Population dynamics and metabolism of *Aequipecten opercularis* (L.) from the western English Channel (Roscoff, France)

Olaf Heilmayer*, Thomas Brey, Daniela Storch, Andreas Mackensen, Wolf E. Arntz

*Alfred Wegener Institute for Polar and Marine Research, P.O. Box 120161, 27515 Bremerhaven, Germany*

Received 17 March 2003; accepted 18 July 2003

**Abstract**

Population dynamics of the queen scallop, *Aequipecten opercularis*, was investigated in the western English Channel off Roscoff. Annual formation of natural growth rings on the shell surface was validated by stable isotope (δ¹⁸O and δ¹³C) analysis. A von Bertalanffy growth function \( H_t = 58.9 \text{ mm} \times (1 - e^{-0.604(t + 0.235)}) \) was fitted to size-at-age data of 249 individuals. Annual somatic and gonad production amounted to 19.74 kJ m⁻² and 0.98 kJ m⁻² y⁻¹, respectively. Total mortality rate \( Z \) was estimated to be 1.716 y⁻¹. Net growth efficiencies (ranging from 45% in 1-y-old to 11% in 6-y-old individuals) were in the same range as in other short-lived scallops. Individual growth, however, was distinctly slower in this population than in other *A. opercularis* populations from similar latitudes, most likely due to a more stressful environment.

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**Keywords:** Bivalvia; Energy budget; Metabolism; Mortality; Pectinidae; Population dynamics

**1. Introduction**

The commercially exploited queen scallop *Aequipecten opercularis* (L.) occurs throughout a wide geographical range on the European continental shelf: from northern Norway to the Mediterranean and from the Adriatic to the west of Ireland (Broom, 1976; Brand, 1991; Waller, 1991). Recruitment success and spatial distribution appear to be highly variable in time, thus inducing major year-to-year changes in extent and position of scallop banks of exploitable size and density, as reported from British waters (Broom, 1976; Brand, 1991) as well as from the eastern part of the English Channel and from northern Brittany (Grainger, 1992; Goulletquer and Heral, 1997). Due to the relatively short time *A. opercularis* can be kept fresh (Ansell et al., 1991), queen scallops support only a small local fishery and market in France, which is about 1/8 of total French scallop landings (Grainger, 1992).

In contrast to the great scallop (*Pecten maximus* L.), interest in population dynamic parameters of *A. opercularis* has hitherto been limited. Paul (1980) investigated salinity and temperature tolerances, Vahl (1972) described the relationship between water transport and oxygen uptake, and McLusky (1973) determined the effect of temperature acclimation on filtration and oxygen consumption rates.
Data on age and growth have been reviewed by Taylor and Venn (1978). Little is known, however, on population biology and production of the queen scallop populations in the western English Channel, besides the growth rates determined by Franklin et al. (1980).

A baseline study on the biology and population dynamics of *A. opercularis* in the western English Channel is presented. In this study growth, mortality, age-specific somatic and gonadal production as well as metabolic rates in individual scallops were calculated and compared with previous findings in the literature.

2. Material and methods

2.1. Study site and general measurements

Queen scallops (*Aequipecten opercularis*) were collected with a small dredge in the western English Channel next to Roscoff (Trezen Vraz, 48°59’ N, 4°53’ W) in 60 to 80 m water depth at three times (February, June and October) in 1999. Water temperature varied between 9.7 °C in January/February and 16.4 °C in August (Fig. 1). The summer phytoplankton bloom consisted of two peaks, with a maximum of 1.53 µg l⁻¹ in April, and a second one amounting to 1.56 µg l⁻¹ by late August. Minimum chlorophyll concentrations occurred in February (SOMLIT-Database). The investigation area, especially the distribution of substrate types and phytoplankton, is described in Sagan and Thouzeau (1998).

After sampling, animals were immediately transported to the “Station Biologique de Roscoff”, where they were maintained in flow-through aquaria until used in experiments or transported to the Alfred Wegener Institute (AWI, Germany). On each occasion shell height (= defined as the maximum distance between the dorsal hinge and ventral margin, Seed, 1980) of a representative subsample was measured to the nearest 0.1 mm using vernier callipers. A size frequency distribution (SFD) was obtained by pooling individuals using class intervals of 1 mm size. For size-mass relationships freshly caught animals were dissected and oven dried at 60 °C for 24 h. Individual gonad and soft tissue dry masses (DM) were weighed to the nearest 0.1 g, linear regression analysis was carried out on log-transformed data, estimating the constants a and b of the allometric equation:

\[ \log(M) = \log(a) + b \cdot \log(H) \]  

where *M* is soft tissue mass (g) of somatic or gonad tissue, respectively, at shell height *H* (mm).

