

## Ctenophore population recruits entirely through larval reproduction in the central Baltic Sea

Cornelia Jaspers, Matilda Haraldsson, Sören Bolte, Thorsten B. H. Reusch, Uffe H. Thygesen and Thomas Kiørboe

*Biol. Lett.* 2012 **8**, 809-812 first published online 25 April 2012  
doi: 10.1098/rsbl.2012.0163

---

### Supplementary data

["Data Supplement"](#)

<http://rsbl.royalsocietypublishing.org/content/suppl/2012/04/24/rsbl.2012.0163.DC1.html>

### References

[This article cites 14 articles, 1 of which can be accessed free](#)

<http://rsbl.royalsocietypublishing.org/content/8/5/809.full.html#ref-list-1>

### Subject collections

Articles on similar topics can be found in the following collections

[developmental biology](#) (51 articles)

[ecology](#) (573 articles)

[taxonomy and systematics](#) (66 articles)

### Email alerting service

Receive free email alerts when new articles cite this article - sign up in the box at the top right-hand corner of the article or click [here](#)

# Ctenophore population recruits entirely through larval reproduction in the central Baltic Sea



Cornelia Jaspers<sup>1,\*</sup>, Matilda Haraldsson<sup>2</sup>,  
Sören Bolte<sup>3</sup>, Thorsten B. H. Reusch<sup>3</sup>,  
Uffe H. Thygesen<sup>1</sup> and Thomas Kiørboe<sup>1</sup>

<sup>1</sup>Centre for Ocean Life, DTU-Aqua, Technical University of Denmark, Kavalergården 6, 2920 Charlottenlund, Denmark

<sup>2</sup>Biological and Environmental Sciences, Gothenburg University, Kristineberg 566, 45178 Fiskebäckskil, Sweden

<sup>3</sup>Helmholtz-Centre for Ocean Research (GEOMAR), Düsternbrooker Weg 20, 24105 Kiel, Germany

\*Author for correspondence ([coja@aqu.dtu.dk](mailto:coja@aqu.dtu.dk)).

**The comb jelly *Mertensia ovum*, widely distributed in Arctic regions, has recently been discovered in the northern Baltic Sea. We show that *M. ovum* also exists in the central Baltic but that the population consists solely of small-sized larvae (less than 1.6 mm). Despite the absence of adults, eggs were abundant. Experiments revealed that the larvae were reproductively active. Egg production and anticipated mortality rates suggest a self-sustaining population. This is the first account of a ctenophore population entirely recruiting through larval reproduction (paedogenesis). We hypothesize that early reproduction is favoured over growth to compensate for high predation pressure.**

**Keywords:** *Mertensia ovum*; comb jelly; reproduction; paedogenesis

## 1. INTRODUCTION

Reproduction before metamorphosis in the larval stage owing to delayed somatic growth (neoteny) or precocious maturation (paedogenesis) is known among amphibians and parthenogenetic insects [1]. In some extreme cases, natural populations consist exclusively of larvae [1].

Marine ctenophores are similarly capable of reproduction in the larval stage [2–5]. While larvae are normally defined as a non-reproductive developmental stage before metamorphosis, larval reproduction in ctenophores has been shown for both metamorphosing (Lobata) and non-metamorphosing (Cydippida) orders in their early life stage less than *ca* 2.7 mm [3–5]. In the cydippid *Pleurobrachia* spp., larval gonad structures are different from adult gonads and first reproduction has been described at a minimum size of 0.4 mm [3]. After larvae of both major orders reach a threshold size, reproduction ceases, animals rapidly grow and then become reproductive again as adults (6–10 mm) [4,5]. While metamorphosis in amphibians generally

involves a habitat shift [6], ctenophore paedogenesis is linked to early age reproduction being favoured over growth [4,5] and has been hypothesized to compensate for high mortality [2–5]. However, this hypothesis has never been confirmed at population level in nature.

