Impact of eutrophication on the life cycle, population dynamics and production of *Ampithoe valida* (Amphipoda) along an estuarine spatial gradient (Mondego estuary, Portugal)

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ABSTRACT: The life cycle, population dynamics and production of *Ampithoe valida* was studied from an intertidal mudflat in central Portugal, close to the northern limit of the species' distributional range in the eastern Atlantic Ocean. Sampling was carried out in eutrophicated areas, where macroalgae blooms of *Enteromorpha* spp. occur usually from January to early summer, and also in non-eutrophicated areas, with *Zostera noltii* meadows. *A. valida* showed a contagious distribution and the population density clearly changed during the study period along the eutrophication gradient. No migratory patterns were detected between the estuary and the sea, but migrations inside the estuary might have occurred. Females were morphologically recognisable at smaller sizes than males. Females reached sexual maturity before males, but males may live slightly longer than females. Females are iteroparous, producing 2, perhaps 3, broods. A 2-generation life cycle involving a short-lived (7 mo), fast-growing summer generation and a longer-lived (9 mo), slower-growing generation that overwinters is hypothesised. Ovigerous females were present year-round. Eggs, depending on the season, increase differently in volume during marsupial development. No correlations were found between fecundity (number of eggs) and the size of females. Along the eutrophication gradient no differences were found regarding the biology of the species. Besides these features, differences were observed between eutrophicated and non-eutrophicated areas with regard to productivity. Growth production (P) of *A. valida* in the most eutrophicated area was 0.098 g m⁻² 18 mo⁻¹ and 0.64 g m⁻² 18 mo⁻¹ in the *Z. noltii* meadows. P/\(\bar{B}\) and E/\(\bar{B}\) ratios (where E is the elimination production and \(\bar{B}\) is the average population biomass) ranged from 1.42 and 3.06 in the most eutrophicated area to 5.98 and 12.41 in the *Z. noltii* beds. To a certain extent, the increase of macroalgae biomass may favour *A. valida* populations, but extensive blooms affecting the whole area of distribution of this species will determine its disappearance.

KEY WORDS: Amphipoda · Life history · Production · Dynamics · Eutrophication · Macroalgae blooms

INTRODUCTION

Most European estuaries are affected by organic pollution and nutrient discharges, which often give rise to eutrophication (Hickel et al. 1993, Yeates 1993, Dijk et al. 1994, Flindt et al. 1997, Marques et al. 1997, Rafaelli et al. 1998). Eutrophication, as a response to nutrient enrichment, commonly causes proliferation of opportunistic green macroalgae, such as *Chaetomorpha*, *Enteromorpha* and *Ulva*, which can cover extensive areas of estuarine intertidal zones. It has been observed that benthic eutrophication in estuaries and coastal lagoons determines a shift from rooted plant communities, dominated by slow-growing species, like the eelgrass *Zostera*, towards free-floating (or partially free-floating) faster-growing macroalgae, like *Enteromorpha* or *Ulva* (Hartog 1994), as well as changes in the biomass and species composition of macroalgae (Lavery et al. 1991) and fauna (Pardal 1998).
Seasonal intertidal macroalgae blooms (mainly of *Enteromorpha* spp.) have been reported in the South Arm of the Mondego estuary (Portugal) for several years (Marques et al. 1993a,b, 1997, Pardal et al. 1993, Pardal 1998, Flindt et al. 1997, Lillebe et al. 1999) due to nutrient enrichment of estuarine waters (Marques et al. 1997, Pardal 1998). As a consequence, *Zostera noltii* beds, which represent the richest habitat with regard to productivity and biodiversity, have been drastically reduced in the South Arm of the estuary, presumably out competed by *Enteromorpha* (Rafaelli et al. 1991, Hodgkin & Hamilton 1993, Pardal 1998).

