Population dynamics of *Venus antiqua* (Bivalvia: Veneracea) in the Bahía de Yaldad, Isla de Chiloé, Southern Chile*

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Abstract

*Venus antiqua* is a dominant member of intertidal and shallow subtidal soft bottom communities in southern Chile, where it is a significant resource for artisanal fisheries. Growth, mortality and productivity of *V. antiqua* were investigated at an intertidal flat in Bahía de Yaldad, Chiloé. Based on growth increment data obtained from a 20 months tagging-recapture experiment, the parameters of the seasonally oscillating von Bertalanffy function were estimated to be \(L_x = 80.0\) mm, \(K = 0.183\) y\(^{-1}\), \(C = 1.211\) and \(t_x = 0.995\). Production of *V. antiqua* is distinctly higher than in other species of the superfamily Veneracea. Annual production amounted to 40.69 g AFDM m\(^{-2}\) y\(^{-1}\) (somatic tissue), 26.42 g AFDM m\(^{-2}\) y\(^{-1}\) (shell organic matter) and 2.00 g AFDM m\(^{-2}\) y\(^{-1}\) (gonad), respectively, corresponding to a P:B ratio of 0.585 y\(^{-1}\). Total mortality \(Z\), natural mortality \(M\) and fishing mortality \(F\) were estimated to be 0.664 y\(^{-1}\), 0.333 y\(^{-1}\) and 0.331 y\(^{-1}\), respectively. The high exploitation rate \(E\), 0.498, indicates serious overexploitation of the stock despite the closure of the fishery.

Key words: Bivalvia; Chile; Population dynamic; Production; Venus antiqua

1. Introduction

The infaunal filter-feeding bivalve *Venus antiqua* (King & Broderip, 1835) inhabits intertidal and shallow subtidal soft bottoms along the Pacific coast of South America from about 12\(^\circ\)S to 54\(^\circ\)S, but also along the Atlantic coast south of 31\(^\circ\)S, including the Malvinas (Falkland Islands) (Osorio et al., 1983). In Chile, the intertidal and subtidal banks of *V. antiqua* are exploited by local fishermen. During the last decade,
annual landings were in the order of 30,000 tons, most of which came from banks around “Isla de Chiloé”. *Venus antiqua* is also a food source for several bird species (mainly seagulls) and the predatory gastropod *Xanthochorus cassidiformis* (de Blainville, 1832). However, despite the significance of this clam for artisanal fisheries only a few investigations on the biology and ecology of *V. antiqua* have been carried out (see Jerez et al., 1991 and references therein).

The purpose of this paper is to examine the population dynamics of *V. antiqua* in the Bahía de Yaldad at the Isla de Chiloé and to evaluate the present state of its exploitation.

2. Methods

2.1. Sampling area

The study was carried out at a *V. antiqua* bank located on a tidal flat at Bahía de Yaldad (43° 07' S, 73° 44' W, Fig. 1), Isla de Chiloé, which is the subject of an interdisciplinary study of the ecology of tidal flats in southern Chile and the impact of artisanal fisheries on the system (see Stead, 1992; Zimmermann et al., 1993; Asencio et al., in press a, b; Navarro et al., 1993). The tidal flat (0.5% slope) has an area of about 1 km² at low tide, and the *V. antiqua* stock is concentrated in a 200 m wide band along MLWN. The amplitude of spring tides just exceeds 5 m, with a semidiurnal regime throughout the year. Sediment in the area of the *V. antiqua* bank is heterogeneous and characterized as gravely muddy sand (according to Folk, 1980), with the gravel fraction composed mainly (> 50%) of broken bivalve shells. Water temperature during the study period ranged from 6.0 °C in winter to 16.5 °C in summer. Salinity fluctuated around 28‰, occasionally dropping below 25‰ after strong rainfalls.

During previous years the *V. antiqua* bank was heavily exploited by local fishermen, with monthly catches reaching 6 tons wet mass and more (information provided by local dealers). However, exploitation was prohibited by the national authorities (Servicio Nacional de Pesca) in 1991, shortly after the start of this investigation, because all clams were below minimum size for legal extraction (55 mm).

2.2. Sampling

Specimens used for the establishment of length-mass relations were collected in December 1990, January, April, August 1991 and January 1992.

To estimate population density and biomass, quantitative samples were collected along a transect reaching from the low water level to the high water level at 13 dates between April 1990 and August 1991. Each transect consisted of five stations, and at each station two sediment cores of 25 × 25: cm area and 20 cm depth were dug out and sieved through 1.5 mm mesh size. All specimens of *V. antiqua* found were counted and measured.

