The effect of temperature on the development of encapsulated embryos of \textit{Concholepas concholepas} along a latitudinal cline

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**ABSTRACT:** Encapsulating species face more constraints than active brooders in adjusting oxygen supply to the needs of the embryos. Therefore, the packing of embryos in gelatinous egg masses or egg capsules is expected to be adjusted to the temperature and oxygen conditions that the embryos are likely to experience. We studied the patterns of embryo packing (number of embryos per unit area) of the gastropod, \textit{Concholepas concholepas}, from 14 sites over an extended geographic area spanning 22° of latitude off the coast of Chile. A clear break in the patterns of embryo packing was found at approximately 29 to 30° S. Capsules collected at the sites located north of this break exhibited significantly fewer embryos per unit area than capsules from southern sites. Embryo packing is correlated with mean temperature shortly before egg deposition. A set of laboratory experiments were conducted to determine if the effects of temperature reside in females (incubating females collected in 2 different sites at a common temperature) or in embryos (incubating capsules at different temperatures). Laboratory experiments showed that temperature does not affect the number of embryos that successfully develop in the capsules within the tolerated range of temperatures, but does influence developmental success at temperatures that are extreme for the sites of the sample population. Our results suggest that packing and protection of embryos in marine invertebrates might be linked to the capacity to supply oxygen to the brood, which can have important consequences for the distribution of brooding and encapsulating species across temperature gradients.

**KEY WORDS:** Brooding · \textit{Concholepas concholepas} · Embryo development · Encapsulated development · Oxygen · Temperature

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**INTRODUCTION**

Over the last 2 decades several studies have shown that oxygen is a limiting factor in the aggregation of embryos in aquatic systems (Strathmann & Strathmann 1995, Cohen & Strathmann 1996, Crump 1996), directly controlling embryonic development (Strathmann & Strathmann 1995, Cohen & Strathmann 1996, Cancino et al. 2003). Oxygen also seems to affect shell calcification in marine gastropods (e.g. Cancino et al. 2000, 2003), which may have enormous consequences on subsequent survival. The problem of oxygen acquisition imposed by aquatic systems can also help to explain the proportional positive relationship between the number of embryos and the surface area of the capsule in marine gastropods (genus Conus, Perron & Corpuz 1982). This evidence suggests that the strong oxygen limitation exhibited by both embryo masses
and capsules in aquatic systems may be an important evolutionary force affecting the capacity of the parents to aggregate the embryos, the patterns of embryo packing and the spread of parental care in aquatic systems in general (Strathmann & Strathmann 1982).

Several studies have shown the effects of oxygen and temperature on embryo oxygen demand and embryo development under laboratory conditions for different types of embryo aggregations (jelly, capsules, crab egg masses; Cohen & Strathmann 1996, Baeza & Fernández 2002, Lardies & Fernández 2002, Brante et al. 2003, Cancino et al. 2003). There is also evidence that parental behavior and the cost of brooding is affected by the oxygen demand of the embryos. For example, female crabs increase the frequency of brooding behaviors that help provide oxygen to the embryos (e.g. abdominal flapping) to compensate for the higher oxygen demand of later embryonic stages, thereby increasing the costs of brooding (energetic cost; Baeza & Fernández 2002, Brante et al. 2003). Similar changes in patterns of brooding behavior and cost in brachyuran crabs are correlated with temperature-dependent embryo oxygen demands (Brante et al. 2003). The positive relationship between temperature and the cost of brooding suggests that the capacity to aggregate embryos in the ocean, and therefore to provide parental care, may be favored at low temperature (Brante et al. 2003). This prediction is strongly based on studies conducted on brachyuran crabs, which always show active brooding behaviors directed towards providing oxygen to the brood (Baeza & Fernández 2002, Brante et al. 2003). However, aggregating embryos is also costly for passive brooding species (e.g. gelatinous embryo masses; Lee & Strathmann 1998). Since passive brooding species face more constraints in adjusting oxygen supply to the needs of the embryos, it is expected that the packing of embryos in gelatinous egg masses or egg capsules is adjusted to the temperature and oxygen conditions that the embryos are likely to experience. We do not disregard other environmental variables that are correlated with temperature and that may also shape embryo packing.