Recently, the Arctic cydippid ctenophore *Mertensia ovum* was discovered in the Baltic Sea possibly as a relict population from the former ice age [7]. While Arctic specimens measure up to 90 mm, *M. ovum* in the northern Baltic remain small (less than 6.5 mm) [7,8]. Here, we describe for the first time that a ctenophore population is recruiting solely through larval reproduction (paedogenesis). The lack of larger-sized specimens is hypothesized to be due to high predation pressure.

## 2. MATERIAL AND METHODS

Zooplankton sampling was conducted (13 monthly cruises) during 2009/2010 in the central Baltic Sea (figure 1), with vertical (90 µm HYDRO-BIOS Kiel, Germany, 0.25 m<sup>2</sup> midi-MultiNet) hauls in five depth-strata from 70, 200 and 180 m for stations 1 to 3. At station 4, the total water column (17 m) was sampled without depth-resolution.

Ctenophore eggs/larvae were measured, either live or after 2 per cent acidified Lugol preservation. Live versus preserved animal sizes (*n* = 1114) were compared. Preserved sizes were 75 ± 12.2% of unpreserved ones and were multiplied by 1.33 to correct for this shrinkage.

Ctenophore DNA from dried samples (*n* = 121) was verified using species-specific primers for the ITS-1 region of the ribosomal-RNA genes for *M. ovum* and invasive *Mnemiopsis leidyi*, respectively [9]. Analyses used standard PCR protocols (electronic supplementary material). All eggs are assumed to be *M. ovum* since no other ctenophore larvae (in the study of Gorokhova *et al.* [7, present study]) have been confirmed for this area.

For reproduction experiments (October 2009, station 2), larvae were collected with vertical 300 µm (MultiNet) tows (90–50 m) and incubated individually in 20 ml tissue-culture trays (Nunc Roskilde, Denmark) with 20 µm filtered water at 7 ± 1°C, salinity of 7. For 24 animals (0.54–1.33 mm), eggs were counted after 48 h with sizes based on averages (before/after the experiment). Regression analysis was performed for estimating size-dependent egg production.

Expected egg abundance at each station was estimated from the observed ctenophore size distribution and the size-dependent egg production, multiplied by the hatching time (*T*, days), excluding station 4 where eggs were never observed. Hatching time for *M. leidyi* eggs is 7 days (7°C, C. Jaspers 2010, unpublished data), and we assume the same for *M. ovum*. From this, expected egg abundance (squared metre) at each station was computed assuming no mortality. With no egg mortality, this computation should yield a larger observed egg abundance than expected at half the stations, on average. To test the hypothesis of zero mortality, the fraction of such stations was compared with 50 per cent using binomial statistics (electronic supplementary material).

## 3. RESULTS

Ctenophores were present throughout the year (figure 1). Molecular analysis confirmed that the individuals sampled were *M. ovum*.

Population sizes peaked during winter with maximum abundances recorded at the deep-water, northernmost station 3. Ctenophores were not observed at the shallow-water station 4 during summer when water temperatures were high (*ca* 17°C). *Mertensia ovum* showed a difference in depth distribution with season, residing deeper during warm seasons, and distributed throughout the water column or in surface waters during cold seasons (electronic supplementary material, figure S1). Hence, the temperature range where larvae were observed was low (−0.3°C to 11.6°C).

Size distributions differed significantly between seasons with the largest sizes during summer (figure. 2*a*). However, throughout the year and area, ctenophores were very small: 95 per cent of the population were

Electronic supplementary material is available at <http://dx.doi.org/10.1098/rsbl.2012.0163> or via <http://rsbl.royalsocietypublishing.org>.