To obtain growth increment data, specimens of *V. antiqua* were sampled, measured, marked with a number on the periostracum (black drawing ink covered with “Loctite Super Bonder Extra”) and returned to the original site at 10 dates between Novem-
Fig. 1. The investigation area. (A) Island of Chiloé, (B) Bay of Yaladad.
ber 1990 and September 1991. At 10 dates between May 15th 1991 and July 3rd 1992, specimens were recovered, measured and returned again to the tidal flat. The sampling schedule is given in Table 1.

Possible effects of repeated recovery on growth were checked by dividing all marked specimens in August 1991 into two groups of 200 animals each. The first group was recovered periodically, whereas the second group was left undisturbed until July 29th 1992, when the complete experimental area was intensely searched for all remaining clams. The growth increments of both groups were compared by ANOVA.

Growth of the newly settled cohort (i.e. age class 0) was followed in 1990/1991 and in 1991/1992 (Stead, 1992). Subsequent mean length data were used as additional size increment data pairs in the growth analysis.

### 2.3. Length-mass relations

Length was measured to an accuracy of 0.1 mm. Shell, somatic tissue and gonad tissue were weighed separately. Ash-free dry mass (AFDM) was determined by drying at 80 °C for 24 h (DM) and subsequent ignition at 500 °C for 2 h (ash). Analysis of covariance [log(M) vs log(L) and vs month] was used to test for differences among different sampling months, and length-mass (L-M) relations of the form

\[ M = a \times L^b \]

<table>
<thead>
<tr>
<th>Date</th>
<th>Newly marked</th>
<th>Re-measured</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 Nov. 1990</td>
<td>83</td>
<td>0</td>
</tr>
<tr>
<td>3 Dec. 1990</td>
<td>88</td>
<td>0</td>
</tr>
<tr>
<td>15 Jan. 1991</td>
<td>37</td>
<td>0</td>
</tr>
<tr>
<td>1 March 1991</td>
<td>28</td>
<td>0</td>
</tr>
<tr>
<td>30 March 1991</td>
<td>25</td>
<td>0</td>
</tr>
<tr>
<td>15 May 1991</td>
<td>36</td>
<td>135</td>
</tr>
<tr>
<td>14 June 1991</td>
<td>32</td>
<td>0</td>
</tr>
<tr>
<td>13 July 1991</td>
<td>55</td>
<td>0</td>
</tr>
<tr>
<td>10 Aug. 1991</td>
<td>164</td>
<td>236</td>
</tr>
<tr>
<td>10 Sept. 1991</td>
<td>59</td>
<td>0</td>
</tr>
<tr>
<td>10 Oct. 1991</td>
<td>0</td>
<td>202</td>
</tr>
<tr>
<td>23 Nov. 1991</td>
<td>0</td>
<td>216</td>
</tr>
<tr>
<td>18 Jan. 1992</td>
<td>0</td>
<td>202</td>
</tr>
<tr>
<td>18 Feb. 1992</td>
<td>0</td>
<td>142</td>
</tr>
<tr>
<td>18 March 1992</td>
<td>0</td>
<td>138</td>
</tr>
<tr>
<td>15 April 1992</td>
<td>0</td>
<td>105</td>
</tr>
<tr>
<td>16 May 1992</td>
<td>0</td>
<td>105</td>
</tr>
<tr>
<td>3 Jul. 1992</td>
<td>0</td>
<td>97</td>
</tr>
<tr>
<td>Total</td>
<td>607</td>
<td>1578</td>
</tr>
</tbody>
</table>
were established by an iterative nonlinear fitting algorithm (SIMPLEX, see Press et al., 1986).

2.4. Growth analysis

Growth was described by a seasonally oscillating von Bertalanffy growth curve according to Pauly & Gaschütz (1979) as modified by Hoenig & Choudary (1982) and Somers (1988):

\[ L_t = L_\infty \times (1 - e^{-K \times (t - t_0)} \times \frac{A}{B}) \]

\[ A = C \times \sin(2\pi \times (t - t_0)) / 2\pi \]

\[ B = C \times \sin(2\pi \times (t_0 - t_0)) / 2\pi \]

where \( L_\infty \) is asymptotic length, \( K \) is the growth constant, \( t_0 \) is age at which length is zero, \( C \) is the amplitude of growth oscillation, and \( t_0 \) is the starting point of the oscillation with respect to \( t = 0 \). The parameters of this function were estimated by fitting a re-arranged function (Appleoorn, 1987) to the size increment data pairs using the SIMPLEX algorithm:

\[ L_2 = L_\infty \times (1 - (1 - L_1/L_\infty) \times e^{-K \times (t_1 - t_0)} \times \frac{A'}{B'}) \]

\[ A' = C \times \sin(2\pi \times (t_1 - t_0)) / 2\pi \]

\[ B' = C \times \sin(2\pi \times (t_2 - t_0)) / 2\pi \]

For \( t_0 \), no estimate can be obtained from growth increment data.

