Migratory Pattern of the Spotted Scat (Scatophagus argus) in the Mangrove Estuary of the Matang Mangrove Forest Reserve, Malaysia, Estimated by Stable Isotope Analysis

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Abstract
Stable isotope ratios of carbon (δ13C) and nitrogen (δ15N) were determined for the spotted scat (Scatophagus argus) in the Matang Mangrove Forest Reserve in Peninsular Malaysia. The ratios from fish samples along with those obtained from samples of their potential food items (i.e., mangrove leaves in the estuary, coastal phytoplankton, microphytobenthos) revealed wide variations, indicating that S. argus utilizes various food items between the coastal area at the river mouth to the inner mangrove estuary. The carbon isotope ratio in the fish body tended to be lower in fish sampled from the inner estuary as compared with fish at the river mouth. The carbon isotope ratio values of the fish body were relatively high in small fish (<20 mm in total length [TL]) but lower in fish of 20-100 mm TL, and relatively higher in fish >100 mm TL. These findings strongly suggest that small-sized S. argus (<20 mm TL) migrate from the outer coastal area into the mangrove estuary and utilize the mangrove estuary habitat as their nursery grounds before eventually returning to the river-mouth area as their growth progresses.

Discipline: Fisheries
Additional key words: stable carbon/nitrogen isotope ratios, mangrove estuary, fish nursery ground, migration

Introduction
The Matang Mangrove Forest Reserve (MMFR) on the west coast of Peninsular Malaysia is one of the world's best-managed mangrove forests. The mangrove estuary there supports a variety of fishes (at least 94 species of 37 families) (Hui 2009). Many euryhaline fishes are reported to spawn offshore of the MMFR, and the developing larvae and juveniles move or are carried by tidal currents to inshore areas or estuarine waters in the mangrove forest area (Blaber 2007, Gillanders et al. 2003, Sarpedonti & Chong 2008, Yamamoto et al. 2010). Among the diverse fish species occurring in the MMFR, the spotted scat (Scatophagus argus) (Perciformes: Scatophagidae) is highly biomass-dominant (Hui 2009), indicating its prominence in the ichthyofauna composition of the area. The species is an important food fish resource and ornamental aquarium species throughout Southeast Asia (Gupta 2016, Musikasung et al. 2006), including the areas surrounding the MMFR. However, limited information is available on its biological aspects, particularly its migratory pattern relative to growth in the area, including its probable dependency on the mangrove estuary as its feeding and nursery grounds. Data on the species’ dynamics in the mangrove estuary could thus be helpful for conservation and management of the S. argus resource.

Stable isotope analysis of carbon and nitrogen can

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provide useful information on the origin and migration paths of an organism if the isotopic compositions of its prey differ following the organism’s movements to a new habitat (Hobson 1999, Tanaka et al. 2011). More specifically, isotope characterization in fish can help to identify long-term migrant individuals from recent ones by using the isotopic ratios of prey species, since the ratio should reflect the isotopic composition of food assimilated over several weeks to months (Herzka 2005). Hence, stable-isotope data can be compared to distinguish between recent immigrants and individuals that have inhabited the sampling site for a relatively long period (Maruyama et al. 2001, Suzuki et al. 2008).

Given the context above, we thus attempted to estimate the migratory pattern of *Scatophagus argus* in the mangrove estuary of the MMFR by analyzing its dependence on the food sources provided in the mangrove habitat by means of stable isotope ratios of carbon and nitrogen of the fish body and several potential food items.