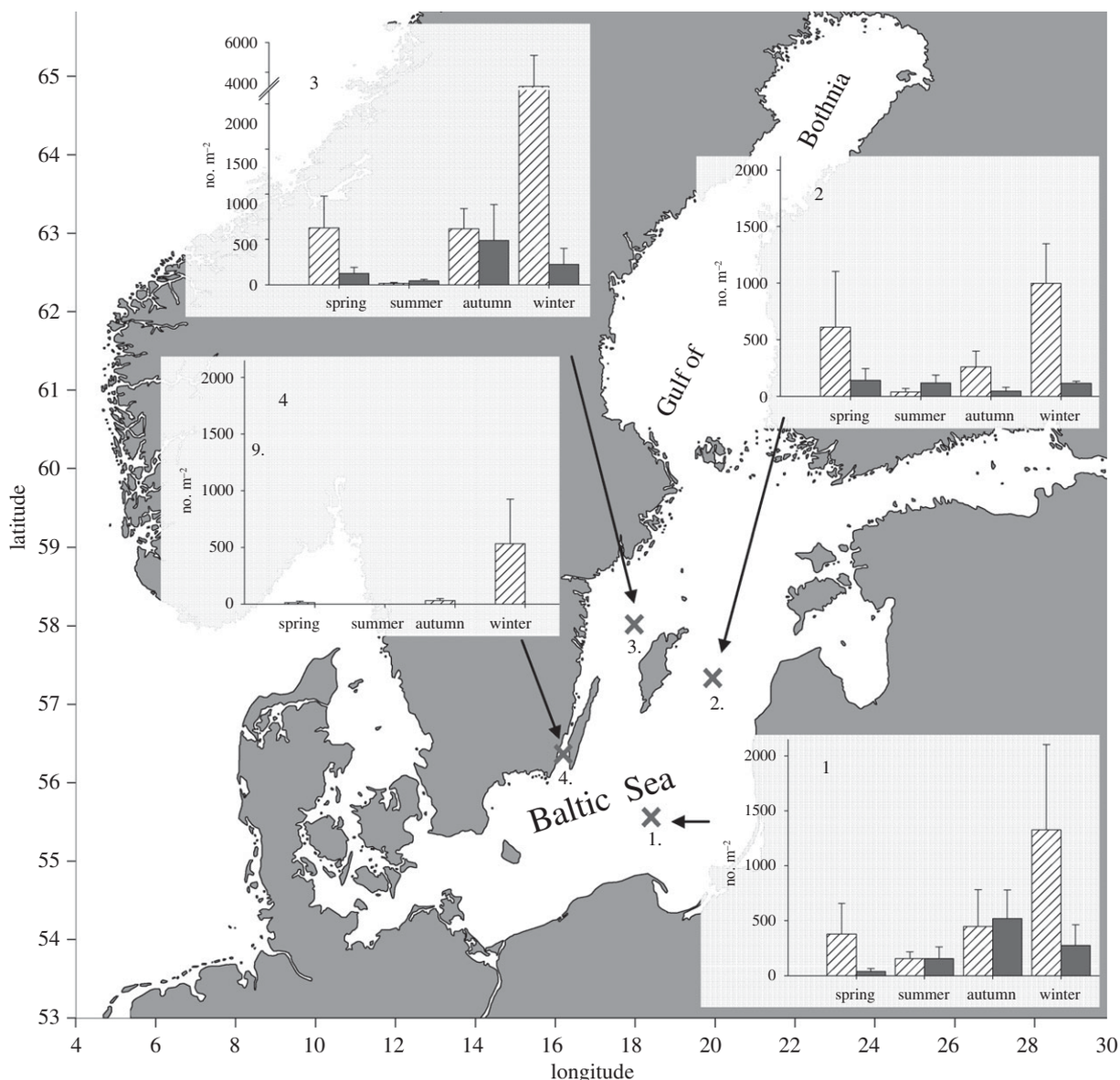


Figure 1. Integrated seasonal *Mertensia ovum* larvae (bars with stripes) and egg (black bars) distribution in the central Baltic Sea, 2009/2010 (pooled for season  $\pm$  s.d.).

less than 1.1 mm and the largest specimen found was 1.6 mm ( $n = 7192$ ).