2.5. Production

Somatic production was calculated by the mass-specific growth rate method (see Crisp, 1984) from (1) an average length-frequency sample based on the pooled samples, (2) the von Bertalanffy growth function and (3) the length-mass relation:

\[ P = \Sigma N_i \times M_i \times G_i \quad \text{[g AFDM} \cdot \text{m}^{-2} \cdot \text{y}^{-1}] \]

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

\[ G_i = b \times K \times ((L_\infty / L_i) - 1) \quad \text{[y}^{-1}] \]

where \( b \) is the exponent of the length-mass relation, \( K \), \( L_\infty \) are parameters of the von Bertalanffy function, and \( L_i \) is the mean length in class \( i \). Annual \( P/B \) ratio was calculated from production \( P \) and biomass \( B \):

\[ B = \Sigma N_i \times (M_i + MG_i) \quad \text{[g AFDM} \cdot \text{m}^{-2}] \]

where \( MG_i \) is annual average individual gonad mass in length class \( i \).
Gonad production was computed from the length-gonad mass relations in the pre-spawning and the post-spawning periods and the pooled length-frequency distributions:

$$PG = \sum N_i \times (MG_{i,pre} - MG_{i,post})$$

where $MG_{i,pre}$ and $MG_{i,post}$ are gonad mass in length class $i$ before and after spawning, respectively.

2.6. Natural, fishing & total mortality and exploitation rate

Natural mortality rate $M$ had to be estimated on an empirical basis. In unexploited populations $M$ equals total mortality $Z$ since fishery mortality $F$ is zero. Allen (1971) showed that in a steady state population the somatic $P/B$ ratio is equal to the total mortality rate $Z$, if mortality can be described by the single negative model and if individual growth follows a von Bertalanffy function. Hence natural mortality of $V. antiqua$ can be estimated from empirical relations between population parameters and annual $P/B$ ratio in unexploited benthic invertebrate populations. We estimated maximum age $A_{Max}$ from an empirical relationship between $A_{Max}$ and the growth constant $K$ of the von Bertalanffy function in mollusc populations (Table 3), and $M$ from empirical relations between $P/B$ ratio and $A_{Max}$ (Table 3).

A size-converted catch curve was used to compute total mortality rate $Z$ of the single negative exponential mortality model:

$$N_i = N_0 \times e^{-Z \times t_i}$$

This curve was calculated from the size-frequency distribution of the pooled quantitative samples and the von Bertalanffy growth function by the ELEFAN (Electronic Length Frequency Analysis) program of D. Pauly and co-workers (see Brey et al., 1988, 1990; Gayanilo et al., 1989):

$$\frac{(N_i - N_i)}{\Delta t} = N_0 \times e^{-Z \times t_i}$$

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

$$\log_e(N_i/\Delta t_i) = a + b \times t_i; \quad Z = -b$$

Fishing mortality rate $F$ and actual exploitation rate $E$ were computed by

$$F = Z - M \quad \text{and} \quad E = F/Z.$$  

3. Results

3.1. Length-mass relations

Neither in ash free dry mass of somatic tissue ($p = 0.07$) and of the shell ($p = 0.55$) nor in the inorganic shell mass (i.e. ash, $p = 0.68$) significant differences in the length-
mass relations among the different months could be detected. Fig. 2 and Table 2 show the length-mass relations for the combined data. Gonad mass was determined in a few animals (≤15) per sampling date only. To increase the reliability of the length-gonad

Table 2
Length-mass relations in *V. antiqua*

(A) Somatic and shell mass

<table>
<thead>
<tr>
<th>Mass</th>
<th>N</th>
<th>a</th>
<th>b</th>
<th>r²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Somatic tissue (mg AFDM)</td>
<td>152</td>
<td>0.030</td>
<td>2.678</td>
<td>0.947</td>
</tr>
<tr>
<td>Shell (mg AFDM)</td>
<td>152</td>
<td>0.006</td>
<td>2.7^-6</td>
<td>0.946</td>
</tr>
<tr>
<td>Inorganic shell (mg Ash)</td>
<td>152</td>
<td>0.117</td>
<td>3.020</td>
<td>0.973</td>
</tr>
</tbody>
</table>