The effects of temperature on embryo packing and clutch size among marine invertebrates are less clear, despite the following evidence for marine invertebrates in general and gastropods that exhibit encapsulated development, in particular. (1) Temperature affects oxygen demand of the embryos (e.g. Brante et al. 2003). (2) The capsule walls limit oxygen diffusion (Cancino et al. 2000, Brante 2006). (3) Intracapsular oxygen conditions affect embryo development and, therefore, clutch size (Cancino et al. 2003). (4) There is a positive relationship between the number of embryos and surface area of the capsule (Perron & Corpuz 1982). This evidence suggests that oxygen conditions could have shaped re-productive attributes, including clutch size, in gastropod species that encapsulate embryos. We studied the patterns of embryo packing of Concholepas concholepas (Mollusca: Muricidae) from 14 sites over an extended geographic area spanning 22° of latitude off the coast of Chile. The response variable was the number of embryos packed per unit area of capsule. This extended geographic region also allowed us to relate temperature to patterns of embryo packing and to draw indirect inferences about the cost of aggregating embryos. A set of laboratory experiments was conducted to determine whether the effects of temperature reside in females, by affecting embryo assignation depending on seawater temperature of the site before egg deposition, or in embryos, by affecting embryo mortality during intra-capsular development. Concholepas concholepas is a good model to address these problems because (1) it shows an extended latitudinal distribution (5 to 54° S), (2) it encapsulates embryos but does not exhibit ovophagia or adelphophagia (Gallardo 1973), which can be alternative determinants of clutch size, and (3) capsules and females can be easily manipulated under laboratory conditions. In addition, C. concholepas is heavily exploited (Bustamante & Castilla 1987) to the point that harvesting negatively affects capsule deposition (Manríquez & Castilla 2001). Therefore, the identification of geographic regions offering different environmental conditions for embryo packing is critical. However, the present study has implications which go beyond the significance for this model species.

MATERIALS AND METHODS

Mature Concholepas concholepas females (>50 mm shell length) collectively deposit capsules in small aggregations in areas characterized by strong water flow in both intertidal and subtidal zones (Castilla & Cancino 1976, Lopez & Varela 1988). Embryos develop within the capsules for approximately 30 d (Gallardo 1973, 1979). After hatching, a free larval phase develops in the plankton for approximately 60 to 90 d (Di-Salvo 1988). Although under laboratory conditions C. concholepas females deposit egg capsules throughout the year, reproduction in nature is seasonal and varies along the coast of Chile (Castilla & Cancino 1976, López & Varela 1988). The capsules contain between a few hundred to more than 14 000 embryos (Gallardo 1973, Castilla & Cancino 1976). The total number of embryos deposited per capsule is positively correlated with capsule size, which in turn is positively correlated with female size (Gallardo 1973, Castilla & Cancino 1976).

Latitudinal pattern of embryo packing. To investigate the patterns of embryo packing in capsules of
Concholepas concholepas between sites in the biogeographic region associated with the Humboldt Current (Strub et al. 1998), capsules were collected during the reproductive season (May to November) from the lowest intertidal zone of wave-exposed areas. Fourteen sites were sampled from northern Chile (Iquique, 20° 53’S) to Chiloé Island (Ancud, 42° 02’S; Fig. 1) between 2003 and 2005. One capsule was collected from each aggregation found and was assumed to have been placed by a single female. The number of capsules analyzed per site is shown in Table 1. After collection, the capsules were preserved in 75% alcohol for later analysis. In the laboratory, the length and width of each capsule was measured to 0.1 mm and the number of embryos in each capsule was counted under a microscope. Embryos were spread evenly on a dish divided into 113 equally sized squares, and all the embryos were counted in 16 randomly selected squares from which the total egg number was estimated. The size of the capsules ranged from 7.5 to 23.0 mm (mean ± SD = 14.8 ± 3.7 mm) in length. Since the number of embryos per capsule was positively correlated with capsule size (length: $r = 0.43$, $n = 399$, $p < 0.0001$), the number of embryos was standardized per unit area (mm$^2$) of the capsule’s surface and used as the response variable. Capsule area was selected because no correlation was found between number of embryos per unit of capsule surface area and capsule length ($r = 0.11$, $n = 283$, $p = 0.27$) and because capsule area is likely to be an indicator of oxygen exchange surface (Perron & Corpuz 1982). Capsule surface area was estimated assuming the shape of a cylinder. Mean number of embryos per mm$^2$ of capsule area was compared across sites using 1-way ANOVA. At most sites we found capsules containing embryos in early developmental stages as well as capsules containing embryos in late developmental stages; however, at some sites capsules containing embryos in only 1 developmental stage were found (Table 1). Although according to Gal-

