Introduction

*Aglantha digitale* is a holoplanktonic hydromedusa distributed through circumpolar arctic waters (Kramp 1959; 1965), and is a common zooplankton component in the subarctic Pacific (Arai & Fulton, 1973; Hirohashi 1988) and Atlantic Oceans (González & González 1996; Søreide et al. 2003). While young *A. digitale* (4–5 mm) ingest diatoms, tintinnids, radiolarians and copepod nauplii, old ones (>5 mm) are typical ambush-predators and feed on copepods and chaetognaths (Smedstad 1972; Hirohashi 1988; Pagès et al. 1996; Colin et al. 2003). Information about predators on *A. digitale* is currently limited; Tsuruta (1963) found this medusa in the stomach of the chum salmon *Oncorhynchus keta*, and Runge et al. (1987) observed ready ingestion of *A. digitale* by the Atlantic mackerel *Scomber scombrus* in the laboratory. Rapid digestion of gelatinous zooplankton in the stomach of fishes may have been masking the importance of this group of animals in the trophodynamics of pelagic systems (cf. Arai et al. 2003). The above results suggest possible trophic importance of *A. digitale* in the pelagic ecosystem of the subarctic Pacific Ocean, but information about their life cycle patterns, as a basis to estimate trophic impacts of this medusa, is currently limited to the population at Station P in the eastern subarctic Pacific (Arai & Fulton 1973).

As part of a research program to evaluate plankton dynamics in the western subarctic Pacific, the present study aims to investigate the abundance, vertical distribution and life cycle patterns of *Aglantha digitale* in the Oyashio region, by analyzing time-series samples collected from 5 discrete depths between 0 and 2000 m. The present results are compared with those in the eastern subarctic Pacific and other regions.

Methods

Monthly depth-stratified sampling was done in the Oyashio region (41°30’ to 42°30’N and 145°00’ to 146°00’E) off southeastern Hokkaido (hereafter referred to as Site H, Fig. 1). Zooplankton were collected with a closing net (60 cm mouth diameter, 100 μm mesh size; Kawamura 1968) equipped with a Rigosha flow-meter in its mouth ring and a TSK depth distance recorder (Tsurumi Seiki Co. Ltd.) or RMD depth meter (Rigosha Co. Ltd.) on its suspension cable. The net was towed vertically at a speed of 1 m s⁻¹, usually through five discrete depths: 0 (surface) to the bottom of the thermocline (BT), BT–250 m, 250–500 m, 500–1000 m and 1000–2000 m (Table 1). During the winter season when the thermocline was not recognized, BT was assumed arbitrarily to be at 100 m depth. After collection, zooplankton samples were immediately preserved in 5% formalin-seawater solution buffered with borax.

**Abstract:** Analyses of monthly depth-stratified (0–2000 m) sampling data in the Oyashio region have revealed that *Aglantha digitale* is distributed largely in the upper 250 m throughout the year. Population biomass ranged from <30 mg DW m⁻² (January) to 1575 mg DW m⁻² (August), with an annual mean of 186 mg DW m⁻². Judging from seasonal sequences of the population structure, the generation length of *A. digitale* is considered to be one year with their spawning season in June–September, which is consistent with that observed at Station P in the eastern subarctic Pacific.

**Key words:** *Aglantha digitale*, vertical distribution, life cycle, Oyashio, North Pacific

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Temperature and salinity profiles were determined with a CTD system at each zooplankton sampling. Chlorophyll a profiles at Site H have been reported elsewhere (Kasai et al. 2001). In the land laboratory, *Aglantha digitale* was sorted from the entire samples and counted under a dissecting microscope. Bell height (BH; mm) was measured under the dissecting microscope to the nearest 0.1 mm. Specimens with “sausage” shaped gonads greater than 10% of BH were designated as “mature”, and those ≤10% of BH as “immature” in this study (cf. McLaren 1969). In order to estimate the population biomass in terms of dry weights (DW; mg), BH data for each specimen was converted to DW data by using an allometry equation: $\log_{10}\text{DW} = 0.454(\log_{10}\text{BH})^2 + 1.883\log_{10}\text{BH} - 2.402$ ($r^2=0.977$, $N=41$, $p<0.001$; Takahashi & Ikeda unpublished data).