(B) Gonad mass

<table>
<thead>
<tr>
<th>Date</th>
<th>N</th>
<th>a</th>
<th>b</th>
<th>r²</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dec. 90, Jan. 91, Jan. 92</td>
<td>44</td>
<td>0.035</td>
<td>2.049</td>
<td>0.109</td>
</tr>
<tr>
<td>April 91, Aug. 91</td>
<td>23</td>
<td>4.66 x 10^-11</td>
<td>7.488</td>
<td>0.577</td>
</tr>
<tr>
<td>All sampling dates</td>
<td>67</td>
<td>0.305</td>
<td>2.678</td>
<td>0.137</td>
</tr>
</tbody>
</table>

Mass = a x length\(^b\).
Units of length are mm.
r\(^2\) = 1 - residual sum of squares/total sum of squares.
mass relations, data from April and August were pooled to represent pre-spawning data, whereas December and January were pooled to represent post-spawning data. These two relations differ significantly ($p = 0.01$, Table 2B).

3.2. Growth

During the investigation period, 607 specimens of *V. antiqua* were tagged and a total of 1578 re-measurements of length were done. The comparison of specimens recovered several times between August 91 and July 92 with those recovered only once showed no significant differences in size increments (ANOVA, $N_1 = 76$, $N_2 = 69$, $p = 0.928$).

All size increment data pairs of $L_1$ and $L_2$ with $\Delta t < 0.3$ years ($N = 1135$) were used for growth analysis. Additionally, 19 data pairs of $L_1$ and $L_2$ from the newly settled cohorts were included (Fig. 3). Fig. 4 shows that the daily length-increments vary among seasons, they are highest in spring and summer, lower in autumn and lowest in winter, indicating strong seasonality in growth.

The parameter values of the seasonally oscillating von Bertalanffy growth function were estimated to $L_x = 71.22$ mm, $K = 0.224$ year$^{-1}$, $C = 1.224$ and $t_x = 0.996$ ($N = 1154$, Residual Sum of Squares = 553.3, $[1 - \text{RSS/TSS}] = 0.997$). Since the obtained value for $L_x$ is below the maximum length observed in subtidal areas at the Bahía de Yaldad (see Discussion), a second fit with $L_x$ fixed at 80.0 mm was performed. It yielded slightly different parameter values of $K = 0.183$ year$^{-1}$, $C = 1.211$ and $t_x = 0.995$, but the goodness of fit was nearly the same (Residual Sum of Squares = 562.4, $[1 - \text{RSS/TSS}] = 0.996$). These parameter values were used for further computations of productivity and mortality. At values of $C > 1$ the oscillating growth curve shows negative

![Fig. 3. Growth of the newly settled cohorts of *V. antiqua* in 1990/1991 and 1991/1992. Vertical bars indicate ± 1 SD (data from Stead, 1992).](image-url)
growth during a certain period due to technical reasons (sinus function), in reality there is zero growth during this time (Figs. 4 & 5).

3.2.1. Productivity

Individual somatic tissue production as well as shell organic matter production increase steadily up to about 310 mg AFDM·ind⁻¹·y⁻¹ and 240 mg AFDM·ind⁻¹·y⁻¹ at 50–55 mm length and then decrease again. Calculated gonad production starts at about 40 mm length and increases exponentially towards the highest length classes, exceeding total non-gonad production above 55 mm length (Fig. 6).

Annual production of the population was computed to be 40.69 g AFDM·m⁻²·y⁻¹ (somatic tissue), 26.42 g AFDM·m⁻²·y⁻¹ (shell organic matter) and 2.00 g AFDM·m⁻²·y⁻¹ (gonad), respectively. Fig. 7 shows the distribution of production among the length classes. Mean annual biomass amounts to 118.06 g AFDM·m⁻² (mean abundance = 227.6 ind·m⁻²), hence annual P/B ratio is calculated to be 0.585 y⁻¹.
Fig. 5. Seasonally oscillating von Bertalanffy growth curve fitted to 1154 data pairs of $L_t$, $L_m$ ($L_m$ fixed at 80 mm, $K = 0.183$, $C = 1.211$, $t_0 = 0.995$; $N = 1154$). Residual sum of squares = 562.4, $[1 - \text{RSS/TSS}] = 0.996$). Superimposed are subsequent growth-increment data of five specimens of $P. antiques$. Each symbol refers to a different specimen. The number of data points differs among specimens due to the different sampling schedules. The x-axis is denoted relative age since age at length zero ($t_0$) is unknown.