2.2. Growth band reading and analysis

Bivalve shell growth bands can result from seasonal oscillations in growth, but also from changes in food availability, spawning events or predation attempts. Identification and interpretation of annual growth bands in *A. opercularis* were based on previous studies in this species (Broom and Mason, 1978; Paul, 1980; Hickson et al., 1999) as well as other scallop species (e.g. Dare and Deith, 1990; Heilmayer et al., 2003). For growth band analysis only the left (upper) valve was used. Prior to analysis the shell was cleaned of organic matter with warm 5% NaOCl solution, washed with 96% ethanol, rinsed with water and dried at 60 °C for 12 h. In each animal, the number of macroscopically visible bands and shell height were recorded. Stable oxygen (*δ¹⁸O*) isotope values in biogenic CaCO₃ reflect ambient water temperature and oxygen isotopic composition of seawater, the latter varying with salinity, while carbon (*δ¹³C*) isotope values reflect primary production at the

![Fig. 1. Sea surface water temperatures in Roscoff (1999: circles; 2000: dots; line: cubic spline through two-year mean values). (Data kindly provided by the Service d’Observation en Milieu Littoral-SOMLIT-INSU-CNRS.)](image-url)
time of carbonate formation (Erez and Luz, 1983; Hickson et al., 1999). Due to low salinity variations at the site of investigation, the oxygen isotopic composition of seawater should be primarily a reflection of ambient temperature variation. Therefore isotope ratio profiles reflect variations in temperature, shell physiology (related to temperature), primary production and a combination of these (see Wefer and Berger, 1991; Owen et al., 2002a). In environments with a seasonal cycle in temperature and/or primary production such isotope ratios can be used to validate seasonal cycle in temperature and/or primary production (e.g. Krantz et al., 1984; Dare and Deith, 1990; Hickson et al., 1999; Lasta et al., 2001; Heilmayer et al., 2003).

For isotope analysis, calcium carbonate powder was sampled from the outer shell layer of four individuals in equally spaced (≈ 1 mm) dorso-ventral series using a small dental drill (bit size 0.5 mm). δ18O and δ13C values were determined in the Stable Isotope Laboratory of the Alfred Wegener Institute with a Finnigan MAT251 mass spectrometer coupled to an automatic carbonate preparation device. The results were reported in ppt (% deviations related to the Vienna Pee Dee belemnite (VPDB) standard through repeated analyses of National Institute of Standards and Technology (NIST) isotopic reference material 19. The precision of measurements was better than ± 0.08% and ± 0.06% for δ18O and δ13C, respectively, based on routine measurements of a laboratory-working standard.

Assuming that stable isotope analysis will prove the annual formation of one shell growth band each winter (see below) individual age in years was taken to equal the number of growth bands X in animals sampled in February. Animals caught in June and October were assumed to be X + 0.33 y and X + 0.67 y old, respectively. A von Bertalanffy growth model (VBGF) was fitted to the resulting size-at-age data pairs by using a non-linear iterative Newton algorithm (for details Brey, 2001):

\[ H_t = H_\infty \cdot \left(1 - e^{-K \cdot (t-t_0)}\right) \quad [\text{mm, y}] \quad (2) \]

where \( H_\infty \) is the mean asymptotic height, \( K \) is the Brody growth coefficient, \( t \) the age and \( t_0 \) is theoretical age at which shell height equals zero.

### 2.3. Production (P)

Age specific individual somatic production (\( P_S \)) was calculated from the increment in dry tissue mass between consecutive age classes \([DM_{t+1} - DM_t]\) (with 1 g dry mass = 20.22 kJ; Brey, 2001).

