11.5 ± 0.5<sup>ad</sup> |
|        | G2    | 151–180           | 10 | −18.2−−−−16.8 | −17.4 ± 0.4<sup>ab</sup> | 11.4−−−−13.0 | 12.1 ± 0.6<sup>a</sup> |
|        | G3    | 181–210           | 11 | −17.2−−−−16.4 | −17.2 ± 0.5<sup>ab</sup> | 10.9−−−−13.9 | 12.3 ± 0.9<sup>ab</sup> |
|        | G4    | 211–240           | 5  | −17.9−−−−16.1 | −17.1 ± 0.6<sup>ab</sup> | 11.5−−−−13.3 | 12.5 ± 0.6<sup>ab</sup> |
|        | G5    | 241–270           | 6  | −17.9−−−−16.1 | −16.8 ± 0.7<sup>b</sup> | 12.1−−−−14.0 | 13.0 ± 0.7<sup>c</sup> |
| April  | G1    | 121–150           | 2  | −18.8−−−−18.5 | −18.8 ± 0.1<sup>c</sup> | 9.1−−−−10.3 | 9.7 ± 0.6<sup>c</sup> |
|        | G2    | 151–180           | 30 | −20.2−−−−17.2 | −18.3 ± 0.7<sup>c</sup> | 8.5−−−−12.1 | 10.6 ± 0.7<sup>c</sup> |
|        | G3    | 181–210           | 20 | −19.4−−−−16.7 | −18.0 ± 8.8<sup>bc</sup> | 10−−−−13.2 | 11.1 ± 0.7<sup>ad</sup> |
|        | G4    | 211–240           | 6  | −19.0−−−−17.6 | −18.4 ± 0.5<sup>c</sup> | 10.5−−−−12.4 | 11.4 ± 0.6<sup>c</sup> |

Different letters indicate significant differences in each group (p < 0.05).
FIGURE 3 | δ¹⁵N and δ¹³C isoscape of Bombay duck in scECS.

(Supplementary Table S2). Modal values of TP generally showed significant higher in November than in April (Figure 5).

**Food Items of Bombay Duck and Its Variation**

Mixing models results (Figure 6 and Table 4) indicated that BD mainly fed on zooplankton and fish, but its prey items varied among different size classes. Generally, BD assimilated less zooplankton but more fish with increased body sizes. In November, zooplankton constituted the major diet of BD in small size classes (median estimate = 57.3%, 90% CR = 45.5–67.9% in G1 and median = 46.5%, CR = 35.7–56.6% in G2), and gradually decreased to 30.5% (CR = 14.6–43.5%) in G5, whereas the fishes’ contribution gradually increased from 25.4% (CR = 6.6–42%) to 47.0% (CR = 22.4–68.6%) with size class. The increase trend in zooplankton and the decrease in fish tended to moderate from groups G3 (181–210 mm) to G4 (211–240 mm). The dietary contributions of shrimp and cephalopod to BD among the body size groups were relatively even (16.6–23.3% with CR among 0–46.5%). The CR values increased with body size, which indicated that the estimations of the proportional contributions of multiple diet sources were characterized by more and more uncertainty. Similar variations in BD’s feeding habit were observed in April, but the increase trend of fish and the decrease trend of zooplankton were still approximately linear from G2 to G4 ($R² = 0.991$ for fish and $R² = 0.959$ for zooplankton, respectively). Comparing the prey contribution of the same size class group in November and April, the zooplankton contribution in April was almost at the same level, while the fish contribution was obviously lower (Table 4). The results of mixing model also supported the variation trend of TP in size class and period.