**Materials and methods**

1. **Study area and sample collection sites**

The Matang Mangrove Forest Reserve (MMFR) (Fig. 1) is the largest single tract of mangrove forest in Malaysia (40,151 ha); it is situated on the northwestern coast of Peninsular Malaysia and measures approximately 52 km in length and 13 km in width. The MMFR is a riverine forest-type mangrove that is inundated during most spring high tides, while the water is mainly confined to channels during neap tides. Figure 1 shows the sample collection sites in the northern part of the MMFR. The depth of the Sangga Besar River basin ranges from 4-6 m (between sites R2 and R6), while the coastal and mudflat areas (at site R1) are shallower (1-3 m deep). The site along the Selinsing River (S1) had a depth of 4-5 m, while the sites in the upstream creek areas (C1-C4) had depths of 2-3 m. Mean water temperature in the study area was 29.6°C (range 26.6-31.1°C). Mean salinity (± standard
deviation) of the bottom water was high as 24.5 ± 1.89 ppt in the outer coastal area (R1), and low as 18.2 ± 7.4 ppt (R5), 22.5 ± 2.1 ppt (S1) and 20.2 ± 2.8 ppt (C4) in the inner estuary area (Okamura et al. 2010) (Fig. 1). Fish were caught using a 2-m-wide otter trawl net or cast net. Fish samples were collected monthly from October 2010 to September 2011 at 11 sites: site R2, situated near the coastal mudflat about 2 km offshore from the mouth of the Sangga Besar River; sites R3 to R6 (distance from R2: ca. 4 to 11 km) located along the Sangga Besar River, the widest river in the estuarine system; site S1 (ca. 6 km) on the Selinsing River where it converges with the Sangga Besar River at the fishing village of Kuala Sepetang; and sites C1 to C4 (ca. 11 to 16 km) located in a creek area. The monthly collections across the sampling sites obtained a total of 1,339 specimens of *Scatophagus argus* (13.0-182.0 mm total length) (Fig. 2). Among these specimens, 132 specimens were used for stable isotope analysis. The total length (TL, mm) and body weight (BW, g) of all specimens were measured before the specific analyses.

### 2. Stable isotope analysis

The 132 specimens above used for stable carbon and nitrogen isotope analysis were frozen immediately after collection. The frozen specimens were thawed and rinsed with distilled water before muscle tissues on the left dorsal side of the body were filleted and then dried in an oven at 60°C for 24-48 h. The tissues were then ground to a fine powder with a mortar and pestle. Fish white-muscle tissue was used for isotope analysis due to its slow turnover rate, which could reflect the isotopic composition of food assimilated by the fish over several weeks to months (Herzka 2005). To eliminate the effect of lipids on the stable carbon isotope measurements, powdered samples were defatted by adding 2:1 chloroform-methanol solution (v/v) and then centrifuged. The defatted samples were oven-dried, and then an aliquot of each sample (~0.8 mg) was put in a tin container for isotope analyses. The carbon and nitrogen isotope ratios (δ¹³C and δ¹⁵N) were determined using a mass spectrometer (DeltaPlus, Thermo Finnigan). Carbon and nitrogen contents of the samples were also determined with an elemental analyzer (Flash EA1112, Thermo Finnigan). The standard reference materials for carbon and nitrogen in the stable isotope analysis were Vienna PeeDee Belemnite (VPDB) and N₂ in atmospheric air, respectively. Results were expressed in standard δ notation, and values were determined based on the following equations:

\[
\delta^{13}C,\text{‰} = \left( \frac{^{13}C}{^{12}C} \right)_{\text{sample}} / \left( \frac{^{13}C}{^{12}C} \right)_{\text{standard, VPDB}} - 1 \times 10^3
\]

\[
\delta^{15}N,\text{‰} = \left( \frac{^{15}N}{^{14}N} \right)_{\text{sample}} / \left( \frac{^{15}N}{^{14}N} \right)_{\text{standard, air}} - 1 \times 10^3
\]

Instrument precision was 0.2‰ for the measurements of both δ¹³C and δ¹⁵N.

### Results

Figure 3 shows dual plots of the organic carbon and nitrogen isotopic compositions of the sampled *Scatophagus argus* together with their potential food items in the MMFR (i.e., phytoplankton from coastal offshore waters, microphytobenthos from sites R2, R5 and C3, mangrove leaves inside the estuary) (data from Okamura et al. 2012). The *S. argus* had an overall mean ± standard deviation (SD) of δ¹³C and δ¹⁵N values of −25.0 ± 2.5‰ and 11.6 ± 1.4‰, respectively, with the

![Fig. 2. Size frequency distribution of *Scatophagus argus* collected from the MMFR](image)

\[n = 1339\]
values ranging widely depending on the sampling site. The differences observed in the δ\textsuperscript{13}C values between the sampling sites are considered to reflect the different carbon sources inherent in the food items available to the fish at each site. Fish sampled from the coastal mudflat area (R2) had the most enriched mean values of δ\textsuperscript{13}C (-19.4 ± 1.0‰) (Fig. 3). In contrast, along the river and in the creek areas (S1/R3-R6 and C1-C4), the δ\textsuperscript{13}C values of S. argus (ca. -19 to -27‰) were between the range covered by the δ\textsuperscript{13}C values of coastal phytoplankton, microphytobenthos, and mangrove leaves (ca. -17 to -30‰), and fish sampled from an upstream creek area (C4) had the most depleted mean value of δ\textsuperscript{13}C (-27.4 ± 2.0‰) (Fig. 3).