The intertidal and subtidal macrobenthic communities of the Mondego estuary were monitored from 1985 to 1990. Amphipods and isopods appeared as abundant groups, most probably playing an important role in the benthic communities (Marques & Nogueira 1991, Marques et al. 1993a,b, 1994, Martins et al. 1997, Zostera noltii meadows Pardal 1998). In the South Arm of the estuary, where increasing processes of eutrophication have been occurring, it is possible to observe the spatial gradient of replacement of *Zostera noltii* beds by green macroalgae. In this intertidal area, *Ampithoe valida* (Amphipoda) is one of the most abundant species.

*Ampithoe valida* is a grazer that can be found on rocky substrata and on muddy bottoms, often in the presence of green macroalgae or macrophytes (e.g. *Zostera noltii*) in salinities higher than 29 psu. This is a species previously found in the Pacific and along Atlantic coasts of North America (Conlan & Bousfield 1982). But even with this wide distribution the species has received little attention, except a study by Borowsky (1983) on the reproductive behaviour under laboratory conditions, and there are only a few studies on the impact of macroalgae blooms on grazer amphipod species population dynamics (Drake & Arias 1995). Such knowledge may definitely contribute to a better understanding of dynamic processes related with shift in primary producers and therefore to the understanding of its theoretical framework.

**MATERIALS AND METHODS**

**Study site and sampling.** The Mondego estuary is a warm-temperate coastal system on the western coast of Portugal (Fig. 1). It consists of 2 arms, north and south, with very different hydrological characteristics. The northern arm is deeper, while the southern arm is largely silted up, especially in upstream areas, which causes most of the freshwater discharge to flow through the northern arm. Consequently, the water circulation in the southern arm is mainly dependent on tidal activity and on the usually small freshwater input of a tributary, the Pranto River, which is controlled by a sluice.

Although, a large part of the southern arm intertidal area still remains more or less unchanged, having sand muddy bottoms covered by *Spartina maritima* marshes and *Zostera noltii* meadows, macroalgal blooms of *Enteromorpha* spp. have been regularly observed over the last 15 yr. This is probably a result of excessive nutrient release into the estuary, coupled with longer persistence of nutrients (nitrogen and phosphorus) in the water column (Flindt et al. 1997, Marques et al. 1997, Pardal 1998). Nevertheless, such macroalgal blooms may not occur in exceptionally rainy years (e.g. 1994) due to low salinity for long periods, as a result of the Pranto River discharge (Pardal 1998).

The population was monitored in the Mondego estuary for 18 mo, from January 1993 to June 1994. Samples of macrophytes, macroalgae, and associated amphipods were taken every 2 wk at 3 different sites during low water tides along an estuarine gradient of eutrophication in the south arm of the estuary (Fig. 1), from a non-eutrophicated zone, where a macrophytic

![Fig. 1. Location of the sampling stations along a spatial gradient of eutrophication in the Mondego estuary, Portugal](image-url)
community (Zostera noltii) was present, up to a heavily eutrophicated zone, in the inner areas of the estuary, from where the macrophytes disappeared while Enteromorpha spp. blooms have been observed there over the last decade. In this area, as a pattern, Enteromorpha spp. biomass normally increases from early winter (January/February) up to July, when an algae crash usually occurs due to anoxia and most of the biomass is washed out to the Atlantic.

Each time at each site 10 cores (with a 143 cm section) were taken to a depth of 15 cm. Each core was placed in a separate plastic bag and sieved within an hour of sampling by washing it in estuarine water through a 500 μm mesh sieve. Given the size of newly hatched juveniles, this mesh should retain all individuals. The residue (sediment, macrophytes or algae and amphipods) was placed into plastic bottles and preserved with 4% formalin in estuarine water.

