Distribution of Thaliacea in SW Taiwan coastal water in 1997, with special reference to *Doliolum denticulatum*, *Thalia democratica* and *T. orientalis*

Kwee Siong Tew¹,², Wen-Tseng Lo¹,*

¹National Sun Yat-sen University, Institute of Marine Resources, Kaohsiung, 804, Taiwan, ROC
²Present address: National Museum of Marine Biology and Aquarium, Pintung, 944, Taiwan, ROC

ABSTRACT: Diel vertical migration of Thaliacea was monitored during different seasonal cruises in 1997 in the southwestern waters of Taiwan using samples collected by multiple plankton sampling nets and a 1 m plankton net. Overall, *Doliolum denticulatum*, *Thalia democratica* and *T. orientalis* were the 3 most dominant species in all seasons, and contributed ≥93% of the numerical total of Thaliacea. These species exhibited about a 2 times higher abundance at night than during the day, if all the data were integrated. In the seasonal multiple net samples, *D. denticulatum* and *T. democratica* exhibited an alternation between sexual and asexual reproduction between different sampling seasons, while aggregate zooids of *T. orientalis*, when compared with solitary zooids, dominated in all seasons and apparently increased from July to December. Different stages of these dominant species had different preferences for water depth in different seasons, suggesting that the vertical distribution of these species might be related to reproduction or food. The abundances of different stages of these thaliaceans were correlated with temperature and salinity. The time-series samples of surface water showed that species richness and abundance of these species varied seasonally, but was always greater at night than during the day.

KEY WORDS: Diel vertical migration · DVM · Thaliacea · Season · Taiwan · Western Pacific

INTRODUCTION

Pelagic tunicates are widely but often patchily distributed in the world’s oceans (Thompson 1948, Deibel 1998). Some thaliaceans are found in dense populations at high latitudes (Bathmann 1988, Huntley et al. 1989), or in productive shelf and coastal waters (Wiebe et al. 1979, Zeldis et al. 1995), particularly during the spring phytoplankton bloom (Gorsky et al. 1991, Ménard et al. 1994, 1997). Some thaliaceans are known to perform extensive diel vertical migration (DVM); for instance, *Salpa aspera* migrates over 800 m in the slope water of the western North Atlantic (Wiebe et al. 1979), while *Pyrosoma atlanticum* migrates from 90 to 760 m with different colony sizes in the Ligurian Sea (Andersen & Sardou 1994). Purcell & Madin (1991), however, mentioned that *Cyclosalpa bakeri* has a migration distance of only 60 feet in the subarctic, and Gibbons (1997) found *Thalia democratica* with little or no DVM in the Agulhas Bank, South Africa. Madin et al. (1996) reported that different species seemed to have different DVM patterns. Some hypotheses proposed that DVM serves to reduce visual predation (Pearre 1973, Ohman et al. 1983), to minimize photodamage (Haney 1988), to optimize feeding on algae (Enright 1977), to reduce metabolic costs (Wiebe et al. 1979), or to increase fecundity (Purcell & Madin 1991).

There have been only a few studies of Thaliacea in the western Pacific (Deibel 1998). Chen et al. (1994) re-
ported 8 unnamed species from their seasonal samplings in the waters around the Nansha (Spratly) Islands in the South China Sea. He et al. (1988) studied zooplankton in the coastal upwelling area of Zhejiang, China, and found that up to 16% of the total wet weight biomass of tunicates, primarily Thalia democra-tica, Dolioolum denticulatum and Dolioletta gegen-bauri, was composed of plankton. They are distributed unevenly, increase clearly in the frontal areas, and are negatively correlated with the abundance of copepods and euphausiids. Tsuda & Nemoto (1992) reported 5 species of pelagic salps; T. democra-tica, Salpa fusi-formis, Traustedtia multitentaculata, Cyclosalpa bak-eri and Ritteriella reteacta accounted for 47% of the wet weight of zooplankton in the upper 200 m of a Kuroshio warm-core ring in summer 1987. They found that all salps appeared mainly in the surface mixed layer (0 to 20 m), with the exception of S. fusiiformis, which was most abundant from 50 to 75 m. Fu et al. (1995) found only 1 thaliacean species, Cyclosalpa pin-nata, with an abundance <1 ind. m⁻³ in zooplankton samples collected from the Pearl River estuary, southern China between June 1991 and March 1992. Among these studies, 2 were carried out in coastal areas (He et al. 1988, Fu et al. 1995), while others were concerned with salps in the open ocean or over continental slope. Relatively few of these studies discussed the DVM of salps and even fewer studies have dealt with different life stages of the same species.