High numbers of ctenophore eggs were observed (figure 1). While larvae abundances were highest in winter, most eggs were present in autumn. Highest egg density was observed for the southernmost station 1, while at the shallow-water station 4 eggs were never found.

Reproduction experiments revealed that *M. ovum* greater than 0.75 mm produced eggs and production increased significantly with size (figure 2*b*). Overall, nine animals produced 22 eggs in  $48 \text{ h}^{-1}$ , while 14 animals (0.54–1.04 mm) did not.

The fraction of stations with lower egg abundance than expected did not deviate significantly from 50 per cent, either for the entire sampling period or within seasons; hence egg mortality can be hypothesized to be zero (figure 2*c*).

#### 4. DISCUSSION

Our data show for the first time that a ctenophore population in nature consists entirely of small, larval

size classes (less than 1.6 mm) throughout the year. We argue that our observations are consistent with a self-sustaining population maintained entirely through larval reproduction.

First, it is unlikely that the population in the central Baltic is supplied by advection from the Gulf of Bothnia in the north. In the north, *M. ovum* individuals are four times larger but their densities are only 1–8% compared with the central Baltic and they occur primarily at depth greater than 50 m [7]. Since the average residence time of water in the Gulf of Bothnia is 6 years and the southward advection mainly consists of surface water [10], drift recruitment is unlikely to be important.

Second, *M. ovum* larvae greater than 0.75 mm produce eggs at significant rates. Minimum size for larval reproduction in *Pleurobrachia* spp. is 0.4 mm [3], and 1.2 mm *P. bachei* produces eight eggs per day at  $15^\circ\text{C}$  [5]. The lobate *M. leidy* (2 mm) produces up to 14 eggs per day at  $22^\circ\text{C}$  at high food concentrations [4]. *Mertensia ovum* (less than 1.3 mm) produces up to two eggs day. Considering the variation in temperature