The δ\textsuperscript{13}C values of Scatophagus argus collected from the coastal mudflat area at the river mouth (R2) showed the highest mean value (-19.4 ± 1.0‰), and the values tended to decrease with increasing distance of the collection sites from the river mouth toward the inner creek areas along the Selinsing River (S1 with a value of -23.0 ± 3.0‰, C1-C4 and SLG2 with ca. -25 to -27‰) (Figs. 1 and 4). Similarly, the δ\textsuperscript{13}C values of fish collected along the Sangga Besar River showed a decreasing trend with increasing distance of the collection sites from the river mouth (R3 with a value of -24.1 ± 1.6‰, R4-R6 with ca. -24 to -25‰) (Figs. 1 and 4). The overall relationship between the δ\textsuperscript{13}C values of fish collected from each sampling site (C) and the distances from the river mouth for each site (D) was significantly correlated with the following formula: $C = -0.39D - 21.12$ ($R^2 = 0.37, n = 132, P < 0.01$) (Fig. 4).

In relation to size, the δ\textsuperscript{13}C values of S. argus were relatively depleted (ca. -26‰) in specimens sized between 20.0 and 100.0 mm TL, while the values were enriched to -22% in the smallest size class (<20.0 mm TL) and tended to increase in fish >100 mm TL with increasing fish size, measuring up to -20% in fish >160 mm TL (Fig. 5). In the three smallest specimens (13.0-15.0 mm TL) analyzed here, the characteristics of the tholichthys larval stage (i.e., protective sheath-like bony plates covering the head) still remained.

**Discussion**

Chew et al. (2012) estimated that 8-44% of nutrition for juvenile fish in the Matang Mangrove Forest Reserve (MMFR) is derived from mangrove sources, with mangrove-derived carbon occurring in the upper estuary becoming more important. In the present study, the δ\textsuperscript{15}N values of Scatophagus argus collected from the river and creek areas (S1/R2-R6, SLG2 and C1-C4) were more enriched (5-7‰) than those of phytoplankton in coastal waters, microphytobenthos (R2, R5 and C3), and mangrove leaves in the estuary (Fig. 3). These values demonstrate that S. argus is a secondary consumer by adapting the generally accepted trophic enrichment

![Fig. 3. Dual plots of stable carbon and nitrogen isotope ratios of Scatophagus argus and its potential prey items (data from Okamura et al. 2012) collected from the MMFR](image).

![Fig. 4. Relationship between values (%) of stable carbon isotope ratios in Scatophagus argus from each collection site and distances from the river mouth (site R2) to each collection site in the MMFR, Malaysia](image)

Open circle: S. argus (with superscripts indicating sample collection sites), Open square: mangrove leaves inside estuary systems, grey square: coastal phytoplankton collected offshore, closed diamond: microphytobenthos collected from site C3, grey diamond: microphytobenthos collected from site R5, open diamond: microphytobenthos collected at the river mouth (site R2), error bars: standard deviations

