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Citation
Journal of the Marine Biological Association of the United Kingdom, 97(2): 387-399

Issue Date
2017-03

Doc URL
http://hdl.handle.net/2115/67082

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Type
article (author version)

File Information
goshimaー1.pdf

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Life history traits and population dynamics of invasive ascidian, _Ascidiella aspersa_, on cultured scallops in Funka Bay, Hokkaido, northern Japan

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ABSTRACT

The European sea squirt, *Ascidiella aspersa* was first found as an alien species in 2008 from Funka Bay, Hokkaido, northern Japan, causing serious damage to the scallop aquaculture industry. We investigated *A. aspersa* on cultured scallops and larval occurrence from July 2010 to June 2014 to clarify life history traits and population dynamics, and consider the relation between the life history of *A. aspersa* and the process of scallop aquaculture. Larvae of *A. aspersa* were found from June to December, and recruitment on cultured scallops occurred mainly between July and October. The ascidians grew well and their weights increased until February. We found that 60–80% of *A. aspersa* that had settled in summer had eggs or sperm in autumn, and 90–100% of *A. aspersa* matured early the following summer. Maturity size in September was 17–20 mm as male, 22–24 mm as female. Scallops in Funka Bay are hung in the spring and harvested from winter to the next spring. *Ascidiella aspersa* settle as larvae in early summer, and grow well until winter, resulting in overgrowth on scallops in the harvest season. The linking of the process of scallop aquaculture and the life history of *A. aspersa* explains why this invasive ascidian has caused serious damage to the aquaculture industry in the bay. In comparison to the earlier descriptions of the native population, *A. aspersa* in Funka Bay has longer reproductive and growth periods, earlier initiation of reproduction, and possibly smaller maturity size.

Keywords: invasive ascidian, *Ascidiella aspersa*, life history traits, population dynamics, aquaculture, scallop
Invasive ascidians have recently become a worldwide issue in coastal waters (Whitlatch & Bullard, 2007; Locke & Carman, 2009). More than 60 non-indigenous ascidians have been recorded in tropical and temperate environments (Shenkar & Swalla, 2011). Non-indigenous ascidians have a rapid growth rate, short life span, and produce large numbers of short-lived non-feeding planktonic larvae. These characteristics, combined with the lack of significant predators, allow ascidians to be successful invaders (Shenkar & Loya, 2009). Ascidians can be strong spatial competitors and, once they become established, often experience population explosions that can develop into dense stands or mats that overgrow and cover available surfaces (Whitlatch & Bullard, 2007). A recent increase in shellfish aquaculture facilities has provided new surfaces (ropes, nets, cages, and shellfishes) for colonisation by invasive ascidians, resulting in overgrowth and smothering of the shellfish (Lambert, 2007). For instance, heavy fouling by cryptogenic species, *Ciona intestinalis* (Linnaeus, 1767), was associated with higher mussel mortality and lower overall size in Nova Scotia (Daigle & Herbinger, 2009). In addition, even if the ascidians have no negative effects on the bivalves directly, removal of the invasive species is costly and requires additional labour by aquaculturists (Carman *et al*., 2010).

The mussel aquaculture industry has been overwhelmed by extremely large numbers of the invasive ascidian *Styela clava* Herdman, 1881 in Prince Edward Island (Bourque *et al*., 2007), resulting in increased production costs estimated at $4.5 million per annum (Shenkar & Swalla, 2011). In Japan, some non-indigenous ascidians have been reported, such as *Molgula manhattensis* (DeKay, 1843) and *Polyandrocarpa zorritensis* (Van Name, 1931) (Tokioka & Kado, 1972; Nishikawa *et al*., 1993).
However, no significant effects of invasive ascidians had been noted on the ecosystems or fisheries prior to the appearance of *Ascidiella aspersa* (Müller, 1776) (The Plankton Society of Japan and The Japanese Association of Benthology, 2009; Kanamori et al., 2012; Nishikawa et al., 2014). The European sea squirt, *A. aspersa*, is a solitary marine and estuarine ascidian that is native from Norway to the Mediterranean (Berrill, 1950; de Kluijver & Ingalsuo, 2004; Mackenzie, 2011). The species has been introduced to North and South America, India, Australia, New Zealand, South Africa, South Korea, and Japan (Brewin, 1946; Kott, 1985; Nagabhushanam & Krishnamoorthy, 1992; Carlton, 2000; Robinson et al., 2004; Tatián et al., 2010; Kanamori et al., 2012; Pyo et al., 2012; Nishikawa et al., 2014). Because there are no efficient predators, *A. aspersa* can form large populations and subsequent high amounts of biomass, which redirects energy to decomposers and not to higher trophic communities (Currie et al., 1998). In addition, colonisation by *A. aspersa* reduces available substrata on which other species recruit successfully (Osman & Whitlatch, 2000). These characteristics have the potential to significantly affect species composition, reducing overall biodiversity (Mackenzie, 2011). *Ascidiella aspersa* also competes directly with other native filter-feeders, including economically important species such as scallops, mussels, and oysters (Currie et al., 1998). Therefore, *A. aspersa* is listed in the Global Invasive Species Database (2010), which is managed by the International Union for Conservation of Nature and Natural Resources, to increase awareness and to facilitate effective prevention and management activities.

