Life history plasticity and reproductive strategy enabling the invasion of *Ligia exotica* (Crustacea: Isopoda) from the littoral zone to an inland creek

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ABSTRACT: A population of the marine isopod *Ligia exotica* was found in an inland habitat subject to unpredictable aridity. We studied the demography of littoral and inland populations in an attempt to define the variations in life history enabling this inland invasion of *L. exotica*. Although the 2 populations have similar steady recruitment and age structures, the patterns of size-specific contribution to yearly reproduction differ. The reproductive event itself may cause mortality in females. Both the net reproductive rate and the intrinsic rate of population growth show that both populations are increasing at similar rates. The longer generation time and the greater minimum age/size at reproduction of the inland population indicate that inland females may postpone the onset of reproduction. This delayed maturity may be an adaptive strategy allowing the inland colonization of *L. exotica*. Inland females are larger at time of reproduction, invest more in reproduction, and produce fewer but larger eggs/juveniles than do littoral females. These differences in the life history traits of *L. exotica* suggest life history plasticity in response to environmental changes. The production of fewer but larger eggs/juveniles of the inland population as a consequence of delayed maturity suggests that increasing offspring size is more important than higher fecundity in enabling *L. exotica* to colonize an inland habitat. The smaller individuals of *L. exotica*, with a lower resistance to desiccation due to high surface-area/volume ratios, have a higher mortality, especially during the juvenile stage. Because of the periodical and unpredictable aridity of the inland habitat, the mortality of inland juveniles is higher than that of littoral juveniles, and the strategy of producing larger offspring may favor inland colonization of *L. exotica*. For the successful invasion from littoral to inland areas, mortality may be the main selective factor determining life history traits: age and body size at maturation may be determined by age/size-specific survivorship, and the mortality of juveniles would seem to be size-dependent.

KEY WORDS: Life history plasticity · *Ligia exotica* · Offspring size · Reproductive strategy

INTRODUCTION

Demography, the key to life history theory, allows us to assess the strength of selection on life history traits. Life histories deal directly with natural selection, fitness, adaptation, and constraints. The evolution of life history traits and their plasticity determines the population dynamics of a species (Stearns 1992). The basic assumption of optimal theory is that, at each stage in its life history, an optimum fraction of an organism’s available material should be diverted to maximizing total reproduction during its lifetime. The energy expenditure of maternal organisms on their offspring focuses on the number of offspring that each maternal organism can produce and the fitness of individual offspring (Smith & Fretwell 1974, Reznick & Yang 1993). Life history theory is concerned with how resources are allocated to different functions of an organism and the circumstances under which different strategies evolve. Organisms belonging to the same species but living in different habitats often display different combinations...
of life history traits, and comparative studies on such variations provide insight into evolutionary processes (Reznick & Endler 1982, Eckelbarger & Watling 1995, Ellers & van Alphen 1997).

*Ligia exotica*, a semelparous isopod, has so far been regarded as a cosmopolitan marine species. It is widely distributed in the littoral zone and can be found on coasts around Taiwan. However, an inland population of *L. exotica* was found colonizing the banks of a hill creek on the Hengchun Peninsula in southern Taiwan (Tsai 1997). The inland habitat of *L. exotica* consists mainly of uplifted reef limestone. The creek bed and banks are composed of reef fragments and are covered by a layer of dense sediment. Most of the Hengchun Peninsula is surrounded by fringing reefs, with scattered sandy beaches carved out of them. A sandy deposit at the inlet area intercepts the creek. Coral reefs in the Hengchun Peninsula, lying on the accretion wedge of the Eurasian Plate, have been uplifted rapidly due to the ongoing arc-continent collision between the Eurasian and the Philippine tectonic plates (Wang & Burnett 1990).

The colonization by *Ligia exotica* in the inland creek may have 2 possible scenarios. The isopods may represent a population which invaded from the littoral area, or a relict population associated with the rapid uplift of land which was isolated from the littoral area. In either case, the inland population of *L. exotica* represents an interesting example for studying the effect of habitat change on life history traits. Environmental conditions differ between the littoral zone and inland creek in many ways (Table 1). The inland habitat of *L. exotica* is subject to unpredictable aridity. The flow of this creek is rain-dependent, and it may occasionally dry out. The features of the watercourse may shift frequently due to flash-floods after precipitation. The microhabitat of inland *L. exotica* is characterized by greater variations in both air temperature and humidity than the littoral zone (Table 1). This inland colonization of *L. exotica* therefore provides an excellent opportunity for studying how life history traits vary in response to environmental changes. In this study, we compare the demography of natural populations of *L. exotica* in the 2 environments in an attempt to determine what combination of life history traits would favor such invasion from the littoral to an inland creek.

