Dynamics of the *Acartia* genus (Calanoida: Copepoda) in a temperate shallow estuary (the Mondego estuary) on the western coast of Portugal

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The purpose of this work was to review the dynamics of the *Acartia* species in the Mondego estuary (a temperate North-Atlantic shallow estuary in Western Portugal) in a genus integrated perspective. The *Acartia* genus is represented in the system by the species *Acartia clausi* and *Acartia tonsa*; the samples were taken between July 1999 and June 2000, with 63 and 125 µm mesh size nets, and between January 2003 and January 2004, with a 335 µm mesh size net, in the downstream and upstream areas of the estuary. Significant differences in abundance were found between months and sampling stations for the two species (ANOVA, P < 0.05). *A. clausi* dominated in the downstream estuary, registering peaks in June (156 ind. m⁻³) and September (73 ind. m⁻³); in the upstream estuary this species showed a maximum of density in September (35 ind. m⁻³). *A. tonsa* dominated in the upstream estuary with peaks of abundance occurring in December (2372 ind. m⁻³) and October (1056 ind. m⁻³) in the downstream estuary this species exhibited higher abundance in August (52 ind. m⁻³). The two species of the genus coexist in time exhibiting a strong spatial segregation behavior in the estuary.

Key words: estuaries, Mondego estuary, Copepoda, *Acartia*, ecology
INTRODUCTION

Copepoda (Crustacea) comprise the most abundant taxa of the marine and estuarine zooplankton (OMORI & IKEDA, 1984; MAUCHLINE, 1998). There they act as an efficient and direct path for energy transfer to higher trophic levels (WILLIAMS et al., 1994; MAUCHLINE, 1998) and contribute substantially to the downward flux of organic material (FEINBERG & DAM, 1998). Abundance and biomass estimates for Copepoda species, in an estuarine ecosystem, provides useful information on the energy flux, and such measurements are essential in ecological studies (VIEIRA et al., 2003b; PASTORINHO et al., 2003).

The copepoda community of the Mondego estuary is dominated by the Acartia genus (Copepoda: Calanoida) (AZEITEIRO et al., 1999, 2000; VIEIRA et al., 2003a). The Acartia genus is represented in the system by the species Acartia clausi (Giesbrecht, 1889) (AZEITEIRO et al., 1999, 2000; VIEIRA et al., 2003a, b) and Acartia tonsa (Dana, 1848) (AZEITEIRO et al., 1999, 2000; PASTORINHO et al., 2003; VIEIRA et al., 2003a). This selected genus is an abundant genus of temperate zooplankton communities (JEFFRIES, 1967; LEE & MCALICE, 1979; BRYLINSKI, 1981; SOBRAL, 1985; GAUDY et al., 2000).

Each one of the two species Acartia was already studied in what concerns morphometric relations, production and turnover-rates (A. clausi: VIEIRA et al., 2003b; A. tonsa: PASTORINHO et al., 2003) using image analysis methods (CHRISTOU & VERRIPOULOS, 1993; JEFFRIES et al., 1998). For zooplankton, the literature reports suitable coefficients, which, starting from linear measurements of body fractions (TACKX et al., 1995) reveal the weight or carbon contents of the specimens studied (SHMELEVA, 1965; PERTSOVA, 1966; KRYLOV, 1968; UYE, 1982).

PASTORINHO et al. (2003) also used histology, histochemistry and biometry in the species A. tonsa to determine whether fecundity is a limiting factor in itself or are zooplankter constrained to respond to an ever-changing environment. Analyses of maturation stage of oocytes in adult ovigerous females demonstrated the presence of all three considered oocytic development stages: immature, vitellogenic and mature, with emphasis in the latter since it indicates a permanent capability for reproduction. Taking in account the relevance of the species, this fact demonstrated the modulating influence of ecological parameters (namely the environmental parameters) in general zooplankton reproductive traits.

