Temperature dependent pelagic larval development: parameterization for *Fissurella latimarginata*

For *F. latimarginata*, the time to complete Planktonic Larval Duration (PLD) was reduced with increasing temperature, it was modeled according to the universal relationship proposed by O’Connor et al. (2007), but using their complete 72 species database published as supplemental information. O’Connor et al. (2007) seem to have used a database of only 69 species. Additionally, we include PLD observations for *F. latimarginata* under laboratory conditions (Chavez 2004) and our laboratory PLD observations for *Loxechinus albus* (see below) under three different experimental temperatures. In consequence, the resulting database that we used included 73 species in total.

We calculated the parameter $\beta_0$ for *F. latimarginata*

The general model has two constants ($\beta_1=-1.3637905$ and $\beta_2=-0.3204597$)

$$\ln(\text{PLD}) = \beta_0 - 1.36 \times \ln \left(\frac{T}{15}\right) - 0.32 \times \ln \left(\frac{T}{15}\right)^2$$

**Equation S1**

$\beta_1$ and $\beta_2$ are constants described by O’Connor et al. (2007) and $T$ is temperature in Celsius degrees and $\beta_0$ ($\beta_0 = 1.81$) is a species-specific parameter (see O’Connor et al., 2007 supplementary information).

We obtained $\beta_0$ considering the random and fixed effects in a linear mixed effects model, as proposed by O’Connor et al. (2007) in their supplemental information file (S1).

A developmental time of 5 days was reported for *F. latimarginata* at 17°C under laboratory conditions (Chavez 2004) and was used in equation S1 to obtain PLDs across a wide temperature range (from 9°C to 20°C), which rendered a PLD range between 4.0 and 11.3 days.

$\beta_0 = 1.809649$

**Linear Model**

In order to model PLD$_{\text{temp}}$ within ICHTHYOP we first transformed PLDs into daily growth rates ($GR$, um/day). We assumed linear growth from a hatching size of 190 microm to a settlement size of 210 microm (Chavez 2004), we named Ontogenetic Growth (OG),

$$GR(T) = OG / \text{PLD}(T),$$

**Equation S2**

and then calculated growth rate $GR(T)$ considering the temperature experienced by the larvae at each time step (i.e. the growth function was updated at each time step), following the same method described in Garavelli et al. 2016. The temperature (T) experienced during each time
step was obtained from the hydrodynamic model and larval size (L in mm) at each subsequent time step (t + dt) was calculated following equation S3.

\[ L_{t+dt} = L_t + GR * (T) * dt \]  \hspace{1cm} \text{Equation S3}

\[ GR = 0.00029073 * T - 0.00099052 \]  \hspace{1cm} \text{Equation S4}

<table>
<thead>
<tr>
<th>Table S1. Length of <em>Fissurella latimarginata</em> larvae, data obtained from Chavez (2004).</th>
</tr>
</thead>
<tbody>
<tr>
<td>Length</td>
</tr>
<tr>
<td>-------------------</td>
</tr>
<tr>
<td>Initial length</td>
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<tr>
<td>Recruitment length</td>
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<tr>
<td>Length variation</td>
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<table>
<thead>
<tr>
<th>Table S2. Planktonic larval duration of <em>Fissurella latimarginata</em> at different temperatures.</th>
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<tbody>
<tr>
<td>Temperature</td>
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<tr>
<td>-------------</td>
</tr>
<tr>
<td>9</td>
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<td>10</td>
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<td>11</td>
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<td>12</td>
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<td>13</td>
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<td>14</td>
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<td>18</td>
</tr>
<tr>
<td>19</td>
</tr>
<tr>
<td>20</td>
</tr>
</tbody>
</table>

Fig. S1. Relationship between temperature and growth rate of *Fissurella latimarginata* larvae.

\[ y = 0.00029073x - 0.00099052 \]

\[ R^2 = 0.99262032 \]
Supplement 2

Temperature dependent pelagic larval development: parameterization for *Loxechinus albus*