The Japanese scallop, *Mizuhopecten yessoensis* (Jay, 1856), is one of the most important seafood species in Japan (Kosaka & Ito, 2006; MAFF, 2015). Funka Bay, located in southwestern Hokkaido,
is one of the main commercially productive areas for scallop culture in Japan, where predominantly suspension culture techniques are used (Kosaka & Ito, 2006). The method for culturing is called ‘Mimi-zuri’ or ear-suspended method: a small hole is drilled at the front-eared beak of the left valve and the scallop is hung on a rope by using artificial strings or plastic clips (Kosaka & Ito, 2006).

In September 2008, *A. aspersa* was first found densely covering cultured scallops in Funka Bay, severely damaging aquaculture activities by causing the facility to sink and the scallops to fall off, and increasing expenses due to the need to dispose of the invasive species (Kanamori *et al*., 2012; Nishikawa *et al*., 2014). The ascidians overgrowing cultured scallops in Funka Bay had been correctly identified as *A. aspersa* through observation of the characteristics of internal morphology, follicle cells of egg, and DNA analysis of mitochondrial cytochrome c oxidase subunit I (Kanamori *et al*., 2012; Nishikawa *et al*., 2014). This is regarded as the first record of *A. aspersa* in the northern Pacific Ocean (Nishikawa *et al*., 2014). In South Korea, Pyo *et al*. (2012) identified many specimens collected in 2010 and 2011 as *A. aspersa* by using morphological and molecular analysis, and concluded that *A. aspersa* was widespread along three coastlines of Korea. However, the relationship between the Japanese and the Korean populations is unknown. In Japan, *A. aspersa* has been found in Hokkaido, Aomori, Iwate, and Miyagi Prefectures, and has become one of the most serious problems for bivalve aquaculture in northern Japan (Figure 1, Kanamori *et al*., 2014). Basic information such as reproductive season, growth patterns, maturity size, and population dynamics of *A. aspersa* in Japanese invasive populations is critical to controlling their impact.

In this study, we examined the recruitment, growth, maturity, and population dynamics of *A. aspersa*.
aspersa on cultured scallops in Funka Bay, and sought to relate the life history of *A. aspersa* with scallop aquaculture, to understand why the invasive ascidian has become a serious problem for the aquaculture industry in the bay. We also compared our results with a past study of native populations by Millar (1952), which is considered the most detailed account of the reproductive cycles of *A. aspersa* (Global Invasive Species Database, 2010), to deepen our understanding of the life history traits of this global invasive ascidian.

**MATERIALS AND METHODS**

**Larval density and seawater analyses**

In preparation for our study, we observed the morphology of larvae and their changes during metamorphosis in the laboratory. Monthly larval surveys were conducted from July 2010 to June 2014 at the sampling station (42°16.208′N, 140°20.568′E, Depth = 32 m, Figure 2) to determine the reproductive period of *A. aspersa*. Larvae were collected in 225-mm or 300-mm diameter plankton nets (NXX13 nylon mesh, opening of 100 μm, RIGO CO. LTD) hauled vertically from the bottom by hand. Our surveys were conducted between 11:30 and 13:30. Samples were fixed with glutaraldehyde (final concentration: 1%), and observed by stereoscopic microscope to count the number of *A. aspersa* larvae.

To determine the environmental factors that affect *A. aspersa* populations, water temperature and salinity were measured at every 1 m by CTD (RINKO-Profiler ASTD102, JFE Advantech Co. Ltd), and 300 mL of seawater was sampled using a Van Dorn sampler (RIGO CO. LTD) at depths of 5, 10,
and 15 m at the sampling station. Each sample was filtered using a glass microfiber filter (GF/F, 47 mm, Whatman, GE Healthcare Life Science), and chlorophyll a (Chl-a) was extracted with 10 mL of $N,N$-dimethylformamide (DMF) (Wako Pure Chemical Industries, Ltd.). The Chl-a content was measured from the change using fluorescence (excitation 436 nm, emission 660 nm) before and after acidification by adding 0.1 mL of 5% HCl in 3 mL of the sample DMF solution. Fluorescence was measured by a fluorescence spectrophotometer (FP6300, JASCO Corp.). The concentration of Chl-a was calculated using Chl-a from chlorella (Wako Pure Chemical Industries, Ltd.) as the standard.

