Reproductive biology of Diopatra neapolitana (Annelida, Onuphidae), an exploited natural resource in Ria de Aveiro (Northwestern Portugal)

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Introduction

The polychaete Diopatra neapolitana Delle Chiaje, 1841 (Onuphidae) inhabits intertidal mudflats and shallow subtidal transitional waters. The geographical distribution records indicate that it is a cosmopolitan species distributed throughout the Mediterranean (Gambi & Giangrande 1986; Dagli et al. 2005), the Red Sea (Fauvel 1923) and the Eastern Atlantic (Fauvel 1923; Lourido et al. 2008) and Indian Oceans (Wehe & Fiege 2002). However, in regions of the world where careful genetic and morphological analysis has been conducted, it was shown that D. neapolitana harbors multiple species. In Europe, four species of Diopatra, D. neapolitana, Diopatra marocensis, Diopatra micrura and Diopatra sp. (not yet described) were identified and distinguished morphologically using characters that have not been used previously (Pires et al. 2010). Such analysis could be applied in other regions, in particular the Red Sea and Indian Ocean.

The species inhabits a tube, has a preference for sediments with mud or a mixture of mud and sand, and grows to about 60 cm (Fauvel 1923; Gambi et al. 1998; Dagli et al. 2005; Rodrigues et al. 2009). The tube consists of a secreted layer, to which sand particles, fragments of solid parts from other animals, such as shells, and algae attach to form a compact tube.

Keywords
Larval development; life history; oocytes; polychaeta; reproduction.

Abstract

Diopatra neapolitana Delle Chiaje, 1841 (Annelida, Onuphidae) is an important economic natural resource in Ria de Aveiro (northwestern coast of Portugal) and throughout Europe. The species is intensively harvested for use as fresh bait. However, there is only limited knowledge about its life cycle derived from a previous study in Mediterranean Sea. Reproduction and development patterns are known to vary biogeographically, making it important to base management decisions on locally appropriate information. This work examines reproduction patterns for populations from the Eastern Atlantic, which have not previously been assessed, with an eye towards drawing Atlantic–Mediterranean comparisons and informing local management strategies. The study was conducted from May 2007 to April 2009 in Ria de Aveiro. The reproductive biology of D. neapolitana was described from the proportional variation of worms with gametes in the coelom and from the progression of the oocyte diameter. Individuals with gametes inside the coelom were found all year round, but the peak reproductive period occurred between May and August, when almost all individuals had gametes in the coelom and females contained more oocytes than at any other time of the year. The overall male:female ratio was close to 1:1 and the oocyte diameter ranged from 40 to 240 µm. In vitro fertilization was performed and the results compared to other studies. Based on the present results, some protection measures are suggested to implement a sustainable exploitation of the species.
Larval development in the Onuphidae is dependent on yolk reserves, with some species being lecithotrophic, feeding only after settlement, and others having a direct development (without larval stages) (Blake 1975; Giangrande 1997). Conti et al. (2005) report that D. neapolitana releases eggs and sperm into the water column and Baud & Cazaux (1987) that it produces planktonic lecithotrophic larvae. Although the spawning of this species has never been observed in nature, artificial fertilization and culture of the larvae was reported by Baud & Cazaux (1987) and Conti & Massa (1998), who described several developmental phases. These authors showed that the larvae were lecithotrophic and free-swimming.

This species is collected to be sold as fish bait and this activity can be locally intense and economically important (Gambi et al. 1994; Conti & Massa 1998). A previous study in Ria de Aveiro, Northwestern Portugal, where the present study was undertaken, indicated an annual harvest of 45,000 kg, valued at over € 325,000 (Cunha et al. 2005). According to Portuguese legislation, bait collection is only allowed by hand gathering or with restricted gear, such as a hoe, operated by licensed personnel (Portuguese legislation: Portaria no 144/2006 2006). No other legislation exists for the Ria de Aveiro and no management or conservation efforts are currently being developed for this species. Its reproductive biology is relatively unknown, as the only field work ever done on this subject was carried out in the Eastern Mediterranean Sea by Dagli et al. (2005).

The present study focuses on the gametes’ characteristics, the larval development, the reproductive period, and the sex ratio of the population of D. neapolitana in Ria de Aveiro. Understanding these life history aspects is important for management and conservation efforts aimed at a sustainable exploitation of the species.

