Population dynamics of the venerid bivalve *Tawera gayi* (Hupé, 1854) in the Ushuaia Bay, Beagle Channel

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Summary

Growth, productivity and potential for exploitation of the clam *Tawera gayi* from shallow waters (3–5 m) of Ushuaia Bay, Beagle Channel were investigated. Mean abundance and biomass in the study area were 1091 ± 737 ind. m⁻² and 901.83 g SFWM m⁻² (shell-free wet mass), respectively. Individual growth was described best by the von Bertalanffy growth model with the parameter values *H₀ = 28.03 mm, K = 0.288 year⁻¹, to = −0.34 (r² = 0.83). Annual production of the population was estimated to be 120.45 g SFWM m⁻² year⁻¹, corresponding to a production-to-biomass ratio (*P/B*) of 0.134 year⁻¹. The single negative exponential mortality model does not fit the population mortality pattern, but predation by gastropods (*Aynemopsis muriciformis, Trophon geversianus, Natica* sp.) appears to be the major cause of mortality. These highly mobile predators together with the comparatively slow growth and low turnover of *T. gayi* in Ushuaia Bay limit its potential for sustainable commercial exploitation.

Material and methods

Sampling

Monthly samples of the hard clams *T. gayi* (n = 4118 specimens) were collected in Ushuaia Bay (54⁰50'18”S, 68’16’25”W), Beagle Channel (Fig. 1) by SCUBA diving in an area about 400 × 200 m at 3–5 m water depth between August 2001 and July 2002. Each month four randomly selected cores of 30 × 30 cm surface area and 15 cm depth were dug out. Specimens were sorted from the unsieved content of the cores. During the study period the monthly mean seawater temperature ranged between 8.2°C in summer and 3.4°C in winter (Fig. 2. C. Schroeder, unpubl. data).

In all individuals collected we measured shell height (H, umbo to the ventral margin), length (L, anterior–posterior axis) and width (W) (precision ± 0.1 mm), and determined total mass (TM), shell mass (SM) and shell-free wet mass (SFWM) (precision ± 0.01 g).

For size–mass relationships,

\[ M = aH^b \] (1)

linear regression analysis was carried out with the log-transformed data, where *M* is the mass, *a* is the intercept and *b* is the slope.

Growth

Individual age was inferred from shell growth bands of 285 specimens randomly selected from all 4118 individuals collected during the 12-month sample period. Each left valve was sectioned along the axis of maximum growth in height (H) and the cross-sections were polished on lapiary wheels using grits of 180, 400, 600 and 1200 grade. Polished shell cuts were examined by stereomicroscope using reflecting light. Acetate peels were made of polished and etched sections from valves embedded in epoxy resin to confirm the pattern found (Rhoads and Lutz, 1980).

A mark-recapture field experiment was carried out to validate the growth pattern found. A group of 96 animals of different size (20–30 mm H) were collected, measured and marked on the surface of the periostracum. Each animal was returned to the subtidal area and allowed to bury in an individual cage consisting of a metal tube 20 cm ø × 20 cm height with perforated walls to permit horizontal water flow through the cage. After 1 year the animals were recovered and annual growth lines were examined as described above.
To confirm during which season translucent and opaque bands (Jones et al., 1990) were formed, the stage of terminal growth in cross-section shells was identified within either the translucent (narrow dark) band or the opaque (wide white) band (see Arnold et al., 1998). Twenty specimens per month were analysed and monthly growth stage data were pooled into seasonal groups.

The number of annual bands and the corresponding height were recorded as age-height data. To describe growth, the von Bertalanffy growth model was used

\[ H_t = H_{\infty} (1 - e^{-K(t-t_0)}) \text{ [mm, year]} \]  

where, \( H_{\infty} \) is asymptotic height, \( K \) is the growth constant, \( t \) is age and \( t_0 \) is age at zero height. The non-linear iterative Newton algorithm was used to fit the model to the data.