**MATERIALS AND METHODS**

**Study site.** The 2 sampling sites are located on the east coast of the Hungchun Peninsula, southern Taiwan (Fig. 1). The littoral population of *Ligia exotica* is distributed in the supralittoral zone of a rocky shore in an area of about 1500 × 20 m. The substratum is emergent limestone, which in some places is covered by boulders. The inland population of *L. exotica* is distributed along the banks of a hill creek covered by muddy sediment and coral fragments. The air temperature and relative humidity of both microhabitats were recorded continually for 5 d bimonthly with a HOBO™ temperature logger and StowAway™ humidity logger (Onsey Instrument Corporation, MA, USA).

**Sampling.** At each site, 9 to 27 quadrats (each 1 × 1 m²) were sampled randomly along the coastline or the bank of the creek. The size of the quadrats was chosen based on a previous study which showed that quadrats larger than 1 × 1 m did not significantly reduce the coefficients of variance among quadrats at both sites (Tsai 1997). Two adjacent quadrats were

<table>
<thead>
<tr>
<th>Feature</th>
<th>Littoral habitat</th>
<th>Inland habitat</th>
</tr>
</thead>
<tbody>
<tr>
<td>Habitat type:</td>
<td>Relatively constant</td>
<td>Unstable; waterways may vary frequently due to flash-floods after precipitation</td>
</tr>
<tr>
<td>Microhabitat:</td>
<td>Surface of boulders, gaps</td>
<td>Crevices, tunnels</td>
</tr>
<tr>
<td>Water resource:</td>
<td>Sea water; regular wave action</td>
<td>Freshwater; unstable; rain-dependent</td>
</tr>
<tr>
<td>Vegetation:</td>
<td>Various seaweeds</td>
<td>Scattered shrubs</td>
</tr>
<tr>
<td>Substratum:</td>
<td>Emergent reef bed covered by large boulders</td>
<td>Coral debris and dense fine sand</td>
</tr>
<tr>
<td>Air temperature of microhabitat:</td>
<td>Relatively constant</td>
<td>Considerable variation</td>
</tr>
<tr>
<td>Min. 22.4°C</td>
<td>Min. 17.3°C</td>
<td></td>
</tr>
<tr>
<td>Max. 38.7°C</td>
<td>Max. 44.5°C</td>
<td></td>
</tr>
<tr>
<td>Average: 25.5°C</td>
<td>Average: 24.7°C</td>
<td></td>
</tr>
<tr>
<td>Humidity of microhabitat:</td>
<td>Relatively high and constant</td>
<td>Relatively low; greater variation than littoral</td>
</tr>
<tr>
<td>Min. 76 %</td>
<td>Min. 55 %</td>
<td></td>
</tr>
<tr>
<td>Max. 100 %</td>
<td>Max. 100 %</td>
<td></td>
</tr>
<tr>
<td>Average: 93 %</td>
<td>Average: 76 %</td>
<td></td>
</tr>
</tbody>
</table>

Table 1. Comparisons of habitat features of littoral and inland populations of *Ligia exotica* in Hengchun Peninsula, southern Taiwan.
Tsai & Dai: Life history plasticity enabling inland invasion of *Ligia exotica*

were then dried in an oven at 100°C for 48 h. All data are presented as dry weight to enable comparison with later studies. A previous study had shown that percentage water content is constant and independent of body size in *L. exotica* (Tsai et al. 1998). The size-age relationship was estimated by the growth rate of each size class. The age of an individual was estimated based on the mean size of juveniles when first released from a brood pouch. The age of any one size class was calculated by size-specific growth rate. The size-age relationship was used for the transformation of body size data into age in subsequent analyses.

Minimum size at first reproduction was defined as the smallest gravid female recorded in the collections, and this minimum size was regarded as the size/age at maturity. Only gravid females with eggs showing no signs of development were measured to reduce the variances caused by different developmental stages and overestimation of the density of gravid females. Body mass, brood mass, fecundity, and egg mass were measured and calculated using some of these gravid females with undeveloped eggs. Reproductive allocation was calculated as the proportion of both brood weight (egg mass) and juvenile biomass to female weight.