Material and methods

Study area and sampling

This study was conducted in Ria de Aveiro, Northwestern Portugal (40°40′01.6″ N, 8°41′39.5″ W; Fig. 1). Ria de Aveiro is a shallow estuarine water system, receiving water from several rivers (Fig. 1), with the Vouga River accounting for more than 50% of the freshwater input, resulting in a complex system of bays, channels and extensive intertidal sand and mud flats (Dias et al. 1999).

Diopatra neapolitana specimens were collected intertidally, monthly from May 2007 to April 2009 with a shovel, at up to 30 cm depth. At least 50 specimens were collected randomly, each month, at the study area.

Laboratory procedures

In the laboratory worms were individually removed from their tubes and washed in sea water. Each specimen was partly dissected to search for the presence of gametes and then fixed in 70% ethanol. Fixed specimens were measured for width at the 10th chaetiger (without parapodia). Total length was measured in entire specimens (about 4% of individuals). These morphological variables were only measured in individuals that were not seen to be regenerating.

The oocytes were extracted from females by dissection of the body cavity. The diameter of each oocyte was measured under a stereomicroscope (resolution 50×) using an ocular micrometer (precision of 0.01 mm). The diameter of 100 oocytes was measured for each female. Different numbers of females were collected each month. During the periods with a larger number of mature individuals (April–August) oocytes were measured in at least 12 females. In the remaining study period, oocytes were measured in all the females collected, as their number was below 12. In some cases, only two to four females with oocytes in the coelom cavity were sampled. In total, oocytes from 332 females were measured. To count the total number of oocytes per female, only complete specimens were used – 12 in total. These were collected between May and December. During the study period, fresh sperm in sea water was observed under a microscope (resolution 1000×).

Fertilization in vitro

Specimens were collected from the study area and kept in the laboratory for at least 2 months. They were maintained at 22 °C and at a salinity range of 30–35. Salinity was measured with a hand-held refractometer and expressed using the practical salinity scale. To study larval development, artificial fertilization was performed following the method described by Conti & Massa (1998) for Diopatra neapolitana. Females and males were cut laterally and left in separated dishes with sea water for 10–15 min to release the eggs and sperm. A portion of sperm was collected and added to the oocytes. The fertilized eggs were cultured at 22 °C and 30–35 salinity. Sea water was changed daily.

The larval development observed in this study was analyzed following the descriptions of Baud & Cazaux (1987) and Conti & Massa (1998).

Once settled, the larvae were fed with homogenized cockles. Four days after fertilization, in the metatrochophore phase, the larvae were moved to an aquarium with fine sediment. The study of larval stages was carried out under an optical microscope.
Data analysis

The relationship between total length ($L$) and the width of the 10th chaetiger ($W$) was studied using second-order polynomial simple regression analysis. This relationship was established from 46 complete individuals, collected over the entire study period, according to the function $L = a + b_1 W + b_2 W^2$, forcing the model through the origin ($a = 0$). SPSS software (version 17) was used to test the overall significance of the model ($F$-test) and of the second-order regression coefficient ($b_2$, $t$-test). The total expected body length of broken specimens was then determined from the measured width of the 10th chaetiger, using the regression function. This relationship was used to determine the expected shortest length of mature individuals.

The mean oocyte diameter (MOD) was calculated per female and per month, and its correlation with total length was assessed using the Pearson coefficient. The variance of oocyte diameter in the period of gametogenesis inactivity (November–January) was statistically compared ($F$-test) with the period of gametes production (March–October).

Results

Relationship between total length and width of the 10th chaetiger

 Entire mature specimens ranged in size from 24 to 725 mm, with the width of the 10th chaetiger varying between 1.9 and 10.88 mm, respectively (Fig. 2). All observed specimens, entire or incomplete, had a 10th chaetiger width of between 1.9 and 13 mm. The regression function relating the body length of the specimens ($L$, in mm) to the width of the 10th chaetiger ($W$, in mm) was statistically significant ($F = 1081.5$; $P < 0.0001$) and was given by the expression $L = 17.955 W + 4.209 W^2$. The regression coefficient associated with $W^2$ ($b_2 = 4.209$) was also found to be significantly different from zero, validating the second-order polynomial ($t = 6.945$; $P < 0.0001$). Under this
regression model, the width of the 10th chaetiger explained 98% of the total length variance ($R^2_{adj} = 0.98$). This regression function was used to estimate the total length of broken specimens. The smallest female observed to be carrying oocytes had $W = 4.2$ mm, corresponding to an estimated body length of 149.7 mm. The smallest male with sperm in the coelom had $W = 4.0$ mm, corresponding to an expected body length of 139.2 mm.