The waters of southwestern Taiwan are a typical subtropical area, with different water masses and monsoons alternating seasonally. During the southwest monsoon season from late spring (May) to early autumn (September), warm and wet winds usually blow from the ocean to the land and cause higher precipitation (>100 mm mo⁻¹). The warmer and fresher South China Sea surface waters also flow northeastward, synchronizing with the southwest monsoon, intruding into the southern Taiwan Strait and dominating in this region. At the end of autumn (November) in the beginning of the northeast monsoon, the Kuroshio Current, a warm and highly saline North Pacific current, which originates from the North Equatorial Current east of the Philippines, moves closer towards the east coast of Taiwan and its branches pass through the Luzon Strait and intrude into the northern South China Sea and the waters off southwestern Taiwan (Wang & Chern 1987, Shaw 1992). This results in a specific water layer between depths of 200 and 300 m in this area with an unusually high salinity and poor nutrients (Li et al. 1997, Xu & Su 1997). However, the northeast monsoon, obstructed by the central mountain range of Taiwan, is generally cool and dry when blowing over the study area and normally makes the mean monthly rainfall <50 mm. The alternations of hydrological and climatic conditions in this region may, therefore, greatly influence the distribution patterns and successions of plankton communities as well as the ecosystems on a seasonal scale.

The area under study in this paper has been the subject of many oceanographic studies, both physical (Wang & Chern 1987, Shaw 1989, Hung & Shy 1995, Tseng 2002) and biological (Fong et al. 1990, Soong & Chen 1991, Mok & Kuo 2001), including zooplankton studies (Lo et al. 2001, 2004, Hwang & Lo 2002), but no attention has been paid to pelagic tunicates. We initiated a project to observe such tunicates, which may play an important role in the ecosystem, with the objective of understanding the seasonal variation in species compositions and life stages of Thaliacea in the study area, and to elucidate the relationship between their patterns of vertical migration and hydrographical conditions.

MATERIALS AND METHODS

The sampling station (TS) was located at the Kaoping Trench, about 15 n miles from the mainland and 8 n miles northwest of Liuchiu Island, with a depth of about 500 m (Fig. 1). Zooplankton samples were collected both during the day (11:00 to 14:00 h) and at night (23:00 to 02:00 h) during the cruises of RV ‘Ocean Researcher III’ on May 2–3, July 7–8, October 17–18 and December 27–28, 1997. We used a rectangular opening and closing multiple plankton sampler (Hydro-Bios), with a mouth area of 0.25 m², a mesh size of 330 µm, and a Hydro-Bios flowmeter fixed at the centre of the mouth of each net. The sampler was first lowered to 200 m and opened the first net at this depth. It was towed horizontally at a cruising speed of about 2 knots for about 30 min. Then, the sampler was quickly pulled up to 150 m to open the second net and close the first net simultaneously, and towed as for the first net. This process was repeated at depths of 100, 50 and 0 to 1 m. In total, 38 samples were collected, with the loss of 2 samples due to a mechanical problem with the second net (150 m) in May. Hydrological variables such as water temperature, salinity and fluorescence were measured with CTD probes (SeaBird Electronics) immediately before the plankton tow. The values of fluorescence were modified to the chlorophyll a concentrations (mg l⁻¹) according to the results of concurrent water samplings.

