Feeding on protists and particulates by the leptocephali of the worm eels *Myrophis* spp. (Teleostei: Anguilliformes: Ophichthidae), and the potential energy contribution of large aloricate protozoa

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SUMMARY: The food sources of the leptocephali of the teleostean superorder Elopomorpha have been controversial, yet observations on the leptocephali of the worm eels, *Myrophis* spp. (family Ophichthidae) collected in the northern Gulf of Mexico indicate active, not passive, feeding. Leptocephali had protists in their alimentary canals. Estimates of the physiological energetics of worm eels indicate that large aloricate protozoa including ciliates could provide substantial energy to these leptocephali toward the end of the premetamorphic and metamorphic stages, given the low energy requirements of metamorphosing leptocephali. Global ocean warming will likely force a shift in oceanic food webs; a shift away from large protozoa toward smaller protists is possible. Such a disruption of the oceanic food webs could further compromise the survival of leptocephali.

Keywords: *Myrophis*, leptocephali, feeding, protozoa, ciliates.

INTRODUCTION

The snake and worm eels, Ophichthidae, are the most diverse family of the true eels, Anguilliformes (McCosker, 1997), inhabiting coastal areas of tropical and warm temperate oceans. Genera of the sub-

families Ophichthinae and Myrophinae live in the Pacific, eastern and western Atlantic, and Indian Oceans, and possibly in the Mediterranean Sea, but the genus *Myrophis* has not been reported in the eastern Atlantic Ocean or Mediterranean (Hureau and Monod, 1973). In the western North Atlantic,
the ophichthids spawn over outer continental shelves or near continental shelf breaks (J.J. Govoni, unpublished observations of egg distributions). Metamorphosing leptocephali of the ophichthids are found near shore (Harnden et al., 1999; Miller and Tsukamoto, 2004).

The larvae of the elopomorph fishes, the leptocephali, are exceptional among fishes in their morphology and in their physiological energetics (Miller and Tsukamoto, 2004). The internal and external morphology of leptocephali is different from the typical vertebrate body plan, with a central acellular core matrix of glycosaminoglycan (GAG; Pfeiler, 1999), and a large fluid filled space separating the central core, visceral organs, nerve cord, and central blood vessels from the musculature and integument (Smith, 1984). The energy sources that fuel metabolism and provide growth of leptocephali have been controversial (Smith, 1989).

Several feeding behaviors and food sources for leptocephali have been hypothesized, including pseudo-parasitism (Moser, 1981), dissolved organic matter (DOM) (Pfeiler, 1986; Hulet and Robins, 1989), and particulate organic matter (Pfeiler, 1986; Smith, 1984). No leptocephalus, however, has been found attached to another organism in a parasitic mode of behavior. While the alimentary canal of leptocephali has been reported to be non-functional and occluded (Hulet, 1978), recent evidence indicates that the alimentary canal is fully functional (Tamura et al., 1993; Otake, 1996; Pedersen et al., 2003), and can serve in the uptake of DOM through the endocytotic absorption of seawater containing DOM by the gut epithelium (Otake et al., 1993). Laboratory experiments using DOM-enriched water (Liao and Chang, 2001), however, indicate that concentrations of DOM in the ocean are unlikely to be high enough to meet the nutritional requirements of leptocephali.

There is growing evidence of active feeding of leptocephali both in the ocean and in the laboratory. In the laboratory, late stage leptocephali feed on plankton (Alikunhi and Roa, 1951), squid paste (Mochioka et al., 1993), and shark-egg paste (Tanaka, 2003). In the ocean, leptocephali were found to have ciliates (Otake et al., 1990), larvacean houses (Otake et al., 1993; Mochioka and Iwamizu, 1996), and particulates including copepod fecal pellets (Otake et al., 1990; Otake et al., 1993; Mochioka and Iwamizu, 1996) in their alimentary canals.

Eel leptocephali have peculiar growth stages that register in differing, periodic energy requirements: a period of increasing total length (TL) after yolk absorption; followed by periods of decreasing TL as leptocephali metamorphose to become glass eels. Using tooth morphology and TL, Leiby (1989) classified the growth of ophichthid leptocephali into engyodontic, eurydontic, and metamorphic stages. Engyodontic and eurydontic stages are periods of increasing TL and body mass, while the metamorphic stage is a period of decreasing TL and decreasing body mass (Bishop et al., 2000).

The energy demand of elopomorph leptocephali is low (Bishop and Torres, 1999) owing to the large amount of body mass occupied by the acellular matrix composed of GAG that is subsequently catalyzed during the metamorphic stage of decreasing TL (Pfeiler, 1996). The speckled worm eel, Myrophis punctatus, exhibits little change in the wet-weight specific metabolic rate through the euryodontic and metamorphic stages (Pfeiler and Govoni, 1993).