In the case of *L. albus*, parameterization of temperature effects on larval development times was obtained through laboratory rearing experiments. To this end, we collected 33 mature urchin individuals (>7 cm) from subtidal areas around Las Cruces, central Chile and took them to the Estación Costera de Investigaciones Marinas where they were injected in the celomic cavity with 3 ml of 0.5M KCl to stimulate gamete release, following the protocol by Bustos et al. (1992). Eggs and sperm were washed with UV filtered seawater and sieved with 500 µm and 170 µm nylon mesh. Then, eggs from each female (n=4) were retained in 55 µm nylon mesh and divided into separate glass petri dishes, one for each female, where they were fertilized in an egg-sperm proportion of 1:100, from sperms of one male. The fertilized embryos were then divided in different batches, which were randomly assigned to experimental temperature treatments. Three temperature treatments were used: 10°C, 13°C and 17°C, encompassing the range observed in the study area (Thiel et al. 2007). The experiments started at the blastula stage and finished at the pre-metamorphic, competent larval stage. We recorded larval arm length under a microscope every 2 days until the 8-arms stage was reached. Thereafter, a reduction in arm length was observed until the pre-metamorphic stage was reached.

Considering the developmental characteristic of echinopluteus larvae, we used the proportion of time larvae remained in each developmental stage, with respect to total developmental time. Therefore, to model temperature effects on development we fitted the following linear function (equation S5) to experimental data:

\[
DP_{t+dt} = DP_t + K \times (T) \times dt
\]

*Equation S5*

where \(DP\) is Development Proportion, \(T\) is temperature in Celsius, \(dt\) is the time increment in days and \(K\) is a coefficient obtained from the relationship between growth rate and temperature \((K=0.38)\) through linear fitting. To incorporate the development proportion into the SEIBM, we followed the degree-day concept applied by Hinckley et al. (1996). At every time step, larvae advanced a fraction of the total development proportion depending on the temperature experienced during the time interval, until completing the entire PLD.
Table S3. Development proportion of each development larval stage of *Loxechinus albus* at three different temperatures.

<table>
<thead>
<tr>
<th>Temperature (T)</th>
<th>Development time (days) (D)</th>
<th>Development time (hours) (D)</th>
<th>Development Stage</th>
<th>Development Proportion (DP)</th>
</tr>
</thead>
<tbody>
<tr>
<td>17</td>
<td>4</td>
<td>96</td>
<td>4-arms</td>
<td>25%</td>
</tr>
<tr>
<td>17</td>
<td>9</td>
<td>216</td>
<td>6-arms</td>
<td>60%</td>
</tr>
<tr>
<td>17</td>
<td>13</td>
<td>312</td>
<td>8-arms</td>
<td>90%</td>
</tr>
<tr>
<td>17</td>
<td>15</td>
<td>360</td>
<td>Competent larvae</td>
<td>100%</td>
</tr>
<tr>
<td>13</td>
<td>5</td>
<td>120</td>
<td>4-arms</td>
<td>25%</td>
</tr>
<tr>
<td>13</td>
<td>13</td>
<td>312</td>
<td>6-arms</td>
<td>60%</td>
</tr>
<tr>
<td>13</td>
<td>19</td>
<td>456</td>
<td>8-arms</td>
<td>90%</td>
</tr>
<tr>
<td>13</td>
<td>21</td>
<td>504</td>
<td>Competent larvae</td>
<td>100%</td>
</tr>
<tr>
<td>10</td>
<td>6</td>
<td>144</td>
<td>4-arms</td>
<td>25%</td>
</tr>
<tr>
<td>10</td>
<td>18</td>
<td>432</td>
<td>6-arms</td>
<td>60%</td>
</tr>
<tr>
<td>10</td>
<td>22</td>
<td>528</td>
<td>8-arms</td>
<td>90%</td>
</tr>
<tr>
<td>10</td>
<td>24</td>
<td>576</td>
<td>Competent larvae</td>
<td>100%</td>
</tr>
</tbody>
</table>

Fig. S2. Relationship between development time and temperatures for each development stage of *Loxechinus albus* larvae.

Linear Model

We obtained a potential growth equation (D=a*T−b) for each development stage. From this equation we determined a development time (D) dependent on temperature (T).

From data of development time for each development stage we obtained a temperature dependent continuous variable (Development Proportion, PD) for each development stage (e.g., DP (4-arms) = 4 *100/15).

We correlate DP with days of development (DP = a*days – b), obtaining one equation for each temperature.
Table S4. Development time for each development proportion for *Loxechinus albus* larvae.