**Sampling, measurement, and maturation level of A. aspersa**

Five scallops, *Mizuhopecten yessoensis*, were collected monthly at 5-, 10-, and 15-m depths from a culture rope near the sampling station between July 2010 and June 2014 (total 15 scallops were collected monthly). In Funka Bay, scallops are produced from a natural population of larvae, from spring to summer. Scallops are reared in cages from autumn to spring, and this is called the intermediate culture. Juvenile scallops, after an intermediate culture, are suspended for hanging culture in spring. Collection of the scallops is initiated after spring (June or July) each year, and completed the following June (from July 2010 to June 2011, from June 2011 to June 2012, from June 2012 to June 2013, and from June 2013 to June 2014). When hanging cultures are started, *A. aspersa* are seldom found on the scallops, which means that the ascidians found on scallops after spring are newly settled. In this study, therefore, the life history traits and population dynamics of *A. aspersa* were surveyed through four generations, the 2010, 2011, 2012, and 2013 cohorts.
Each scallop was placed in a zippered plastic bag to prevent the ascidians from falling off and carried to the laboratory in a cooler box. The surface of scallop was examined by direct observation and under a stereoscopic microscope. *Ascidiella aspersa* were removed using forceps. The number of individuals per each scallop was counted to assess seasonal variation in abundance, and the wet weight of *A. aspersa* was measured to assess seasonal variation in biomass. The weight of each scallop was quantified to compare it with the weight of the ascidians attached to it. Body length of each ascidian was determined within 0.1 mm using digital vernier calipers to examine size structure and growth. For small individuals (body length < 5 mm) found using a microscope, body length was measured from images captured using a Digital Sight Ds-Fi1 camera with NIS-Elements software (Nikon Corporation). More than 50 *A. aspersa* were randomly chosen from all depths in September, December, March, and June in 2010, 2011, and 2012, and fixed in 5–10% formalin seawater. After measuring body length, the specimen was dissected and genital ducts examined for eggs and sperm to evaluate the maturity. The 2013 cohort was not examined in terms of maturity. Sizes during maturity as male and female in September were analysed using generalised linear model (GLM) with a binomial error distribution. The response variable was whether eggs or sperm were in the ducts; the explanatory variable was body length, by using the statistical software R version 3.01 (R Development Core Team, 2013).
RESULTS

Larval density and environmental factors

The larvae of *A. aspersa* appeared in July–December 2010, July–November 2011, June–December 2012, and June–December 2013 (Figure 3), and were not found in the samples from January to May each year. Densities (individuals/m$^3$) were the highest between July and October. The highest density in each year was 74.3 in October 2010, 95.5 in August 2011, 37.7 in September 2012, and 22.6 in July 2013. Data were not collected in December 2012 because the plankton net was broken during the survey.

Water temperature reached its peak in August or September, except at 15-m depth in 2010 (Figure 4A). During summer 2010, a strong thermocline developed, in which the water temperature in August at 5-m depth was 23.9°C, whereas at 15-m depth, it was only 12.9°C. After the thermocline dissipated, the maximum water temperature at 15-m depth was 17.5°C, recorded in October. Water temperature was the lowest in February or March at all depths, in the range of 2.0–3.2°C. The seasonal fluctuation in salinity was stable in comparison with that of water temperature (Figure 4B). From spring to summer, the salinity was relatively low, fluctuating from 31.0 to 33.0 in part because of the inflow of the Oyashio Current, with low salinity, and in part because of the discharge of land water, including snowmelt runoff (Ohtani *et al.*, 1971a). From autumn to winter, the salinity fluctuated from 33.0 to 34.0 because of the inflow of the Tsugaru Warm Current, with high salinity (Ohtani *et al.*, 1971b). There were no obvious differences in salinity between depths, except in August–September 2010, when the thermocline developed intensely. A strong increase in Chl-a, a
spring bloom, occurred between February and April every year, and the concentration of Chl-a peaked at 6–8 μg/L (Figure 4C). After the spring bloom, the concentration remained low in summer and had an annual variability in autumn. A difference in Chl-a concentration between depths was not noted, and the average concentrations in 5-, 10-, and 15-m depths through the survey period were nearly the same at 1.53, 1.49, and 1.51 μg/L, respectively.