Production

Somatic production was calculated by the mass-specific growth rate method (Crisp, 1984; Brey, 2001) from the size–frequency distribution obtained from all pooled samples, the von Bertalanffy growth function and the size–mass relation:

\[ P = \Sigma N_i M_i G_i \text{ [g SFWM m}^{-2} \text{ year}^{-1}] \]  

where, \( N_i \) and \( M_i \) are the average number of animals (N m\(^{-2}\)) and mean individual body mass in size class \( i \), and \( G_i \) is the mass-specific growth rate:

\[ G_i = bK((H_{\infty}/H_i) - 1) \text{ [year}^{-1}] \]  

where, \( b \) is the exponent of the size–mass relation, \( K, H_{\infty} \) are parameters of the von Bertalanffy function, and \( H_i \) is the mean size in class \( i \).

Production-to-biomass ratio (\( P/B \)) was computed by dividing \( P \) by the biomass \( B \) of the size-frequency sample.

Mortality

Total mortality rate \( Z \) was estimated from the overall size–frequency distribution and the von Bertalanffy growth function parameters by a size-converted catch curve (Pauly, 1984a,b):

\[ N_i/\Delta t = N_0 e^{-Zt_i} \]  

where, \( N_i \) is the number of individuals in size class \( i \), \( \Delta t_i \) is the time required to grow through this size class and \( t_i \) is the relative age in the mid-size of class \( i \). Total mortality \( Z \) is computed by the linear regression:

\[ \ln(N_i/\Delta t_i) = a + bt_i; Z = -b \]  

Results

Significant exponential size–mass relationships between whole animal TM, SM, SFWM as dependent variables and H as the independent variable were found (Table 1).

Growth

Under reflecting light, polished shell cuts showed a pattern of alternating broad opaque and narrow translucent bands, which were also confirmed by acetate peels (Fig. 3). In general, the translucent bands showed a pink to purple colour, similar to the internal part of the shell.

Regarding the mark-recapture field experiment, only 10 animals (10.4%) were recovered alive. A further 18 (18.8%) were recaptured dead; all of them had gastropod drill holes in their shells. Analysis of the surviving specimens confirmed that one translucent growth band and one opaque growth band were formed each year. Seasonal growth terminal stage data (Table 2) showed that the translucent band was formed during autumn/winter (higher percentage of specimens forming translucent band), i.e. seasons with lowest sea temperature, and the opaque band was formed during spring/summer.

Table 1

<table>
<thead>
<tr>
<th>Dependent variable</th>
<th>Independent variable X</th>
<th>( a )</th>
<th>( b )</th>
<th>( r^2 )</th>
<th>( n )</th>
</tr>
</thead>
<tbody>
<tr>
<td>( \log(\text{TM}) )</td>
<td>( \log(\text{H}) )</td>
<td>-3.6201</td>
<td>3.2428</td>
<td>0.99</td>
<td>1913</td>
</tr>
<tr>
<td>( \log(\text{SM}) )</td>
<td>( \log(\text{H}) )</td>
<td>-3.4819</td>
<td>2.9965</td>
<td>0.99</td>
<td>1723</td>
</tr>
<tr>
<td>( \log(\text{SFWM}) )</td>
<td>( \log(\text{H}) )</td>
<td>-4.3728</td>
<td>3.3751</td>
<td>0.96</td>
<td>1723</td>
</tr>
</tbody>
</table>

\( L \), shell length; \( H \), shell height; \( \text{TM} \), total body mass; \( \text{SM} \), shell mass; \( \text{SFWM} \), shell-free wet mass.
Hence, individual age corresponds to the number of shell growth bands. The 285 size-at-age data obtained from internal ring readings were fitted best by the von Bertalanffy growth function (Fig. 4):

\[ H_t = 28.03 \text{ mm} \left( 1 - e^{-0.288(t+0.34)} \right); n = 285; r^2 = 0.83 \]

The clams can reach a maximum age of 15 years. Mean density was 1091 ± 737 ind. m\(^{-2}\). The annual shell height frequency data showed a peak at 2–6 mm and another peak at 23–28 mm (Fig. 5). A distinct peak of specimens smaller than 12 mm height occurred between December and March with smaller pulses the rest of the year (Fig. 6).