The mass-specific growth rate method could not be used to compute somatic production-to-biomass ratio (\( P/S \)) and production of the population, because animals < 35 mm H were strongly under-represented in our size frequency distribution owing to the comparatively large 25 mm × 25 mm mesh size of the dredge. This problem was circumvented by using the annual mortality rate \( Z \) (see below) as a proxy for the somatic production-to-biomass ratio \( P_S/B \), as \( Z \) and \( P_S/B \) are equal in steady-state (see Allen, 1971) and the catch curve approach to estimate \( Z \) is quite robust against the effects of gear selectivity (Pauly and Munro, 1984). The queen scallop bed at Trezen Vraz is not ephemeral but has been present for at least several decades, as anecdotal evidence (local fishermen reports) and multi-year scientific research (e.g. Sagan and Thouzeau, 1998) indicate. Hence, the assumption of steady-state appears to be justified.

Consequently annual somatic production of the population was estimated by:

\[ P_S = Z \cdot M_S \cdot N \quad [\text{kJ m}^{-2} \text{y}^{-1}] \quad (3) \]

where \( M_S \) is the average individual somatic body mass in our sample and \( N \) is the mean abundance of 0.82 ind m\(^{-2}\) provided by Franklin et al. (1980) for the western English Channel.

Gonad production (\( P_{G,i} \)) for each age class was estimated from the decline of gonad mass during spawning in scallops of given age (determined from the von Bertalanffy equation describing shell height as a function of age), determined from logarithmic regressions of gonad mass to size before \((M_{G,i} \text{ before})\) and after spawning \((M_{G,i} \text{ after})\). June) (1 g gonad DM = 23.4 kJ; average from: MacDonald and Thompson, 1985; MacDonald et al., 1991):

\[ P_{G,i} = M_{G,i} \text{ (before)} - M_{G,i} \text{ (after)} \quad [\text{kJ ind}^{-1} \text{y}^{-1}] \quad (4) \]
Individual gonad production is negligible below 30 mm H (see Results). Therefore population $P_G/B$ ratio was computed from gonad production of total sample and biomass of total sample:

$$P_G/B = (\Sigma N_i \cdot P_{G,i})/(\Sigma N_i \cdot M_{S,i}) \quad [y^{-1}]$$

(5)

where $M_{S,i}$ is the average individual body mass in class $i$. Hence population gonad production ($P_G$) can be estimated by:

$$P_G = (P_G/B) \cdot M_S \cdot N \quad [kJ \ ind^{-1} y^{-1}]$$

(6)

2.4. Mortality rate ($Z$)

The total mortality rate $Z$ was estimated from the SFD and the von Bertalanffy growth curve by a size-converted catch curve (Pauly and Munro, 1984; Brey, 2001):

$$\ln(N_i/\Delta t_i) = a + b t_i \quad \leftrightarrow \quad Z = -b$$

(7)

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 at the mid-size of class $i$.

2.5. Metabolism ($R$)

Oxygen consumption rates of unfed (deprived of food for at least 3 days), unstressed and inactive animals were used as a proxy of standard metabolic rate as defined by Bayne and Newell (1983) (a detailed description of animal handling is given in Heilmayer and Brey, 2003). Measurements of scallops sampled in June (summer) and October (autumn) were done at ambient temperatures within 5–20 days after sampling. Winter rates were measured on animals sampled in October 1999 and maintained under simulated winter conditions (reduced food, dimmed light) in temperate aquaria with recirculating seawater ($10 \pm 0.5 \, ^\circ C, 33–34 \, psu$). A size range of animals as large as possible was used on each sampling occasion.

Oxygen content was assessed using an intermitted flow system and oxygen microoptodes connected to a Microx 8-array, as previously described in Gatti et al. (2002). Individual metabolic rates ($VO_2$) were obtained by comparison with control chambers (no animals). $VO_2$ was expressed as a function of soft tissue dry mass ($DM$):

$$\log(VO_2) = a + b \cdot \log(DM)$$

(8)

where $a$ is a constant and $b$ the mass scaling exponent. The model was fitted by least squares linear regression after logarithmic transformation of both variables. In order to compare oxygen consumption among seasons, rates were standardised for body mass (Packard and Boardman, 1999) by:

$$VO_{2,S} = VO_2 \cdot (DM_S/DM_E)^b$$

(9)

where $VO_{2,S}$ is the metabolic rate of a standard-sized scallop of 1 g dry mass ($DM_S$), and $VO_2$ and $DM_E$ are observed oxygen rate and body mass of the experimental scallop. Oxygen consumed was converted to energy by 1 ml O$_2$ = 20.1 J (cited in Brey, 2001).