Seasonal variation in size, weight, and maturity of *A. aspersa* on scallops

In June, few *A. aspersa* were found on cultured scallops, and the average number per scallop at all depths was 0–0.9 individuals. In July, the average number increased to 0.9–7.8 individuals per scallop, and *A. aspersa* was observed at all depths except at the 5-m depth in 2013. After July, the number of *A. aspersa* per scallop increased and reached its peak between August and October. The average number of *A. aspersa* per scallop at each depth in each year is shown in Figure 5. The maximum number per scallop on average for all depths in each year was 117.4 individuals in October 2010, 39.2 individuals in August 2011, 22.9 individuals in September 2012, and 45.7 individuals in August 2013. During the time the numbers were increasing, as water depth increased, the number of *A. aspersa* also increased. After that, their numbers decreased, with an especially rapid rate of decrease at the 15-m depth. Because of this trend, in winter, the difference in number between the 10-m and 15-m depths became small. The number of *A. aspersa* at the 5-m depth was relatively low throughout the survey. June 2011 abundance data are not represented because only five scallops were collected, without the depth information.
No clear variation in size structure of *A. aspersa* on cultured scallops was noted between depths. However, seasonal variation in the size frequency was noted when all depths were combined, as shown in Figure 6. Juvenile ascidians (body length < 5 mm) dominated during the period of increasing abundance. For the 2010 cohort, many juvenile ascidians were found from August to October, whereas for the 2011, 2012, and 2013 cohorts, juvenile ascidians were found mainly from July to August. Figure 7 shows the seasonal variation in the body length of *A. aspersa* on cultured scallops at all depths. *Ascidiella aspersa* grew well until February following each season, when their body length remained unchanged or decreased slightly from February to March or April.

The biomass of the scallops increased steadily in each year. The biomass of *A. aspersa* on scallops increased, with fluctuations, until February, and after that, changes were less clear (Figures 8, 9). For the 2010 cohort, the average weight of *A. aspersa* at all depths exceeded that of the scallops even in November, and was three to seven times heavier in harvest season, from December to April, meaning that the weight of *A. aspersa* accounted for 75–90% of the total weight of the harvest. For the 2011 cohort, the average weight of the ascidians was less than that of the scallops except in February and March. For the 2012 cohort, the average weight of the ascidians was always less than that of the scallops. The weight of the ascidians in the 2013 cohort was more than that of the scallops in and after November. June 2011 weight data are not represented because only five scallops were collected, without the depth information.

*Ascidiella aspersa* with eggs and sperm in the ducts were found as late as September 2010, 2011, and 2012 (Figure 10). Because there were many juvenile ascidians in September 2010, the ratio of
ascidians having gametes was low (15%) at that time. On the other hand, in 2011 and 2012, the ratios were high, at 72.2% and 62.3%. Although there were few larvae and juveniles in December and March, many ascidians had eggs or sperm in the ducts. The ratios of individuals having gametes in December 2010, 2011, and 2012 were 52.0%, 81.6%, and 78.0%, respectively, and, in March 2011, 2012, and 2013, the ratios were 54.6%, 84.4%, and 87.5%, respectively. In June 2011, 2012, and 2013, the ratios were 92.1%, 100%, and 100%, respectively. In September, estimated 50% maturity size as male was 17–20 mm, and as female, 22–24 mm (Figure 11). The maturity size as female was approximately 5 mm larger than that as male, and in December and March, there were many A. aspersa with no gametes, even if the body length exceeded the 50% maturity size estimated in September. In the GLM analysis of maturity related to size as male and female in September, all of the estimated coefficients for body length were significant (Table 1, Wald test P < 0.001).

DISCUSSION

Life history traits and population dynamics of A. aspersa in Funka Bay

In Funka Bay, the larvae of A. aspersa appeared between June and December, and the highest density was observed between July and October. In addition, juvenile ascidians were found on cultured scallops mainly between July and October. Therefore, the reproductive period of A. aspersa is thought to be from June to December, and the main breeding season, from July to October. A study conducted from 1991 to 1997 in Long Island Sound, New England, showed that recruitment of A. aspersa started between June and July and, on average, initiation of recruitment was estimated to
occur on 1 July (Stachowicz et al., 2002). The onset of recruitment of *A. aspersa* in Funka Bay corresponds to that in Long Island Sound. The reproductive season of ascidians usually coincides with the period of maximum food production (Lambert, 2005). However, this idea does not apply to *A. aspersa* in Funka Bay because it is between February and April that the bay has a spring bloom and conditions for filter-feeders are good. *Ascidiella aspersa* grew well until February following the reproductive season. Their body length remained stagnant from February to March or April, when the bay has high production. The Oyashio Current, a subarctic current, introduces cold water to the bay and water temperatures fall below 4°C in February and March. Hence, the growth of *A. aspersa* would be depressed by low water temperature.