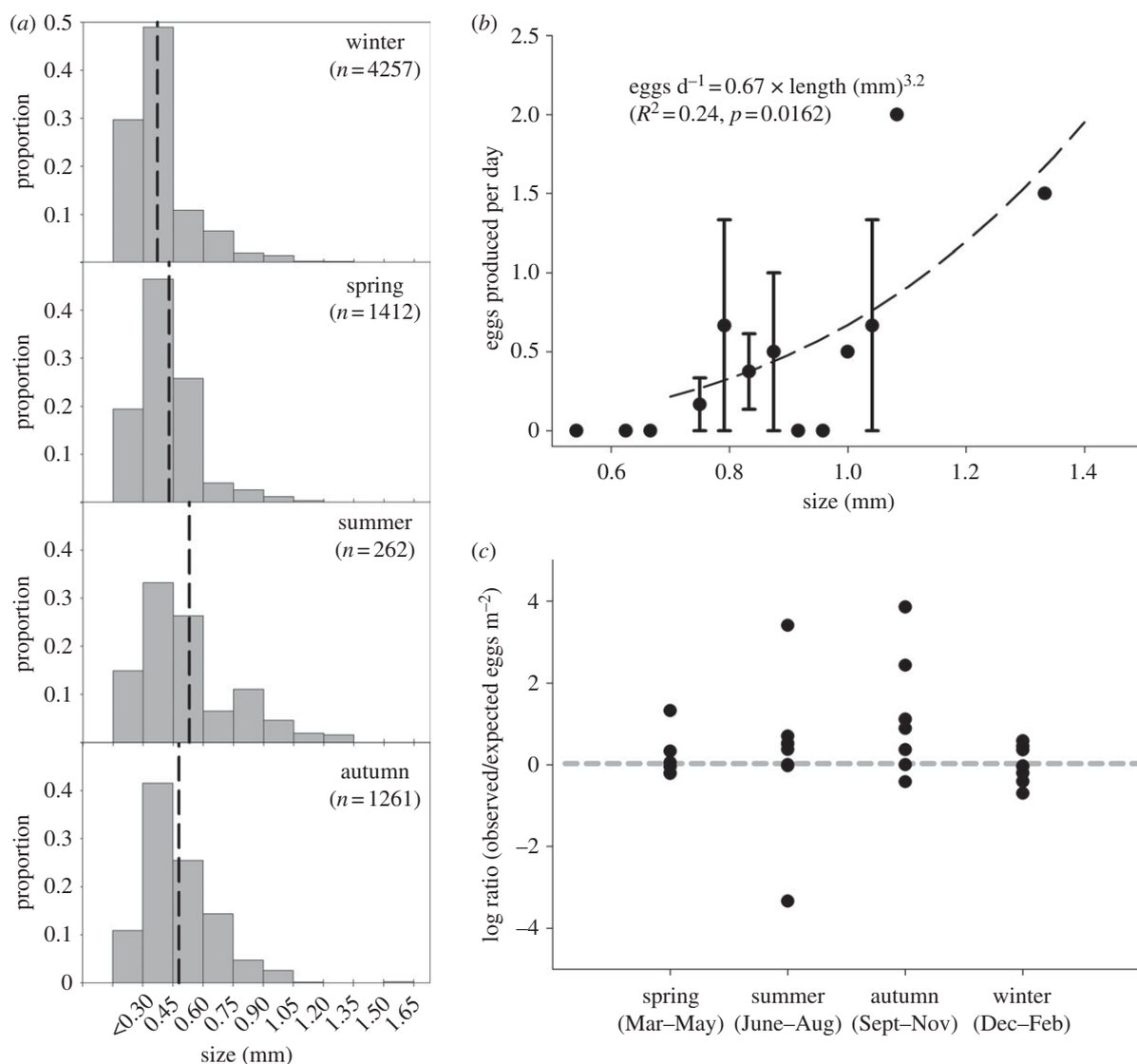


Figure 2. (a) Seasonal size distribution of *Mertensia ovum* in the central Baltic (0.15 mm size bins, averages (dashed lines), log-transformed lengths significantly differed with season (one-way ANOVA  $F_{3,7845} = 821$ ,  $p < 0.0001$ ) (b) size-dependent egg production at 7°C, October 2009 (average sizes  $\pm$  s.e.) (c) log ratio of observed to expected egg abundance with zero observations substituted by 0.1 (log ratio = 0, no egg mortality). The fraction of stations where expected egg abundance exceeds observed egg abundance does not differ significantly from 50% ( $p = 0.36$ ).

[11] and sizes, larval reproductive rates are similar for the different species and confirm our observation of reproducing larvae in the Baltic.

Finally, the observed egg production is sufficient to maintain the population in the face of likely mortality rates. The requirement for population maintenance is that the net reproductive rate  $R_0 \geq 1$  for simultaneous hermaphrodites. Assuming an egg hatching time of  $T_1$ , that it takes time  $T_2$  to grow to reproductive size whereupon growth is replaced by egg production at a constant rate ( $m$ ), and that mortalities in the egg ( $\delta_1$ ) and post-hatch ( $\delta_2$ ) stages are constant, then  $R_0 = \exp(-\delta_1 T_1)(m/\delta_2)\exp(-\delta_2 T_2)$  [12]. Using temperature-corrected maximum growth rates reported for *M. leidy* larvae [11,13], it would take *M. ovum* about 8 days to reach 1 mm size at 7°C. If we insert the numbers relevant to our October reproduction experiments ( $\delta_1 = 0$ ,  $T_2 = 8$  d,  $m = 0.7$  d $^{-1}$ ), then  $R_0 > 1$ , as long as the mortality is less than 0.174 d $^{-1}$ . Faster growth and reproduction during summer allows for higher

mortality, and conversely during winter. In general, mortality rates of similarly sized pelagic organisms are similar or less [14] and the observed larval fecundity is thus consistent with a self-sustaining population.