The purpose of this work was to describe and interpretate the dynamics of the Acartia genus in the Mondego estuary with the specific published data available and new collected data.

MATERIALS AND METHODS

Study Site

The Mondego estuary, located in the Portuguese west coast (North Atlantic Ocean) (40º08’N; 8º50’W), has an area of 3.3 km² and a volume of 0.0075 km³. The hydrological basin of the Mondego, with an area of 6670 km², provides an average discharge of 8.5x10⁹ m³s⁻¹ (Fig. 1).

Fig. 1. Location of zooplankton sampling stations in the Mondego estuary
The circulation in the South arm of the estuary depends on the tides and, in much smaller amount on the freshwater discharge from a tributary – the Pranto River, which is controlled by a sluice located 3 km from the confluence with the Mondego River. The sampling stations were located along the southern arm of the estuary (Fig. 1): station 1 near its mouth and station 2 further inland in the inner area of the southern arm (1.7 m deep in high tide).

Sampling program

Samples were taken monthly, from July 1999 to June 2000 (63 and 125 µm taxocenosis) (VIEIRA et al., 2003a) and from January 2003 to January 2004 (335 µm taxocenosis).

Determination of environmental parameters, phytoplankton and zooplankton

All samples were analyzed in situ for salinity, temperature, dissolved oxygen and pH. Samples were also analyzed in the laboratory (in triplicate) for their content in chlorophyll a concentration (BACELAR-NICOLAU et al., 2002, 2003; VIEIRA et al., 2002).

Sub-surface (20 cm depth) phytoplankton samples (horizontal hauls) were collected with a 25 µm mesh size net (VIEIRA et al., 2002).

Sub-surface (20-40 cm depth) zooplankton samples (horizontal hauls) were collected using 63 and 125 µm (VIEIRA et al., 2003A) and 335 µm mesh size nets.

Production study methodology

Each one of the two species was already studied in what concerns morphometric relations, production and turnover-rates using image analysis methods (A. clausi: VIEIRA et al., 2003b and A. tonsa: PASTORINHO et al., 2003).

Data analysis

ANOVA and Regression analysis were applied to find an explanatory model for the dynamics of the two species.

Table 1a. Environmental results (temperature, salinity, pH, oxygen dissolved -saturation% and chlorophyll a values) from monthly annual sampling cycle in the south arm of the Mondego estuary, in both sampling stations, between July 1999 and June 2000

<table>
<thead>
<tr>
<th>Month</th>
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<td>S2</td>
<td>S1</td>
<td>S2</td>
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<tr>
<td>18.9</td>
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<td>20.0</td>
<td>25.0</td>
<td>19.1</td>
<td>20.7</td>
<td>16.9</td>
</tr>
<tr>
<td>Sal (psu)</td>
<td>24.5</td>
<td>18.1</td>
<td>23.0</td>
<td>26.0</td>
<td>25.0</td>
<td>31.0</td>
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<tr>
<td>pH</td>
<td>8.2</td>
<td>7.9</td>
<td>8.1</td>
<td>7.6</td>
<td>8.3</td>
<td>8.3</td>
</tr>
<tr>
<td>DO2 %</td>
<td>80.0</td>
<td>72.0</td>
<td>94.0</td>
<td>85.0</td>
<td>86.0</td>
<td>48.0</td>
</tr>
<tr>
<td>Chl a(mg m⁻³)</td>
<td>0.605</td>
<td>1.445</td>
<td>0.415</td>
<td>2.275</td>
<td>0.250</td>
<td>2.270</td>
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<td>S1</td>
<td>S2</td>
<td>S1</td>
<td>S2</td>
<td>S1</td>
<td>S2</td>
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<tr>
<td>11.9</td>
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<td>14.2</td>
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<tr>
<td>Sal (psu)</td>
<td>23.2</td>
<td>23.2</td>
<td>29.7</td>
<td>29.4</td>
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<td>-</td>
</tr>
<tr>
<td>pH</td>
<td>7.8</td>
<td>7.9</td>
<td>7.9</td>
<td>7.6</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>DO2 %</td>
<td>89.5</td>
<td>91.0</td>
<td>99.4</td>
<td>72.4</td>
<td>-</td>
<td>-</td>
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<tr>
<td>Chl a(mg m⁻³)</td>
<td>0.190</td>
<td>0.810</td>
<td>0.340</td>
<td>2.040</td>
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</table>
RESULTS