Fig. 1. Study area on the coast of Chile, which encompasses 22° of latitude, showing the 14 study sites

Table 1. Concholepas concholepas. The 14 study sites where capsules containing early and late stage embryos were collected. Latitude (lat.) of the site, mean water temperature (MWT) in °C (±SE), number of capsules (early and late) analyzed for each embryo stage and total no. of capsules are given; nc: no capsules found

<table>
<thead>
<tr>
<th>Site</th>
<th>Lat. (°S)</th>
<th>MWT (±SE)</th>
<th>Early stage</th>
<th>Late stage</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Iquique</td>
<td>20°53'</td>
<td>16.51 (0.81)</td>
<td>18</td>
<td>5</td>
<td>23</td>
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<tr>
<td>Punta Chacaya</td>
<td>22°58'</td>
<td>15.58 (0.61)</td>
<td>15</td>
<td>5</td>
<td>20</td>
</tr>
<tr>
<td>Juan Lopez</td>
<td>23°28'</td>
<td>15.42 (0.47)</td>
<td>14</td>
<td>5</td>
<td>19</td>
</tr>
<tr>
<td>Radison</td>
<td>23°40'</td>
<td>14.19 (0.19)</td>
<td>14</td>
<td>11</td>
<td>25</td>
</tr>
<tr>
<td>El Way</td>
<td>23°42'</td>
<td>14.22 (0.54)</td>
<td>4</td>
<td>5</td>
<td>9</td>
</tr>
<tr>
<td>Huasco</td>
<td>28°24'</td>
<td>13.48 (0.63)</td>
<td>9</td>
<td>20</td>
<td>29</td>
</tr>
<tr>
<td>Punta Choros</td>
<td>29°15'</td>
<td>13.75 (1.16)</td>
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<td>nc</td>
<td>58</td>
</tr>
<tr>
<td>Temblador</td>
<td>29°28'</td>
<td>13.33 (0.45)</td>
<td>7</td>
<td>7</td>
<td>14</td>
</tr>
<tr>
<td>Caleta Hornos</td>
<td>29°39'</td>
<td>11.52 (0.25)</td>
<td>nc</td>
<td>34</td>
<td>34</td>
</tr>
<tr>
<td>Totoralillo</td>
<td>30°03'</td>
<td>13.49 (0.67)</td>
<td>24</td>
<td>6</td>
<td>30</td>
</tr>
<tr>
<td>Montemar</td>
<td>32°57'</td>
<td>13.24 (0.19)</td>
<td>8</td>
<td>6</td>
<td>14</td>
</tr>
<tr>
<td>El Quisco</td>
<td>33°23'</td>
<td>12.61 (0.32)</td>
<td>nc</td>
<td>16</td>
<td>16</td>
</tr>
<tr>
<td>Concepción</td>
<td>36°48'</td>
<td>12.01 (0.32)</td>
<td>13</td>
<td>nc</td>
<td>15</td>
</tr>
<tr>
<td>Ancud</td>
<td>42°02'</td>
<td>10.50 (0.34)</td>
<td>nc</td>
<td>17</td>
<td>17</td>
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</tbody>
</table>
lardo (1973), the number of encapsulated embryos in *Concholepas concholepas* does not vary during embryo development, the comparisons were conducted independently for each embryo stage as well as with pooled samples of capsules containing early and late stage embryos.