**Table 1.** Zooplankton sampling data at Site H in the Oyashio region (N: nighttime, D: daytime, Os: TS “Oshoro Maru”, Ho: TS “Hokusei Maru”, Hs: RV “Hokusin Maru”, Hk: “Hokko Maru”, Ts: RV “Tansei Maru”)

<table>
<thead>
<tr>
<th>Sampling date</th>
<th>Time</th>
<th>Ship</th>
<th>Discrete sampling depths (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>19 Sep. 1996</td>
<td>N</td>
<td>Ho</td>
<td>0–30, 30–250, 250–500, 500–1000, 1000–1500</td>
</tr>
<tr>
<td>8 Dec. 1996</td>
<td>D/N</td>
<td>Hs</td>
<td>0–80, 80–250, 250–500, 500–1000, 1000–1700</td>
</tr>
<tr>
<td>13 Jan. 1997</td>
<td>D</td>
<td>Hk</td>
<td>0–50, 10–500, 300–1700</td>
</tr>
<tr>
<td>20 Feb. 1997</td>
<td>N</td>
<td>Hs</td>
<td>0–100, 100–250, 250–500</td>
</tr>
<tr>
<td>17 Mar. 1997</td>
<td>N</td>
<td>Hk</td>
<td>0–100, 100–250, 250–500, 500–1000, 1000–2000</td>
</tr>
<tr>
<td>11 Apr. 1997</td>
<td>D/N</td>
<td>Ho</td>
<td>0–100, 100–250, 250–500, 500–1000, 1000–1500</td>
</tr>
<tr>
<td>7 May 1997</td>
<td>N</td>
<td>Hk</td>
<td>0–100, 100–250, 500–1000, 1000–2000</td>
</tr>
<tr>
<td>4 Jun. 1997</td>
<td>N</td>
<td>Os</td>
<td>0–80, 80–250, 250–500, 500–1000</td>
</tr>
</tbody>
</table>

**Fig. 1.** The Oyashio region in the western subarctic Pacific Ocean (a) and the sampling site (“Site H”, shaded)(b). Depth contours (200, 1,000, 3,000, 5,000 and 7,000 m) are superimposed in (b).
Results

Site H in this study is located near the southern end of the southwestward alongshore flow of the Oyashio Current. Over the study period, surface temperatures ranged from 2°C (March to April 1997) to 18°C (September to October 1996 and 1997) (Fig. 2). Oyashio Current water, characterized by salinities from 33.0 to 33.3 and temperatures below 3°C (Ohtani 1971), occurred in the upper 150 m from February to April 1997. After April, less saline, seasonally warmed water (possibly originating from the Okhotsk Sea; T. Kono pers. comm.) occupied the upper 50 m of the water column. Surface temperatures above 10°C were observed in September to November 1996 and in June to October 1997, when the thermocline was well established at 20 to 50 m in the water column. Effects of warm core rings originating from the Kuroshio extension were seen in September in both 1996 and 1997, and from December 1996 to January 1997, as judged by temperature at 200 m (>4°C) and salinity in the 0 to 200 m layer (>33.5) (Fig. 2). The temperature and salinity in the 200 to 1500 m layer were stable and nearly constant at 2 to 3°C and 33.5 to 34.5, respectively, throughout the year. Phytoplankton biomass, estimated in terms of chlorophyll a concentrations, showed a marked seasonality (Fig. 2). Chlorophyll a at the surface was around 0.4 mg m⁻³ from August 1996 to the end of February 1997, and increased rapidly to >1 mg m⁻³ in mid March forming an annual peak of >9 mg m⁻³ in May 1997. The surface chlorophyll a concentrations had decreased to 2 mg m⁻³ by the end of June and to 0.4 mg m⁻³ toward the end of 1997. Chlorophyll a concentrations were <0.4 mg m⁻³ below 100 m depth throughout the year. *Aglantha digitale* occurred throughout the year, with a mean abundance of 211 specimens m⁻² over the entire study period (Fig. 3). The abundance during the period of December 1996 and May 1997 was low (55–259 specimens m⁻²), showing a consistent decreasing pattern with progress of the seasons. After that, the abundance increased rapidly, with an annual peak (896 specimens m⁻²) in August 1997. The seasonal change in population biomass paralleled that of abundance. The maximum was 1575 mgDW m⁻² in August, and the minimum <30 mgDW m⁻² in January, with a mean biomass of 186 mgDW m⁻².