Time-series of bi-hourly samples were taken at the same station (TS), over a 24 h period on March 14–15 and October 26–27, 1997. A 1 m net (1 m diameter opening, 4.5 m long, with a 330 µm mesh size) with a Hydro-Bios flowmeter was towed horizontally near surface water (1 to 5 m) with a cruising speed of about
2 knots for 10 min. After collection, all plankton samples were immediately preserved on board in 5% borax-buffered formalin in seawater. Identification and enumeration of specimens were performed in the laboratory under a dissecting microscope (Askania model GSZ 2) and, when necessary, under a light microscope (Olympus model BX50F-3). Doliolids were identified to species and 3 life stages: gonozooids, phorozooids and oozooids (nurse stage, the older phase of oozooids, was included in oozooids), and salps were identified to species and solitary and aggregate stages. The species identification was based on Thompson (1948), Yount (1954), Chihara & Murano (1997) and Godeaux (1998).

The sampling data, including the filtered water volume, subsample size and number of thaliaceans counted in each sample, are listed in Table 1. The filtered water volume varied from 71 to 671 m$^3$ due to the direction of tow along or against the ocean currents and 3 to 449 thaliacean specimens were counted in each sample. Data of numerical abundance were standardized to ind. m$^{-3}$ (except ind. per 10$^3$m$^3$ in Table 2) before analysis. The weighted mean depths (WMD) (Roe et al. 1984) were calculated to show the preferred depths of different generations of the 3 most dominant species and to evaluate the distance they migrated between day and night. The equation of computation was:

$$WMD = \frac{\Sigma(n_i \times z_i)}{\Sigma n_i}$$

where $n_i$ was abundance (ind. m$^{-3}$) of species at depth $z_i$.

Forward stepwise regression and partial correlation were used to evaluate the relationships between the abundance of different generations of the 3 most dominant thaliacean species and the hydrological variables. To compare the thaliacean abundance during different diel periods, a data matrix of time versus species abundance was built. A Student’s t-test was used to examine the significance of differences in the total abundance and also of the 3 most dominant thaliacean species between day (07:00 to 17:00 h) and night (19:00 to 05:00 h).

**RESULTS**

**Hydrographic conditions**

Vertical profiles of temperature, salinity and fluorescence at the sampling site are shown in Fig. 2. Surface temperature varied slightly among seasons, from 26.0°C in December to 28.8°C in July. The depth of the mixed layer was about 70 m in October and December, and about 20 m in July. No clear thermal stratification was present in May. Salinity in surface water ranged from 33.9 to 34.6‰ during the study period, except in July when it was 32.7‰. In October, there was an unusual decline in salinity at a depth of 70 m, coupled with a sudden peak of chlorophyll $a$ (0.2 µg l$^{-1}$) and the beginning of a thermocline. The chlorophyll $a$ concentration was generally low, except for 1 high value (0.57 µg l$^{-1}$) found near the surface zone in July.

**Day and night vertical distribution**

There were 16 thaliacean species (14 species of salps and 2 species of doliolids) found in the multiple net collections (Table 2). *Thalia democratica* and *T. orientalis* were the most dominant salp species, while the majority of the doliolids were *Doliolum denticulatum*. These 3 dominant species were present in all samples, and contributed to 93% of the numerical total of Thaliacea. Seasonal changes in numerical abundance and species number of Thaliacea were notable, with a peak value, both in numerical abundance and species richness, in October and least abundance in May. Lowest species richness was found in July (Fig. 3). Abundance of total Thaliacea and the 3 dominant species at night was about 2 times as high as during the day when all the data were integrated (Table 2). No apparent difference was observed in species richness between day and night in all sampling seasons, with the exception of October, which showed higher species richness at night (Fig. 3).
Table 1. Sampling data for the multiple plankton net and the 1 m net. FWV: filtered water volume; SS: subsample; AC: no. of animals counted in the subsample; N, D: night and day; –: no data

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Fig. 2. Profiles of environmental parameters at station TS near Liuchiu island, southwestern Taiwan
Doliolum denticulatum was the most dominant species among the thaliaceans at this study site, with an average abundance of 0.7 ± 0.2 ind. m\(^{-3}\), and comprised 62% of the total count of Thaliacea in the 200 m water column. It exhibited apparent seasonal change in abundance, with a peak abundance in October (>1 ind. m\(^{-3}\)), and least abundance in May (Fig. 4). Different life stages of *D. denticulatum* showed different seasonal trends during the sampling periods. The phorozooid stage was the most abundant of all life stages. Its abundance increased from May to October but abruptly declined in December. The abundance of the oozooid stage increased in May, in the subsequent season, and became the most dominant stage (comprising 60% of the total count of this species) in December. The gonozooid stage was the least abundant of all life stages and generally contributed to less than 10% of the total number. Its abundance decreased from May to December.