To assess the effects of temperature on the number of encapsulated embryos per unit area of capsule along the study region, temperature data loggers were installed at all study sites 30 d before the capsules were collected. Data loggers were always placed in the lowest intertidal zone of exposed rocky platforms. Within this region no latitudinal trends were found in aerial exposure (Finke et al. 2007); therefore, the pattern of mean temperature was not expected to be biased by aerial or water temperature throughout the latitudinal gradient. Mean temperature was calculated using all data points obtained during this period (Table 1). Correlation analyses (Pearson) were conducted to assess the relationship between temperature and number of eggs per mm$^2$ of capsule area for each embryo stage and all capsules (pooling both embryo stages).

**Role of temperature in clutch size determination in *Concholepas concholepas***. Since temperature is correlated with the number of embryos assigned by females mm$^{-2}$ of capsule area in nature (see ‘Results’), laboratory experiments were conducted to determine whether the effect of temperature resides in females — through the effects of environmental temperature before egg deposition on patterns of embryo packing — or in embryos — through the effects on embryo mortality during intracapsular development. Two experiments were conducted to identify the relative importance of each mechanism.

**Effects of temperature on females**: Since our field studies show that local temperature at the study site a month before egg deposition is a good predictor of the number of embryos that females deposit per unit area of capsule (see ‘Results’), a laboratory experiment was designed with the goal of removing the effect of temperature of the site of origin by acclimating females for between 4 and 6 mo to a common constant temperature (12°C). Female *Concholepas concholepas* were collected from 2 sites within our study area, one situated in the northern region (at 29° S), where females deposit fewer embryos per unit area of capsule, and the second in the southern region (at 42° S). Experimental animals were transported to the Laboratorio Costero Lenga (36° 44′ S, 73° 11′ W) shortly after collection, where females from different sites of origin were placed in 5001 aquaria. Animals were fed ad libitum with clams (*Protothaca thaca* and *Semece solidia*) and were maintained at constant temperature and salinity (33 ± 1‰) under air flow.

After the acclimation period we collected the capsules that females deposited in the aquaria. Recently deposited capsules (<36 h) were removed and assigned to 5 temperature treatments: 6, 9, 12, 15 and 18°C, to assess the effects of temperature on developmental success. These temperatures are within the temperature range to which capsules are exposed under natural conditions. The capsules from the different sites of origin were placed in small containers (0.5 l). Four containers per site of origin and incubation temperature were used, and approximately 20 capsules from different females were incubated in each container. Thermostats maintained constant temperature. Filtered seawater was replaced every other day during the incubation period. The capsules were maintained with constant aeration. Embryo development was monitored regularly until the pre-veliger stage was reached, at which time 1 capsule was sampled from each treatment combination. Most of the capsules were sampled during the experiment, but only 1 per container was used to record (1) length and width of the capsule, and (2) number of embryos (see protocol previously described). The remaining capsules were only used to monitor development. At some temperatures (6, 9 and 18°C) embryo mortality occurred, preventing us from conducting a full-factorial ANOVA. For this reason, 2 independent analyses were conducted. First, 1-way ANOVAs were conducted for each site of origin to compare the effect of temperature on the mean final number of embryos per unit area of capsule among temperature treatments in which development occurred. Since development at the different temperatures depended on the site of origin, different incubation temperatures were compared for the northern (12, 15 and 18°C) and southern origins (9, 12, 15°C). A 2-way ANOVA was then used to compare the mean number of embryos per unit area of capsule between sites of origin and temperatures for the 2 temperatures at which development was observed (12 and 15°C). Data were not transformed because the assumptions of the model were met. Tukey’s HSD tests were used to assess differences among treatments.