In autumn, *A. aspersa* had eggs or sperm. *Ascidiella aspersa* are known to be hermaphroditic, although the male sex organs develop first (Millar, 1952). In Funka Bay, the maturity size as males was estimated to be 17–20 mm, and as females was estimated to be 22–24 mm. Ascidians that reached these sizes in autumn had gametes and were expected to start reproduction. In December and March, there were many immature ascidians whose body length was greater than the maturity size in September. This indicated that factors other than body length influenced the accumulation of gametes. Because larvae and juvenile ascidians were scarcely found in winter and spring, the ascidians having gametes in December and March are thought to be the animals that reach maturity size in autumn and continue to have gametes after the reproductive season. In June, most of the *A. aspersa* had eggs and sperm, showing that the conditions needed for the maturity are fulfilled between March and June. Temperature is correlated with the timing of reproduction in many ascidian
species (Millar, 1971; Goodbody, 2004; Shenkar & Loya, 2008; Rius et al., 2009). The average temperatures found at 5–15-m depth in September, December, March, and June were 21.3°C, 8.0°C, 2.6°C, and 11.2°C, respectively. Hence, _A. aspersa_ stopped gamete accumulation when water temperature decreased from 21.3°C to 8.0°C, and started it again when water temperature increased from 2.6°C to 11.2°C. From this, we speculate that _A. aspersa_ have a critical temperature to start or stop the gamete accumulation, estimated to be between 8 and 11°C.

The number of _A. aspersa_ on the cultured scallops increased sharply after July and the number of juvenile ascidians increased with increasing water depth. In most cases, larval behaviour is a good predictor of adult distribution of ascidians (Svane & Young, 1989). At first, the larvae of _A. aspersa_ exhibit positive phototaxis and negative geotaxis; however, the reactions are reversed at later stages (Niermann-Kerkenberg & Hoffman, 1986). The reaction of larvae of _A. aspersa_ to environmental factors may explain the difference in quantity of ascidians at varying depths in our results. In our survey, the number of juvenile ascidians did not increase in autumn 2011, 2012, and 2013, although the generation from the previous year would continue reproduction; moreover, the recruits in summer would start spawning in autumn. _Ascidiella aspersa_ and other fouling animals settled over the surface of scallops in summer, and they may have prevented larvae of _A. aspersa_ from settling on scallops in autumn.

In 2010, the increase in ascidians was the greatest from August to September; however, in other years, it was from July to August. In Funka Bay, warm and less saline water is found in the surface layer from spring to summer, and a strong seasonal thermocline is formed (Ohtani et al., 1971a). The
thermocline dissipates by atmospheric influences and inflow of the Tsugaru Warm Current from summer to autumn (Ohtani et al., 1971b). The average air temperature of northern Japan in summer 2010 was the highest it had been since 1946 (Japan Meteorological Agency, 2010a), and in autumn, the temperature continued to be higher than that in an average year (Japan Meteorological Agency, 2010b). In addition, the inflow of the Tsugaru Warm Current was delayed, and not observed until mid-September (Hakodate Fisheries Research Institute, 2010). Under these conditions, the strong thermocline developed for a long time, and water temperatures in the depths below 15 m did not increase in summer. The low water temperature at deeper zones in summer 2010 may have influenced the reproduction of *A. aspersa* populations, resulting in the delay of *A. aspersa* increasing on cultured scallops.

During our survey, the Great East Japan Earthquake and the subsequent tsunami occurred on 11 March 2011. Funka Bay is approximately 500 km away from the centre of shock. Even so, the waves (maximum 1.6-m high) repeatedly struck the bay, damaged the facilities for scallop aquaculture, and affected coastal fauna (Japan Meteorological Agency, 2012; Natsuike et al., 2014). Most of the *A. aspersa* on the scallops at 5-m depth disappeared in and after March 2011 because the tsunami caused ascidians to drop off scallops in the shallow water. The effect on the ascidian population at 10–15-m depth appears small. The facilities damaged by the tsunami were removed and new facilities were established between 2011 and 2012 (Hokkaido Government, 2012), and consequently, many ascidians attached to the facilities were also removed. Facilities of aquaculture are considered important habitats for invasive ascidians (Lambert 2005; Howes et al. 2007; Carman...
et al. 2010). The tsunami and the removal of damaged facilities may explain why the numbers of A. aspersa on the scallops decreased in 2011 and 2012.