Reproduction of *Diopatra neapolitana*

The presence/absence of gametes was analyzed in 1163 specimens, of which 320 were males, 332 females and 511 undetermined (with no gametes in the coelom). No external morphological differences were noticed between males and females. However, during the main reproductive period males turned a cream color and females became greenish, mainly due to the gametes in the coelom.

The overall male:female sex ratio was close to 1:1 from April to September. For the other months, very few individuals with gametes were captured and the sex ratio was not determined.

The reproductive cycle of *D. neapolitana* can be inferred from the proportional variation of worms with gametes in the coelom, from the development of the size of oocytes and from the number of oocytes in complete females (Figs 3 and 4; Table 1). Individuals with gametes inside the coelom were always found, but the percentages of males, females and of individuals without gametes varied (Fig. 3) and showed a consistent pattern in the two consecutive years. In February 2008 and February 2009, a single specimen with oocytes and a single specimen with sperm were found, respectively, whereas in April–August a larger proportion of individuals with gametes (varying from 39.22–54.29% in females and 35.14–50.0% in males) (Fig. 3) were found.

The smallest oocyte found in a female’s coelom had a diameter of 40 $\mu$m and the largest a diameter of 240 $\mu$m, with the mean for all specimens being 164.4 ± 40.8 $\mu$m. Small oocytes (<140 $\mu$m) were present in almost every month. The number of small oocytes reached a peak in March and April, decreasing until September. Oocytes were absent in some autumn/winter months (October, November and December) (Fig. 4). The decrease in the number of small oocytes paralleled the increase of larger oocytes (Fig. 4). The mean oocyte diameter was not correlated with the size of the females, measured by the width of the 10th chaetiger ($r = -0.011$; $P > 0.05$). Mean oocyte diameter increased rapidly from March to May, and continued to increase slowly until January (Fig. 4). The variance in oocyte size was significantly larger from March to October ($s^2 = 1354$) than during the winter months, from November to January ($s^2 = 264$; $F = 5.1$; $P < 0.001$). This can be appreciated in Figs 4 and 5.
Females from November to January contained mainly large oocytes of between 140 and 240 μm (Fig. 5). Nurse cells were observed in oocytes with a diameter of up to 160 μm (Fig. 6A). They were attached to the immature oocytes with two strings measuring up to 230 μm in length (mean = 177.5 ± 35.4 μm) and containing up to 39 cells (mean 29.4 ± 4.5 μm) 12 μm in diameter. Oocytes larger than 160 μm did not have nurse cells attached (Fig. 6B).

Sperm had a spherical, short and rounded head with a long tail and were grouped in capsules in the coelom. When sperm were observed under the microscope, between May and August, the majority of the males contained spermatozoa with a mobile flagellum, moving actively in sea water. From October to January, spermatozoa had tails but reduced mobility. Sperm were immobile during the other months.

The first chaetiger with gametes varied. In females where the oocytes were observed they were located between the 35th and the 70th chaetiger. In males, sperm were found from chaetigers 50 to 70. The mean location of the chaetiger where gametes first appeared was 52.7 ± 8.6 for oocytes and 59.3 ± 7.3 for sperm. No significant correlation was found between the first chaetiger bearing gametes and the size of the individuals (r = 0.01).

In May and June, the months where it was possible to count the total number of oocytes per female (complete females), females had the highest number of oocytes in the coelom (Table 1).

Fertilization in vitro

Table 2 presents the main characteristics of larval development of *Diopatra neapolitana* in this and in other studies (Bhaud & Cazaux 1987; Conti & Massa 1998).

Larval development was followed up to the age of 7 days. Seven hours after fertilization the embryo had cilia and swam in the water column, becoming a free-swimming protrochophore larva after 19 h. The protrochophore larvae were sub-spherical, with an apical tuft and were almost completely covered by cilia 210 μm in length. At 2–3 days after fertilization, the metatrochophore larvae had a length of between 240 and 280 μm and were segmented in three chaetigers with chaetae; the prostomium was ciliated and with two red eyes. After 3 days, the metatrochophore larvae lost the apical tuft, had four chaetigers and a length of 300 μm. On the 4th day, some metatrochophore larvae swam slowly in the water column, and others started to sink to the bottom and aggregate detritus around them. At this phase, larvae were moved to an aquarium with fine sediment and fresh sea water to allow the larvae to create a wrapping and protecting niche, and later permit the construction of the tube.