**Effects of temperature on embryos**: Since the experiment described in the previous section does not distinguish the effect of temperature on females (before capsule deposition) from the effect on the embryos (survival during intracapsular development), a second laboratory experiment was conducted to determine whether the number of embryos deposited by females per unit area of capsule (initial number) changed during intracapsular development (final number) and to assess whether the changes were temperature dependent. Recently deposited capsules of *Concholepas concholepas* were collected from the intertidal zone of Las Cruces (33° S) in July 2003. Immediately after collection the capsules from different females were transported to the laboratory and
haphazardly assigned to either the 12 or the 15°C temperature treatment. Thermostats maintained constant temperature. Five and six 1 l containers were used for 12 and 15°C respectively; 6 capsules were assigned haphazardly to each container. One capsule was removed from each container at both the beginning of the experiment (initial time) and the end of the experiment (final time). One container from the 15°C treatment was lost during the experiment. The experiment was carried out for 60 and 40 d at 12 and 15°C, respectively. The capsules were maintained with filtered seawater and constant aeration. Water was exchanged every other day throughout development. Embryo development was monitored regularly. Before hatching (final time), 1 capsule was sampled per container. Length and width (to 0.1 mm) of the capsule and total number of embryos were recorded for each capsule sampled for each container, temperature and time. The number of embryos was estimated as described in ‘Latitudinal pattern of embryo packing’ above. After calculating capsule area (see ‘Latitudinal pattern of embryo packing’ above) the number of embryos mm⁻² of capsule area was estimated. A 2-way ANOVA was used to assess the effects of temperature and developmental time (initial and final) on the mean number of embryos per unit area of capsule. The data met the assumptions of the ANOVA model. Tukey’s HSD tests were used to assess differences among treatments.

RESULTS

Latitudinal pattern of embryo packing

Significant differences in the mean number of embryos per unit area of capsule were detected among sites ($F = 50.9$, df = 13, 310, $p = 0.001$), showing a clear break at approximately 29 to 30° S (Fig. 2A). Capsules collected at the sites located north of 29° 28' S exhibited significantly fewer embryos per unit area than capsules from sites south of 29° 39' S ($p < 0.05$). No differences were detected among the 8 sites sampled north of 29° 28' S or among the 6 sites sampled south of 29° 39' S ($p > 0.05$). The pattern described above was not affected by the fact that some sites had capsules containing only early or late stage embryos (Table 1) because a break in the mean number of embryos per unit area of capsule was still found between 29 and 30° S when early and late stage embryos were analyzed separately. The mean number of embryos per unit area of capsule was significantly lower in sites north of 29 to 30° S than in the southern sites for both embryo stages (early: $F = 18.05$, df = 10,173, $p = 0.001$; late: $F = 27.13$, df = 11,126, $p = 0.001$; Fig. 2B). No significant differences were detected in the mean number of early (or late) embryos per unit area of capsule either among sites located south of 29 to 30° S ($p > 0.05$) or among sites sampled north of 29 to 30° S ($p > 0.05$). Although mean capsule size was significantly different among sites ($F = 76.84$, df = 13,310, $p < 0.0001$), the patterns of embryo packing were not related to capsule size (see ‘Materials and Methods’ for the analysis of the response variable). The sites showing the largest capsules (Punta Choro and Ancud) exhibited opposing patterns of embryo packing. Contrasting patterns of embryo packing were also found between the 2 sites showing the smallest capsules (Iquique and Totorallillo).

The observed trend in the mean number of embryos per unit area of capsule was correlated with the mean temperature at the study site before egg deposition. The mean number of early stage embryos per unit area of capsule was negatively correlated with temperature (early stage: $r = 0.87$, $n = 11$, $p < 0.001$; Fig. 3A), suggesting that females pack embryos differently depending on mean temperature or a correlate of temperature. The same trend was observed when capsules containing late stage embryos were analyzed (late stage: $r = 0.7$, $n = 11$, $p < 0.001$; Fig. 3A). Pooling all capsules, the relationship between mean temperature at the study site and mean number of embryos per unit area of capsule was also significant and negative (Fig. 3B).
Role of temperature in clutch size determination