3. Results

3.1. General measurements

The pooled size-frequency distribution of the year 1999 (Fig. 2) was polymodal and dominated by...
(>70%) by large individuals (>41 mm). An ANCOVA showed significant (p < 0.001) seasonal changes in the regressions of somatic and gonad dry mass against shell height (Table 1, both log-transformed). Both somatic dry mass and gonad dry mass of a standard-sized 45 mm scallop were lowest during the summer period (soma in g DM: 0.371 in June vs. 0.583 in February and 0.847 in October; gonad in g DM: 0.0122 in June vs. 0.0465 in February and 0.0204 in October).

3.2. Age and growth

\[ \delta^{18}O \] and \[ \delta^{13}C \] profiles of the two shells examined showed more or less synchronous cyclical patterns reflecting environmental seasonality (Fig. 3). Over the whole lifetime, \[ \delta^{18}O \] values obtained ranged from \(-0.3\) to \(+1.1\) \% (Fig. 3a) and \(-0.4\) to \(+1.3\) \% (Fig. 3b). \[ \delta^{13}C \] values ranged from \(-0.5\) to \(+0.4\) \% (Fig. 3a), and from \(-0.5\) to \(+0.5\) \% (Fig. 3b). There is a distinct spatial coincidence between macroscopically identified growth bands and relative \[ \delta^{18}O \] and \[ \delta^{13}C \] maxima of the isotope profile (i.e. temperature as well as primary production minima), thus indicating a formation of one growth band each winter.

Size-at-age data of 249 specimens were fitted best by the von Bertalanffy equation:

\[ H_t = 58.9 \text{ mm} \cdot (1 - e^{-0.604 \cdot (t + 0.235)}); \ r^2 = 0.827 \] (Fig. 4). Accordingly, the largest scallop (\( H = 58.9 \) mm) found at Roscoff was about six years old.

<p>| Table 1 |
| Allometric relationships of tissue mass and shell height in <em>Aequipecten opercularis</em> collected in 1999 from Trezen Vraz |</p>
<table>
<thead>
<tr>
<th>February</th>
<th>June</th>
<th>October</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Soma</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>a</td>
<td>-4.151</td>
<td>-4.921</td>
</tr>
<tr>
<td>b</td>
<td>2.369</td>
<td>2.716</td>
</tr>
<tr>
<td>( r^2 )</td>
<td>0.41</td>
<td>0.76</td>
</tr>
<tr>
<td>N</td>
<td>11</td>
<td>27</td>
</tr>
<tr>
<td><strong>Gonad</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>a</td>
<td>-12.024</td>
<td>-10.078</td>
</tr>
<tr>
<td>b</td>
<td>6.467</td>
<td>4.938</td>
</tr>
<tr>
<td>( r^2 )</td>
<td>0.18</td>
<td>0.89</td>
</tr>
<tr>
<td>N</td>
<td>11</td>
<td>27</td>
</tr>
</tbody>
</table>

Regression are of the form \( \log(M) = a + b \cdot \log(SH) \), where \( M \) is the predicted soft tissue dry mass (g) of the somatic tissue or of the gonad, \( SH = \) shell height (mm), and \( a \) and \( b \) are fitted parameters. \( N \) is number of data, \( r^2 \) is the determination coefficient. All equations are statistically significant at \( p < 0.001 \).
3.3. Production and mortality

Individual somatic production increased to a maximum of 10.66 kJ yr\(^{-1}\) (0.527 g DM yr\(^{-1}\)) at an age of 2 years and decreased thereafter, whereas individual gonad production increased exponentially with shell height, reaching 4.17 kJ yr\(^{-1}\) (0.177 g DM yr\(^{-1}\)) at an age of 6 years (Fig. 5a). Mean annual biomass of *A. opercularis* was 13.67 kJ m\(^{-2}\) (0.67 g DM m\(^{-2}\)), total mortality rate \(Z\) amounted to 1.761 yr\(^{-1}\) (Fig. 6). Correspondingly, annual somatic production \(P_s\) was estimated at 19.74 kJ m\(^{-2}\) yr\(^{-1}\) (0.976 g DM m\(^{-2}\) yr\(^{-1}\)). Annual gonad productivity \(P_G/B\) and production \(P_G\) amounted to 0.087 yr\(^{-1}\) and 0.98 kJ m\(^{-2}\) yr\(^{-1}\) (0.042 g DM m\(^{-2}\) yr\(^{-1}\)), respectively. Hence, total annual production for the 1999 season amounted to 20.72 kJ m\(^{-2}\) yr\(^{-1}\) (1.02 g DM m\(^{-2}\) yr\(^{-1}\)), corresponding to an annual \(P_{tot}/B\) ratio of 1.848 yr\(^{-1}\). The turnover ratio \(P/B\) calculated for each age class declined with age from 1.45 to 0.14 (Fig. 5b). Similarly \(P/R\) ratio and net growth efficiency \(K_2 = P/A = P/(P + R)\) decreased but levelled off in the oldest age classes (Fig. 5b).