Variation in environmental variables, phytoplankton and zooplankton

The environmental parameters, relatively to the 99/00 annual cycle (Table 1a), description and analysis were published by BACELAR-NICOLAU et al. (2002, 2003) and VIEIRA et al. (2002). Relatively to the 03/04 annual cycle the data is presented in Table 1b.

In the 99/00 annual cycle in sampling station 1 the average temperature was of 16.1 ºC, varying between 20.0 ºC in August 2000 and 11.8 ºC in December 1999. The salinity varied along the year with a minimum of 9.5 psu in April 2000, and a maximum of 31.0 psu in September 1999. pH suffered a little fluctuation during the studied period, varying between a maximum value of 8.4 in December 1999 and a minimum value of 7.5 in September 1999, with an average value of 7.9. Dissolved oxygen concentration presented a minimum value of 48.0 % in September and November 1999, a maximum value of 91.0 % in January 2000 and an annual average of 69.4 %. Chlorophyll a concentration presented higher average values than in station 1 (1.747 mg m\(^{-3}\)), with a maximum of 2.730 mg m\(^{-3}\) in May 2000, and a minimum of 0.810 mg m\(^{-3}\) in January 2000 (BACELAR-NICOLAU et al., 2002,
In the phytoplankton, the most abundant groups were Bacillariophyceae, Cyanoprokaryota, Dinophyta, Euglenophyta and Chlorophyta. The composition of phytoplankton differed between the two sampling stations. Bacillariophyceae dominated in sampling station 1 during all the year. In station 2, Bacillariophyceae dominated September, December, February and May. Cyanophyta dominated in August and June. Chlorophyta presented important percentages in January and March (VIEIRA et al., 2002).

In the 03/04 annual cycle temperature ranged from 8.7 to 23.9 °C, reaching minimum and maximum values during winter and summer, respectively, showing a typical seasonal pattern for temperate Atlantic estuaries. Significant differences among the two stations and months were observed (ANOVA, P < 0.05). The spring-summer months exhibited an overall warming trend, especially, in downstream sampling station. Salinity values during the sampling period varied between 0.6 psu observed in May and 33.4 psu observed in August. Dissolved oxygen levels ranged from 45.0 % in January 2003 and 125.0 % in February. A fluctuation in these values was observed during the study period. During this study chlorophyll a concentration ranged from 0.74 mg m⁻³ in January to 21.89 mg m⁻³ in June. The maximum chlorophyll a concentrations were observed at the stations S2. Peaks of concentration occurred in June and August.

In the 99/00 annual cycle the zooplankton community, in terms of species composition and distribution, was dominated by estuarine and estuarine/marine copepods, adults and copepodits (VIEIRA et al., 2003A). In the 63 µm taxocenosis A. tonsa occurred with low densities throughout the year, at station 1, registering a peak in April (1860 ind. m⁻³). At station 2, this species showed a peak in October (29070 ind. m⁻³) and important densities in August (3793 ind. m⁻³), November (2121 ind. m⁻³), December (2724 ind. m⁻³) and April (1966 ind. m⁻³). A. clausi presented low abundances all year round, registering a peak in April (560 ind. m⁻³), at station 1, and in November (1160 ind. m⁻³), at station 2 (VIEIRA et al., 2003A).