Temperature did not affect the mean number of embryos that developed in the 2 populations studied (northern and southern origin; Table 2). However, temperature tolerance of embryos differed with latitude of maternal population (Fig. 4). Embryo development did not occur at the 2 lowest experimental temperatures in capsules deposited by females collected in the northernmost location (29°S), nor was any development noted at the lowest (6°C) and highest temperatures (18°C) in capsules deposited by females collected in the southern site. Regardless of the site of origin, 100% mortality was observed at 6°C. Although temperature did affect the mean final number of embryos per unit area of capsule within each region, site of origin significantly influenced the number of embryos that reached the pre-veliger stage ($F = 40.08, df = 1,12, p < 0.0001$), when the 2 temperatures at which development occurred in both sites of origin (12 and 15°C) were compared. The mean final number of embryos incubated at 12 or 15°C was significantly lower in the capsules deposited by females collected in the northern (34.5, SD = 1.91) than in the southern region (39.4, SD = 1.97; $p < 0.05$). The 2 experimental temperatures at which development occurred did not affect the mean number of embryos that reached pre-veliger stage per unit area of capsule ($F = 4.51, df = 1,12, p = 0.07$). The interaction term was not significant ($F = 1.15, df = 1,12, p = 0.306$). Remarkably, the mean number of embryos mm$^{-2}$ of capsule that attained pre-veliger stage did not show a difference between capsules sampled in the northern region and capsules deposited in the laboratory by females collected in the north ($t = 0.97, df = 44, p = 0.33$). The same trend was found for the

Table 2. *Concholepas concholepas*. Results of the 1-way ANOVAs conducted to assess the effects of incubation temperatures (12 and 15°C) of the capsules on the mean number of embryos per unit area (mm$^2$) of capsule that successfully developed for 2 sites of origin located to the north (<30°S) and the south (>30°S) of the break identified in our study. Since development at the different temperatures depended on the site of origin, different incubation temperatures were compared for individuals of northern origin (12, 15 and 18°C) and southern origin (9, 12 and 15°C).

<table>
<thead>
<tr>
<th>Site of origin</th>
<th>Source of variation</th>
<th>df</th>
<th>$F$</th>
<th>$p$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Northern origin (29° S)</td>
<td>Incubation temperature</td>
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<td>1.172</td>
<td>0.353</td>
</tr>
<tr>
<td>Southern origin (42° S)</td>
<td>Incubation temperature</td>
<td>2</td>
<td>2.758</td>
<td>0.116</td>
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</table>

Fig. 3. *Concholepas concholepas*. Relationship between seawater temperature (°C) and mean number of embryos per unit area (mm$^2$) of capsule estimated (A) for each stage separately (O: capsules containing early stage embryos; •: capsules containing late stage embryos) and (B) by pooling capsules containing early and late stage embryos ($n = 14$). Regression parameters are given in (B).

Fig. 4. *Concholepas concholepas*. Mean number of embryos per unit area (mm$^2$) of capsule developed at the end of the incubation experiment (pre-veliger stage) in capsules at 5 temperatures (6, 9, 12, 15 and 18°C). Experimental capsules were deposited by females collected at 2 sites (29° S and 42° S, northern and southern populations, respectively) and maintained at a common (acclimation) temperature of 12°C for 4 to 6 mo before capsules were laid; nd = no embryo development at that temperature. Vertical bars indicate 1 SE.
southern region \((t = 1.86, df = 27, p = 0.27)\). Temperature did not affect the survival of *Concholepas concholepas* embryos during intracapsular development, since the mean number of embryos \(mm^{-2}\) did not change between initial and final time (Table 3, Fig. 5). The interaction between the 2 factors was not significant (Table 3).

**DISCUSSION**

The consistent patterns observed in our field sampling and laboratory experiments allowed the following conclusions. (1) Embryo packing (number of embryos packed per unit area of capsule) in capsules of *Concholepas concholepas* depends on the site of origin. (2) Temperature does not affect the number of embryos that successfully develop in the capsules, but (3) temperature is correlated with mean number of embryos per unit of area (embryo packing) and affects development, although the effect of temperature on development depends on the site of origin of females. The break identified in the patterns of embryo packing and the effect of temperature on embryo developmental success suggests differences between populations, although we cannot distinguish whether this pattern is related to genetic differences or environmental responses.