**Production**

Individual production of total soft tissues peaked at 19.5 mm shell height (0.42 g SFWM ind.\(^{-1}\) year\(^{-1}\), Fig. 7), whereas SM production was highest at 18.25 mm shell height (0.96 g SM ind.\(^{-1}\) year\(^{-1}\)). Annual production of the population was 120.45 g SFWM m\(^{-2}\) year\(^{-1}\), corresponding to an annual \(P/B\) ratio of 0.134 year\(^{-1}\) (mean biomass, \(B = 901.83\) g SFWM m\(^{-2}\)). Specimens between 21.5 and 28 mm shell height contributed most to population production (Fig. 5).

**Mortality**

The poor fit of the single negative exponential mortality model to the population data (Fig. 8) indicated that the mortality rate \(Z\) changes with age. Hence, an overall value representative for

<table>
<thead>
<tr>
<th>Season (month)</th>
<th>Percentage of animals in translucent growth stage</th>
<th>n</th>
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</thead>
<tbody>
<tr>
<td>Winter (July, August, September)</td>
<td>79</td>
<td>60</td>
</tr>
<tr>
<td>Spring (October, November, December)</td>
<td>65.66</td>
<td>60</td>
</tr>
<tr>
<td>Summer (January, February March)</td>
<td>33.08</td>
<td>60</td>
</tr>
<tr>
<td>Fall (April, May, June)</td>
<td>68.33</td>
<td>60</td>
</tr>
</tbody>
</table>

Shells were sectioned from the umbo to the ventral margin (axis of maximum growth) and terminal bands were examined by stereomicroscope using reflecting light.

Hence, individual age corresponds to the number of shell growth bands.

The 285 size-at-age data obtained from internal ring readings were fitted best by the von Bertalanffy growth function (Fig. 4):

\[ H_t = 28.03 \text{ mm} \left( 1 - e^{-0.288(t+0.34)} \right); n = 285; r^2 = 0.83 \]
shell height in *T. gayi*. Values for \( P_{\text{med}} \) in size classes above \( H_w = 28.03 \) were set to the value of \( P_{\text{max}} \) of the size class 27.75 mm the whole population cannot be computed. The tagging-recapture data – 18 of 18 recovered dead clams drilled by snails – indicate that predation by gastropods is a major source of mortality.

**Discussion**

The shell growth pattern of *T. gayi* in Ushuaia Bay – alternating narrow translucent and broad opaque bands (Fig. 3) – resemble growth patterns observed in other species of the genus *Tawera* (Luckens, 1990, 1991; Isla and Gordillo, 1996) as well as in other venerid bivalves such as *Eurhomaeda exalbida* (Lomovasky et al., 2002a,b), *Mercenaria campithiensis* (Jones et al., 1990; Arnold et al., 1996, 1998), *M. mercenaria* (Jones et al., 1990; Arnold et al., 1991, 1996, 1998) and *Venus verrucosa* (Arnieri et al., 1998).

The presence of growth rings indicates periods of very slow or even halted shell growth (Rhoads and Lutz, 1980) possibly caused by either (i) low metabolic rates related to low environmental temperatures (Grizzle and Lutz, 1988; Jones et al., 1990; Lomovasky et al., 2002a); (ii) a lack of food (Green, 1973; Beukema and Desprez, 1986; Beukema and Cadée, 1991; Kibe et al., 1996; Arneri et al., 1998); or by (iii) a diversion of metabolic products away from shell growth into gamete production (Morton, 1978; Ankur, 1980; Peterson and Fegley, 1986; Ramón et al., 1995; Arneri et al., 1998; Lomovasky et al., 2002a). In *T. gayi*, the translucent bands corresponding to slow or halted growth are formed in autumn/ winter, i.e. temperature (Fig. 2) and/or food availability are most likely to affect shell growth.