Why does *M. ovum* population in the central Baltic consists of only larvae that are 1–2 orders of magnitude smaller in length (3–6 orders in mass) than in the Arctic [7,8]? The temperature–size rule predicts that individual sizes within a species increase with decreasing temperatures, but a temperature difference as observed here of 10°C predicts only a 30 per cent difference in individual masses [15]. Similarly, marine species are often smaller in brackish water systems like the Baltic [16], but not to the degree observed here. Also, the largest *M. ovum* were observed in the least saline Gulf of Bothnia [7]. Therefore, neither temperature nor salinity can explain the observed differences.

Life-history theory predicts that the optimum age of maturation decreases with increasing juvenile mortality

[17]. The trade-off is between maturing early at a small size and low fecundity but high chance of surviving to maturity versus maturing late at a large size and high reproduction but lower chance of reaching maturity. Our observation of early reproduction in *M. ovum* is consistent with the suggestion of high juvenile mortality. We substantiate this by a simple calculation. If juvenile growth in mass is a power function of time with an exponent of  $c = 2$  as in the initial phase of the von Bertalanffy growth model [14], egg-production rate is proportional to individual mass (or length<sup>3</sup>, figure 2*b*), and growth ceases subsequent to start of egg production [4,5], then the maturation time yielding the highest  $R_0$  is  $c/\delta_2$  [12]. Assuming  $\delta_2 = 0.17 \text{ d}^{-1}$  as has been shown for less than 19 day larvae of the ctenophore *Pleurobrachia bachei* [18] and as calculated above, implies a development time of 12 days, close to the estimated time required to reach reproductive size. Higher temperatures may imply shorter maturation time but probably also higher mortality, and vice versa for lower temperatures.

Why would mortality be higher in the Baltic than in the Arctic where *M. ovum* are much bigger? We have no strongly substantiated explanation of this but note that planktivorous fish, potential predators on *M. ovum*, are abundant in the central Baltic Sea [19], and, more generally, that the relative significance of pelagic versus demersal fish increases from Arctic to temperate and tropical ecosystems [20]. The Baltic relict population of *M. ovum* [7] has had several thousand years to adapt to the local predation pressure.

In amphibians, a wide range of life-history strategies exist including facultative and obligate metamorphosis [1], and neoteny has been shown to be a response to density-dependent processes in newts with adult and larval reproduction occurring at the same age [6]. In extreme cases, natural populations are 'trapped' in the larval stage [1]. In insects, paedogenesis has been linked to optimum utilization of food patches, leading to short generation times in response to food availability [1]. In contrast, *M. ovum* paedogenesis is suggested to be caused by high predation pressure. However, very little is known about the potential predators on ctenophores in the Baltic, and this suggestion should be tested, e.g. through diet analyses and experiments with and without the presence of predators.

The project was funded by BONUS (BAZOOCA: project no. 210-2008-1882/-1889).

- 1 Gould, S. J. 1977 *Ontogeny and phylogeny*, 1st edn. London, UK: Belknap.
- 2 Chun, C. 1892 *Die Disoogenie, eine Form der geschlechtlichen Zeugung*. In *Festschrift zum siebzigsten Geburtstag* (ed. R. Leuckarts), pp. 77–108. Leipzig, Germany: Engelmann.