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values (2.4% for δ¹³N) (Sugisaki et al. 2013). Although organisms as primary consumers were not analyzed in the present study, S. argus is known to be an omnivore (Gandhi 2002) feeding on various food items including such primary consumers as zooplankton, protozoan, and mysids (Thimdee et al. 2004, Wongchinawit & Paphavasit 2009, Gupta 2016), thus indicating the species niche as a secondary consumer. In addition to the observations above, the δ¹³C values of S. argus observed here covered a wide range, with higher δ¹³C values (ca. −19‰) found in fish collected from the river mouth area (R2) to lower δ¹³C values in fish from the inner mangrove estuary (<−27‰ at C4) (Fig. 3). Moreover, the trend in changes of δ¹³C values by distances of the fish collection sites from the river mouth (Fig. 4) strongly suggests that the δ¹³C values of fish reflected the δ¹³C values of the food items occurring at each collection site, and this consideration is also explainable by the trend of δ¹³C values of potential prey items being higher in coastal/ river mouth areas than in inner estuary areas (Figs. 1 and 3). In addition, Tanaka et al. (2011) reported that the δ¹³C values of mysids (Mesopodopsis, Acanthomysis, and Rhopalophthalmus) and decapods (Acetes spp.) as primary consumers and food items of S. argus were higher in the river mouth area, and that the values became lower towards the upstream of the estuary system in the MMFR. Furthermore, Okamura et al. (2010) revealed that the δ¹³C values of surface sediments, which are the traces of potential food organisms for S. argus, showed a similar trend in the MMFR. This information also suggests that the δ¹³C values of the fish body are dependent on the δ¹³C values of the food items by collection sites with different carbon source origins.

Relatively higher δ¹³C values in fish <20 mm TL and >100 mm TL, and lower values in fish of 20-100 mm TL were observed (Fig. 5). Given that the mature specimens occurred offshore (Gandhi et al. 2014) and were scarcely present in the MMFR (S. Morioka, unpubl. data), S. argus is considered to breed offshore. Planktonic larvae of S. argus reportedly feed on phytoplankton/ zooplankton in surface water (Boonruang et al. 1994, Musikasung et al. 2006, Wongchinawit & Paphavasit 2009), and subsequently the feeding area shifts from surface water to the bottom area during development at the juvenile/adult stages (Gandhi 2002, Gupta 2016). In the present study, the three smallest specimens of 13.0-15.0 mm TL still had the characteristics of the tholichthys stage, that is, pelagic/planktonic larvae with a size range of 6-12 mm TL (Barry & Fast 1992). Given this context, the higher δ¹³C values in fish <20 mm TL (Fig. 5) are thus considered a signature of such prey as coastal phytoplankton having higher δ¹³C values observed in this study (Fig. 3). Since S. argus is consistently a bottom omnivorous feeder after reaching the juvenile stage (Sivan & Radhakrishnan 2011), lower δ¹³C values in fish of 20-40 mm TL (Fig. 5) are considered to be caused by the shift in feeding habitats from offshore surface water to bottom areas in the estuary system as part of the ontogenetic development of fish, and subsequent low δ¹³C values in fish of 20-100 mm TL reflect their food items in the inner estuary having lower δ¹³C values [Fig. 3 and Okamura et al. (2010)]. Moreover, higher δ¹³C values are observed in fish >100 mm TL (Fig. 5), although the trend is slower than that of the decreasing trend in fish of 20-40 mm TL, suggesting that the habitat shift of fish migrating from the inner estuary towards the river mouth area as being due to fish growth. The turnover of isotopic compositions is known to be more rapid in younger (smaller-size) specimens and more extended in larger ones (Hoffman et al. 2007, 2011, Xia et al. 2013). The slower increasing trend in δ¹³C values in larger fish >100 mm TL than the decreasing trend in fish <40 mm TL (Fig. 5) is thus likely due to the slower turnover of isotopic compositions in larger fish during the habitat shift from the inner estuary to the river mouth area.

**Conclusion**

The sizes at sexual maturity of Scatophagus argus were reported to be >140 mm TL for females and >120 mm TL for males, and mature specimens occurred offshore (Gandhi et al. 2014). However, the limited portions of specimens reaching those sizes [4.9% (65 fish) >140 mm TL as female maturation size and 11.4% (152 fish) >120 mm TL as male maturation size] were observed in the MMFR (Fig. 2). In addition, the larvae of
many euryhaline tropical fishes are known to move or be carried into the mangrove estuary of the MMFR from offshore (Gillanders et al. 2003, Blaber 2007, Sarpedoni & Chong 2008). Hence, the above information and findings obtained in the present study suggest the ontogenetic migration of \textit{S. argus}; that is, the species breeds in offshore waters, the planktonic larvae (tolichthys larval stage) are considered to subsequently migrate into the mangrove estuary from outer coastal areas (probably carried by the tidal current), grow in the estuary as a nursery grounds, and eventually migrate back to the offshore water for breeding after reaching maturation size. To confirm this migratory pattern of the species, analysis of gonadal development of \textit{S. argus} is to be made.

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