The size-fecundity relationship was established by a simple linear regression model, and was used in further estimations of size-specific fecundity for a given size class. Gravid females with embryos in later stages were raised in the laboratory, and the number and size of young released were measured. The proportional loss ($v_x$) arising from brood pouch mortality was assessed as the difference between the mean number of eggs and number of young released. The estimation of yearly reproduction ($N_x$) followed the method of Willows (1987). The average density of females ($D_x$), and the proportion of gravid females in each size class ($p_x$) were assessed. Yearly reproduction is calculated as:

$$N_x = D_x \times 2m_x \times p_x \times (1 - v_x)$$

where $2m_x$ is the weight-specific fecundity calculated from the size-fecundity regression under the assumption that the male:female ratio is 1.

**Demographic analysis.** Average survivorship curves and life tables were constructed based on the density estimates for each size class on each sampling occasion with different intervals. The size range which individuals of each size class should have attained at the subsequent sampling was calculated by the size-specific growth rate. Only those females within the expected size range at the subsequent sampling were considered to belong to the same size class, and were used for assessing survivorship. This approach is justified given the stable size distribution and limited variation in population density between years (Willows 1987).
The net reproductive rate, $R_0$, the intrinsic rate of population growth, $r$, and the generation time, $T$, were calculated by the equations (Birch 1948):

$$R_0 = \sum_{x=0}^{\infty} 1_x \times m_x$$

$$\sum_{x=0}^{\infty} e^{-r x} 1_x \times m_x = 1$$

$$R_0 = e^{rT}$$

Reproductive value, $V_x$, was calculated as:

$$V_x = \frac{1}{1_x} \sum_{x>x} 1_x \times m_x$$

(Fisher 1930, Stearns 1976).

RESULTS

Recruitment and population structure

In both the littoral and inland populations of *Ligia exotica*, daily growth rates varied between size classes. Growth rates of juveniles increased quickly after their release from the brood pouch, reached similar high levels in the size class ranging from 15 to 35 mg dry wt, and then gradually decreased (Fig. 2). The turning point of the growth rate curve corresponded roughly to the age/size at first reproduction. This suggests that an energy trade-off exists between individual growth and reproduction.

In general, both populations showed a similar size-age relationship. The results of a Kolmogorov-Smirnov test showed that the age-histograms of the 2 populations were similar among sampling months (p > 0.05) and between years (p > 0.05). In both populations, juveniles were released at a steady rate, and no obvious recruitment peak could be identified throughout the sampling period. Within each population, no significant difference in the quantity of recruitment could be found either among sampling months (littoral: p > 0.05; inland: p > 0.05) or between years (littoral: p > 0.05; inland: p > 0.05). However, the dry weight of newly released inland juveniles was significantly heavier (p < 0.001) than of those from the littoral. This indicates that inland females release larger offspring than littoral females.

Sex ratio and size-fecundity relationship

The results of 2 × 9 contingency table tests showed that female/male ratios among size classes differed in both populations (littoral: $\chi^2 = 759.19$, p < 0.001; inland: $\chi^2 = 490.49$, p < 0.001). For those individuals lighter than 35 mg dry weight, the sex ratios of both populations were approximately 1:1 (Table 2). However, for individuals heavier than 35 mg dry weight (i.e. approaching body size at reproduction), the number of males significantly exceeded that of females (Table 2). This suggests that reproduction may cause more mortality in females than in males. In both populations, the relationship between female size and fecundity fit a simple linear regression model well (Fig. 3). The slopes of the size-fecundity regression lines of both populations were similar (p > 0.05), but the intercepts differed (p < 0.001). The size-fecundity relationships were then used to calculate the size-specific fecundity of each population.

Yearly reproduction, survivorships and life tables

Yearly reproduction was estimated for both populations (Table 3). The midpoint of each size class and mean weight of brood-carrying females were used to calculate the average female fecundity ($2m_f$) based on the size-fecundity regression models. Although the total reproduction of littoral and inland populations were similar, the distribution patterns of young released by each maternal size class differed. A contribution index of each...
size class to total reproduction was calculated by dividing the reproduction of each size class by total natality. Total reproduction of the littoral population was contributed to mainly by size classes below 60 mg dry weight, while the contribution of larger size classes (>60 mg) to total reproduction declined rapidly (Fig. 4a). The total reproduction of the inland population was contributed to mainly by larger size classes, with a maximum contribution by the 60 to 70 mg size class (Fig. 4b).