Our study showed a break in embryo packing of the carnivore *Concholepas concholepas* between 29 and 30°S. We hypothesize that a break in environmental variables or a break in the genetic structure of local populations could explain our results. Several lines of evidence provide support for the first argument. The clear break in embryo packing coincides with a clear break in eddy kinetic energy and equator-ward wind stress at 30°S (Hormazábal 2004) and with 2 contrasting regimes in chlorophyll concentration in coastal areas and offshore (Yuras et al. 2005). Moreover, contrasting patterns of recruitment and abundance of several key intertidal species occurred on each side of the break (Camus 1998, Broitman et al. 2001, Navarrete et al. 2005). Among the environmental variables that may affect the pattern of embryo packing, local temperature and oxygen concentration have been studied in aquatic systems (Strathmann & Strathmann 1995, Cohen & Strathmann 1996, Brante et al. 2003, Cancino et al. 2003). In our study area both variables can be influenced by upwelling persistence and strength, which breaks at 29 to 30°S (Hormazábal 2004), although the expected effects of oxygen concentration may occur only in subtidal areas (Helly & Levin 2004). Our results clearly show that the number of embryos per unit area of capsule in intertidal zones is determined by local temperature before egg deposition, which, in turn, is strongly correlated with temperature during embryo development (\(r = 0.75, n = 13, p < 0.04\)). We think that latitudinal patterns of embryo packing might respond to oxygen demand during embryonic development, as temperature affects embryo oxygen consumption (e.g. Brante et al. 2003) and oxygen is generally considered a limiting factor in embryo aggregation (Cohen & Strathmann 1996, Crump 1996). Limits in the acquisition of oxygen in aquatic embryo aggregations have also been proposed as an explanation of the positive relationship between number of embryos and surface area of the capsule in marine gastropods (Perron & Corpuz 1982). In our model species, limited oxygen supply through capsule walls (Brante 2006), which is expected to increase at higher temperatures due to the increase in embryo oxygen demand, may determine clutch size. This hypothesis is supported by evidence in the literature showing lower clutch sizes at higher temperatures in brachyuran crabs, which seems to be related to the higher cost of oxygen provision (Brante et al. 2003). Our results also

Table 3. *Concholepas concholepas*. Results of the 2-way ANOVA conducted to compare the mean initial number of embryos deposited by females per unit area of capsule and the final number of embryos that successfully developed between the 2 experimental temperatures 12 and 15°C

<table>
<thead>
<tr>
<th>Source of variation</th>
<th>df</th>
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<tr>
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<tr>
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<td>Error</td>
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</tbody>
</table>

Fig. 5. *Concholepas concholepas*. Mean number of embryos per unit area of capsule \(mm^{-2}\) in capsules collected at 33°S on the coast of Chile (southeastern Pacific Ocean) and incubated at 2 different temperatures (12 and 15°C) until hatching. Initial and final numbers of embryos per unit area of capsule are shown. Vertical bars indicate 1 SE
suggest that the cost for *C. concholepas* of packing embryos increases with temperature, since fewer embryos can be packed per unit of extra embryonic material (capsule wall). Increases in encapsulation and brooding cost with temperature might affect the distribution of brooding species at low latitudes, which is in line with the higher prevalence of encapsulating and brooding species at low temperatures compared with those at high temperatures (Thorson 1950, Gallardo & Penchazadeh 2001). These results suggest that different strategies in the way marine invertebrates pack and protect embryos might be shaped by the capacity to supply oxygen to the brood, and this depends on temperature (Strathmann & Strathmann 1982). However, we do not disregard the effects of other environmental factors that might be correlated with temperature, such as metabolites produced by the embryos or development rates of protozoa or bacteria (Cancino et al. 2000).