3.4. Metabolism

Metabolic rates were measured in June (\(T = 12.4^\circ C\)), October (\(T = 14.4^\circ C\)) and December (\(T = 10.3^\circ C\)) over the complete size range of scallops available at Roscoff (June: 0.06–1.81 g DM, October: 0.47–1.46 g DM, December: 0.13–0.71 g DM). The temperature range examined approximates the annual range encountered by queen scallops in Roscoff waters. The allometric relationships between body mass and metabolic rates (\(\text{VO}_2\)) of *A. opercularis* were:

The combined effects of body mass and temperature on metabolic rate can be described by the multiple linear model:

<table>
<thead>
<tr>
<th>Month</th>
<th>(\log(\text{VO}_2)) Formula</th>
<th>(r^2)</th>
<th>N</th>
<th>Animals</th>
</tr>
</thead>
<tbody>
<tr>
<td>June</td>
<td>(2.253 + 0.654 \cdot \log(\text{DM}))</td>
<td>0.483</td>
<td>72</td>
<td>34 animals</td>
</tr>
<tr>
<td>October</td>
<td>(2.408 + 1.014 \cdot \log(\text{DM}))</td>
<td>0.496</td>
<td>101</td>
<td>34 animals</td>
</tr>
<tr>
<td>December</td>
<td>(2.288 + 0.707 \cdot \log(\text{DM}))</td>
<td>0.662</td>
<td>101</td>
<td>22 animals</td>
</tr>
</tbody>
</table>
\[
\log(\text{VO}_2) = 2.093(\pm 0.059) + 0.774(\pm 0.029) \\
\cdot \log(\text{DM}) + 0.02(\pm 0.005) \cdot T;
\]
\[N = 274, r^2 = 0.74, F = 381.311,\]
\[p < 0.001, \text{temperature in} ^\circ C.\]

Standard metabolic rates (\(\text{VO}_{2,\text{S}}\)) were indistinguishable in June (188.62 ± 60.32 \(\mu\) O2 g DM\(^{-1}\) h\(^{-1}\)) and December (204.17 ± 67.9 \(\mu\) O2 g DM\(^{-1}\) h\(^{-1}\)), but significantly higher (ANOVA: \(p < 0.001\)) in October (264.59 ± 69.05 \(\mu\) O2 g DM\(^{-1}\) h\(^{-1}\)). This indicated a 30% seasonal factorial rise over the lowest winter metabolic rates.

4. Discussion

4.1. Age and growth in *Aequipecten opercularis*

Partitioning of oxygen isotopes between molluscan shells and seawater has historically been assumed to closely approximate in isotopic equilibrium (Witbaard et al., 1994; McConnaughey et al., 1997; Hickson et al., 1999; Schoene et al., 2002; Schoene, 2003). On the other hand more recently Owen et al. (2003a, b) showed in a thorough laboratory study a positive deviation from equilibrium (+ 0.6 \(\delta^{18}O\) in *Pecten maximus*. Although changes in the \(\delta^{18}O\) and \(\delta^{13}C\) profiles cannot be used for an exact environmental reconstruction the amplitude still can be used as a proxy for relative seasonal oscillation of ambient temperature. The maximum \(\delta^{18}O\)-amplitudes in the two shells shown in Fig. 3 correspond to a relative temperature range of 5.6 and 6.8 \(^\circ\)C (according to Craig’s palaeo-temperature equation as given by Erez and Luz, 1983), respectively, which is close to the observed annual range of 6.7 \(^\circ\)C (Fig. 1). The close spatial coincidence between shell growth band position and both the \(\delta^{18}O\) and \(\delta^{13}C\) peaks in the isotope profiles (Fig. 3) indicate sufficient evidence that bands are formed during times of lowest temperature and low primary production, i.e. one band is formed each winter (for review, see Richardson, 2001). The same pattern was found in populations from Plymouth (December-March: Pickett and Franklin, 1975; Broom and Mason, 1978) and the North Sea (December-February: Ursin, 1956).