**Laboratory procedures.** Amphipods were identified, counted, measured and sexed. Due to the difficulties in accurately measuring total body length (\(T_L\)) in such small organisms that are typically comma-shaped, an alternative and more accurate length was measured: the cephalic length (\(C_L\)) (measured between the extremity of the rostrum and the base of the head). We measured 261 individuals (males females and juveniles) for both lengths, and an equation for \(C_L - T_L\) conversion was determined (\(T_L = -0.1355 + 9.4233C_L\), \(n = 262, r = 0.965\)). Length-weight relationships were determined for production estimates. Since length ash-free dry weight (AFDW) relationships did not show significant seasonal differences based on analysis of variance (\(p \leq 0.05\)), a data set obtained throughout the study was used to provide a single regression equation (\(W [\text{weight}] = 0.00085 \times C_L^{0.631}\), \(n = 95, r = 0.983\)). Individuals were dried at 60°C for 72 h and weighed to the nearest 0.01 mg. Small individuals were pooled to obtain measurable values. AFDW was assessed after combustion of samples for 8 h at 450°C. The same procedure was used to quantify the macroalgae and Zostera noltii biomasses.

The determination of sex was based on the presence or absence of oostegites and/or broods (males), and of genital papillae (males). Animals without these features were considered to be juveniles. When broods were present, eggs were counted (to estimate the fecundity), measured and examined to determined the development stage. Taking into consideration several similar criteria (Steele & Steele 1969, Goedmakers 1981, Skadsheim 1982, Marques & Nogueira 1991, Marques et al. 1994) 5 stages were considered: (1) newly laid, eggs grouped and resembling a gelatinous mass; (2) eggs well separated, internally homogeneous; (3) embryo comma-shape, pereopods starting to be visible; (4) constriction of the comma clearly visible, appendages segmented and looking slender, eyes visible, cephalorax orange-red; (5) hatched and free juveniles, which may stay for some days in the brood pouch. Since eggs become oval in shape during development, an average of their length and width was taken as measurement of their size.

**Statistical analysis.** An index, \(I = S^2/X/X\) where \(S^2\) is the variance of abundances and \(X\) is the mean abundance; Elliot (1977) was used to analyse the population spatial dispersion. Stepwise, least squares multiple regression models for Ampithoe valida were developed to examine the relationship between the density and biomass of A. valida and the biomass of macroalgae, salinity and water temperature. All variables except temperature and salinity were submitted to a ln(x + 1) transformation prior to analysis in order to minimise heteroscedasticity. Only variables with partial effects significant at the \(p < 0.01\) level were retained in the regression model.

**Growth.** Growth rates were estimated like in previous papers of Marques & Nogueira (1991), Marques et al. (1994), Pardal (1998), and Lillebø et al. (1999) by tracking recognisable cohorts with size frequency distributions from successive sample dates.

Growth rates are not constant throughout the year. Thus to express field growth rates, we used a model that takes into consideration seasonal changes (Guschütz et al. 1980), expressed as

\[
L_t = L_0 \{1 - e^{-kD(t - t_0)} + C(1 - e^{D/2}) \cos 2\pi(t - t_1)/D\}
\]

where \(L_t\) length of the organism at a given moment \(t\); \(L_0\): maximum possible length of the organism; \(t_0\): instant when the organism would have a length = 0; \(t_1\): time interval between start of growth (when \(t = 0\)) and the first growth oscillation; growth is expressed by a sine curve with a period of 1 yr (the fact that the sine function has a period of 1 yr addresses normal seasonal oscillation of growth rates with temperature. Animals under study may live less [or more] than 1 yr, but their growth rate oscillation will be expressed by the sine curve function. The model of course is calibrated with real data from field observations); \(k\): intrinsic growth rate; \(C\): constant, whose values can change from 0 to 1; and \(D\): parameter that expresses metabolic deviations from the Von Bertalanffy \(\frac{2}{2}\) rule.

**Production.** Production was estimated like in previous papers of Marques & Nogueira (1991), Marques et al. (1994), Pardal (1998), and Lillebø et al. (1999), and was based upon cohort recognition. Growth increments or net production (\(P\)) and elimination production (\(E\)) were estimated with a method derived from Allen (1971), as described in Dauvin (1986).