Differences in day/night vertical distribution patterns of *A. digitale* on 4 September and 8 December 1996, and 11 April and 5 October 1997 (cf. Table 1) were all not significant (Kolmogorov-Smirnov tests, p>0.1, data not shown). On this basis, only the night data for these sampling days were used to analyze seasonal changes in vertical distribution patterns (Fig. 4). All mature *A. digitale* were found in the top 250 m throughout the year. Part of the immature population resided below 250 m in early September (41%) and December (41%) in 1996, and January (92%) and October (66%) in 1997. It is noted that the broad distribution exceeding 1000 m depth in January 1997 may be an artifact due to the failure of stratified samplings (cf. Table 1).

The entire range of BH (1 to 23 mm) of *Aglantha digitale* was divided into 1 mm increments (Fig. 5). The population structure of *A. digitale* was characterized by the predominance and salinity in the 200 to 1500 m layer were stable and nearly constant at 2 to 3°C and 33.5 to 34.5, respectively, throughout the year. Phytoplankton biomass, estimated in terms of chlorophyll a concentrations, showed a marked seasonality (Fig. 2). Chlorophyll a at the surface was around 0.4 mg m⁻³ from August 1996 to the end of February 1997, and increased rapidly to >1 mg m⁻³ in mid March forming an annual peak of >9 mg m⁻³ in May 1997. The surface chlorophyll a concentrations had decreased to 2 mg m⁻³ by the end of June and to 0.4 mg m⁻³ toward the end of 1997. Chlorophyll a concentrations were <0.4 mg m⁻³ below 100 m depth throughout the year. *Aglantha digitale* occurred throughout the year, with a mean abundance of 211 specimens m⁻² over the entire study period (Fig. 3). The abundance during the period of December 1996 and May 1997 was low (55–259 specimens m⁻²), showing a consistent decreasing pattern with progress of the seasons. After that, the abundance increased rapidly, with an annual peak (896 specimens m⁻²) in August 1997. The seasonal change in population biomass paralleled that of abundance. The maximum was 1575 mgDW m⁻² in August, and the minimum <30 mgDW m⁻² in January, with a mean biomass of 186 mgDW m⁻².

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nance of small (<4 mm BH) immature specimens in September-December 1996, and by medium-sized (4–11 mm) immature specimens in January–May 1997. In June 1997 onward, large (>11 mm) mature specimens increased and occupied the major portion of the population. In October 1997, the large mature specimens disappeared and the small specimens occurred again. The minimum and maximum BH of the specimens recorded in this study was 1.4 mm and 22.5 mm, respectively.

**Discussion**

The annual range of the abundance of *Aglantha digitale* recorded at Site H (55–896 specimens m$^{-2}$) is in the same order of magnitude to that (10–500 specimens m$^{-2}$) observed at Station P (Arai & Fulton 1973), although the latter was based on the sampling of 0–150 m water column and specimens less than 5 mm were not taken into account. In the central subarctic Pacific, isolated abundance data (150–1500 specimens m$^{-2}$, 0–150 m depth) of *A. digitale* in June-August have been reported by Hirohashi (1988). In terms of population biomass, information comparable to the present results (annual mean: 186 mg DW m$^{-2}$) is only that (225 mg DW m$^{-2}$) for *A. digitale* in Toyama Bay studied by Ikeda & Imamura (1996).

The vertical distribution of *Aglantha digitale* was largely limited to the upper 250 m at Site H throughout the year with a few exceptions of extended distribution down to 1000 m (Fig. 4). Recent observations by manned submersibles in the Kurile-Kamchatka region (near Site H) in July-August are consistent with the present results (e.g. the occurrence from the upper 200–250 m) (Vinogradov & Shushkina 2002). In adjacent seas of the subarctic Pacific, the bathymetric distribution of *A. digitale* has been reported as in the upper 200 to 700 m, depending on the season, in the Japan Sea (Zenkevitch 1963; Ikeda & Imamura 1996), and the upper 200 m (no sampling below 200 m depth) in the summer of the southern Bering Sea (Hirohashi 1988). In the Northeastern Atlantic, Williams & Conway (1981) studied the vertical distribution of this medusa over one year, and found that most of both smaller (1–3 mm BH) and larger specimens (3–12 mm) were in the top 100 m in May to August, and the latter extended distribution down to 500 m in July (no sampling was made at >500 m). These regional similarities and dissimilarities in the vertical distribution patterns of *A. digitale* may be interpreted as a result of complex attributes of physical conditions of the upper layers, decrease of food zooplankton abundance and possibly passive sinking behavior of this medusa in response to these environmental conditions (cf. review by Arai 1992).