Non-linear growth patterns as derived from the VBGF can be compared indirectly only, e.g. using composite indices of overall growth performance (OGP) (e.g. \(P = \log(K) + \log(M_{\infty})\); Pauly and Munro, 1984). According to Brey (1999), this index is more or less consistent for closely related species. However, in suspension feeding bivalves food supply and/or temperature are the main factors controlling patterns of life history, whereas the genome commits the range of possible adaptations (Steams, 1976; Widdows et al., 1984; Bayne and Newell, 1983). Hence, the attainable data of Pectinids populations (\(N = 134\)) were divided into four broad groups regarding to (i) their hypothetical longevity (estimated as 0.95 \(\cdot\) Age at \(H_{\infty}\) derived from the appropriate VBGF), and (ii) the average annual water temperature of the study sites. An auximetric plot of \(\log(K)\) versus \(\log(M_{\infty})\) is given in Fig. 7. The mean OGP of the four groups are with one exception (see below) significantly different (ANCOVA with Bonferroni-Dunn post-hoc \(p < 0.001\)). OGP of *Aequipecten opercularis* falls in the range of other temperate short-lived scallops (TSL, mean = 1.92 ± 0.26, \(N = 31\), Fig. 7). While previous OGP estimates of *A. opercularis* range between 1.9 (Lee, 1973; Taylor and Venn, 1978) and 2.1 (Rolfe, 1973; Allison, 1994), values of the Roscoff population (1.74) appear to be comparatively low, indicating a potentially more stressful environment (i.e. lower food availability, increased depth).

Cold-water long-lived species showed quite similar values (CLL, mean = 1.81 ± 0.23, \(N = 19\), Fig. 7) and hence are statistically not distinguishable from the TSL group, albeit having lower K but higher \(M_{\infty}\) values. The TSL group consists mostly of species belonging to the supragenera *Aequipecten* and *Mimachlamys* (suggested by Waller, 1991), while in the CLL all species belong to the supragenus *Chlamys*. All species of this genus are close relatives, all of them have been considered as subspecies of *C. islandica* at some time of their nomenclatural history (see Waller, 1991). OGP in both warm water short-lived (WSL, mean = 2.89 ± 0.28, \(N = 25\), Fig. 7) and temperate long-lived species (TLL, mean = 2.55 ± 0.18, \(N = 59\), Fig. 7) is distinctly higher.
In general, phylogenetically ‘younger’ species of the supragenera belong to temperate groups, thus supporting the general idea (e.g. Arritz and Fahrbach, 1991) that recent scallops evolved from warm water ancestors, i.e. within the supragenus Pecten. Amusium spp. belong to the WSL group, while Pecten maximus belongs to TSL. This remains, however, pure speculation until modern molecular methods (e.g. Canapa et al., 2000) will clarify the hitherto rather uncertain phylogeny of pectinids.

4.2. Metabolic rates

Metabolic rate (R) reflects the energy expenditure of an organism, and hence is an important factor in estimating energy flow through both populations and the individual organism (Bayne and Newell, 1983). Rates measured in this study are 2–3 times lower than values for the same species published by McLusky (1973); most likely the latter study used too short an acclimation time to the holding systems (10–24 h).

Our results indicate a significant seasonality of metabolic activity in A. opercularis, as previously observed for many boreal scallops (MacDonald and Thompson, 1985; Bricelj et al., 1987; Shumway et al., 1988). Such seasonal cycles in metabolism result from complex interactions of environmental and endogenous variables acting on the metabolism (for review, see Bayne and Newell, 1983). Body size, environmental temperature and the energetic demands of gametogenesis, are considered to have a profound influence on respiratory activity (Shafee, 1982; Bayne and Newell, 1983; Bricelj et al., 1987; MacDonald and Bourne, 1987; MacDonald and Thompson, 1988).

