INTRODUCTION

Studies of coral reef fish assemblages have often missed or ignored many of the small and/or cryptic species that numerically dominate this habitat (Ackerman & Bellwood 2000). Until recently, this has resulted in their virtual exclusion from investigations of reef ecology. Newly developed evidence, however, suggests that their contribution to reef trophodynamics is far from negligible. Although the so-called cryptobenthic fishes make up the majority of reef fishes, their small size means that they have very low standing biomass (Ackerman & Bellwood 2000, Depczynski et al. 2007). Despite this, they account for a large portion of overall biomass production and overwhelmingly dominate the length production of reef fishes, i.e. the ‘overall’ growth in body length of all fish (as determined in cm m⁻²) per unit time (Depczynski et al. 2007).

Several features of the life history of these small fishes are responsible for this significant contribution to overall productivity. Body size is negatively correlated with several life history features, including mortality, and positively correlated with such traits as lifespan and time of maturation (Blueweiss et al. 1978, Calder 1984, Roff 1992). Recently, Depczynski & Bellwood (2006) characterized the entire life cycle of 3 members of the cryptobenthic fish fauna of the Great Barrier Reef: the pygmy gobies *Eviota sigillata*, *E. melasma* and *E. queenslandica*. These incredibly short-lived fish (*E. sigillata* is the shortest-living vertebrate known, with a maximum recorded age of 59 d) exhibited rapid, linear growth, produced several generations over their exceedingly brief lifespans, and experienced high daily mortality (Depczynski & Bellwood 2006). Species of the Gobiidae make up a large portion of the cryptobenthic community (Depczynski & Bellwood 2003, Depczynski et al. 2007),...
and several other species in this family have short lifespans and similarly high mortalities (e.g. *Istigobius decoratus*, Kritzer 2002; *Valenciennea strigata*, Reavis 1997). These factors combine to produce a rapid turnover of the cryptobenthic community, resulting in its disproportionately large contribution to overall production.

The genus *Trimma* is a speciose group of small (mostly <30 mm standard length), colourful, Indo-Pacific coral reef gobies. Some 58 valid species have been described to date, with a further 30 to 40 remaining undescribed. In Palau, the nasal pygmygoby *Trimma nasa* is similar in size to *Eviota sigillata*, and is one of several western Pacific species in the genus that hover in large, loose schools adjacent to the steep or vertical walls of the outer reef. Members of these schools usually adopt a ‘head-up’ posture and probably pick zooplankton in the water column (Saeki et al. 2005, Winterbottom 2005). As such they are best described as epibenthic, though epi- and cryptobenthic are part of a continuum of reef fish behaviour (Miller 1979). These tiny fish are good candidates for life history analysis because if they show the rapid growth and turnover of other small gobies, their consumption of (probably allochthonous) plankton and presumable role as a food source for larger reef fishes may put them at an important trophic intersection.

This study investigated the age–length relationship, approximate lifespan, timing of settlement and mortality of *Trimma nasa* using information obtained from otoliths. These data will provide a foundation for future work on population structure and biomass production in this and other small but abundant reef fish species.

**MATERIALS AND METHODS**

**Fish collection.** Fish were collected on 10 May 2007 at the Mutremdiu wall, Uchelbeluu Reef (Short Drop Off), in Palau at a depth of 15 m using the anaesthetic quinaldine and a hand-net. Captured fish were immediately placed in 80% ethanol. Within 3 wk of capture, all 110 specimens were sexed based on external genital morphology and their standard lengths were measured to the nearest 0.1 mm. Genital morphology is a good indicator of sex in these types of gobies, but does not necessarily indicate sexual maturity (K. Cole pers. comm.); thus, while we assumed fish with undifferentiated genitalia were sexually immature, not all differentiated individuals had necessarily reached maturity.

**Otolith preparation.** Sagittal otoliths were extracted from all specimens, cleaned, dipped in 70% ethanol and stored dry in plastic culture plates. Each sagitta was mounted, sulcus side down, on a microscope slide with the thermoplastic glue Crystalbond 509™ with the anterior end (but not the core) hanging over the edge. The anterior end was ground on 9 and 3 µm lapping film until level with the edge of the slide, and progress was monitored under a dissecting microscope. The glue was then reheated so the otolith could be repositioned with the ground surface face down. The posterior end was ground to provide a thin section showing the entire sequence of increments from the primordium to the outer edge.