<table>
<thead>
<tr>
<th>Development Proportion (DP)</th>
<th>Development time (days) (D)</th>
<th>Development time (days) (D)</th>
<th>Development time (days) (D)</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>25</td>
<td>4</td>
<td>5</td>
<td>6</td>
</tr>
<tr>
<td>60</td>
<td>9</td>
<td>13</td>
<td>18</td>
</tr>
<tr>
<td>90</td>
<td>13</td>
<td>18</td>
<td>23</td>
</tr>
<tr>
<td>100</td>
<td>16</td>
<td>20</td>
<td>25</td>
</tr>
</tbody>
</table>

Fig. S3. Relationship between development proportion and development time in three different temperatures for *Loxechinus albus* larvae.
Finally, we estimated the slope of the equation of each experimental temperature, and plotted the slope versus temperature.

Table S5. Parameters of the equation of the relationship between development proportion and development time for *Loxechinus albus* larvae for three different temperatures.

<table>
<thead>
<tr>
<th>Temperature (ºC)</th>
<th>b</th>
<th>Slope (a)</th>
</tr>
</thead>
<tbody>
<tr>
<td>10</td>
<td>0.6896</td>
<td>3.8777</td>
</tr>
<tr>
<td>13</td>
<td>0.7625</td>
<td>5.0634</td>
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<tr>
<td>17</td>
<td>0.2971</td>
<td>6.5725</td>
</tr>
</tbody>
</table>

Fig. S4. Relationship between slopes (showed in table S5) and temperature.

The slope of this linear model is the parameter $K$ used in equation S5

$$DP_{t+dt} = DP_t + K \times (T) \times dt$$

**Literature cited**


Supplement 3
Comparative results and discussion of model scenarios

Recruitment success

For *F. latimarginata* average PLD ranged between 5.83 and 5.70 days under fixed PLD scenarios (Table 2), and between 8.03 and 8.10 days under PLD\textsubscript{Temp} scenarios (Table 2) showing an extension of precompetency time of 39% under PLD\textsubscript{Temp} scenarios. Conversely for *L. albus* average PLD varied between 22.92 and 22.67 days under fixed PLD scenarios (Table 2), and between 23.28 and 22.23 days under PLD\textsubscript{Temp} scenarios (Table 2) not showing differences of precompetency time between PLD scenarios, even a reduction of 1% under PLD\textsubscript{Temp} scenarios.

Significant among-year differences were observed in terms of recruitment success for both species ($X^2$=20809.9, df=3, p<0.01 for keyhole limpet and $X^2$=5516.0, df=3, p<0.01 for red sea urchin). The highest mean recruitment success in both species was observed in 2012 (Fig. S5A, B), while the lowest in 2011 for keyhole limpets (16.95%, Fig. S5A) and 2013 for red sea urchins (Fig. S1B). Larval DVM had significant effects on recruitment success in both species (behavior: $X^2$= 18609.9, df= 1, p<0.01, for keyhole limpet and $X^2$=9609.0, df= 1, p<0.01 for red sea urchin), generally increasing onshore recruitment in both species (Fig. S5A, B). However, the effect of larval behavior on red sea urchin recruitment disappeared in 2011 (Fig. S5B). Consequently, the interaction term (year: behavior) was significant for red sea urchin ($X^2$= 1972.0, df= 3, p< 0.01) and not significant for keyhole limpets ($X^2$= 153.8, df= 3, p= ns), indicating that the effect of DVM depends on the year only for long PLD species (red sea urchin). Having a temperature dependent larval development time (PLD\textsubscript{Temp}) significantly reduced recruitment success with respect to fixed development time in keyhole limpets and sea urchins ($X^2$= 29910.6, df= 1, p<0.01, for keyhole limpet and $X^2$=278.5, df= 1, p<0.01 for red sea urchin), but the effect was much smaller and constant among years in red sea urchins (Fig. S5B). Therefore, the interaction term (year: PLD) was significant for keyhole limpet ($X^2$= 2874.9, df= 3, p< 0.01) and not significant for red sea urchin ($X^2$= 95.0, df= 3, p= ns), suggesting that temperature interactions with PLD vary more among years for short PLD species than long PLD species.