The survival rate of inland juveniles in the first several months after release was lower than that of littoral juveniles (Table 4, Fig. 5). Before the onset of reproduction, males and females had similar life expectancies in both populations. However, the life expectancy of females was lower than that of males after the onset of reproduction (Fig. 5). The maximum longevity of littoral males was similar to that of inland males (about 25 mo), but the longevity of inland females (about 22 mo) was longer than that of littoral females (about 18 mo). This indicates that the reproductive event may affect the longevity of females.

**Fitness, life histories and reproductive traits**

The net reproductive rates were similar, and greater than 1 ($R_0 = 1.56$ and 1.54 for littoral and inland populations, respectively). The intrinsic rates of population growth were slightly greater than 0 ($r = 0.037$ and 0.035 per generation for littoral and inland populations, respectively). This indicates that both populations were increasing at similar rates. The estimated generation time of the inland population (13.87 mo, Table 5) was longer than that of the littoral population (11.32 mo, Table 5). This suggests that the inland population may postpone the onset of reproduction, and consequently a larger female size would be attained at reproduction over that of the littoral population.

Comparisons of life history traits between littoral and inland populations showed that the inland population had a longer life span, a later maturity, a lower brood pouch mortality, and a longer generation time than the littoral population (Table 6). The 2 populations also showed significant differences in several reproductive traits (Table 7). Inland females had a larger body size at reproduction ($p < 0.01$), invested more materials in reproduction ($p < 0.001$), and produced heavier broods ($p < 0.001$) with fewer (p < 0.001) but larger offspring ($p < 0.001$) than littoral females.

**Table 2.** *Ligia exotica.* Percentage of females in each size class in the 2 populations, and results of $\chi^2$ test on sex ratio departure from 1:1. ns: not significant at $p > 0.05$; **significant at $p < 0.01$; ***significant at $p < 0.001$

<table>
<thead>
<tr>
<th>Size class (mg)</th>
<th>Total individuals</th>
<th>Littoral population % of females</th>
<th>$\chi^2$</th>
<th>p</th>
<th>Total individuals</th>
<th>Inland population % of females</th>
<th>$\chi^2$</th>
<th>p</th>
</tr>
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<tr>
<td>5</td>
<td>688</td>
<td>49.74</td>
<td>0.01</td>
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<td>0.09</td>
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<td>15</td>
<td>1006</td>
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<td>0.19</td>
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<td>25</td>
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<td>50.10</td>
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<tr>
<td>35</td>
<td>756</td>
<td>44.08</td>
<td>10.73</td>
<td>**</td>
<td>964</td>
<td>46.00</td>
<td>3.16</td>
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<td>45</td>
<td>563</td>
<td>39.10</td>
<td>26.87</td>
<td>***</td>
<td>856</td>
<td>40.50</td>
<td>15.47</td>
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<tr>
<td>55</td>
<td>458</td>
<td>29.28</td>
<td>78.82</td>
<td>***</td>
<td>732</td>
<td>37.97</td>
<td>21.47</td>
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<td>75</td>
<td>514</td>
<td>12.60</td>
<td>286.88</td>
<td>***</td>
<td>440</td>
<td>24.45</td>
<td>60.97</td>
<td>***</td>
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<tr>
<td>85</td>
<td>392</td>
<td>0.25</td>
<td>388.01</td>
<td>***</td>
<td>320</td>
<td>3.13</td>
<td>180.23</td>
<td>***</td>
</tr>
<tr>
<td>Total</td>
<td>7267</td>
<td>36.38</td>
<td>257.66</td>
<td>***</td>
<td>6818</td>
<td>39.00</td>
<td>166.84</td>
<td>***</td>
</tr>
</tbody>
</table>

Fig. 3. *Ligia exotica.* Size-fecundity regression of littoral ($y = 8.80 + 0.782x$, $F_{1, 98} = 795.7$, $p < 0.001$) and inland ($y = 0.243 + 0.788x$, $F_{1, 80} = 1116.2$, $p < 0.001$) populations.
DISCUSSION

Life history variations

Neither the littoral nor inland populations of *Ligia exotica* were at equilibrium densities ($R_0 > 1$ and $r > 0$). However, this did not affect the stability of the population structure, since the proportion of organisms in each age class was relatively stable. Each population might achieve a steady recruitment of young through different combinations of life history traits such as proportions of gravid females, fecundity, age of breeding, and size/age-specific mortality. Although the *L. exotica* populations in the 2 habitats had similar net reproductive rates and intrinsic rates of increase, the inland population had a longer generation time. In this case, the net reproductive rate ($R_0$) proved an incomplete measure of fitness because it did not consider the...
The differences in the life history traits of littoral and inland populations may indicate life history plasticity of *Ligia exotica* in response to habitat change. The capability to modify their life history traits in different environments is possibly an adaptive strategy that is common among isopods or other crustaceans (Davis 1984). Geographic differences in life history traits have been reported in size at maturity and number of eggs in isopods (Sastry 1983). In California, the isopod *Armadillidium vulgare* breeds after 1 yr at a small body size (Paris & Pitelka 1962, Lawlor 1976a,b), but in other regions, it does not initiate breeding until after 2 yr, at a larger body size (Heeley 1941).