Total values of \(P\) and \(E\) for the population are expressed as:

\[
P = \sum_{n=1}^{N} P_{cn}
\]
\[
P, \text{ and } E, \text{ are the growth and elimination of cohort } n.
\]

\[
P/B \text{ and } E/B \text{ ratios were determined. } B \text{ (average population biomass) is expressed as:}
\]

\[
\bar{B} = \frac{1}{T} \sum_{n=1}^{N} (B_n t)
\]

where \(T\): period of study; \(N\): number of successive cohorts in the period \(T\); \(B_n\): average biomass of cohort \(n\); \(t\): duration of cohort \(n\).

**RESULTS**

**Macrophyte and macroalgae biomasses**

As expected, *Zostera noltii* biomass exhibited a clear seasonal variation. During autumn and winter total biomass depended essentially on the rhizomes, while during spring and summer total biomass increased due to the growth of leaves (Fig. 2A).

The specific composition of macroalgae biomass changed very much along the eutrophication gradient. Red macroalgae, essentially *Gracilaria* sp., were relatively abundant at the *Zostera noltii* meadows, decreasing along the eutrophication gradient, while green macroalgae showed exactly the opposite pattern (Fig. 2).

In the intermediate eutrophicated area small amounts of green macroalgae were present in both spring seasons, but the biomasses estimated could not be considered a typical bloom.

In the most eutrophicated area *Enteromorpha* sp. exhibited a typical spring bloom in 1993, with biomass reaching 413 g m\(^{-2}\) (AFDW) in April. In early summer an algae crash occurred, causing a severe impact on the macrofauna in the area attained by the phenomenon (Figs. 2 & 3). In 1994 no macroalgae bloom occurred because it was a very rainy year in the region which increased freshwater discharges (by the sluice of Pranto River), keeping salinity low. Such conditions inhibited *Enteromorpha* growth, since salinity was below 10% for several months.

**Spatial distribution and abundance**

*Ampithoe valida* showed an aggregated spatial distribution in the 3 study areas along the eutrophication gradient. Estimated values of \(I\) were always higher than 1 (Elliot 1977). Population density changed throughout the period of study but the pattern of variation was not the same at the 3 sampling stations (Fig. 3). In the most eutrophicated area density was consistently higher during the algae bloom, with a peak in April (2026 ind. m\(^{-2}\)). After the algae crash density declined sharply and remained low, or organisms even disappeared throughout the study (Fig. 3A).

In the intermediate eutrophicated area the variation in density was correlated with the variation in macroalgae biomass (\(r = 0.87, n = 26\)), but density never reached abundances as high as in the previous case (maximum peak, 154 ind. m\(^{-2}\)) (Fig. 3B). With abundances so low, most of the statistical analyses that we performed in the present paper were not possible in this area. At the *Zostera noltii* meadows the population...
Nevertheless, in the *Zostera noltii* meadows no significant correlation was observed between these population parameters and leaf biomass, salinity or temperature of the water.

**Reproduction, sex ratio, and fecundity**

The *Ampithoe valida* population was sexually active throughout the year in the south arm of the Mondego estuary. However, taking into consideration the percentage of ovigerous females over the total female population (Fig. 4), sexual activity was higher from late spring to late summer. The variation of the percentage of juveniles in the population was clearly related to increases in recruitment.

Females were usually more abundant than males overall, and no significant seasonal changes of this pattern were observed (Fig. 5). However, some inversions of the values might be observed, due to the death of older females after recruitment (June and August of 1993 and April/May of 1994). Males appeared to live longer than females. Also these maxima in sex-ratio

**Fig. 3. Abundance of *Ampithoe valida* along the eutrophication gradient. (A) Most eutrophicated area; (B) eutrophicated area; (C) *Zostera noltii* meadows**

Density increased during summer and fall of 1993, reaching a maximum in November (887 ind. m$^{-2}$), and also during spring of 1994 (maximum peak, 710 ind. m$^{-2}$) (Fig. 3C).