The multiple-regression approach used in the present work attempts to quantify the simultaneous effects of body size and seasonal temperature. The equation obtained explains 74% of total variation recorded in metabolic rate. Because no interactive effects have been considered (for the sake of clarity), the two variables tested can be assumed to account almost completely for the seasonal fluctuations in metabolism. The metabolic mass exponent obtained by this equation ($b = 0.774, S.E. = 0.03$) is slightly below the mean overall value of $0.807 (S.E. = 0.02)$ derived from a large number of scallop studies ($N = 80, 13$ species; Heilmayer et al., unpubl. data).
4.3. Individual energy budget model

The partitioning of ingested or metabolizable energy into maintenance and production and possible trade-offs between growth and reproduction are important to characterise the life-history of species (e.g. Sibly and Calow, 1986; Wieser, 1994). The changing pattern of individual energy expenditure during the 6-y lifespan of *A. opercularis* clearly illustrates (i) an increasing share of maintenance requirements (expressed as respiration) in the absorbed energy (Fig. 5a, b), and (ii) a progressive transition of production from somatic growth to gonad output with increasing age (Fig. 5a), as previously observed for other scallop species, too, e.g. *Mimachlamys varia* (Shafee, 1982), *Mizuhopecten yessoensis* (Fuji and Hashizume, 1974) and *Placopecten magellanicus* (MacDonald and Thompson, 1985, 1988; Claerembout and Himmelman, 1996). The general decrease of somatic growth with increasing age (Calow, 1977) is not necessarily caused only by a shift between *P_s* and *P_g* (Fig. 5a) or the declining ability of the older animals to convert assimilated food into new tissue (Calow and Townsend, 1981), but may also be caused by a shorter growing season of older individuals, as already shown for *Chlamys islandica* (Vahl, 1981) and *Pecten maximus* (Chauvaud and Strand, 1999).

Net growth efficiency (*K_2*) values for each year of life, which is a measure of the ability to turn assimilated energy into production, decreases during life time from 45% to 11% in *A. opercularis* (Fig. 5b). These data are among the lowest recordings for populations of suspension and deposit feeding bivalve molluscs so far (Bayne and Newell, 1983; Shumway, 1991).

4.4. Population production and productivity

With a total P/B ratio of 1.848 y⁻¹ annual somatic production of *A. opercularis* is estimated at 20.717 kJ m⁻² y⁻¹ in Trezen Vraz and hence in the same order of magnitude as previous estimates from other populations (Broom and Mason, 1978; Taylor and Venn, 1978; Richardson et al., 1982).

The *A. opercularis* population from Roscoff waters invests only around 5% of its total annual production into gametes. This is clearly less than in other short-lived scallops (16% in *Mimachlamys varia*, Shafee and Conan, 1984) and far below the investment of either long-lived scallops (≥50% in *Placopecten magellanicus* and in *Patinotopecten caurinus*; MacDonald and Thompson, 1985; MacDonald and Bourne, 1987) or cold-water scallops (20–30% in *Chlamys islandica* and in *Adamussium colbecki*; Vahl, 1981; Heilmayer et al., 2003). We may, however, have underestimated gonad productivity by assuming one single spawning event during the reproductive season. This view is supported by Aravindakshan (1955) and Ursin (1956), but Taylor and Venn (1978) reported repeated spawning in populations from the Clyde Sea. Conclusive information is lacking so far.

The high *P_{Tot}/B* ratio is in slight contradiction to the observed low OGP value of the population. This might be explained by the comparatively low abundance of large individuals (>60 mm), which is either a consequence of spatial differences in settlement from year to year (Brand, 1991) or caused by size-selectivity of commercial fisheries.

In conclusion, despite its 6-y lifespan, *A. opercularis* shows some typical features of r-strategy orientated quasi-opportunists, e.g. fast growth and comparatively low reproductive effort. This life strategy combined with environmental variability causes a highly dynamic pattern of occurrence in time and space with typical high-density patches of limited continuance. Locally these patches of *A. opercularis* are likely to play a significant role in system energy flow. Although the sampling extent may not be considered adequate for a precise estimate of population energy parameters, it is still possible to arrive at some conclusions on the individual energy level, which may be considered as a first approximation for future research in this field.

Acknowledgements

The help of the ‘Mya’ crew and the colleagues from the logistics of the ‘Station Biologique de Roscoff’ for fishing and aquarium-maintenance of *A. opercularis* is greatly acknowledged. We are grateful to Günther Meyer and Katrin Blancke (AWI) for technical support in stable isotope analysis. We also thank three anonymous reviewers for constructive comments that greatly improved the manuscript.


References