Fig. 6. Relation between length (mm) and individual somatic and gonad production (mg AFDM·y⁻¹). ⬜, somatic tissue; Δ, shell organic matter; □, gonads. See chapter "methods" for computation procedures.
3.3. Mortality and exploitation rate

Maximum age $A_{\text{Max}}$ of $V. \text{antiqua}$ at Yaldad was estimated to be 10.4 years by the empirical relation between $A_{\text{Max}}$ and $K$ (Table 3), which is a reasonable estimate with respect to our growth curve (Fig. 5). Subsequently, natural mortality rate $M$ was estimated to be 0.333 yr$^{-1}$ (mean of 0.341, 0.423, 0.236 from equations in Table 3).

Total mortality $Z$ was estimated to be 0.664 yr$^{-1}$ by the size converted catch curve shown in Fig. 8. Hence, fishing mortality $F$ was 0.331 yr$^{-1}$ and exploitation rate $E$ was 0.498.

Table 3

<table>
<thead>
<tr>
<th>Author</th>
<th>$X$</th>
<th>$Y$</th>
<th>Intercept $a$</th>
<th>Slope $b$</th>
<th>Number of data</th>
<th>$r^2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Brey (unpubl.)$^1$</td>
<td>$K$</td>
<td>$A_{\text{Max}}$</td>
<td>0.450</td>
<td>-0.768</td>
<td>89</td>
<td>0.566</td>
</tr>
<tr>
<td>Brey (unpubl.)$^1$</td>
<td>$A_{\text{Max}}$</td>
<td>$P/B$</td>
<td>0.682</td>
<td>-1.130</td>
<td>219</td>
<td>0.767</td>
</tr>
<tr>
<td>Hoening (1983)</td>
<td>$A_{\text{Max}}$</td>
<td>$P/B$</td>
<td>0.625</td>
<td>-0.982</td>
<td>134</td>
<td>0.820</td>
</tr>
<tr>
<td>Warwick (1980)$^2$</td>
<td>$A_{\text{Max}}$</td>
<td>$P/B$</td>
<td>0.538</td>
<td>-0.112</td>
<td>85</td>
<td>0.567</td>
</tr>
</tbody>
</table>

$log_\text{e}(Y) = a + b \times log_\text{e}(X)$; $^1$ Molluscs only; $^2$ $log_\text{e}(Y) = a + b \times X$. 
4. Discussion

4.1. Growth

The computed oscillating von Bertalanffy growth curve (Fig. 5) shows an excellent fit to the data with the exception of specimens \( \leq 10 \) mm length (age class 0), where growth seems to be generally overestimated by the above function. This indicates that a growth function with inflexion point such as the Gompertz function may fit the data better in this particular size range. Our first estimate of \( L_x = 71.22 \) mm, underestimates the true \( L_x \), since specimens up to 80 mm length are known from subtidal areas of the Bahia de Yaldad as well as from other areas of southern Chile (Bustos et al., 1981; Jerez et al., 1991). In the intertidal of Yaldad, however, these large specimens do not occur due to the impact of artisanal fishermen. Hence, they are missing from our growth increment data set (Fig. 4), leading to an under-estimation of \( L_x \), as indicated by the almost equally good fit of the growth curve obtained with \( L_x \) fixed at 80 mm.

Our growth parameter values differ clearly from those estimated for a population of \( V. \) antiqua in the Bahia de Ancud, \( \approx 150 \) km north of Yaldad. From growth ring readings, Bustos et al. (1981) estimated a non-oscillating von Bertalanffy growth curve with \( L_x = 78.7 \) mm and \( K = 0.421 \) y\(^{-1}\). Jerez et al. (1991) estimated \( L_x = 102 \) mm and \( K = 0.421 \) y\(^{-1}\) by a method based on the analysis of length-frequency samples. Bustos et al. (1981) assumed \( V. \) antiqua to form two growth rings per year as stated by Estay et al. (1979), but did not validate this assumption. There is new evidence from recent
investigations that *V. antiqua* in the sublitoral of the Bahía de Ancud forms only one growth ring per year during autumn/winter (C. Rojas, pers. comm.), as observed in our experiments in Bahía de Yaldad, too. Consequently, Bustos et al. (1981) may have overestimated the growth constant *K* by about 100%. We therefore conclude that *V. antiqua* around Isla de Chiloé will reach size at first reproduction (46–48 mm, Lozada & Bustos, 1984) at an age of 4–5 years, whereas size of permitted legal extraction (55 mm) will be reached at 5–6 years of age.