The interactive effects of larval traits on recruitment success was apparent for the short PLD species (keyhole limpet) (Fig S1A), which rendered significant the behavior: PLD interaction term ($X^2$= 7026.2, df= 1, p< 0.01). For longer PLD species (red sea urchin) the interaction behavior: PLD was not significant ($X^2$= 90.4, df= 1, p= ns), indicating that larval behavior had similar effects on recruitment in all years, regardless of the small fluctuations in time of PLD induced by temperature (Fig. S1B). Despite among year variation in average recruitment, the second order interaction term (year: behavior: PLD) was not significant in either species ($X^2$= 207.2, df= 3, p= ns, for keyhole limpet and $X^2$= 97.4, df= 3, p= ns for red sea urchin).

Some larval traits, but not all of them, had significant effects on connectivity patterns and, these changes affected some but not all indicators of retention recruitment in keyhole limpets and/or red sea urchins. A significant effect of temperature dependent PLD was observed on self-recruitment (SR) and LR in keyhole limpet (Table S5 and Fig. S6), which can be visually appreciated comparing panels C and D of Fig. 4. Higher larval LR and SR were observed under fixed PLD (Table 2) than when PLD was allowed to vary with temperature, probably due to increased dispersal distance (see below) and larval waste. No significant effect of behavior was detected on SR or LR in keyhole limpet (Fig. S6), nor did behavior modify the effect of type of PLD (interaction term behavior: PLD was not significant, Table S5). In
contrast, significant effects of larval behavior were observed on larval LR and SR in the case of the red sea urchin (Fig. S6), while the type of PLD and the interaction with behavior were not significant (Table S5). Interestingly, generally lower red sea urchin LR and SR were observed under DVM (Table 2). AR was not affected by larval PLD or behavior in both species (Table S5).

**Dispersal distance**

As expected, dispersal distances were generally longer for red sea urchins than keyhole limpets, but on some years (2013) such a pattern reversed, with red sea urchins mean dispersal distances being lower than those of keyhole limpets across all larval traits (Fig. S5C, D, average of 47.39 km for keyhole limpet and 39.52 km for red sea urchins). Consequently, we observed that release year had a significant effect on dispersal distance in both species ($X^2=26214.9$, df=3, $p<0.01$, for *F. latimarginata* and $X^2=42066$, df=3, $p<0.01$ for *L. albus*). Averaging across the four years of simulations showed increased mean dispersal distances in the much shorter PLD species, keyhole limpets, under some larval trait scenarios (Table 2).

The year with highest mean dispersal distance was different between species; 2011 for keyhole limpet (79.85 km) and 2010 for red sea urchin (83.11 km) (Fig. S5C and D). Larval behavior had a significant effect on dispersal distance in both species ($X^2=463.3$, df=1, $p<0.01$ for *F. latimarginata* and $X^2=6720$, df=1, $p<0.01$ for *L. albus*), but in the opposite direction and generally larger effect on red sea urchins than keyhole limpets (Fig. 4C, D, Table 2). Consistently longer mean distances under DVM scenarios were observed for red sea urchin (Fig. S5D), increasing average dispersal distance across years from 49.41 to 76.98 under fixed PLD and from 56.96 to 70.29 km under PLD$_{temp}$ (Table 2).

In the keyhole limpet, DVM generally and slightly decreased mean dispersal distance across years (Fig. S5C, average 75.84 km in LAG and 64.09 km under DVM; Table 2).

When averaging across years, the type of PLD had significant main effect on dispersal distance only for keyhole limpet ($X^2=14901.7$, df=1, $p<0.01$, for *F. latimarginata* and $X^2=2$, df=1, $p=\text{ns}$ for *L. albus*), increasing dispersal when PLD was temperature dependent (Fig. S5C, Table 2). In this short PLD species, an extension of two days of planktonic development increased dispersal distance from 57.10 to 75.84 km under LAG scenarios and from 54.92 to 64.09 km under DVM scenarios (Table 2). In contrast, the effect of PLD type on the red sea urchin was small and varied in direction depending on larval behavior: a slightly negative effect under DVM and slightly positive effect under LAG behaviors (Fig. S5D, Table 2). The interaction term (year: behavior) was statistically significant on dispersal distance for red sea urchin but not for keyhole limpet ($X^2=32.6$, df=3, $p=\text{ns}$, for *F. latimarginata* and $X^2=2523$, df=3, $p<0.01$ for *L. albus*). The interaction term (year: PLD) was statistically significant on dispersal distance for both species ($X^2=283.8$, df=3, $p=0.01$, for *F. latimarginata* and $X^2=49$, df=3, $p<0.01$ for *L. albus*). The behavior: PLD interaction term had a statistically significant influence on dispersal distance for red sea urchin and not for keyhole limpet ($X^2=102.8$, df=1, $p=\text{ns}$, for *F. latimarginata* and $X^2=530$, df=1, $p<0.01$ for *L. albus*). The third level interaction term (year: behavior: PLD) was not significant in either species ($X^2=33.9$, df=3, $p=\text{ns}$, for *F. latimarginata* and $X^2=15$, df=3, $p=\text{ns}$ for *L. albus*).