Life history of A. aspersa and the process of scallop aquaculture

The surface of newly suspended scallops is clean because they rub against netting or other scallops in the cage during intermediate culture; thus, they become a suitable substrate for sessile organisms, especially species that begin reproduction in early summer, such as A. aspersa. Harvest season for cultured scallops in the bay is mainly from December to April in order to avoid the shellfish toxin period and competition with other areas of production (Imai et al., 2014). Consequently, there is enough time for A. aspersa that have settled in summer to grow prior to scallop harvesting, and hence the harvest and shipment must be conducted after the weight of ascidians become several times heavier than that of scallops. The linking of “hang in spring and harvest in winter” of the cultured scallops and “recruitment after spring and rapid growth until winter” of A. aspersa results in serious problems in the aquaculture industry in Funka Bay (Figure 12). Effects of invasive organisms on an aquaculture industry depend on the relationship between the life history of the invasive species and the process of aquaculture in the introduced area. It is important to understand the life history and adaptations of invasive species in order to evaluate the risk of introduction to fisheries activities.

Comparison of life history of A. aspersa in Funka Bay and native area

The article by Millar (1952) is considered to be the most detailed account of the life history of A.
aspersa (Global Invasive Species Database, 2010), and the description in the literature and many databases are based on this significant work (e.g. Global Invasive Species Database, 2010; Mackenzie, 2011). Millar (1952) studied the reproductive cycle and population dynamics of A. aspersa throughout 1950 and 1951 in Ardrossan, southwestern Scotland, which is their native habitat and we summarise his findings here.

Larvae settle in the summer (July–August) and grow until the end of September. Ascidiella aspersa grow again after winter or spring. The life span is on the order of 18 months, extending approximately from the middle of one summer until the winter of the following year. Ascidiella aspersa have only one spawning season, and that is in the year after A. aspersa settled as larvae. Ascidiella aspersa is hermaphroditic and protandrous, in which the male reproductive organs come to maturity before the female reproductive organs. Sexual maturity is dependent on size; sperm development occurs when the animals are about 25-mm long, while eggs are found in the oviduct when the animals are about 30-mm long (Millar 1952). Most of the life history traits of A. aspersa in Funka Bay seem to be essentially identical to that summarised by Millar (1952). However, there are some clear differences.

The estimated reproductive period (June–December) and the main breeding season (July–October) in Funka Bay is longer than the recruitment season in Ardrossan (July–August). Ascidiella aspersa grow well until February in Funka Bay, and the average water temperature at 5–15-m depth fluctuates between 4 °C and 21°C from July to February. In Ardrossan, A. aspersa grow until late in September. From the information in Saltcoats, a town near Ardrossan, the peak water temperature is
14°C in August, and the lowest is 7 °C in March (World Sea Temperatures, 2015). This suggests that factors other than water temperature influenced the differences in growth period of *A. aspersa* between Funka Bay and Ardrossan. In Funka Bay, 60–70% of *A. aspersa* settled in summer have eggs or sperm in September, and *A. aspersa* would start to reproduce. From January to May, *A. aspersa* stop reproduction, and start spawning again in June. In contrast, *A. aspersa* in Ardrossan is regarded as the typical annual species, which has only one spawning season in the year after it has settled. Further, the extra generation of *A. aspersa* does not occur in the native population on the west coast of Norway (Dybern, 1969). The natural distribution of *A. aspersa* includes European low latitudes, such as the Mediterranean, but we have no information about the reproduction of *A. aspersa* in these areas. *Ascidiella aspersa* populations in the warmer temperature of the native range perhaps start to reproduce in the recruitment year as seen in Funka Bay. There is a possibility that the voltinism and reproductive traits of *A. aspersa* population is directly influenced by the habitat temperature, as discussed in the case of peracarida crustaceans (e.g. Vincente & Sorbe, 2013). Study of the life history and population dynamics of native *A. aspersa* population in warmer habitats is required to understand the life history strategy of this species.

The maturity size of *A. aspersa* in Funka Bay is approximately 5–8 mm smaller than that in Ardrossan. In Millar’s study, the samples were fixed after they were narcotised with menthol; in our study, the samples were directly fixed, which may have led to an underestimation of the body length. The test of *A. aspersa* is firm, and their siphons are short. Consequently, the difference in body length between individuals narcotised and those not narcotised was small, up to 3.5 mm, when the body
length was from 10.3 to 44.6 mm (N = 30, examined by MK on 14 September 2015). The differences in method of fixation would not fully account for the disagreement of maturity size between Funka Bay and Ardrossan. Millar (1952) also described that ascidians in Loch Sween, Argyll, western Scotland, became mature at a smaller body size than did those in any of the samples from Ardrossan. Further analysis is required to determine whether maturity size is different between Funka Bay and native ranges.