In the 03/04 annual cycle the copepods dominated the zooplankton community throughout the year. The species A. clausi and A. tonsa had contributed about 11% and 75%, respectively, of the total copepod abundance. Significant differences in abundance were found between months and sampling stations for the two species (ANOVA, P ≤ 0.05). A. clausi (Fig. 2a) dominated in the downstream station (S₁), registering peaks in June (156 ind. m⁻³) and September (73 ind. m⁻³); at station 2 this species showed a maximum of density in September (35 ind. m⁻³). A. tonsa (Fig. 2b) dominated in the upstream station (S₂) with peaks of abundance occurring in December (2372 ind. m⁻³) and October (1056 ind. m⁻³). At station 1 this species exhibited higher abundance in August (52 ind. m⁻³).

Weight-length relationships

The following biomass/length relationship was estimated for specimens of A.clausi and A. tonsa, respectively: Y=0.11e³.14x, with a r² of 0.59 (VIEIRA et al., 2003b) and Y=0.15e³.04x, with a r² of 0.62 (PASTORINHO et al., 2003).

Production and P/B ratio

Length-weight relationships were used to estimate production taking into account cohort growth and mortality. The annual production
was calculated for specimens of *A. clausi* and *A. tonsa*, respectively: 63.44 mgCm\(^{-3}\)yr\(^{-1}\) (VIEIRA et al., 2003b) and 43.12 mgCm\(^{-3}\)yr\(^{-1}\) (PASTORINHO et al., 2003). The production/biomass (P/B) ratio was estimated, respectively, at 25.50 and 10.56 (PASTORINHO et al., 2003; VIEIRA et al., 2003b).

**Multiple regression analysis**

Multiple regression analysis between environmental parameters and zooplankton total density showed that zooplankton increases with increasing salinity, temperature and chlorophyll \(a\) \((r=0.859; r^2=0.737)\) (VIEIRA et al., 2003a).

In the 125 µm taxocenosis for the two species *A. clausi* and *A. tonsa*, respectively, \(r=0.990; r^2=0.981\) (P<0.001) (VIEIRA et al., 2003b) and \(r=0.964, r^2=0.929\) (P<0.001) (PASTORINHO et al., 2003), the abundance increased, respectively, with increasing salinity and temperature (VIEIRA et al., 2003b) and dissolved oxygen and temperature (PASTORINHO et al., 2003). The regression analysis, performed with the 335 µm taxocenosis, revealed no significant effect of the environmental factors (salinity, temperature, dissolved oxygen and chlorophyll \(a\)).

*Fig. 2. Spatio-temporal distribution of salinity, temperature and chlorophyll \(a\) and Acartia species between January 2003 and January 2004*
DISCUSSION

The distribution patterns of the environmental parameters observed are mostly in agreement with previous studies (AZEITEIRO & MARQUES 2000; AZEITEIRO et al., 2002; BACELAR-NICOLAU et al., 2002, 2003; VIEIRA et al., 2002) defining a clear spatial unidirectional salinity gradient and secondarily a temperature-chlorophyll a temporal gradient (AZEITEIRO & MARQUES 2000; AZEITEIRO et al., 2002; BACELAR-NICOLAU et al., 2002, 2003; VIEIRA et al., 2002). The estuarine and marine phytoplanktonic communities are frequently dominated by dinoflagelate and diatoms species. Both stations presented diatoms and dinoflagelates as the most abundant phytoplankton species. There were also found Chlorophyta, Euglenophyta and Cyanoprokaryota (VIEIRA et al., 2002). This flora conditioned the primary consumers, not only for the form of the cells and the associations that they establish with each other, but also because of the nutritional quality of the different species. Several studies have shown the importance of food quality on fertility within the Acartia genus (DAM et al., 1994; PAGANO & SAINT-JEAN, 1994b; JONASDOTTIR, 1994; JONASDOTTIR & KIORBOE, 1996). The development of Microcystis aeruginosa blooms induces mortality within A. clausi assemblages (PAGANO & SAINT-JEAN, 1994a) and some diatoms species inhibits the fertility in A. clausi individuals (JONASDOTTIR, 1994; IANORA et al., 1996). However the nutritional quality requirements, ecological population interactions and other ecological interactions were not studied.