**Fig. S5.** Average recruitment success (A and B) and boxplots showing dispersal distance (C and D) for *Fissurella latimarginata* (A and C) and *Loxechinus albus* (B and D) in each year simulated under the four scenarios: Diel Vertical Migration behavior and fixed Planktonic Larval Duration (PLD) (DVM + PLD$_{fixed}$), DVM and PLD based on a species-specific temperature dependent PLD (DVM + PLD$_{temp}$), passive Lagrangian transport (LAG) behavior and fixed PLD (LAG + PLD$_{fixed}$) and LAG and PLD based on temperature.
dependent PLD (LAG + PLD_{Temp}). The black diamonds show the average across all four experiments in a given year.
Table S5. Analysis of deviance for the Generalized Linear Models (GLMs) with quasipoisson error structure applied to the potential connectivity metrics calculated for self-recruitment, local retention, relative local retention and allochthonous recruitment for *Fissurella latimarginata* and *Loxechinus albus*.

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<tbody>
<tr>
<td></td>
<td><em>Fissurella latimarginata</em></td>
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<tr>
<td><strong>Self-recruitment</strong></td>
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<tr>
<td>Deviance explained:</td>
<td>1.82%</td>
<td></td>
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<td></td>
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<tr>
<td></td>
<td><strong>Df</strong></td>
<td><strong>Deviance</strong></td>
<td><strong>Resid. Df</strong></td>
<td><strong>Resid. Dev.</strong></td>
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<tr>
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<td></td>
<td>555</td>
<td>26.34</td>
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</tr>
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<td>554</td>
<td>26.30</td>
</tr>
<tr>
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<td>0.39</td>
<td>553</td>
<td>25.92</td>
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<td>0.06</td>
<td>552</td>
<td>25.86</td>
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<td><strong>Df</strong></td>
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<td>4.63*10^-2</td>
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<td>553</td>
<td>11.02</td>
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<tr>
<td>Behavior * PLD</td>
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<td>1.80*10^-3</td>
<td>552</td>
<td>11.02</td>
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<td><strong>Relative local retention</strong></td>
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<tr>
<td>Deviance explained:</td>
<td>1.52%</td>
<td></td>
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<tr>
<td></td>
<td><strong>Df</strong></td>
<td><strong>Deviance</strong></td>
<td><strong>Resid. Df</strong></td>
<td><strong>Resid. Dev.</strong></td>
</tr>
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<td>555</td>
<td>27.51</td>
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</tr>
<tr>
<td>Behavior</td>
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<td>4.28*10^-3</td>
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<td>27.50</td>
</tr>
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<td>PLD</td>
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<td>0.41</td>
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<td>27.09</td>
</tr>
<tr>
<td>Behavior * PLD</td>
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<td>8.80*10^-4</td>
<td>552</td>
<td>27.09</td>
</tr>
<tr>
<td><strong>Allochthonous recruitment</strong></td>
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<tr>
<td>Deviance explained:</td>
<td>0.00%</td>
<td></td>
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<td></td>
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<tr>
<td></td>
<td><strong>Df</strong></td>
<td><strong>Deviance</strong></td>
<td><strong>Resid. Df</strong></td>
<td><strong>Resid. Dev.</strong></td>
</tr>
<tr>
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<td>555</td>
<td>345.37</td>
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<tr>
<td>Behavior</td>
<td>1</td>
<td>9.74*10^-4</td>
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<td>345.37</td>
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<tr>
<td>PLD</td>
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<td>7.28*10^-3</td>
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<td>345.36</td>
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<td>Behavior * PLD</td>
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<td>345.36</td>
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<tr>
<td></td>
<td><em>Loxechinus albus</em></td>
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<td></td>
</tr>
<tr>
<td><strong>Self-recruitment</strong></td>
<td>~ Behavior * PLD, family= quasipoisson</td>
<td></td>
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Fig. S6. Relationship between self-recruitment (SR) and latitudes for *Fissurella latimarginata* (A and C) and *Loxechinus albus* (B and D), and between Local Retention (LR) and latitudes for *F. latimarginata* (E and G) and *L. albus* (F and H). The upper plots show the comparison between the two PLD scenarios, PLD fixed (red dots and solid line) and temperature-dependent PLD (blue dots and dashed line). The bottom plots show the comparison between behavior scenarios, passive Lagrangian transport (LAG) (red dots and solid line) and Dial Vertical Migration (DVM) (blue dots and dashed line).
Discussion