**Age determination.** Sections were viewed at 400× magnification on a computer monitor by means of a Spot Flex model 15.2 camera attached to a Leitz Dialux 22 compound microscope using Spot Diagnostic Instruments software (v4.5.9.5). Increments were counted from the core outward along the longest axis (Fig. 1). These increments were assumed to be deposited daily, as recent studies have confirmed daily increment deposition in 9 species of reef-associated goby (Hernaman et al. 2000, Shafer 2000, Depczynski & Bellwood 2006) as well as several tropical estuarine (Hernaman et al. 2000) and temperate marine (e.g. Iglesias et al. 1997) gobies. In addition, the first increment was assumed to be deposited at hatching. The hatch check, as the first increment is often called, may coincide with hatching, mouth opening or absorption of the yolk sac (Neilson 1992). In several species of *Trimma* and in the
related *Priolepis nocturna*, these events all occur within 24 h (Sunobe 1995, Wittenrich et al. 2007). The location of the settlement mark, an abnormal increment corresponding to the transition from larval to settled life, was also noted when possible. Each otolith was counted twice; if counts differed by >10% of the mean, the otolith was re-counted and the average of the 2 closest counts was used, provided that they agreed within 10%.

**Data analysis.** Linear and power regression lines were fitted to the age-at-length data. A mixed model of the effects of age, sex, and age × sex interaction on standard length was performed using SAS statistical software to investigate sex-specific growth. Hoenig’s (1983) equation was used to estimate the total daily mortality rate from the maximum age of the fish examined as follows:

\[
\ln Z = 1.46 \times 1.01 \ln T_{\text{max}}
\]

where \( Z \) = instantaneous mortality rate and \( T_{\text{max}} \) = maximum age.

**RESULTS**

The mean increment number versus standard length data were fitted with linear and power regression lines (Fig. 2), with resulting coefficients of determination of \( r^2 = 0.8458 \) and 0.8464, respectively. The maximum age was 87.5 d. In the mixed model, the effects of age, sex and age × sex interaction were all significant (Table 1). Tukey’s HSD post hoc test revealed that males were significantly smaller than females and juveniles (\( p < 0.05 \); Fig. 3). The male age versus length relationship was significantly different from those of females and juveniles (\( p = 0.0218 \) and 0.0066, respectively; Fig. 4).

In 60% of otoliths examined, a change in increment morphology that was interpreted as a settlement mark was observed (Fig. 1). It consisted of an abrupt decrease in increment width accompanied by a change in increment contrast and focal plane. Based on the position of this settlement mark, the mean pelagic larval duration (PLD) was found to be 33.9 ± 3.7 d (Fig. 2); this constitutes 38.7% of the maximum recorded lifespan and corresponds to 53.6 d of post-settlement life. Daily total mortality (\( Z \)) obtained from Hoenig’s equation was 4.7%.

**DISCUSSION**

Linear and power regression lines fit the age-at-length data equally well; therefore, more sampling at the extremes of the size spectrum could determine the growth trajectory more accurately. We conducted an ANOVA to look for effects of sex as well as of age; both factors, plus an age × sex interaction term, had significant effects on length. Interestingly, males were smaller than females when corrected for age. Males
also had significantly slower growth than females and undifferentiated juveniles, which suggests that their growth is not strictly linear but instead fits a ‘broken stick’ or ‘breakpoint’ model in which growth slows after maturity (Toms & Lesperance 2003); however, accurate determinations of sexual maturity status based on gonad histology will be necessary to refine our description of the growth trajectory.

Sex-based differences in growth patterns are known in other gobies and often can be explained by the mating system. For example, Hernaman & Munday (2005) reported larger, faster-growing males compared with females in 3 polygynous gobies, and no difference between male and female growth in 2 monogamous species. In addition, Reavis (1997) reported no overall size difference between males and females in the monogamous goby Valenciennea strigata, although pairs tended to be assorted by size.

Trimma nasa’s mating system is not currently known; however, its congener, T. okinawae, is a harem, serial sex-changer (Sunobe & Nakazono 1990). T. nasa females are larger and faster-growing than males, but the sex ratio is only slightly female-biased (1 male : 1.6 females). At present, there is no obvious evidence that T. nasa is hermaphroditic; no individuals with intermediate gonad morphology have been found. The lack of a strongly biased sex ratio and the presence of larger females suggest that polygyny is unlikely, but there is little clear evidence for any particular mating system. Further study may be worthwhile to see whether small body size and short lifespan in fish place constraints on the mating system.