The production values estimated to the two species are in accordance with values reported by other authors to the Acartia genus and other copepods (Table 2).

The P/B values obtained had given expected modal turnover rates (VALIELA, 1995). Those P/B values mean that although the biomass of small sized species (e.g. copepods) may be small, the higher specific production makes them important producers. A. clausi is an r-strategy-type species, characterized by a high productivity (HIRCHE, 1992). The two species differences in production values and turnover rates are due to

<table>
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<tr>
<th>Species</th>
<th>Reference</th>
<th>Study site</th>
<th>Methods</th>
<th>P (annual)</th>
<th>P/B</th>
</tr>
</thead>
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<tr>
<td>Acartia spp.</td>
<td>(Hirst et al., 1999)</td>
<td>Coastal station, Solent UK</td>
<td>weight</td>
<td>17,62 mg C m⁻³ yr⁻¹</td>
<td></td>
</tr>
<tr>
<td>Acartia bifilosa</td>
<td>(Irigoien and Castel, 1995)</td>
<td>The Gironde (SW France)</td>
<td>size</td>
<td></td>
<td>28</td>
</tr>
<tr>
<td>Acartia spp.</td>
<td>(Escaravage and Soetaert, 1995)</td>
<td>Westerschelde estuary, Nether</td>
<td>growth rate methods</td>
<td>5 g C m⁻² y⁻¹</td>
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<tr>
<td>Copepoda</td>
<td>(Hirst et al., 1999)</td>
<td>Coastal station, Solent UK</td>
<td>weight</td>
<td>32,2 mg C m⁻³ y⁻¹</td>
<td></td>
</tr>
<tr>
<td>Eurytemora spp.</td>
<td>(Escaravage and Soetaert, 1995)</td>
<td>Westerschelde estuary, Nether</td>
<td>growth rate methods</td>
<td>6 g C m⁻² y⁻¹</td>
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<tr>
<td>Paracalanus parvus and Pseudocalanus elongatus</td>
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<td>1,67 mg C m⁻³ y⁻¹</td>
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<td>Temora longicornis</td>
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<td>weight</td>
<td>4,77 mg C m⁻³ y⁻¹</td>
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<tr>
<td>Acartia tonsa</td>
<td>(Pastorinho et al., 2003)</td>
<td>Mondego estuary, Portugal</td>
<td>cohorts</td>
<td>43,12 mg C m⁻³ y⁻¹</td>
<td>11</td>
</tr>
<tr>
<td>Acartia clausi</td>
<td>(Vieira et al., 2003b)</td>
<td>Mondego estuary, Portugal</td>
<td>cohorts</td>
<td>63,44 mg C m⁻³ y⁻¹</td>
<td>26</td>
</tr>
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</table>
the ecological conditions that the two species experienced: *A. clausi* more productive in a less stable environment (a tide variable environment) and predator pressure (by medusa, siphonophores, chaetognaths, large copepods and fishes) and *A. tonsa* less productive in a more stable environment with a direct food competitor the species *Calanipeda aquae-dulcis* (YELIZARENKO, 1992) that co-exists with *A. tonsa* (AZEITEIRO et al., 1999).