As described above, compared with the native population in Ardrossan, *A. aspersa* in Funka Bay has a longer reproductive and growth period, earlier initiation of reproduction, and possibly smaller maturity size. The vigour and success of invasive species has been explained by favourable environments where they are introduced and by release from natural enemies and the adaptation or evolution of increasing competitive ability (Blossey & Nötzold, 1995, Keane & Crawley, 2002; Colautti et al., 2004). Further studies that assess environmental factors, such as temperature and food conditions, and enemies regulating the population in native regions, are necessary to compare life history traits of the global invasive species, *A. aspersa*, in native and introduced ranges.
ACKNOWLEDGEMENTS

We thank Professor Teruaki Nishikawa of Toho University for the invaluable information on identification of ascidians. We are also grateful to Mr. Daisuke Achiya of the Yakumo Town Fisheries Cooperative, Associate Professor Isao Kudo of the Hokkaido University, and the students in Research Group of Marine Environmental Science, Graduate School of Fisheries Sciences, Hokkaido University for their helpful assistance in the field samplings. We appreciate Dr. Joan Cartes and the anonymous reviewer for their valuable suggestion and comments.

FINANCIAL SUPPORT

This study was conducted as a part of the contract researches from the Hokkaido Scallop Fisheries Promotion Association in 2010, 2011, 2012, and 2013.
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Fig. 1. Cultured scallops, *Mizuhopecten yessoensis*, overgrown by the invasive ascidian, *Ascidiella aspersa*, in Funka Bay, Hokkaido, northern Japan: (A), (B) a cultured rope with scallops hung by using plastic clips; (C) a cultured scallop held in the hand, having shell length of approximately 90 mm. More than 30 ascidians were attached to the scallop in (C) when the photos were taken on 18 May 2015.

Fig. 2. Maps showing Funka Bay, Hokkaido, northern Japan and a sampling station (42°16.208′ N, 140°20.568′ E, Depth = 32 m). Recording of environmental conditions and plankton surveys were conducted at the sampling station. Cultured scallops were collected around the sampling station to investigate the attached *Ascidiella aspersa*.

Fig. 3. Seasonal variation in larval density of *Ascidiella aspersa* at a sampling station (42°16.208′ N, 140°20.568′ E, Depth = 32 m), Funka Bay, Hokkaido, northern Japan from July 2010 to June 2014. J, S, N, J, M, M: July, September, November, January, March, May.

Fig. 4. Seasonal variation in (A) water temperature, (B) salinity, and (C) chlorophyll a concentration at a sampling station (42°16.208′ N, 140°20.568′ E, Depth = 32 m), Funka Bay, Hokkaido, northern Japan from July 2010 to June 2014. J, S, N, J, M, M: July, September, November, January, March,
Fig. 5. Seasonal variation in the number of *Ascidiella aspersa* on cultured scallops. (first J on the horizontal axis is June of the year presented on the graph; last J is June of the following year). Average and standard error of number of *A. aspersa* on a scallop in each depth are shown: (A) 2010 cohort from July 2010 to May 2011; (B) 2011 cohort from June 2011 to June 2012; (C) 2012 cohort from June 2012 to June 2013; and (D) 2013 cohort from June 2013 to June 2014. For June 2012 and 2013, cultured scallops hung in the previous year and the year were collected. Scales of vertical axes are different.

Fig. 6. Seasonal variation in size frequency of *Ascidiella aspersa* on cultured scallops at all depths.

Fig. 7. Seasonal variation in the body length of *Ascidiella aspersa* on cultured scallops at all depths (first J on the horizontal axis is June of the year presented on the graph; last J is June of the following year). The medians are shown as representative values. Bars indicate 25th and 75th percentiles: (A) 2010 cohort from July 2010 to June 2011; (B) 2011 cohort from July 2011 to June 2012; (C) 2012 cohort from June 2012 to June 2013; and (D) 2013 cohort from July 2013 to June 2014.

Fig. 8. Seasonal variation in biomass of *Ascidiella aspersa* on cultured scallops (first J on the horizontal axis is June of the year presented on the graph; last J is June of the following year).
Average and standard error of wet weight (w.w. in grams [g]) of *A. aspersa* per month at each depth is shown: (A) 2010 cohort from July 2010 to May 2011; (B) 2011 cohort from June 2011 to June 2012; (C) 2012 cohort from June 2012 to June 2013; and (D) 2013 cohort from June 2013 to June 2014. For June 2012 and 2013, cultured scallops hung in the previous year and the year were collected. Scales of vertical axes are different.