**DISCUSSION**

**Isotopic Variations and Trophic Position of Bombay Duck**

Spatiotemporal stable isotopic variations have been extensively studied in different waters, such as lagoons, estuaries, and coastal waters (Goering et al., 1990; Vizzini and Mazzola, 2003; Doi et al., 2005; Zheng et al., 2019). It has been proven that the stable
isotope composition varies in consumers living in contrasting environments, such as L. polyactis living in near-shore and offshore waters or the macrozoobenthos and fishes living in lagoons in different seasons due to different compositions of organic matter sources (Ji et al., 2011a; Zheng et al., 2020). In inshore areas easily susceptible to runoff, the mixing of terrestrial and marine-originated carbon might result in variations in $\delta^{13}C$ inshore and offshore, while $\delta^{15}N$ may be affected by enrichment in the ocean or proximity to nitrogen land sources (Sherwood and Rose, 2005). In this study, stable isotopic composition of BD displayed a gradient change from north to south on the whole (Figure 3), though completely opposite trend was observed in two seasons. It to some extent might reflect the temporal change of marine environment in the scECS. The scECS is seasonally affected by two water masses, the Zhejiang and Fujian coastal current (ZFCC) characterized by low temperature and salinity, as well as the Taiwan warm current (TWC) characterized by high temperature and salinity. Under the influence of the northerly winds the ZFCC can be extended to the southern Taiwan Strait in winter and spring, while the TWC controlled most of the scECS in summer and autumn. Different marine environmental characteristics can cause differences in stable isotope ratio of food sources since the origins of carbon sources and other nutrients varied (Uzundumlu and Büyükates, 2019), and then affect the stable isotope composition of consumers. Our observation shows that the seasonal variation of water mass is quite consistent with the variation of $\delta^{13}C$. Generally, the $\delta^{13}C$ value in the area affected by TWC is relatively stable, while the area affected by ZFCC is more complicated due to the joint influence of terrestrial-derived carbon and marine-derived carbon. For $\delta^{15}N$, its distribution pattern is similar to $\delta^{13}C$, but the spatial difference is relatively insignificant. This is consistent with other studies in the ECS, for example, L. polyactis and Pampus argenteus (Ji, 2011). This might be related to the similar composition of prey in the scECS. On the other hand, BD mainly fed on zooplankton and fish, while isotope ratios of pelagic consumers are generally more stable (Sherwood and Rose, 2005).

Our results showed that the $\delta^{13}C$ and $\delta^{15}N$ values varied with body size, suggesting ontogenetic variability in BD, as in other fish species (Currin et al., 2003; Rooker et al., 2006; Ji et al., 2011a; Kimirei et al., 2013; Zhang et al., 2015). Such variation in size could be explained by shifts in dietary preference. For example, it has been proven that the consumption of euphausiids and copepods decreases during the growth of small yellow croaker, whereas the consumption of large prey items, such as decapods and fish, increases (Xue et al., 2010; Ji et al., 2011a). Similar results were also found in a stomach content analysis of BD, which may be responsible for the variations in stable isotope composition (Pan and Cheng, 2011). We also noticed the correlation between the body size and $\delta^{15}N$ of BD was a little weak, although significant. That might be due to the spatial variation of BD’s $\delta^{15}N$ (Figure 3). The difference would weaken
the regression relationship. Also, a capacity of transtrophic feeding of BD revealed by wide amplitude of trophic position within the same body size (Figure 5), corresponding to food availability, also intensified the weak link. The δ¹⁵N values of BD would fluctuate largely in the same body class due to the different diet composition. When high-quality food items was lacking, BD will turn to low-quality food as a supplement. Similar phenomenon also exist in other fish species in the ECS, such as P. argenteus (R² = 0.24) and Argyrosomus argentatus (R² = 0.175) (Ji et al., 2011b).

Trophic Position of Bombay Duck

Due to differences in baseline organisms, TEF and assessment methods, it is difficult to directly compare the TP of BD among studies (Huang et al., 2008; Pan and Cheng, 2011; Yang et al., 2013; Du, 2018; Shi, 2018). However, many studies have indicated that the TP of fishes in the East China Sea are gradually decreasing, including BD (Pan and Cheng, 2011; Du, 2018; Ma et al., 2019). With the overfishing of fishery resources in this area, the trend of miniaturization and younger age of fish species is obvious, and this may be the internal reason for the decline in the TP of BD (Lin et al., 2006; Du, 2018). However, more likely, it may be that BD’s prey has changed dramatically over time and seasonally. It has been proven that the main prey of BD changed from fish species of the genera Engraulis and Sardinella to some smaller species that mainly prey on zooplankton, such as B. pterotum and B. mcelwendy (Pan, 2011; Ghosh, 2014; Du, 2018). In this study, we also find significant seasonal variation in TP (Figure 5) due to the increased contribution of shrimp and cephalopod, while decreased of fish (especially for G2 and G3) in April (Figure 6). Considered there is only half of the biomass for nektonic BD prey caught in April (Table 1), this shift of prey contribution is perhaps caused by the reduction in nektonic prey availability. This temporal fluctuation in food availability would force BD to feed more on lower-quality resource in its ontogenic stages in April, leading to a decreasing in TP.