Juveniles were observed 7 days after fertilization and had five chaetigers, five small antennae on the prostomium, and a pair of small anal cirri in the pygidium. Tube formation was not observed, although individuals with particles around the body were seen.

Regenerating specimens

During the study period, about 5% of the specimens were regenerating the anterior end of the body, from two to 13 chaetigers. A minor proportion of specimens, about 0.3%, were regenerating the posterior end and a much larger number of chaetigers (56 to >100). Specimens regenerating the anterior end were found in almost all the sampling occasions, and represented between 1.4% and 17.0% of the sampled population. Individuals regenerating the posterior end were rare and only observed in 5 sampling occasions, randomly scattered throughout the sampling period. The majority of the regenerating specimens did not contain gametes, with the exception of some females with small oocytes.

Discussion

The study of the reproductive biology of *Diopatra neapolitana* showed that this species contained gametes in the coelom in all months of the year, but had the highest proportion of individuals with gametes from May to August. These results are similar to those of Dagli *et al.* (2005) in Izmir Bay, Turkey, where individuals with gametes were reported all year round, except in January. The number of oocytes in the females’ body cavity was higher in May–August, but was decreasing during this period. In the month of October and December, it was similar numbers of oocytes were found in the body cavity.
Fig. 5. Size-frequency distribution of oocytes of *Diopatra neapolitana* during the study period (n = number of females observed).
of females, suggesting that no oocytes were released over this period. A number of small oocytes (<140 μm) was present almost every month, showing a peak in March and April, and then decreasing until September. The absence of small oocytes between November and January indicates that the females were not producing oocytes. The large oocytes probably were not expelled during spawning and remained in the coelom cavity. The decrease of small oocytes was paralleled by an increase in the number of larger oocytes.

Conti et al. (2005) described mature sperm of *D. neapolitana* as having a long tail attached to a spherical, short and rounded head, which is similar to our findings for sperm from May to August. The sperm also had the highest mobility during this period. These results indicate that the beginning of gametogenesis should be in March/April, the spawning period from May to August, and gametogenic inactivity from November to February.

The oocyte diameter varied between 40 and 240 μm, with a mean of 164.4 ± 40.8 μm. Oocytes should be released from the coelom into the water column with a diameter of about 200 μm (Dagli et al. 2005). In fact, in the present study only a small percentage of the oocytes remaining in the coelom had a larger diameter. This is in agreement with our observations of the artificial fertilization, and with Bhaud & Cazaux’s (1987) results, as the fertilized eggs had a diameter between 210 and 215 μm.

![Fig. 6. Oocytes of *Diopatra neapolitana*. (A) Immature oocyte with nurse cells attached. (B) Mature oocyte.](image)

### Table 2. Principal characteristics of larval development of *Diopatra neapolitana* in this study and in comparison with Bhaud & Cazaux (1987) and Conti & Massa (1998).

<table>
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<tr>
<td>P 19 h. Shape sub-spherical. Almost completely covered by cilia. Apical tuft. Larvae swimming actively in water column</td>
<td>24 h. Shape sub-spherical to piriform. Length 215 μm. Almost completely covered by cilia. Apical tuft. Larvae swimming in water column</td>
<td>5 h. Larvae swim free in the water column</td>
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<tr>
<td>M 2–3 days. Length 240–280 μm. 3 chaetigers. Prostomium ciliated. 2 red eyes</td>
<td>3 days. Length 240 μm. 3 chaetigers. Prostomium ciliated. 2 red eyes</td>
<td>24 h. Larvae present. Positive phototropism</td>
</tr>
<tr>
<td>M 3 days. Length 300 μm. 4 chaetigers. Loss of apical tuft. 2 red eyes. 4 days. 380 μm. 4 chaetigers. 2 red eyes. Some of them swimming in water column and others on the bottom, with detritus around them (starting the tube construction)</td>
<td>4 days. Length 390 μm. 4 chaetigers. Loss of apical tuft. 2 red eyes. 4–5 days. Larvae sink to the bottom and produce mucus where particles will aggregate</td>
<td>3 days. Larvae sink to the bottom. 4 days. Black jaws visible through body cavity. 3 chaetigers. Larvae start to agglutinate diverse detritus</td>
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<tr>
<td>E 5–6 days. 500 μm. 5 chaetigers, 2 red eyes. 5 large buds in prostomium. Rudimentary anal cirri. Black jaws visible through body cavity</td>
<td>6 days. Length 550 μm. 5 chaetigers. 5 large round antennal buds at the front of the prostomium. Black jaws visible through body cavity, rudimentary anal cirri</td>
<td>Not described</td>
</tr>
<tr>
<td>J 7 days. Length 540 μm. 5 chaetigers. 2 red eyes. 5 small antennae on the prostomium. 2 anal cirri. Juvenile present positive phototropism</td>
<td>7 days. Parapodia more developed. 5 antennae. 16 days. Length 1250 μm. 7 chaetigers. Parapodia and antennae more developed</td>
<td>1 month. 25 chaetigers. First branchia appears in 5th parapodia. 1 month and 20 days. Second branchia appears in 6th parapodia. 3 months. Length 15 mm</td>
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P, protrocophore; M, metatrocoephore; E, erpochaeta; J, juvenile.
Nurse cells were observed in oocytes with a diameter equal or less to 160 μm, as reported by Dagli et al. (2005) and reached up to 39 cells, 230 μm long, and 12 μm wide, which is larger than observed by Dagli et al. (2005). Nurse cells are common in the Onuphidae family, and they probably transport nutrients taken up from the coelomic fluid to the developing oocytes. Usually, larger oocytes had few or no nurse cells attached, probably because nutrients will not be absorbed by the mature oocytes (Blake 1975).