Stepwise multiple regression analysis indicated that abundance (A) and biomass (B) of *Ampithoe valida* in the most eutrophicated area was positively correlated with *Enteromorpha* sp. biomass (ENT) and *Gracilaria* sp. biomass (Grace):

$$\ln(1 + A) = 0.350 + 1.142 \ln(1 + \text{ENT})$$

$$\ln(1 + B) = -0.030 + 0.030 \ln(1 + \text{ENT}) + 0.096 \ln(1 + \text{Grace})$$

**Fig. 4. *Ampithoe valida*. Biological features of the population. (A) variation of females in different physiological stages in relation to the female population; (B) variation of ovigerous females in the total female population and of juveniles in the population**
were normally followed by an increase in the proportions of unreproductive females.

The fecundity of females of similar size was quite variable, and no correlation between the number of developing embryos and female size was found. Measurements of eggs in different developmental stages (Table 1) showed that egg volume increase changes seasonally. For instance an increase of 361% was estimated in summer, 452% in spring and 492% in autumn. It was also possible to recognise that for the same developing stage egg volume was greater in colder periods, while the average number of eggs per female was smaller (Table 1).

The difference in time between peaks of eggs in the first stage of development (A) and the last one (E) provided us with an estimation of the duration of the embryogenic development (Fig. 6). During spring and summer the embryogenic development was faster (30 to 45 d) than in autumn or winter (60 d).

**Growth and lifespan**

Size frequency polymodal distributions were analysed for recognisable cohorts (Fig. 7), allowing the identification of significant differences in the population structure on the *Zostera* beds and in the most eutrophicated area. Individuals reached larger dimensions at the most eutrophicated area in comparison with the other 2 areas (*Z. noltii* meadows and intermediate eutrophicated areas). Nevertheless, in the first months the recruitment pattern was the same all along the eutrophication gradient. Following the algae crash the population disappeared from the most eutrophicated area and cohort tracking became only possible at the *Z. noltii* meadows. In January 1993, 3 cohorts were identified, and 6 new ones were recognised during the study period ($\chi^2$ and G not significant, $p \leq 0.05$). Minimum average $C_L$ of cohorts ranged from 0.327 (spring) to 0.397 mm (winter), corresponding to 2.946 and 3.606 mm of total length.

Growth was continuous through life (Fig. 8). Nevertheless, growth rates were higher during spring and summer and decreased during winter. Lower growth rates during winter were probably a function of lower temperatures and lower biomasses of macroalgae (food resources) in the estuary.

Growth data from cohort C6 were used to calibrate a growth model proposed by Gashutz et al. (1980) (Fig. 9). The model parameters were estimated as follows: $L_m$, which was fixed at a value of 1.6 mm $C_L$, slightly larger than the maximum value we observed; $k = 1.401$; $T_0 = -0.116$; $t_0 = 0.625$; $C = 0.175$; $D$, since metabolic rates are not known, it was assumed that there were no deviations from the Von Bertalanffy's $\frac{2}{3}$ rule, and therefore $D$ was fixed at 1.

Growth data fitted the model very well ($r = 0.994$). Based on the model it is clear that growth rates decreased during cold months. During the rest of the year growth rates were higher.

Lifespan were estimated at 191 ± 30 to 238 ± 15 d for summer and spring cohorts and at 242 ± 30 for the autumn cohort. Similarly, the age at which females and males matured was lower in spring and summer
cohorts than in the autumn-winter cohorts (Table 2). Nevertheless, during the yearly cycle, males always reached maturity before females.

**Production estimates**

Length-weight relationships previously established were used in production estimates in the most eutrophicated area and in the *Zostera noltii* meadows (Fig. 10, Table 3). Taking into account the whole period of study, results clearly showed that growth productivity (P), elimination productivity (E) and the relations P/B and E/B were much higher in the *Z. noltii* meadows. But in the most eutrophicated area during the algae bloom P, E and population density (Fig. 4) were clearly higher than in *Z. noltii* meadows (Fig. 10).

**DISCUSSION**

**Abundance**

Population density clearly changed along the eutrophication gradient as a result of the macroalgal annual dynamics, as previously observed for *Microdeutopus gryllotalpa* (Drake & Anas 1995). In fact, in the most eutrophicated area, the occurrence of a macroalgal bloom in the first year prepared the conditions for the presence of large densities of *Ampithoe valida*. The algae crash, which acted as a catastrophic event (Marques et al. 1997), and the non-occurrence of an algae bloom in the second year had a strong effect on the population’s density in the inner parts of the estuary. Consequently, since in the second spring (1994) there were no macroalgal habitats available *A. valida* individuals seemed to occur only in the *Zostera noltii* meadow.