**Delayed maturity and its consequences**

Age at maturity is a key factor determining the sensitivity of each life history trait in response to selection pressure (Meats 1971, Green & Painter 1975, Snell 1978). Both earlier and later maturity may lead to increased fitness. In some cases, earlier maturity is the

![Diagram](image1)

**Fig. 4. Ligia exotica. Contribution of each size class to total reproduction of littoral and inland populations**

effects of different generation times (Lande 1982, Stearns 1992). Both the longer generation time and the larger minimum size/age at reproduction of inland females indicate that the life cycle of inland females had been delayed. Littoral offspring had a higher brood pouch mortality (i.e. mortality occurring in the brood pouch during the development period from egg to juvenile). However, after being released, the littoral juveniles had a higher survival rate than did inland juveniles. In the brood pouch, the survivorship of offspring is mainly related to the internal environment afforded by the gravid female. Once the juveniles are released from the brood pouch, the external environment becomes the main factor affecting their survivorship. Inland females with larger body sizes may provide better internal conditions (perhaps available space/resources) for the egg/juvenile development in the brood pouch. However, the desiccation stress of the inland habitat may lead to higher mortality of juveniles after release.

![Diagram](image2)

**Fig. 5. Ligia exotica. Survivorship curves of male and female of littoral and inland populations**
simplest and most effective way of increasing fitness by increasing the chance of survival to maturity (Caswell & Hastings 1980, Roff 1992, Stearns 1992). Individuals that mature earlier may have higher fitness because their offspring are born earlier and reproduce sooner (Hamilton 1966, Bell 1980). When the timing of maturity is earlier than the occurrence of the highest mortality in the life cycle, the benefit of earlier maturity would be theoretically maximal. Thus, the benefit of earlier maturity may depend on age/size-specific mortality. However, the high mortality of the early juvenile stages of Ligia exotica seems unlikely to favor evolution toward earlier maturity, and the very low fecundity associated with small maternal size is also disadvantageous. Stearns (1992) suggested 2 important factors that may cause delay in maturity. First, if delaying maturity permits further growth and increasing fecundity with increasing maternal size, then delay in maturity would lead to higher fecundity. Second, if delaying maturity means that the quality of the offspring produced is higher, then delayed maturity would reduce the juvenile mortality of the offspring. In the case of the present study, the delayed maturity of inland L. exotica is possibly due to the second reason. Additionally, the longer generation time of the inland population does not seem to result in a significant increase in mortality (Fig. 5). Thus, due to the unpredictable aridity of the inland habitat and the high juvenile mortality of L. exotica, a strategy of delayed maturity may be advantageous: life cycle delays can increase individual fitness in strongly variable environments.

Selection pressures on offspring size

Body size may affect survival probabilities, reproductive output, and individual fitness in many organisms (Twombly & Tisch 2000). Age-specific mortality
possibly subject to stronger selective pressure than capability of (Tsai 1997). Tsai et al. (1998) demonstrated that the fourth month after release displayed lower mortality than that of the littoral population in the present study. However, in given arid conditions (27°C, 85% relative humidity) in the laboratory, inland juveniles in the habitability occurs during the first few months after release, and juvenile mortality of the inland population was higher than that of the littoral population in the present study. However, in given arid conditions (27°C, 85% relative humidity) in the laboratory, inland juveniles in the fourth month after release displayed lower mortality (Tsai 1997). Tsai et al. (1998) demonstrated that the capability of L. exotica to resist desiccation was closely related to body size: large individuals with lower surface-area/volume ratios had higher desiccation resistance than smaller ones. The high juvenile mortality of L. exotica may be due to the small body size of its newly released juveniles. This may also explain why attaining a larger body size, especially in the juvenile stage, is important for inland L. exotica in reducing mortality due to desiccation stress. For the successful invasion from littoral to inland areas, mortality may be the main selective force driving life history traits. It is interesting to note that age and body size at maturation seem to be determined by age/size-specific survivalship, while morality of juveniles seems to be size-dependent. The larger maternal size of the inland population, as a consequence of delayed maturity, may be advantageous in producing more offspring to compensate for juvenile mortality or in producing larger juveniles. In the present study, inland females produced fewer but larger eggs instead of increasing their fecundity, due to the trade-off in egg size and number (Tsai & Chen 1997). This indicates that in the inland population of L. exotica, offspring size is possibly subject to stronger selective pressure than the fecundity, and that the inland habitat, with its periodical and unpredictable periods of desiccation, may favor females that produce larger juveniles. In many cases, offspring size is closely related to fitness (e.g., Parker & Begon 1986, Lloyd 1987; Bauer 1994). Under the constraint of reproductive investment, inland females may allocate more material/energy to increasing offspring size rather than number. The variations in the life history traits between the 2 populations of L. exotica indicate life history plasticity in response to environmental change; this plasticity may allow them to exploit various habitats.