During the study period, some individuals were regenerating the anterior end of the body. The regenerative capacity of the anterior segments has already been observed for Diopatra species, including D. neapolitana (Beli 2006). The majority of individuals that were regenerating the anterior end had no gametes in the coelom, except for some females that contained small oocytes. It is thus possible that the individuals in regeneration concentrate all their energy on this process.

The first chaetiger with gametes ranged in females from the 35th to the 70th and in males from the 50th to 70th, usually after the chaetigers with branchiae. These results are very similar to those obtained by Dagli et al. (2005), who reported the appearance of oocytes in chaetiger 55 ± 0.9 (mean) and of sperm in chaetiger 51 ± 0.9 (mean). In agreement with those authors, our study confirmed the absence of a significant correlation between the first appearance of the oocytes and the width of the 10th chaetiger, chosen as a measure of the total length of the specimens.

In the present study, the smallest mature male and female were 139.2 and 149.7 mm long, respectively. These values are higher than those obtained by Dagli et al. (2005), who reported minimum length in females of 125 and in males of 110 mm. However, in their study, the largest entire worm had a length of 347 mm, whereas in our study we found larger individuals. During our collecting period we harvested 46 complete individuals with a total length between 24 mm and 725 mm. Only specimens of about 600 mm long are reported in the literature.

According to Paxton (1993), there are four reproduction patterns in Diopatra: Group I – species that breed in the parental tube, Group II – species with direct development in a cocoon, Group III – species that attach their eggs to the parental tube and present direct development, and Group IV – species with broadcast spawning with a free-swimming larval stage. Diopatra neapolitana belongs to group IV, as no eggs were observed inside the tubes or any gelatinous mass containing eggs attached to their distal end. This conclusion is supported by the artificial fertilization experiment, where only free-swimming lecithotrophic larvae were observed. This is also in agreement with the conclusions of Dagli et al. (2005). In our fertilization experiment, free-swimming prototrochophore larvae were obtained 19 h after fertilization, at 22 °C. In Conti & Massa (1998), this phase appeared 5 h after fertilization (25–32 °C) and Bhaud & Cazaux (1987) only observed the prototrochophore larva 24 h after fertilization. The larvae developed faster in Conti & Massa’s (1998) study compared with our study and Bhaud & Cazaux’s (1987) description. Morphologically, the results of our experiment were similar to Bhaud & Cazaux’s (1987) larval description, but the time of development was different, as in our study, larvae developed faster (about 1 day) until the metatrochophore phase.

These differences could be explained by the temperature, 22 °C in our case, and 25–32 °C in Conti & Massa (1998); temperature was never mentioned in Bhaud & Cazaux (1987). The number of larvae or the size of the containers could influence larval development, but we do not have information about these features. Larvae of D. neapolitana observed in our experiments and in Bhaud & Cazaux (1987) were morphologically different to those described by Choe (1960) in Japan as being D. neapolitana. This supports Paxton’s (1993) suggestion that the species mentioned by Choe (1960) was not D. neapolitana (Paxton 1993).