During the macroalgal bloom the population density was much higher (2026 ind. m\(^{-2}\) by 22 June 1993) than in the *Zostera noltii* meadows in the same period. This may be related to habitat protection against potential predators (crabs, fishes and birds) and food resources (Greeze 1968, Duffy & Hay 1991) since they can feed directly on the algae. Next, during the algae crash, when anoxic conditions occurred, the population suffered a sudden and drastic reduction to 56 and 0 ind. m\(^{-2}\) (6 July and 5 August). After the algae crash the population was never able to recover since no food resources were available. In contrast, in the *Zostera noltii* meadows the population increased in number after the algae crash. A possible movement of adult individuals inside the estuary, moving to avoid the extreme conditions of anoxia, is the most probable explanation, together with new recruitments taking place in this area. During this specific period the movement of adult individuals was identified based on the observation that the older cohorts increased greatly in number of individuals with no other possible explanation available. It seems therefore that this species has a large potential for dispersion inside the estuary, avoiding as much as possible the situations of stress caused by macroalgae blooms in early summer. It seems reasonable to conclude that to a certain extent the development of macroalgae biomass favours *Ampithoe valida* populations, but extensive blooms affecting the whole area of distribution of this species will determine its disappearance.

**Reproduction, growth and life cycle**

Data showed that females almost always outnumbered males, which is a common feature in amphipod populations (Hastings 1981, Dauvin 1988a,b, Marques & Nogueira 1991, Morritt & Stevenson 1993, Covi & Kneib 1995, Sudo & Azeta 1996). The values were close to 1:1 only after recruitments, as a function of the death of older females after reproduction. Moreover, young males developed secondary sexual characteristics earlier than females (Table 2).

The duration of the embryogenic development in amphipods from high latitudes is very long and can reach 6 mo (Bregazzi 1972). In contrast, in temperate coastal areas it is much faster, taking usually only a few weeks (Steele & Steele 1973, Moore 1981, Powell & Moore 1991). In our case embryogenic development

<table>
<thead>
<tr>
<th>Season</th>
<th>Maturation (d)</th>
<th>Males</th>
<th>Females</th>
</tr>
</thead>
<tbody>
<tr>
<td>Winter</td>
<td>44</td>
<td>61</td>
<td></td>
</tr>
<tr>
<td>Spring</td>
<td>24</td>
<td>28</td>
<td></td>
</tr>
<tr>
<td>Summer</td>
<td>35</td>
<td>37</td>
<td></td>
</tr>
<tr>
<td>Autumn</td>
<td>41</td>
<td>57</td>
<td></td>
</tr>
</tbody>
</table>

Table 2. *Ampithoe valida*. Annual cycle of maturation in the Mondego estuary

<table>
<thead>
<tr>
<th>Season</th>
<th>P/B</th>
<th>E/B</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Z. noltii</em></td>
<td>0.64</td>
<td>5.98</td>
</tr>
<tr>
<td>Most eutrophicated area</td>
<td>0.098</td>
<td>1.42</td>
</tr>
</tbody>
</table>

Table 3. *Ampithoe valida*. Production estimates at *Zostera noltii* meadows and at the most eutrophicated area, taking into account the whole period of study (18 mo)
Fig. 7  *Amphitrite valida*. Size-frequency polymodal distribution at *Zostera noltii* meadows and at the most eutrophicated area. Arrowheads indicate average cephalic length of numbered cohorts; N: number of individuals.
Zostera noltii meadows

Most eutrophicated area

Fig. 7. (continued)
they contain higher nutritive reserves. Such volume increase was higher than the one observed by Moore & Wong (1996), although they were similar to the ones calculated by Marques & Nogueira (1991) for *Echinogammarus marinus*. Moreover, in spring and summer sexual maturity was reached at smaller sizes in comparison with autumn cohorts, which agrees with previous observations on other amphipods (Moore 1981, Powell & Moore 1991, Sudo & Azeta 1996).