*D. denticulatum* was a typical epipelagic species that stayed mostly within the upper 50 m of the water column (Fig. 5). Its abundance was greater at night than during the day, with a night:day ratio of 1.9 when the data from the upper 200 m of the water column were integrated (Table 2). If different zooids were considered separately, it was apparent that phorozooids and gonozooids mainly inhabited the surface water, while most oozooids stayed at a depth of 50 m (Fig. 5).

Both solitary and aggregate zooids of *Thalia democratica* showed a clear seasonal change in abundance,
with the greatest abundance in December and the lowest abundance in July 1997 (Fig. 4). Solitary zooids comprised 76% of the total catch of the species in May, and about 27% in December. Aggregate zooids steadily increased from May to December, contrasting with the decline in solitary zooids over the same period. As far as numerical abundance is concerned, both solitary and aggregate zooids increased from July to December, aggregate zooids decreased from December to May, while solitary zooids increased over the same period. Furthermore, the night/day vertical distribution of *T. democratica* varied seasonally but the salps were mainly distributed in the upper 100 m of the water column, both during the day and at night (Fig. 6). The day/night vertical distribution of solitary and aggregate zooids showed different preferences for depths. Solitary zooids were mostly near surface water, particularly in May and December, while aggregate zooids were mainly at a depth of 50 m and had a higher abundance at night.

*Thalia orientalis* showed a similar seasonal distribution to that of *T. democratica* (Fig. 4), having a peak abundance in December, and lowest abundance in July. Aggregate zooids dominated in all samples, accounting for >80% of the species in all seasons. Both zooids of this species were mainly found in the upper 100 m of the water column, and were relatively scarce at depths below 150 m; however, solitary zooids were mainly found at depths between 50 and 100 m, while aggregate zooids were found over the whole sampling depth (Fig. 7). Furthermore, this species showed seasonal differences in the night/day vertical distribution, with higher daytime abundance in May, no clear day/night difference in July, and peak nighttime abundance at depths of 100 and 50 m in October and December, respectively.

Based on the average weighted mean depths (WMD), the 3 most dominant thaliacean species seemed to stay at shallower depths at night, while remaining deeper in the water column during the day (Table 3). Both zooids of *Thalia orientalis* showed significant day/night differences (paired t-test, p < 0.05) in WMD, but no significant differences were found in the other 2 species. This confirms results that the majority of *Doliolum denticulatum* oozooids and solitary zooids of *T. orientalis* inhabit deeper water (59 to 104 m), whereas gonozooids and phorozooids of *D. denticulatum* remain in shallower water (18 to 36 m) (Table 3, Figs. 5 to 7). The night/day ratio of the 3 most dominant species, which averaged about 1.9, implied that the nighttime abundance of these species was about twice that of the day.

The abundance of the 3 dominant species was significantly correlated with the hydrological variables and varied with species and stages (Table 4). The abundances of *Doliolum denticulatum* gonozooids and *Thalia democratica* solitary zooids were negatively and significantly correlated with depth; the abundance of *D. denticulatum* oozooids, and aggregate zooids of
T. democratica and T. orientalis were significantly influenced by temperature. The abundance of D. denticulatum oozooids and phorozooids showed significant correlation with salinity but with a reverse trend, while no significant relationship was found between thaliacean species and fluorescence.

**Time-series samplings within 24 h**

Thirteen species of salps and 2 species of doliolids were identified in the 1 m net samples (Table 2). Doliolum denticulatum, Thalia democratica and T. orientalis were again the 3 most dominant species in both sampling seasons, and together contributed to 93% of the numerical total of Thaliacea. D. denticulatum was the most dominant species in both seasons, with an overall mean abundance of 1.5 ± 0.3 ind. m⁻³ and comprised 55.1% of total Thaliacea. T. democratica and T. orientalis ranked second and third, comprising 19.5 and 18.7%, respectively. Local sunrise occurred at about 06:00 h and sunset at 18:00 h in both seasons; thus, we defined nighttime as 19:00 to 05:00 h and daytime as 07:00 to 17:00 h. We found nighttime abundances of the total thaliacean species to be 3 times more than daytime abundances when all the data were integrated.