Female growth, unlike male growth, is not significantly different from that of undifferentiated juveniles. Females maintain their pre-reproductive growth rates throughout their lives, seemingly disregarding the usual trade-off between somatic growth and reproduction. This, and their faster growth compared with males, is probably due to selection for larger female size and, thus, greater fecundity (see e.g. Depczynski & Bellwood 2006), but the presence of this selective pressure has not yet been tested.

The settlement mark appeared as a transition from narrower to wider increments, accompanied by a change in increment contrast and focal plane. This transition can be classified as a Type IV settlement mark (Wilson & McCormick 1999). Although we have not verified that this mark corresponds to settlement, Sponaugle & Cowen (1994) and Shafer (2000) reported that a similar mark was associated with settlement in 3 other gobies: Coryphopterus glaucofraenum and Gnatholepis thompsoni in the Caribbean, and Bathygobius coalitus in Hawaii.

The 33.9 d larval life fits within the typical 3 to 6 wk duration reported by Brothers et al. (1983) for 12 reef fish families including the Gobiidae. However, as a percentage of the total lifespan, the PLD of Trimma nasa is remarkably long, i.e. 38.7% of the maximum recorded age. The vast majority of reef fishes for which data exist spend less than 1% of their lives as pelagic larvae (Depczynski & Bellwood 2006). Four small (<30 mm standard length), cryptic reef fish species—3 gobies and 1 tripterygiid—exceed 20%, with Eviota sigillata spending 41.8% of its total life in the plankton (Longenecker & Langston 2005, Depczynski & Bellwood 2006). Just as an increasing amount of data indicate that many reef fishes have short lifespans with linear growth, it is likely that further investigation of PLD compared with total lifespan in small reef fishes will uncover many more species for which the PLD represents a large proportion of the total life. There may be a lower limit on PLD length due to a minimum level of development required for settlement onto the reef.

Trimma nasa experiences 4.7% daily mortality, similar to the values reported for Eviota melasma and E. queenslandica by Depczynski & Bellwood (2006). This value represents natural mortality (M) because T. nasa is not exploited by humans, and furthermore, mortality is expected to be primarily due to predation in small (<50 mm total length) species. Theory predicts selection for early maturity and high reproductive effort when mortality is high (Miller 1996). Experimen-
tal work, such as that of Reznick & Endler (1982), has shown that increased predation selects for earlier maturation and faster growth in fish. Extensions of this work to *T. nasa* and other small reef fishes will help us understand the role mortality plays in shaping their life histories.

In addition to the questions about the consequences of small body size for life history that it raises, this study lays a foundation for further ecological work on small gobies. *Trimma nasa* and its relatives potentially play an important trophic role because, if their food source is primarily allochthonous plankton, they may be importing energy to the reef. While these fishes account for only a small part of the standing biomass (Depczynski et al. 2007), their generation times are probably very short, as in *Eviota* (Depczynski & Bellwood 2006), leading to rapid turn-over and, thus, higher biomass production. A precise, histological determination of age at female maturity — the factor on which generation time mainly depends — has yet to be done for *T. nasa*, but if we use the estimate that female maturation occurs at about 65% of maximum length in fish generally (Charnov 1993), age at maturity is less than 45 d, hinting at a minimum of 5 generations per year. Such a rapid turn-over combined with their potential status as a trophic intermediate and their large numbers on suitable parts of the reef suggest that *T. nasa* may significantly affect reef trophodynamics, out of proportion to their small body size. This has implications for coral reef conservation, arguing for spatial management rather than management on a species-by-species basis.

*Trimma nasa* has many of the life history characteristics of small reef fish: short lifespan, relatively long pelagic larval duration, rapid post-settlement growth and high mortality. Its sex-based differences in growth, apparently unusual among gobies, may be products of its presently unknown mating system. As one of several gobid fish at the extreme small end of the vertebrate body length scale, the evolutionary consequences of its size on its life history and mating system are worth investigating. As an epibenthic planktivore, it may be an important trophic intermediate, importing energy from the pelagic plankton to the reef. As part of the diverse assemblage of small fish that numerically dominate the Uchelbeluu Reef ecosystem in Palau, its influence on reef ecology may be disproportionately large.

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