Ovigerous females of *Ampithoe valida* were found throughout the year in the Mondego estuary, and the population is multivoltine with a maximum of 3 generations yr⁻¹. Continuous sexual activity during the year is a common feature in amphipods (Franz 1989, Marques & Nogueira 1991, Covi & Kneib 1995, Drake & Arias 1995, Moore & Wong 1996). Unfortunately the life cycle of this species at different latitudes was not studied, but our results are consistent with the generalised notion that ecosystems with warmer temperature regimes allow earlier maturation and higher voltinism (Birklund 1977, Marques & Nogueira 1991, Sudo & Azeta 1996).

Summer and spring growth rates were much higher than during colder months, which was also observed for other amphipod species in temperate areas (Hastings 1981, Dauvin 1988a,b, Franz 1989, Marques & Nogueira 1991, Uitto & Sarvala 1991, Drake & Arias 1995, Wilson & Parker 1996), and our estimated growth rates were similar to other speci-

Like other amphipod species, *Ampithoe valida* is a semi-annual with short-lived spring and summer generations (7 to 8 mo) and longer-lived autumn and winter generations (9 mo) (Moore 1981, Powell & Moore 1991, Beare & Moore 1994, Sudo & Azeta 1996).

*Ampithoe valida* appeared as an r strategist, with iteroparous females (except summer generation), a multivoltine cycle, high individual fecundity and recruitment throughout the year. This is the most common pattern in epifaunal species (van Dolah 1980) from physically controlled communities according to the stability-time theory (Sanders 1969). In environmentally stressed systems like estuaries we should expect the evolution of opportunistic adaptive strategies to take place.

**Production estimates**

Taking into account the whole period of study, *P* (growth productivity) and *E* (elimination productivity) as well as *B* (standing stock) showed significantly higher values in the *Zostera noltii* beds than in the most eutrophicated area. Nevertheless, this difference does not reflect the entire reality. In fact we must look carefully at the dynamics of the 2 areas. For a short period (during the macroalgae bloom) in the most eutrophicated area *P*, *E* and *B* were higher than in the *Z. noltii* meadows. This means that during the macroalgae bloom the most eutrophicated area was clearly the preferential habitat for *Ampithoe valida* in the Mondego estuary probably due to resource availability (Fig. 10).

The results clearly suggest that *Ampithoe valida* population dynamics and spatial distribution may change as a function of environmental changes through adaptive behavioural mechanisms. Nevertheless, it appears that macroalgae may mainly represent a good additional resource for populations with a stable habitat like *Zostera noltii* beds. But the macroalgae mats alone cannot sustain stable populations of *A. valida* due to their own dynamics. A conclusion would be that the total replacement of *Z. noltii* by macroalgae mats would negatively affect this kind of grazer.

As in other intertidal areas of Europe and North America, *Ampithoe valida* in the Mondego estuary is quite accessible to aquatic predators. High *P/B* and *E/B* ratios found in the *Zostera noltii* meadows, as well in the most eutrophicated area during the macroalgae bloom (Table 3, Fig. 10), suggest that this species may play a more important role in the trophic dynamics than one might expect from its standing stock biomass.
As pointed out by Fredette & Diaz (1986), our results show that in warm shallow marine habitats relatively low biomass of benthic invertebrates can result in large production estimates, because a lack of extreme cold temperatures might allow prolonged reproduction periods (Kalejta & Hockey 1991).

As a whole, the present work reinforces the generalised notion that estuaries are highly productive systems, and reflect the important role of amphipods in the productivity of habitats colonised by them. In the Mondego estuary, although macroalgae seem to be favourable for this amphipod population, if we look at the following effects of the algae crash it is reasonable to conclude that more extensive blooms which affect the whole area of distribution of the species will, on the contrary, have a strong negative impact and could imply its disappearance.

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