In the 24 h time-series collections, 7 species of salps were found in March and 11 species in October 1997, plus 2 species of doliolids in both seasons (Table 2, Fig. 8). In March 1997, total thaliacean abundance steadily increased at night and reached a peak (~7 ind. m⁻³) before midnight, and then decreased to <1 ind. m⁻³ near noon (11:00 h). Lowest abundance was found at 17:00 h. A similar trend was also found in species number (Fig. 8). However, the average abundance of total thaliacean species was not significantly different in terms of diel changes (Table 5). In October 1997, peak abundance was found close to dawn (05:00 h) and the highest species number was found at 03:00 h. This was in contrast to the lowest values, which
were observed at 09:00 h. Average nighttime abundance was significantly higher than during the day (p < 0.05), but species number showed no significant difference between night and day.

The diel period distribution patterns of the 3 most dominant thaliacean species, Doliolum denticulatum, Thalia orientalis and T. democratica were similar; they fluctuated in March and peaked at 05:00 and 03:00 h, respectively in October. There was no significant difference (p > 0.05) in abundance between March and October in these dominant species. The abundance of total thaliaceans between day and night was significantly different in October but not in March. The abundance of D. denticulatum gonozooids and phorozooids in March was significantly different (p < 0.05) between day and night; phorozooid and oozooid stages in October showed significant changes in day/night abundance (p < 0.05, Table 5). Except for the aggregate zooids of T. democratica in October, both solitary and aggregate generations of T. orientalis and T. democratica were generally not significantly different in abundance between day and night.

Fig. 7. Thalia orientalis. Seasonal variation in night/day vertical distribution at station TS near Liuchiu island, southwestern Taiwan

Table 3. Doliolum denticulatum, Thalia democratic and T. orientalis. Average weighted mean depths (WMD) ± SE for both day and night, and mean amplitude of the migration (dm) at Station TS near Liuchiu island in 1997. **, *: day and night differences (paired t-test) in WMDs are significant at the 0.01 and 0.05 levels, respectively. G: gonozooid; O: oozooid; P: phorozooid; S: solitary; A: aggregate

<table>
<thead>
<tr>
<th></th>
<th>WMD (day)</th>
<th>WMD (night)</th>
<th>dm</th>
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<tbody>
<tr>
<td>Doliolum denticulatum</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>G</td>
<td>36 ± 21</td>
<td>23 ± 8</td>
<td>13</td>
</tr>
<tr>
<td>O</td>
<td>78 ± 14</td>
<td>59 ± 5</td>
<td>19</td>
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<tr>
<td>P</td>
<td>24 ± 8</td>
<td>18 ± 3</td>
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<tr>
<td>Thalia democratic</td>
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<tr>
<td>S</td>
<td>53 ± 14</td>
<td>34 ± 18</td>
<td>19</td>
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<tr>
<td>A</td>
<td>59 ± 14</td>
<td>34 ± 15</td>
<td>25</td>
</tr>
<tr>
<td>Thalia orientalis</td>
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<tr>
<td>S</td>
<td>104 ± 4</td>
<td>59 ± 15</td>
<td>45**</td>
</tr>
<tr>
<td>A</td>
<td>59 ± 10</td>
<td>43 ± 13</td>
<td>16*</td>
</tr>
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Fig. 8. Diel period distribution of abundance and species number of total thaliacean species in surface water at station TS near Liuchiu island in 1997. SS: sunset; SR: sunrise
DISCUSSION

Day-night vertical distribution and abundance

Studies on Thaliacea in the western North Pacific Ocean are very sparse; only a few of studies have been published in China and Japan (He et al. 1988, Tsuda & Nemoto 1992, Chen et al. 1994, Fu et al. 1995). The present study was the first to describe species composition and DVM of salps and doliolids in the surrounding water of Taiwan. It is difficult to compare our data with others, partly due to the difference in sampling strategies, and the scales of spatial and temporal variations. However, our results confirm previous studies that both Thalia democratica and Doliolum denticulatum are dominant and widely distributed in the western North Pacific Ocean and adjacent shelf waters. The abundance of Thaliacea in this study was similar to a previous study (Lo & Hwang 2000) in the northern South China Sea, and also to studies by Tsuda & Nemoto (1992) and Fu et al. (1995) in waters adjacent to our study area. The abundance was, however, lower than in the studies reviewed by Deibel (1998) in other geographical areas.