Interestingly, in addition to ontogenetic variation, BD within the same size classes also fluctuates greatly in TP (Figure 5). This indicates a high individual variation of BD and a wide-ranging prey composition for this species, from low to high trophic levels. This may be partly related to its morphological characteristics. A large mouth split makes BD more unique than other carnivorous fishes, giving it more choice in the size and items of prey and resulting in a high trophic plasticity of this species.

Feeding Habits of Bombay Duck and Trophic Plasticity

Zooplankton and fish were the main prey items of BD, which was consistent with previous stomach content analysis (Lin et al., 2010; Pan and Cheng, 2011; Ghosh, 2014). However, the dietary contribution of fish varied between the two methods: median estimate less than 50% with a broad CR 6.3–68.6% for the stable isotope method and 80% for stomach content analysis. In our case, the small size of BD may be responsible for the difference: the body size of BD in this study ranges from 120 to 270 mm, and zooplankton contribute more in its ontogenetic stages. However, stomach contents can only reflect the information of instantaneously captured food items in a short time rather than assimilated food items, which can be captured by the stable isotope method. It is difficult to quantitatively analyze zooplankton because it is more easily digested and absorbed compared to fishes, and the results may be biased toward some indigestible food items (Li et al., 2005).

The diet of BD varied with body size, and such preference has also been proven in other fish, such as L. polyactis, Trachurus japonicus, and T. lepturus (Guo et al., 2010; Yan et al., 2010; Jiang et al., 2013). These shifts are usually associated with body size, depending on feeding ability and energy requirements, and many studies have also confirmed that changes in feeding organs are one of the main factors that cause diet to change with increasing body size (Wang, 2001; Post, 2003; Pan, 2011). For many juvenile fish, zooplankton are the main prey to support their growth. As body size grows, fish gradually develop feeding organs (such as the mouth split), which makes the juvenile more powerful in hunting. At this stage, predators tend to prey on species of large individuals to ensure sufficient energy sources, which is supported by the “optimal foraging theory” (Gerking, 1994). This shift could expand the dietary resources of the population, meet the nutrition demand at different ontogenetic stages, and alleviate competitive pressure when food resources are limited. The body size thresholds that correspond to dietary changes in BD vary in different studies (Pan, 2011; Du, 2018). In our case, the dietary conversion of BD did not occur suddenly but shifted continuously with age class. It is characterized by the consumption of gradually reduced zooplankton and increased fish associated with increased body size. Interestingly, the fish’s contribution to BD stabilized in November when the body size grew to 211–240 mm (G4), it is similar to the result of dietary conversion by stomach content analysis (250 mm; Pan and Cheng, 2011). But fish’s contribution will continue to increase in April, this means that BD developed a diet heavily based on nekton in November,
but low availability forced a change in April. This might be related to cannibalism habit of BD. Studies show that the prey of BD is mainly composed of small BD when its body size is 250–300 mm (Pan and Cheng, 2011), and BD might have increased feeding of its own species in the deficient of other nektonic prey. This might also explains the shrimp
and cephalopod contribution sudden decrease in G4 in April (Figure 6 and Table 4).

A body of evidence has shown that the spatial and temporal differences in the feeding habits of consumers can be affected by the abundance and type of available prey (Sedberry, 2000; Zheng et al., 2014, 2020; Rolla et al., 2020). When the biomass of the main prey items in the habitat decreases drastically, the small-sized BD with poor predatory ability are under greater food competition pressure, and this pressure can be alleviated by strengthening the predation of zooplankton, shrimp and cephalopod. It also further proves that BD has strong trophic plasticity. High trophic plasticity enables consumers to exploit broad food resources and succeed at interspecific competition. For example, marked differences in the diet and trophic position of top mouth gudgeons (Pseudorasbora parva) have been found in neighboring systems only a few kilometers away, and plasticity in food selection makes them a successful invader in European waters (Rolla et al., 2020).