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<table>
<thead>
<tr>
<th>Reproductive traits</th>
<th>Littoral population</th>
<th>Inland population</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean no. of eggs per brood</td>
<td>50.4</td>
<td>47.4</td>
<td>***</td>
</tr>
<tr>
<td>Mean no. of young per brood</td>
<td>48.3</td>
<td>46.6</td>
<td>***</td>
</tr>
<tr>
<td>Brood-pouch mortality</td>
<td>0.042</td>
<td>0.016</td>
<td>***</td>
</tr>
<tr>
<td>Mean brood weight</td>
<td>10.4 mg</td>
<td>11.8 mg</td>
<td>***</td>
</tr>
<tr>
<td>Mean maternal somatic weight</td>
<td>53.18 mg</td>
<td>57.74 mg</td>
<td>**</td>
</tr>
<tr>
<td>Mean egg weight</td>
<td>0.207 mg</td>
<td>0.243 mg</td>
<td>***</td>
</tr>
<tr>
<td>Mean juvenile weight(^a)</td>
<td>0.32 mg</td>
<td>0.35 mg</td>
<td>***</td>
</tr>
<tr>
<td>Reproductive investment(^b)</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>based on egg weight</td>
<td>19.52 %</td>
<td>20.58 %</td>
<td>***</td>
</tr>
<tr>
<td>based on juvenile weight(^a)</td>
<td>30.33 %</td>
<td>33.32 %</td>
<td>***</td>
</tr>
</tbody>
</table>

\(^a\)After release
\(^b\)(Brood wt/maternal wt) \times 100

Table 7. Ligia exotica. Comparison of reproductive traits of littoral and inland populations in Hengchun Peninsula, southern Taiwan. ***significant at level of \(p < 0.001\); **significant at \(p < 0.01\)

LITERATURE CITED


Lawlor LR (1976a) Parental and offspring fitness in terrestrial isopod Armadillidium vulgare (Latr.) (Crustacea: Oniscoidea). Evolution 30:775–785


may be an important factor in deciding life history traits (e.g., Schaffer 1974, Reznick & Endler 1982, Roff 1992). In the life cycle of Ligia exotica, highest mortality is related to body size: large individuals with lower surface-area/volume ratios had higher desiccation resistance than smaller ones. The high juvenile mortality of Ligia exotica may be due to the small body size of its newly released juveniles. This may also explain why attaining a larger body size, especially in the juvenile stage, is important for inland Ligia exotica in reducing mortality due to desiccation stress. For the successful invasion from littoral to inland areas, mortality may be the main selective force driving life history traits. It is interesting to note that age and body size at maturation seem to be determined by age/size-specific survivorship, while mortality of juveniles seems to be size-dependent.

The larger maternal size of the inland population, as a consequence of delayed maturity, may be advantageous in producing more offspring to compensate for juvenile mortality or in producing larger juveniles. In the present study, inland females produced fewer but larger eggs instead of increasing their fecundity, due to the trade-off in egg size and number (Tsai & Chen 1997). This indicates that in the inland population of Ligia exotica, offspring size is possibly subject to stronger selective pressure than the fecundity, and that the inland habitat, with its periodical and unpredictable periods of desiccation, may favor females that produce larger juveniles. In many cases, offspring size is closely related to fitness (e.g., Parker & Begon 1986, Lloyd 1987; Bauer 1994). Under the constraint of reproductive investment, inland females may allocate more material/energy to increasing offspring size rather than number. The variations in the life history traits between the 2 populations of Ligia exotica indicate life history plasticity in response to environmental change; this plasticity may allow them to exploit various habitats.

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