The 3 most dominant thaliacean species, Thalia democratica, T. orientalis and Doliolum denticulatum in this study had higher abundances at night than during the day in the upper 200 m of the water column. They were mostly epipelagic and did not show clear DVM. Different life stages, however, might inhabit different layers of water within the upper 100 m and their abundance varied seasonally (Figs. 5 to 7).

Gonozooids and phorozooids of D. denticulatum mainly lived in shallower water (59 to 78 m), which coincided with lower chlorophyll a concentrations (<0.1 µg l⁻¹). This might be a predator avoiding strategy (Ohman et al. 1983) or a strategy to conserve energy through a decrease in metabolic rate (Wiebe et al. 1979). The distribution patterns of T. democratica in this study agree with previous observations in the Agulhas Bank, where the species displays no vertical migration (Gibbons 1997), as well as in a Kuroshio warm-core ring, where both generations of this species dominate in the upper 20 m of the water column and show no DVM (Tsuda & Nemoto 1992). T. democratica is thought to spawn when chains of aggregates are liberated from solitary zooids in early morning (Heron 1972).

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The shallower occurrence of the solitary zooids of T. democratica...
observed in the present study might be linked to sexual reproduction, similar to observations from the same species in Agulhas Bank (Gibbons 1997) and from *Cyclosalpa bakeri* in the subarctic Pacific (Purcell & Madin 1991). *T. orientalis* has been rarely reported as a dominant species in other studies. Our data show that this species occurs throughout the upper 100 m of the water column and that its solitary zooids are less abundant and inhabit deeper water than aggregate zooids (Table 3, Fig. 7). According to Godeaux et al. (1998), *Thalia* generally has a very short generation time, in the order of days. The dominance of *T. orientalis* aggregate zooids in our samples was probably a result of random samplings during its asexual reproduction. 

Doliolids are mostly epipelagic in the ocean (Paffen-höfer et al. 1995), and are found in or above the thermocline (Ménard et al. 1997). The distribution of *Doliolum denticulatum* is limited by water temperature when it is less than 15°C (Berner & Reid 1961), whereas in the present study the water temperatures above 200 m were higher than 15°C in all seasons; thus, temperature is not likely to be a limiting factor for this species in our study area. There was an indication from our samples that different stages of doliolids were located at different depths. Large proportions of gonozooids (83%) and phorozooids (89%) of *D. denticulata* were caught above the thermocline (~60 m), and mainly in the surface water regardless of the season and time of day (Fig. 5). A similar result was reported by Paffen-höfer et al. (1995) for *Doliioletta gegenbauri* on the middle continental shelf off Georgia and South Carolina: namely, that more than 50% of the gonozooids assembled in the upper part of the water column. They found, however, that the oozooid stage of this species is more abundant (51%) at around 50 m in all seasons and has an overall WMD of 71 m during the day and 59 m at night.

The high abundance and species richness in October in multiple-net collections (Fig. 3) suggested that autumn, at the end of the wet season and during the transition period between southwest and northeast monsoons, might be a more favorable season for most thaliacean species to live and reproduce in this region. However, Ménard et al. (1994) found that *Thalia democratica* and *Salpa fusiformis* were more abundant from April to June in the Mediterranean Sea, and proposed that this might be a result of the spring phytoplankton bloom. Ménard et al. (1997) reported that doliolids in the bay of the northwestern Mediterranean Sea were abundant mostly from July to December and seemed to be temperature-related, while Gibson & Paffen-höfer (2000) further revealed that the feeding and growth rates of doliolids are a function of food concentration and temperature. Our results show that there are more thaliaceans, particularly *Doliolum denticulatum* in October, and their abundance is likely affected by temperature and salinity in this study area (Table 4).