BD was not regarded as a fish species of high economic value in the past, but in the context of the severe decline of traditional fishery resources, the increasing catch of BD re-asserted its commercial value (Lin, 2009; Luo et al., 2012). BD in the ECS is mainly divided into two populations: one is distributed in the northern ECS (30°30′–33°30′ N, 122°30′–126°00′ E), and the other is distributed at 27°00′–28°30′ N, 121°30′–123°00′ E and is regarded as the southern coastal group featuring less abundance than the northern group (Pan, 2011; Luo et al., 2012). However, a survey of fishery resources in recent years showed that BD in the southern coastal group is becoming the dominant species and has an extensive distribution (Song et al., 2012; Zhong et al., 2016; Du, 2018). Short generation and high fecundity have been considered important factors for population expansion (Rolla et al., 2020). BD possesses a series of ecological traits that make it dominant in the southern ECS, such as growing and maturing rapidly and spawning persistently (Pan, 2011). Additionally, the high trophic plasticity of BD may partly explain the success of this species in competition with other fish. Under the multiple influences of environmental changes and fishing pressures, the community structure of the ECS has undergone tremendous changes. It has been proven that some species with higher TLs in the ECS are gradually replaced by some small species that feed on zooplankton (Lin et al., 2006; Dai et al., 2020). In this survey, a variety of small fish species were abundant, such as B. pterotum and Bregmaceros macclellandi, and only T. lepturus, as a potential competitor to BD, has a relatively higher biomass (unpublished data). This variation not only alleviates the pressure on BD from competitors but also expands the range of sources of its prey, which guarantees the necessary space and dietary environments for the population extension of BD. The capacity of BD to quickly adjust its food items in its ontogenetic stages and within the same size class enables BD to be more competitive in their habitats and ensures its nutritional requirements at different ontogenetic stages or environments.

### DATA AVAILABILITY STATEMENT

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

### ETHICS STATEMENT

Animals involved in this research were collected through scientific surveys and were not exposed to any unnecessary pain, injuries or suffering, and no sensitive or endangered species involved in this study. The capture and handling of animal was approved by the Fujian Provincial Department of Ocean and Fisheries (collecting permit LS-100014).

### AUTHOR CONTRIBUTIONS

P-QS, YL, and L-SL contributed to the conception and design of the study. P-QS and H-SZ conducted the field work with sample collection and wrote the first draft of the manuscript. YL and Y-YL analyzed the field survey data. P-QS, X-QZ, and H-SZ analyzed the stable isotopic data. All authors contributed to the manuscript revision and approved the submitted version.

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**Table 4** | Estimates of prey species in Bombay duck diet based on Bayesian mixing models using δ¹⁵N and δ¹³C values.

| Period | Prey item    | G1       | G2       | G3       | G4       | G5       |
|--------|--------------|----------|----------|----------|----------|----------|
|        | Median 5% 95%| Median 5% 95%| Median 5% 95%| Median 5% 95%| Median 5% 95%| Median 5% 95%|
| November | zooplankton | 57.3 45.5 67.9 | 46.5 35.7 56.6 | 39.4 26.4 51.6 | 37.1 20.4 52.1 | 30.5 14.6 43.5 |
|         | fishes      | 25.4 6.6 42.0 | 32.0 11.5 50.2 | 38.2 13.6 58.1 | 40.8 15.3 63.0 | 47.0 22.4 66.6 |
|         | Shrimp-cephalopod | 16.6 0.0 38.3 | 21.2 0.0 43.7 | 22.6 0.0 46.3 | 22.5 0.0 46.5 | 23.3 0.0 46.4 |
| April   | zooplankton | − − −   | 49.1 35.9 63.0 | 39.2 24.6 55.3 | 34.5 0.7 54.9 | − − −   |
|         | fishes      | − − −   | 19.5 6.3 36.9 | 30.6 14.9 48.1 | 38.6 16.6 59.8 | − − −   |
|         | Shrimp-cephalopod | − − −   | 30.8 8.2 52.8 | 30.2 5.9 51.5 | 27.5 0.0 49.5 | − − −   |

Median, 5 and 95% values are credible interval outputs from Bayesian isotope mixing models provided.
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SUPPLEMENTARY MATERIAL

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