The observed strong seasonal pattern of growth in *V. antiqua* with highest growth rates in spring and summer and lowest growth rates during winter may be related to the similar annual pattern of primary production in southern Chile waters (Navarro et al., 1993).

4.2. Productivity

Annual somatic P/B ratio of *V. antiqua* in the intertidal of Yaldad, 0.572 yr⁻¹, is in the upper range of values referring to various species of the superfamily Veneracea, if the relation between population P/B ratio and mean body mass is taken into account (Fig. 9). The above-average P/B ratio may be explained best by increased mortality due to human exploitation, since P/B ratio and mortality rate are more or less equivalent (Allen, 1971). Under natural conditions, P/B ratio is expected to be equal to natural

![Graph](image-url)

**Fig. 9.** Somatic productivity of *V. antiqua* at Yaldad compared to other species of the superfamily Veneracea. The relation between P/B ratio and mean body mass (converted to kg) is described by log(P/B) = -0.738 - 0.729 × log(Mean Body Mass), N = 14, r² = -0.746 1. *Chione cancellata*, Florida/USA, Moore & Lopez (1969); 2, *Mercenaria mercenaria*, Southampton/UK, Hibrert (1976); 3, *Mercenaria mercenaria*, Georgia/USA, Walker & Tenore (1984); 4, *Venerupis aequa*, Southampton/UK, Hibrert (1976); 5, *V. ovata*, Morlaix Bay/Spain, Dauvin (1985); 6, *V. striatula*, Bristol Channel/UK, Warwick et al. (1978); [V] *V. antiqua*, Yaldad, this study.
mortality rate $M$, which has been estimated to 0.331 y$^{-1}$. This figure would fit much better in the plot of log(P/B) vs log(body mass), even if mean body mass is expected to increase due to reduced mortality.

Annual somatic production, 42 g AFDM·m$^{-2}$, is three times higher in *V. antiqua* than in that species coming closest to this figure, the population of *Mercenaria mercenaria* at Southampton, 14 g AFDM·m$^{-2}$ (recomputed from Hibbert, 1976). Two factors are likely to be significant for this high production: the comparatively long period of high primary production at Bahía de Yaldad, $\approx$10 wk; and the high levels as well as high nutritional value of suspended organic matter at this particular tidal flat (Navarro et al., 1993). Especially bacteria appear to provide a significant additional food source for *V. antiqua* throughout the year (Zimmermann et al., 1993).

### 4.3. Mortality and state of exploitation

In that part of the *V. antiqua* population at Yaldad living in the intertidal, fishing mortality $F$ (0.331 y$^{-1}$) equals natural mortality $M$ (0.331 y$^{-1}$), and the resulting exploitation rate of 0.5 indicates a high fishing pressure on the stock, despite the closure of the fishery since 1991. Gulland (1971) suggested $E = 0.5$ (i.e. $F = M$) to represent optimum exploitation conditions in fish stocks. However, Beddington & Cooke (1983), Caddy & Csirke (1983), Francis (1974) and others showed that this assumption generally overestimates potential yield by a factor of 2 or even higher, i.e. $E = 0.5$ is far above optimum exploitation conditions. These considerations are likely to be valid for benthic invertebrate stocks, too, since there are no differences in the underlying principles of population dynamics and fisheries. Hence, *V. antiqua* in the Bahía de Yaldad seems to be in a state of overexploitation, comparable to the Bahía de Aucel, where *V. antiqua* was found to be exploited by a rate 40% above optimum level (Jerez et al., 1991).

It is obvious from this study that the measures taken against overexploitation by the authorities are ineffective to control the fishery. Unfortunately, these are not isolated phenomena. Although total landings of *V. antiqua* in southern Chile keep on rising (Anonymous, 1991), the productivity to yield ratio is descending rapidly and average size of extracted specimens is also diminishing. These observations indicate continuous overexploitation of the traditional fishing areas around Isla de Chiloé (Reyes et al., 1991). Hence, there is an urgent need for effective management of the *V. antiqua* stocks and the Venus fishery to prevent the coastal ecosystem from permanent damage and to maintain the significance of this species for the local artisanal fisheries.

### 5. Acknowledgements

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6. References