Two dominant thaliaceans, *Thalia democratica* and *Doliolum denticulatum*, displayed alternation of asexual-sexual reproduction in different seasons in this study. In *T. democratica*, a solitary-dominated popula-

![Fig. 9. Doliolum denticulatum, Thalia democratic and T. orientalis. Diel period distribution at station TS near Liuchiu island in 1997. SS: sunset, SR: surese](image-url)
aggregates of

tralia and found that juvenile solitary and mature

growth rate of salps in the Indo-West Pacific near Aus-

ter (80%) (Fig. 4). Heron & Benham (1985) studied the

gate-dominated population in autumn (75%) and win-

tion (80%) in late spring was replaced by an aggre-

gate-dominated population in autumn (75%) and win-

Further studies are needed to examine whether this

change was simply a result of random sampling or

whether it was due to other hydrological effects. The

horozooid stage of D. denticulatum was by far the

most common life stage (~80%) in summer and autumn,

suggesting that asexual reproduction pre-

vailed during these seasons. Oozooids were found in

larger proportions (~55%) in winter, indicating that it

was possibly a suitable season for sexual reproduction

of this species. Similar results were recorded by Bra-

connot (1963) (cited in Deibel 1998) in the Mediter-


in the waters surrounding the Great Barrier Reef, and

Paffenhöfer et al. (1995) off the southeastern USA dur-

ing winter.

Time-series observations

As far as we know, few studies in the literature

address the diel time scale of the migration of Thali-

acea. Atkinson et al. (1978) documented the abun-

dance of doliolids increasing from 7 to 10-fold between

midnight (minimum) and noon (maximum) at a station

off Augustine, Florida, with 3 to 12 h time-interval

samplings in 3 consecutive days. However, their

results did not discuss the species or stage levels and

the mechanism of this diel variability. Andersen et al.

(1992) noted that Pyrosoma atlanticum showed a clear

diurnal symmetry from a time-series of horizontal

hulls around sunrise (02:00 to 08:00 h) and sunset

(15:00 to 20:00 h) at a depth of 230 m in the NW

Mediterranean Sea. The bulk of the population crosses

this depth before sunrise during its descent and after

sunset during its upward migration. From our observ-

ations of 24 h time series samplings, most stages or

generations of these 3 thaliacean species showed

higher abundance at night, but their peak abundance

occurred at different times between 23:00 and 05:00 h,

and varied seasonally. We speculate that these thali-

aceans are epipelagic species that migrate vertically

within a small range to and from the surface on a diel

basis. It is further evident from our results of multiple-

net samples that most of these thaliaceans have shal-

low WMDs both during the day and at night. The

potential metabolic advantage could be important for a

species such as Salpa aspera, which migrated through a

depth of 800 m with a temperature gradient of over

20°C (Wiebe et al. 1979). Metabolic advantages for the

species reported in this study, which migrates through a

temperature range of less than 5°C, if present at all,

would be minimal. Some salmonids, clupeids and

anchovies, which are known to prey on salps, might

affect the DVM pattern of salps (Kashkina 1986, Harbi-

son 1998, Mianzan et al. 2001). However, we did not

have information on any predator in our study area

and were, therefore, unable to assess potential preda-

tion pressures on the behavior of salps. We cannot

eliminate the possibility of predator-caused descent of

salps and doliolids during the daytime in this area.

In conclusion, the 3 dominant thaliacean species,

Doliolum denticulatum, Thalia democratica and T. or-

rentalis, were typically epipelagic in this study site.

They mostly stayed in the upper 100 m of the water col-

umn, both during the day and at night, throughout the

year. Different life stages of these species tended to

stay at different depths of the water column, but most

of them showed no clear DVM despite their abun-

dances at night being usually higher than during the

day. Most thaliacean species in this study showed sea-

sonal changes in abundance, and perhaps alternation

of generation during the year. We speculate, based on

the low thaliacean abundance and chlorophyll a con-

centration in this study area, that these distribution

patterns might be food-limited and reproduction-

related. Furthermore, significant correlations between

thaliacean abundance and water temperature and salinity
could have been influenced by hydrographic

conditions (i.e. alternate intrusions of the Kuroshio

Branch Current and South China Sea Warm Waters) and

southwest and northeast monsoons.

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