MATERIALS AND METHODS

Collections and leptocephalus taxonomy

Leptocephali were removed from collections of larval fishes taken from the continental shelf of the northern Gulf of Mexico (Govoni et al., 1983; 1985; 1986; Govoni and Chester, 1990). Day and night collections were taken at discrete depth intervals and preserved in a 5% (volume to volume) mixture of 35% saturated formaldehyde solution in seawater buffered with sodium borate. Leptocephali were identified using the morphological and meristic characters provided by Leiby (1989) and Miller and Tsukamoto (2004).

Alimentary canal content analysis

Complete alimentary canals were removed and cut open, and the contents washed into standard depression slides. The resulting slurries were drawn into cross-sectionally square and optically flat capillary tubes, which were examined and photographed with transmission and interference contrast light microscopy. Slurries containing large particulates were also examined with scanning electron microscopy following Turner (1984). Organisms were frequently difficult to identify owing to their semi-digested state and cellular rupture or distortion owing to preservation (Leakey et al., 1994; Stoecker et al., 1994).
Aloricate ciliates were identified using photographs taken by Lynn and Montagnes (1988a,b), Lindholm and Mörk (1990), Lynn et al. (1991) and Crawford (1993).

RESULTS

Leptocephalus taxonomy

Seven of the nine leptocephali examined were identified as the leptocephali of the speckled worm eel; two leptocephali were unidentifiable to species level, because of deviations in meristic counts from those given in Leiby (1989). One leptocephalus was late euryodontic; the remaining leptocephali were metamorphic.

Alimentary canal contents

Six of the nine leptocephali collected contained chime (amorphous, semi-digested food (sensu Govoni et al. (1983)), or unidentified particulates in their guts (Table 1). Three leptocephali contained identifiable organisms or products of digestion. Two leptocephali contained dinoflagellates. Two contained single, double or clumped ovoid cells, 6 μm in diameter (Fig. 1a), tentatively identified as foraminifers. The following cells were found in different individual leptocephali: unidentified, nucleated cells, 13 μm in diameter (Fig. 1b); a nucleated cell, 20 μm along the long axis, tentatively identified as a cryptophyte (Fig. 1c); an amoeboid cell, 10 μm (Fig. 1d); nucleated and flagellated cells, 20 μm along the long axis (Fig. 1e, f); a ciliated cell, 100 μm along the long axis, tentatively identified as a scuticociliatid ciliate (Fig. 1g); a tintinnid Codonellopsis spp.; and a rotifer. Another leptocephalus contained many apparent polykenetids (Lynn et al., 1991) of possible oligotrich or choreotrich ciliates (Fig. 1h, i). The exact number of aloricate ciliates is uncertain because of their semi-digested state. Another leptocephalus

Table 1. – Contents of the guts of Myrophis spp. leptocephali collected from the northern Gulf of Mexico (IC denotes identification with interference contrast microscopy; SEM denotes identification with scanning electron microscopy).

| Total length (mm) | Taxon          | Examination method | Alimentary canal contents                      |
|------------------|---------------|--------------------|------------------------------------------------|
| 75               | M. punctatus  | SEM                | Unidentified particulates                      |
| 64               | M. punctatus  | IC                 | Unidentified particulates                      |
| 64               | M. punctatus  | IC/SEM             | Unidentified ovoid cells with large nuclei     |
| 58               | M. punctatus  | IC                 | Unidentified ovoid cells                        |
| (incomplete specimen) | M. punctatus  | IC/SEM             | Ciliate polykenetid                            |
| (incomplete specimen) | Myrophis spp. | IC/SEM             | Dinoflagellates                                |
|                  |               |                    | Amoeboid                                       |
|                  |               |                    | Unidentified particulates                      |
|                  |               |                    | 2 ciliate polykenetid                          |
|                  |               |                    | 2 dinoflagellates                              |
|                  |               |                    | Unidentified ovoid cells with oral groove      |
|                  |               |                    | Unidentified ciliate polykenetid               |

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contained a copepod fecal pellet that packaged many bacilli, fragments of centric and chain-forming diatoms (one a fragment of a possible *Ceratium* spp.) and a dinoflagellate tentatively identified as *Gyrodinium* (Fig. 2).