Judging from the occurrence of mature specimens followed by smaller specimens (Fig. 5), the major spawning
The season of *Aglantha digitale* is deduced to be June-September at Site H. At Site H, June-September corresponds to the period of high abundance of young copepods of grazing copepods such as *Metridia pacifica* (Padmavati et al. 2004), *Neocalanus* spp. (Kobari & Ikeda 1999, 2001a, b), and *Eucalanus bungii* (Shoden et al. 2005), all of which grew rapidly in the upper layers through the phytoplankton bloom in mid March-June. Assuming that *A. digitale* is repeating the same life cycle pattern every year, the present results (Fig. 5) show that the new generation of *A. digitale* created in a given year overwinters and reaches maturity to spawn in June-September of the next year (e.g. annual life cycle), although tracing growth trajectory of the overwintering population is not very clear because of scatter in the size frequency data.

The comparison of the present results for *Aglantha digitale* at Site H with those at Station P in the eastern subarctic Pacific is of special interest to investigate possible regional variation in life cycle patterns of this medusa, as was made for large/medium grazing copepods such as *Neocalanus* spp. (Kobari & Ikeda 1999, 2001a, b), *Eucalanus bungii* (Shoden et al. 2005) and *Metridia* spp. (Padmavati et al. 2004). The annual range in surface temperatures at Station P is narrower (6–14°C; Figure 6 in Miller et al. 1984) than that at Site H (2–18°C, cf. Fig. 2). Station P is warmer than Site H in general; temperatures as low as 3°C throughout the year are seen only below 1000 m in the latter. At Station P, phytoplankton biomass does not show clear seasonality throughout the year, with peaks below 200 m in the latter. At Station P, phytoplankton biomass in the 0–2000 m water column mentioned above. According to Araki & Fulton (1973), this wet biomass of epipelagic zooplankton in the Oyashio region is equivalent to 2,070 mg DW m⁻² (115×0.12×150) in the top 150 m water column, which is 11 times greater than that (186 mg DW m⁻²) of *A. digitale* in the 0–2,000 m water column mentioned above. According to Araki & Fulton (1973), *A. digitale* at Station P spawn in June-September, and resultant offspring overwinter and spawn the next year (annual life cycle). The maximum size specimens attained was 20 mm BH. Excepting this smaller maximum size (20 mm BH as compared with 22.5 mm BH of Site H specimens), the life cycle patterns of *A. digitale* at Station P are remarkably similar to those at Site H mentioned above.

Despite dissimilar environmental conditions between Site H and Station P mentioned above, similar life cycle patterns seen for *A. digitale* living in these two habitats may be interpreted as a result from opposite effects of temperature and food abundance. That is, the negative effect of lower temperatures on the development of *A. digitale* at Site H is compensated by the positive effect of higher food concentrations in the same region. As with *A. digitale*, no regional differences have been reported in the life cycle and associated ontogenetic vertical migration patterns of some grazing copepods (*Neocalanus* spp.) inhabiting Site H and Station P, but the body sizes of specimens at Station P were smaller than those at Site H (Kobari & Ikeda 1999, 2001a, b) as was the case for *A. digitale* mentioned above.

One generation per year of *Aglantha digitale* at Site H and Station P in the subarctic Pacific is also the case for the populations in the North Sea (Kramp 1927) and in the Northeast Atlantic (Williams & Conway 1981). However, more diverse generation numbers have been documented for local *A. digitale* populations; including two generations per year in Toyma Bay (Ikeda & Imamura 1996), Ogac Lake of Baffin Island, (McLaren 1969) and the Strait of Georgia (Araki & Mason 1982), or even three to four generations per year in a Norwegian fjord (Smedstad 1972) and off Plymouth (Russell 1938). Not only the number of generations, but the size at maturity for *A. digitale* has also been reported to be highly variable between different habitats; 4.3 mm for the population in Ogac Lake, 19 mm for those living its neighboring open sea (McLaren 1969), and 17 mm in Toyma Bay (Ikeda & Imamura 1996). While temperature and food abundance were considered to be environmental parameters affecting the life cycle features of *A. digitale* in the subarctic Pacific in this study, broad analyses across all the diverse habitats of this medusa should be done to confirm these and identify new environmental parameters of consequence.

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