While location of origin and spawning date played major roles on dispersal distance and successful onshore recruitment, environmental conditions affecting larval development and larval behavior did have significant, albeit lower effects on these important population variables. Much has been speculated about the effect of increased sea surface temperature on larval dispersal and resulting recruitment and population connectivity, both in the context of latitudinal variation in mean temperatures (Thorson 1950, Bradbury et al. 2008, Ayata et al. 2010, Leis et al. 2013) and through climate change (O’Connor et al. 2007, 2012, Munday et al. 2009, Lacroix et al. 2018). Indeed, the dominant effect of temperature on metabolic and development rates (Gillooly et al. 2002, Brown et al. 2004) leads to highly predictable and universal reduction in larval development times (Connor et al. 2007), which is expected to generally reduce effective dispersal distances. However, the actual thermal regimes experienced by developing larvae in a complex realistic ocean might depart very significantly from laboratory controlled conditions. Therefore, it is critical to examine temperature effects under realistically variable thermal conditions and in different regions of the world. In our model, temperature dependent development had an important effect on recruitment success through changes in PLD, but surprisingly, the impact was greater on the short PLD species (keyhole limpets), than on longer PLD species (red sea urchin). For red sea urchin, the small effect of temperature on developmental time had marginal effects on recruitment success (Fig. SSB) and no consistent effect on dispersal distances (Fig. S5D). For *F. latimarginata* temperature dependent development increased the average time to reach the settlement size by two days, which represents 39% increase from the nominal development time. However, we used 5 days of precompetency period for both species although it might be extreme for *F. latimarginata*. Hence in this species we observed increased larval waste leading to lower recruitment (Fig. S5A) and slight but very consistent increase in average dispersal distances (Fig. S5C). It is remarkable that the relative impact of temperature dependent development is stronger on species exhibiting shorter larval developmental times. This is not because development in this species is more sensitive to temperature, but because of the differences in the actual thermal regimes and temperature ranges experienced in a realistic ocean, even when larvae are released at the same time. The longer the PLD, the more variable the thermal environment experienced through development. Long PLDs will necessarily extend beyond the warm water season and be exposed to colder waters that could have important effects on developmental times. Lacroix et al. (2018) predicts a PLD increase of 22% in a flatfish species in the North Sea in a climate change scenario, but our results suggest a complex interaction between the effect of temperature on developmental times and across regions since variability in temperature depends on the oceanographic conditions determining temperature (e.g., dominance of upwelling). Moreover, a rise in ocean temperature is expected to not only decrease larval duration, other consequences warrant consideration such as changes in spawning period (Fincham et al. 2013), changes in reproductive output (Shama 2015) and changes in larval mortality (Garavelli et al. 2016, Madeira et al. 2016). Thus, we submit that predicting consequences of increased ocean temperature on dispersal and recruitment can be counter intuitive and, is much more complex than simply projecting mean ocean temperature increases (Connor et al. 2007, Byrne & Selvakumaraswamy 2011, McLeod et al. 2015, Lacroix et al. 2018).