These production results indicate that the *Acartia* genus may play a significant role in transferring energy to higher trophic levels in the estuary. Although production by nauplii is not included in the present study, this does not typically exceed 25% of copepod total production (MULLIN, 1988; LIANG & UYE, 1996; LIANG et al., 1996). Most copepods reproduce throughout all the year. The cohorts represent the maximum of generations possible and the duration of the cohorts represents the longevity of the generations (BINET, 1977). Copepoda life cycle in temperate regions have an average of 25 to 45 days. In temperate regions the adult longevity exceed rarely two months (GAUDY, 1972) with variable annual growth generations (Table 3) function of the latitude of the systems analyzed. *Acartia* genus biological cycle depends on the local of study. Annual generations depend on latitude and trophic availability. Both temperature and food availability are known to play a significant role in the copepod production activity (LEE & MCALICE, 1979; KLEIN & GONZALEZ, 1988). Temperature together with food should explain the *Acartia* genus biological cycles features in the Mondego estuary.

Histological observation of the positioning and distribution of cells and different cellular structures, qualitative evaluation of chemical contents, through histochemistry, pointing existence of differences between development stages in terms of the accumulation of reserve substances (nominately glycoproteins and, by indirect extrapolation, lipids), measurement of cellular size and determination of the C/N ratio, were applied to *A. tonsa* by PASTORINHO et al. (2003) in order to obtain a clear and convincing answer the environmental determination in the *Acartia* genus biological cycles. These methods and techniques allowed the identification of three very distinct stages of maturation of oocytes and hence the elaboration of a functional scales, numerically based and statistically validated, allowing comparisons between samples. It was verified that all three maturation stages were always present. The presence in the gonadic masses of the females of stage III oocytes (mature) mean that these females are fully capable of reproduction and only fecundation has to occur, since these cells possess all that is necessary for the juvenile to be viable (PASTORINHO et al., 2003). The simultaneous existence of fully functional reproductive cells in registered high density epochs and in minimum density periods of the reproductive cycle of the organisms, indicate that the main modulating influence over these cycles comes out of ecological abiotic parameters which notoriously

<table>
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<td>Ocidental Atlantic</td>
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<td>(Vučetić, 1957)</td>
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<tr>
<td>4</td>
<td>Roscoff</td>
<td>(Razouls, 1965)</td>
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<td>4</td>
<td>Long Island Sound</td>
<td>(Conover, 1956)</td>
</tr>
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<td>5</td>
<td>Plymouth</td>
<td>(Digby, 1950)</td>
</tr>
<tr>
<td>6,7</td>
<td>Sebastopol</td>
<td>(Greze &amp; Baldina, 1972)</td>
</tr>
<tr>
<td>8</td>
<td>Black Sea</td>
<td>(Porumb, 1968)</td>
</tr>
<tr>
<td>9</td>
<td>Karadag</td>
<td>(Tchaianova, 1950)</td>
</tr>
<tr>
<td>Continuous</td>
<td>Mediterranean</td>
<td>(Bernard, 1958)</td>
</tr>
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</table>
superimpose over the continuous reproductive capabilities of zooplanktonic organisms, namely the genus *Acartia* as referred by HEINLE (1966), creating the well described fluctuating patterns (KLEPPEL, 1992). As a corollary one could conclude that seasonal variability in zooplankton densities reflects advantageous or disadvantageous assemblages of external factors acting over a latent reproductive potential.

Temperature is an important factor controlling seasonal distribution of copepods (HALSBAND-LENK *et al*., 2002), namely the *Acartia* species (GONZALES, 1974; SULLIVAN & MCMANUS, 1986). The thermal sensitivity of both species (CERVETTO, 1985), in the studied system, could explain their continuous distribution year around, at temperatures ranging between 11 and 25 ºC. Seasonal fluctuations of both species are described by several authors (DEEVEY, 1948, 1960; CONOVER, 1956; HEINLE, 1966; JEFFRIES, 1962, 1967; SAGE & HERMAN, 1972; LEE & MICALICE, 1979; SOBRAL, 1985; SULLIVAN & MCMANUS, 1986; GAUDY *et al*., 2000) where *A. clausi* and *A. tonsa* co-exist and reach great abundances. All these investigations have indicated a well defined relationship between seasonal changes and the temperature.