Although we suggest that the latitudinal patterns of embryo packing in *Concholepas concholepas* are linked to local temperature and oxygen requirements of the embryos, other strategies to adjust clutch size in order to assure oxygen provisioning to the embryos according to their requirements may occur. For instance, ovophagia and adelphophagia have been reported in many species of marine invertebrates that encapsulate embryos (i.e. Kohn & Perron 1994, Collin 2003). It is not clear yet if intracapsular cannibalism is a plausible mechanism to reduce sibling competition for oxygen under adverse and unpredictable environmental conditions; however, the number of developing embryos increases with oxygen partial pressure in *Acanthina monodon*, suggesting that intrasibling consumption increases (Lardies & Fernández 2002). *C. concholepas* embryos do not exhibit ovophagia or adelphophagia; this is clearly shown by the lack of change in the number of embryos during development in our laboratory experiments. Therefore, the clear response of females to temperature is critical, since clutch size does not seem to be regulated after deposition. More information on the relationship between embryo packing and environmental variables in species exhibiting ovophagia or adelphophagia during encapsulated development is needed to fully understand the set of mechanisms determining clutch size in species that encapsulate the embryos and to generalize our findings.

The alternative hypothesis to explain the 35% change in the number of embryos per unit of capsule size with a small change in temperature (<1°C) is a change in the genetic structure of *Concholepas concholepas* populations on each side of the break. Although a break in the genetic structure over the short distance in which changes in embryo packing was observed is unlikely in a species exhibiting a long-lived larval stage (Palumbi 2003, Shanks et al. 2003), the break identified in the mean number of embryos packed per unit area of capsule coincides with the break in the genetic structure of *C. concholepas* reported between populations north and south of Coquimbo (about 30°S; Guiñez et al. 1992, Fig. 1). Moreover, no genetic differences were detected among populations sampled south of the break, which also showed a homogeneous pattern of embryo packing, or in comparisons of northern populations among themselves (Guiñez 1992, Gallardo & Carrasco 1996). However, ongoing studies question the existence of genetic differences among *C. concholepas* populations along the coast of Chile (L. Cardenas pers. comm.). Nevertheless, our results show a clear break in reproductive patterns of *C. concholepas* north and south of 30°S latitude. First, females from different sites of origin, but incubated at a common temperature, packed embryos as at their site of origin, which could suggest different behaviors among populations, regardless of temperature. Second, there was a clear effect of the origin of females on embryo developmental success at low and high temperatures. It is interesting to note that capsules deposited by females collected in the northern region showed no embryo development at the 2 lowest temperatures, which are extreme for the site of origin. Similarly, embryo development did not occur at 18°C in capsules deposited by females collected in the southern region. Although these results might be linked to genetic differences among populations, we cannot assume that 4 to 6 mo of acclimation was enough to delete the stable and predictable environmental signal of the site of origin (Hormazábal et al. 2004, Yuras et al. 2005). Therefore, more evidence on genetic structure of *C. concholepas* populations is needed to determine the influence of maternal genetic effects and environmental factors in shaping female behavior, as well as the interaction between these factors. Nevertheless, the existing evidence suggests that the reproductive behavior of populations of *C. concholepas* differs north and south of 30°S, which may have important implications for the dynamics of the larval pool and the population dynamics of *C. concholepas* and, therefore, for management of this species.

**Acknowledgements.** We are grateful to M. Aguilera, I. Albornoz, A. Brante, M. Cifuentes, C. Inostroza, P. Pappalardo, R. Soto and F. Vélez for their help. We also thank R. Finke, J. Holl and S. Kimberlin for comments on the manuscript. Four anonymous reviewers made helpful comments and interesting suggestions to improve this manuscript. This study was fully supported by the Fondecyt 1020860 (M.F.) and 1990451 (J.M.C.). We also thank the Humboldt Foundation for their donation of equipment. M.F. acknowledges the FONDAP-Fondecyt grant 1501-0001 to the Center for Advanced Studies in Ecology and Biodiversity and the International Laboratory Dispersal and Adaptation in Marine Species (DIAMS).
LITERATURE CITED


Editorial responsibility: Otto Kinne (Editor-in-Chief), Oldendorf/Luhe, Germany


Submitted: November 8, 2006; Accepted: May 9, 2007

Proofs received from author(s): September 21, 2007