Laval behavior, especially the ability to perform diel vertical migration, has been pinpointed as the most important missing information in studies of larval dispersal (Levin 2006, Metaxas & Saunders 2009, Morgan 2014). Our results showed that DVM enhances recruitment success in both model species and across all years modeled. DVM increased recruitment success twice as much for red sea urchin than for the keyhole limpet, suggesting that DVM
seems to be more important in species exhibiting longer PLDs probably because larvae may experience baroclinic circulation patterns for longer periods of time. While the effect of DVM on spatial and temporal variability in recruitment was rather minor, it is remarkable that it was consistently positive across all conditions and for both species. In many modeling and observational studies conducted in upwelling regions DVM has been proposed as a mechanism promoting onshore larval retention (Marta-Almeida et al. 2006, Morgan et al. 2009, Aiken et al. 2011, Morgan 2014). Offshore currents at the surface and onshore currents at deeper depths during upwelling events (Strub et al. 1998, Kirincich et al. 2005, Morgan et al. 2009) allow organisms performing DVM to migrate below the Ekman layer, and reduce offshore transport (Marta-Almeida et al. 2006, Queiroga et al. 2007, Aiken et al. 2011). This general mechanism may play a role in the observed increased recruitment rates in our model. However, in contrast to the idea that DVM may enhance recruitment through reducing dispersal and increasing larval retention nearshore during development, DVM had positive effects on dispersal distance (see also Ospina-Alvarez et al. in press), negatively impacting local retention and self-recruitment, particularly of red sea urchins. In a previous study conducted in Monterey Bay, Carr et al. (2008) observed that DVM did not substantially lead to nearshore retention as the daytime return flow did not compensate offshore nighttime transport. In our study, for red sea urchin we also observed that allochthonous recruitment under DVM was mainly northward, compared to LAG where more than 20% was southward (Fig. 6). This result reinforces the idea that larvae that vertically migrate were advected northward and then a fraction returned to nearshore locations, possibly by the mechanisms described above. Thus, our results suggest that for species with moderate PLD (20 to 25 days), like the red sea urchins, in the upwelling ecosystem of central Chile, vertical migration is not a behavior promoting local retention, nor self-recruitment, but it increases the probability of successful onshore transport of competent larvae and, probably increases coastal alongshore dispersal distances in a coastal band across the region. Increased onshore recruitment and reduced dispersal have been suggested as one factor that may favour evolution of DVM in invertebrate and fish larvae (Batchelder et al. 2002, Marta-Almeida et al. 2006, Morgan 2014). DVM may provide an adaptive advantage as it leads to higher recruitment in our simulations for both species across all scenarios, but this comes at the expense of increased dispersal distances. Further studies are needed to determine if increased recruitment can be a sufficiently strong selective force leading to DVM in competent larvae.

**Literature cited**


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Fig. S7. Spatial variability on recruitment success and dispersal distance along the study region under different simulated scenarios. The map shows the study area where the recruitment (A to I) and release (H and J)-locations are represented by nearshore latitudinal bands of 2 km (location). Black bars show mean recruitment success (number of larvae that successfully reach the coastal zone) per latitudinal band and red lines indicate mean dispersal distance of successfully recruited larvae at a given latitudinal band. Recruitment success is shown for different simulation scenarios, and for *Loxechinus albus* and *Fissurella latimarginata* respectively, are: (A and F) passive Lagrangian transport and fixed Planktonic Larval Duration (PLD), (B and G) passive Lagrangian transport with a species-specific temperature-dependent PLD, (C and H) particles with Diel Vertical Migration (DVM) behavior and fixed PLD, and (D and I) particles with DVM and PLD based on temperature-dependent PLD. The spatial distribution of larval released, measured as the contribution (percentage) of oocytes/m² (potential egg production) of each location to the regional (study area) production, is shown for *L. albus* (E) and *F. latimarginata* (J). Dispersal distance is orthodromic distance from the spawning (release) to the successful coastal recruitment location, in km.
Fig S8. Temporal variability on the number of particles recruited (upper plot panel) and on dispersal distance traveled by recruited particles (lower plot panel) under different simulated scenarios. Black circles show passive Lagrangian transport (LAG) and white circles Diel Vertical Migration (DVM) behavior. Solid lines show fixed Planktonic Larval Duration (PLD$_{\text{fixed}}$), 5 days for $F$. latimarginata and 20 days for $L$. albus and dotted line)s temperature-dependent PLD (PLD$_{\text{Temp}}$).