- 3 Garbe, A. 1901 Untersuchung über die Entstehung der Geschlechtsorgane bei den Ctenophoren. *Z. wiss. Zool.* **69**, 472–491.
- 4 Martindale, M. Q. 1987 Larval reproduction in the ctenophore *Mnemiopsis mccradyi* (order Lobata). *Mar. Biol.* **94**, 409–414. (doi:10.1007/BF00428247)
- 5 Hirota, J. 1972 Laboratory culture and metabolism of the planktonic ctenophore, *Pleurobrachia bachei*. In *Biological oceanography of the North Pacific* (ed. A. Y. Takenouti), pp. 465–484. Tokyo, Japan: Idemitsu-Shoten.
- 6 Harris, R. N. 1987 Density-dependent paedomorphosis in the salamander *Notophthalmus viridescens*. *Ecology* **68**, 705–712. (doi:10.2307/1938476)
- 7 Gorokhova, E., Lehtiniemi, M., Viitasalo-Frosen, S. & Haddock, S. H. D. 2009 Molecular evidence for the occurrence of ctenophore *Mertensia ovum* in the northern Baltic Sea and implications for the status of the *Mnemiopsis leidyi* invasion. *Limnol. Oceanogr.* **54**, 2025–2033. (doi:10.4319/lo.2009.54.6.2025)
- 8 Matsumoto, G. 1991 Functional morphology and locomotion of the Arctic ctenophore *Mertensia ovum* (Fabricius) (Tentaculata: Cydippida). *Sarsia* **76**, 177–185.
- 9 Reusch, T. B. H., Bolte, S., Sparwel, M., Moss, A. & Javidpour, J. 2010 Microsatellites reveal origin and genetic diversity of Eurasian invasions by one of the world's most notorious marine invader, *Mnemiopsis leidyi*. *Mol. Ecol.* **19**, 2690–2699. (doi:10.1111/j.1365-294X.2010.04701.x)
- 10 Myrberg, K. & Andrejev, O. 2006 Modelling of the circulation, water exchange and water age properties of the Gulf of Bothnia. *Oceanologia* **48**, 55–74.
- 11 Hansen, P. J., Bjornsen, P. & Hansen, B.W. 1997 Zooplankton grazing and growth: scaling within the 2–2,000- $\mu\text{m}$  body size range. *Limnol. Oceanogr.* **42**, 687–704. (doi:10.4319/lo.1997.42.4.0687)
- 12 Kjørboe, T. & Hirst, A. 2008 Optimum development time in pelagic copepods. *Mar. Ecol. Prog. Ser.* **367**, 15–22. (doi:10.3354/meps07572)
- 13 Baker, L. & Reeve, M. 1974 Laboratory culture of lobate ctenophore *Mnemiopsis* with notes on feeding and fecundity. *Mar. Biol.* **26**, 57–62. (doi:10.1007/BF00389086)
- 14 Kjørboe, T. 2008 *A mechanistic approach to plankton ecology*, 1st edn. Oxford, UK: Princeton University Press.
- 15 Forster, J., Hirst, A. & Atkinson, D. 2011 How do organisms change size with changing temperature? *Funct. Ecol.* **25**, 1024–1031. (doi:10.1111/j.1365-2435.2011.01852.x)
- 16 Kautsky, N. & Tedengren, M. 1992 Ecophysiological strategies in Baltic Sea invertebrates. In *Proc. 12th Baltic Marine Biologists Symp. 25–30 August 1991, Helsingør, Denmark*, pp. 91–96. Fredensborg, Denmark: Olsen & Olsen.
- 17 Stearns, S. C. 1992 *The evolution of life histories*, 1st edn. Oxford, UK: Oxford University Press.
- 18 Hirota, J. 1974 Quantitative natural history of *Pleurobrachia bachei* in La Jolla Bight. *Fish. Bull.* **72**, 295–335.
- 19 Casini, M., Lovgren, J., Hjelm, J., Cardinale, M., Molinero, J. C. & Kornilovs, G. 2008 Multi-level trophic cascades in a heavily exploited open marine ecosystem. *Proc. R. Soc. B* **275**, 1793–1801. (doi:10.1098/rspb.2007.1752)
- 20 Petersen, H. & Curtis, M. 1980 Difference in energy flows through major components of subarctic, temperate and tropical marine shelf ecosystems. *Dana* **1**, 53–64.