**DISCUSSION**

Aloricate protozoa are not easily recognized and counted in the alimentary canals of fish larvae and are consequently under-recognized as energy sources for fish larvae including leptocephali. Many of the organisms found in the alimentary canal of worm eels (i.e., diatoms, armored dinoflagellates, tintinninds and rotifers) are common to the diets of other larval fishes in the northern Gulf of Mexico, and these organisms are easily recognized by their remnant hard parts that are refractory to digestion. Ciliates have been reported in the guts of a leptocephalus of *Anguilla japonicus* (Otake et al., 1990), but these were identified using transmission electron microscopy and were indicated solely by the microfibrillar intracellular supports of their cilia in the digestive residue; they were not enumerated.

There is evidence from larval fishes collected in the ocean (Ohman et al., 1991; Fukami et al., 1999; Figueiredo et al., 2005; Pepin and Dower, 2007) and from laboratory experiments (Hunt von Herbing and Gallager, 2000) that oligotrich or choreotrich ciliates might be important food sources for larval fishes. Aloricate ciliates are preferred food of fish larvae in the laboratory (Hunt von Herbing and Gallager, 2000). Larval fishes are known to selectively feed by discrete feeding strikes on planktonic prey that they can effectively detect with their sensory systems and that they can easily capture given their behavioral responses to prey and their swimming abilities (Govoni et al., 1986). The leptocephali examined here were from the same collections as other larval fishes examined from the northern Gulf of Mexico; consequently, the availability of ciliates as food for leptocephali was the same. Leptocephali are fully capable of sensing (Hulet, 1978; Okamura et al., 2002; Døving and Kasumyan, 2008) and capturing (Pfeiler, 1989) protozoa, including aloricate ciliates. Ciliates are likely to be easy prey for leptocephali in the ocean and they are eaten readily by other larval fishes (Figueiredo et al., 2007). Aloricate ciliates are a preferred food of fish larvae in the laboratory (Hunt von Herbing and Gallager, 2000).

The energy contribution of large protozoa to the physiological energetics of worm eel leptocephali may be substantial. Two leptocephali examined contained food organisms recognized as aloricate protozoa that were larger, ca. 100-200 µm, and of higher nutrient (carbon and nitrogen) and energy (Finlay and Uhlig, 1981; Stoecker and Govoni, 1984; Crawford and Stoecker, 1996)) content per cell than are dinoflagellates, diatoms, and small particulates.

Without knowledge of daily feeding periodicity and the evacuation rate of the alimentary canal, an accurate estimate of the daily ration of leptocephali is impossible. Examination of the gut contents indicates alimentary canal contents at a moment in time, whereas feeding and digestion are continuous over some unknown period. A rudimentary estimate of
the energy contribution of protozoa is, nonetheless, possible. For example, the 64 mm TL leptocephalus contained large (ciliates and an amoeba) and small (ovoid cells and dinoflagellates) protists. The assumption of five ciliates and one amoeba (large protozoa), and five ovoid cells and two dinoflagellates (small protists), is justified and conservative. With this assumption, and with enthalpy conversions given by Finlay and Uhlig (1981), Stoecker and Govoni(1984), and Crawford and Stoecker (1996) – 44 J mg C⁻¹ and 5 x 10⁻⁶ mg C cell⁻¹ for ovoid cells, 43 J mg C⁻¹ and 3 x 10⁻⁶ mg C cell⁻¹ for dinoflagellates, and 47 J mg C⁻¹ and 1.25 x 10⁻⁴ mg C cell⁻¹ for the ciliates and the amoeba – the total energy available from the gut contents of this leptocephalus would be 0.04 J. Large protozoa, the amoeba and ciliates, account for most of the energy consumed at this one point in time. This energy contribution, while small when compared with the total, daily energy requirement of leptocephali, indicates that these large protozoa can be physiologically substantial for several reasons: because they are likely to be consumed continuously throughout the feeding period of leptocephali; because the overall energy demand of metamorphic worm eels is low; because energy (matter) for growth is provided by the transformation GAG into morphological (structural) components; and because GAG supplies much of the energy for metabolism. The importance of large protozoa, including ciliates that lack lorica, in the diets of leptocephali and other larval fishes, should be assessed comprehensively (Govoni, 2005).

The survival of eels through early life might be compromised by any decline in the availability of large protozoa as food for elopomorph leptocephali. Ciliates are common, abundant and widely distributed (Stoecker et al., 1989; Lessard and Murrell, 1996), and are key components of oceanic food webs (Pierce and Turner, 1992). Ciliate abundance and availability might change owing to global climate change and its impact on the world’s oceans (Miller et al., 2009). A shift in the complexity of primary producers driven by increased temperatures from large diatoms to small dinoflagellates, and consequent shifts in the abundance of heterotrophs and mixotrophs, could change food webs (Friedland et al., 2006; Bonhommeau et al., 2008) in ways that could compromise the availability of large protozoa as food for leptocephali. This would influence negatively the survival of eels through their early life history (Desaunay and Guerault, 1997).

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