Larval development has also been studied in other Diopatra species, namely Diopatra cuprea (Allen 1959), which has a developmental pattern similar to D. neapolitana, and Diopatra marocensis (Fadlaoui et al. 1995), which breeds in the parental tube. The first larval stage observed in this study in D. neapolitana, the prototrochophore, was similar to that described for D. cuprea (Allen 1959) and D. marocensis (Fadlaoui et al. 1995). This stage is characterized in the three species by the presence of the apical tuft and ciliation around the body. Diopatra neapolitana and D. cuprea are active swimmers and had red eye spots during the initial development stages. Diopatra cuprea starts to settle to the bottom 3 days after fertilization, with four chaetigers, producing mucus to build the tube (Allen 1959). This was similar to what was observed in this study for D. neapolitana. Five antennae and anal cirri were observed at the 5th-chaetiger stage in D. cuprea and D. neapolitana, and at the 6th-chaetiger stage for D. marocensis. In D. marocensis, the ceratophores appear at the 12th-chaetiger stage and in D. cuprea at the 5th-chaetiger, with one to two rings, whereas in D. neapolitana the ceratophores still had no rings at the 50th-chaetiger stage (Conti & Massa 1998). According to these authors, the first branchiae appear at the 25th-chaetiger stage on the 5th chaetiger in D. neapolitana, and at the 18–20th chaetiger stage in D. marocensis, also on the 5th chaetiger (Fadlaoui et al. 1995).

In Ria de Aveiro, D. neapolitana is intensively exploited as live fish bait. No management or conservation regulations are currently set for the species and there is very...
little legislation. In the case of the Sado estuary, located about 350 km south of Ria de Aveiro, harvesting of *D. neapolitana*, *Marphysa sanguinea* and *Hediste diversicolor* is not allowed from 1 November until 30 April (Portuguese legislation: Portaria no 576/2006 2006). That period is reported in the legislation as coinciding with spawning and juvenile growth. However, this is not supported by the present or other studies. The main reproductive period for *H. diversicolor* in the Sado estuary was from April to August/September (Garcês, unpublished data). In the Southwestern coast of Portugal (Odeceixe, Aljezur and Carrapateira) the same species was reported as reproducing throughout the year, with important peaks in September and May (Fidalgo e Costa 2003). In Ria de Aveiro, the species also showed two important reproductive periods, in March and September (Abrantes *et al.* 1999). The reproductive period of *Marphysa sanguinea* was mainly from March/April to October/November in the Sado estuary (Garcês, unpublished data) and a peak spawning period in April–May was reported from the Venice Lagoon (Italy) by Prevedelli *et al.* (2007). The main reproduction peak for *D. neapolitana* in the present study and in that of Dagli *et al.* (2005), in Izmir Bay (Eastern Mediterranean), was from May to August.

In Portugal, with the exception of the resting period established for the Sado estuary, the exploitation of polychaetes occurs all year, being more intense in warm months. Cancela da Fonseca & Fidalgo e Costa (2008) observed that the capture of these species has increased in recent years and that the mean size of harvested individuals is smaller. Dagli *et al.* (2005) reported that *D. neapolitana* occurred in the past in high densities in Inciralti (Mediterranean Sea), and by the time they did their study, the species was only present in their study area. They also observed that each digger needed 10 h to collect about 2000 specimens, whereas 10 years before they collected the same number in only 2 h.

The digging activity has negative impacts on the entire ecosystem. The benthic community is affected as a whole, as are the species which depend on it for food (mainly birds and fishes). In addition, the biogeochemical cycles could be affected and the release of nutrients and bio-availability of metals enhanced (Cancela da Fonseca & Fidalgo e Costa 2008). All of this emphasizes the urgent need for a sustainable exploitation of these natural resources, not only in Ria de Aveiro but in all coastal areas. The use of scientifically supported legislation coupled with control in the allocation of bait-digging licenses with regular monitoring of the impacted areas should be implemented. Mitigation measures could be applied either by restricting the harvest (in the case of *D. neapolitana* in Ria de Aveiro, the most suitable period seems to be April until September) or by establishing yearly rotating resting areas. The rotation system has been suggested as an effective solution to minimize the negative impacts of this kind of resource exploitation by Fowler (1999) and Cancela da Fonseca & Fidalgo e Costa (2008) and is being used in Korea, which is one of the largest exporters of bait polychaetes in the world (Choi 1985).

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**References**