The growth parameters values of the von Bertalanffy function were estimated to be \( H_w = 28.03 \) mm (\( L_w = 34.30 \) mm; see Table 1 for conversion), \( K = 0.288 \) and \( t_0 = -0.34 \) (Fig. 4). Urban and Tesch (1996) determined distinctly different parameter values for *T. gayi* from Laredo Bay (51°58’S), Magellan Strait, \( L_w = 21.7 \) mm, \( K = 0.462 \) and \( t_0 = -0.13 \). These values, however, are most likely affected by biased sampling, as Urban and Tesch (1996) collected a ‘qualitative’ sample that only contained animals <21 mm shell height and that were under 5 years of age.

*Tawera gayi* from Ushuaia Bay (54°50’S; this paper) can reach a maximum age of 15 years, similar to *M. mawsoni* from Macquarie Island (54°29’–55°01’S), New Zealand (Luckens, 1990). Other maximum age estimates for *T. gayi* appear to be unreliable owing to non-representative sampling as well as non-validated ageing methods (Isla and Gordillo, 1996; Urban and Tesch, 1996).

The annual somatic production of *T. gayi* was 120.45 g SFWM m\(^{-2}\) year\(^{-1}\) (=456.51 kJ; converted by 3.79 kJ g\(^{-1}\) SFWM; Brey, 1999) with a mean biomass of 901.83 g m\(^{-2}\) (=3417.94 kJ m\(^{-2}\)), corresponding to a \( P:B \) ratio of 0.134 year\(^{-1}\). This value is as low as the \( P:B \) ratio reported for a sympatric population of another venerid, *E. exalbida*, 0.119 year\(^{-1}\) (Lomovasky et al., 2002a) and is among the lowest ever reported for venerid populations such as *Anomalocardia brasiliana* (Monti et al., 1991), *Callista brevisiphonata* (Selin and Selina, 1988), *Chamelea gallina* (Ramón and Richardson, 1992), *Chione cancellata* (Moore and Lopez, 1969), *Dosinia hepatica* (Hanekom, 1986), *M. mercenaria* (Hibbert, 1976; 1977a,b, Walker and Tenore, 1984), *Tapes philippinarum* (Yap, 1977), *Venerupis decussata* (Guelorget et al., 1980), *V. aurea* (Hibbert, 1976; Guelorget et al., 1980), *V. antiqua* (Clasing et al., 1994), *V. ovata* (Dauvin, 1985) and *V. striatula* (Warwick et al., 1978).

Recruits (animals <12 mm shell height, <1 year old) were present year-round (Fig. 6), with a major peak occurring in summer. The preliminary analysis of the reproductive cycle of *T. gayi* (E. Morriconi, pers. comm.) indicates that the observed recruitment pattern corresponds to the continuous presence of mature gonads throughout the year, with a major spawning event during November.

Mortality patterns within this population of *T. gayi* appear to be complex. Obviously, the single negative exponential mortality model does not apply and hence we cannot provide a population mortality estimate. The size-converted catch curve plot indicates, however, that mortality is quite high in small specimens and lower in larger specimens, which to some extent may be able to outgrow their predators. The dead shell assemblage along the coastline (about 95% of all shells with drill holes; pers. comm.) as well as our field experiment strongly indicate that gastropods such as *Xymentopsis mariniformis*, *Trophon geverianus* and *Natica* sp., are the major predators of *T. gayi* from Ushuaia Bay, a situation similar to the one observed in *T. mawsoni* from Macquarie Island, New
Zealand (Luckens, 1990). Tawera gayi, as other species of the genus Tawera from Australasian (Luckens, 1990, 1991), are usually found close the sediment surface. Thus, they are easier prey for snails than are the deeper-dwelling venerids such as E. exalbida (Lomovsky et al., 2002a). These highly mobile predators, together with the comparatively slow growth and low turnover of T. gayi in Ushuaia Bay, limit its potential for sustainable commercial exploitation.

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