**Fig. 9.** Seasonal variation in biomass of *Ascidiella aspersa* and cultured scallop, *Mizuhopecten yessoensis* (first J on the horizontal axis is June of the year presented on the graph; last J is June of the following year). Average wet weight (w.w. in grams [g]) of *A. aspersa* and *M. yessoensis* per month at all depths is shown: (A) 2010 cohort from July 2010 to May 2011; (B) 2011 cohort from June 2011 to June 2012; (C) 2012 cohort from June 2012 to June 2013; and (D) 2013 cohort from June 2013 to June 2014. Scallops were hung in spring each year. For June 2012 and 2013, cultured scallops hung in the previous year and the year were collected. Scales of vertical axes are different.

**Fig. 10.** Size frequency and the presence of sperm and eggs in the ducts of *Ascidiella aspersa*: (A) 2010 cohort; (B) 2011 cohort; and (C) 2012 cohort. Ascidians having neither eggs nor sperm in their ducts are regarded as immature.

**Fig. 11.** Relation between body length and maturity of *Ascidiella aspersa* in September. Maturity is assessed by the presence of gametes in the ducts. The best-fit logistic curves are shown. Maturity
size (M₅₀) indicates the size at which 50% of *A. aspersa* mature, estimated according to the logistic curves.

**Fig. 12.** Life history of *Ascidiella aspersa* and basic process of scallop culture in Funka Bay, Hokkaido, northern Japan. Scallops hung in spring become suitable substrate for *A. aspersa*, which start their reproduction in early summer. The rapid growth and weight gains of *A. aspersa* from summer to winter cause serious problems for the scallop-harvesting season.
Fig. 1
Fig. 2
Fig. 3

Larval density of A. aspersa (Number/m$^3$)
Fig. 4
Fig. 5
Fig. 6
Fig. 7
Fig. 8

A 2010 Cohort

B 2011 Cohort

C 2012 Cohort

D 2013 Cohort

Weight on a scallop (w.w.g.)

Month
Fig. 11
<table>
<thead>
<tr>
<th>Winter</th>
<th>Spring</th>
<th>Summer</th>
<th>Autumn</th>
<th>Winter</th>
<th>Spring</th>
<th>Summer</th>
</tr>
</thead>
<tbody>
<tr>
<td>Life history of A. aspersa</td>
<td>Recruitment (Peaks; Jul.-Sep.)</td>
<td>Growing (Rapidly; Jul.-Feb.)</td>
<td>Reproduction</td>
<td>Reproduction</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Process of scallop culture

- Intermediate culture (Oct.-Feb.)
- Ear-hanging (Mar.-May)
- Hanging culture (Jun.-Nov.)
- Harvesting (Dec.-Apr.)

Many ascidians attach to the newly cultured scallops. Weight gains of the ascidians cause the problem for harvesting.
Table 1. Results of generalized linear model (GLM) analysis for the maturity of *Ascidiella aspersa* collected in September. All of the coefficients for body length are significant (P < 0.001, Wald test). The maturity size indicates the size at which 50% of *A. aspersa* mature.

<table>
<thead>
<tr>
<th></th>
<th>Explanatory variable</th>
<th>50% Maturity size (mm)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Intercept ($\beta_0$)</td>
<td>Body length ($\beta_1$)</td>
</tr>
<tr>
<td></td>
<td>Coef.    SE  z    p</td>
<td>Coef.    SE  z    p</td>
</tr>
<tr>
<td>Male</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2010</td>
<td>-13.612  3.563 -3.820 &lt;0.001</td>
<td>0.681  0.185 3.678 &lt;0.001</td>
</tr>
<tr>
<td>2011</td>
<td>-8.850   2.382 -3.715 &lt;0.001</td>
<td>0.474  0.121 3.901 &lt;0.001</td>
</tr>
<tr>
<td>2012</td>
<td>-5.471   1.087 -5.003 &lt;0.001</td>
<td>0.320  0.059 5.431 &lt;0.001</td>
</tr>
<tr>
<td>Female</td>
<td></td>
<td></td>
</tr>
<tr>
<td>2010</td>
<td>-13.962  3.338 -4.183 &lt;0.001</td>
<td>0.583  0.145 4.016 &lt;0.001</td>
</tr>
<tr>
<td>2011</td>
<td>-10.238  2.359 -4.340 &lt;0.001</td>
<td>0.473  0.107 4.415 &lt;0.001</td>
</tr>
<tr>
<td>2012</td>
<td>-8.450   1.483 -5.697 &lt;0.001</td>
<td>0.351  0.065 5.4506 &lt;0.001</td>
</tr>
</tbody>
</table>