*A. tonsa* is a brackish water euryhaline and eurythermic species; in low concentrations of dissolved oxygen retard the development of eggs and results in death of all copepod stages of *A. tonsa* (Oxyphilic species) (SAZHINA, 1987). In the Mondego estuary adults occur in the inner and upstream estuary in a shallow, low salinity water stable area; *A. tonsa* performs both horizontal and daily vertical migrations that ensures the residence in the area; young individuals undertake migrations only within surface layers (BRODSKY, 1950) what explains the 63 and 125 µm taxocenosis distribution. The eurythermic behavior explains why *A. tonsa* copepods were found in the plankton year round. *A. clausi* occurs in the mouth of the estuary in a high salinity/tidal area. The correlations given by the regression analysis reflected the species ecological preferences clarifying the distribution found in previous descriptive zooplankton studies in the estuary (AZEITEIRO *et al*., 1999, 2000), and also the genus biology (KLEIN & GONZALES, 1988) and ecology (ALCARAZ, 1983; SOBRAL, 1985), namely in the Mondego estuary (AZEITEIRO *et al*., 2000). In Atlantic temperate systems the species of the genus (namely *A. tonsa* and *A. clausi*) sometimes co-exist at the same time (LEE & MICALICE, 1979); in the Mondego estuary they co-exist (AZEITEIRO *et al*., 1999, 2000; VIEIRA *et al*., 2003a) but exhibiting a strong spatial segregation. This two species have different salinity (and temperature) preferences (LANCE, 1963; SAGE & HERMAN, 1972): the two species do not co-exist in the space due to the strong spatial unidirectional salinity gradient (AZEITEIRO *et al*., 2000). The *A. tonsa* failure to develop in more saline waters (coastal marine water) is due to the sensitivity to the salinity factor (CONOVER, 1956; TESTER & TURNER, 1991; CERVETTO *et al*., 1999; GAUDY *et al*., 2000) and, according to PAFFENHOFER & STEARNS (1988) to the low concentrations of appropriate food in sea water comparatively to estuaries. The absence of *A. clausi* population in the estuary middle zone and upstream could result from the predation by the dense resident *A. tonsa* population (CONOVER, 1956; ANRAKU & OMORI, 1963; ANRAKU, 1964; LONSDALE *et al*., 1979; GIFFORD & DAGG, 1988; WHITE & ROMAN, 1992) or, according to LUTZ *et al*. (1992), a failure of hatching of their eggs due to environmental disadvantageous conditions (anoxic conditions in the bottom layer); the negative impact of anoxia on the viability of subitaneous eggs of calanoid copepods as well as the impact of the benthic-pelagic coupling on their life cycle has already been evidenced in shallow coastal aquatic systems (MARCUS & LUTZ, 1994; MARCUS & BOERO, 1998).
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Dinamika roda Acartia (Calanoida: Copepoda) u temperiranom plitkom estariju (ušće rijeke Mondego) na zapadnoj obali Portugala

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SAŽETAK

Cilj ovog rada je dati pregled dinamike vrsta Acartia u estuariju Mondego (temperirani Sjeveroatlantski plitki estuarij u zapadnom Portugalu) za sve nađene vrste toga roda. Rod Acartia je prisutan s tri vrste: Acartia clausi, Acartia bifilosa var. inermis i Acartia tonsa; uzorci uzeti od srpnja 1999 do lipnja 2000 su dobiveni pomoću mreže veličine oka 63 i 125 µm, a od siječnja 2003 do siječnja 2004 pomoću mrežice veličine oka 335 µm, oboje u nizvodnom i uzvodnom dijelu estuarija. Nađene su znatne razlike u abundanciji dviju vrsta (ANOVA, P < 0.05). Acartia clausi je bila dominantna u nizvodnom estuariju s maksimumom u lipnju (156 ind. m⁻³) i rujnu (73 ind. m⁻³); u uzvodnom estuariju ova vrsta je pokazala maksimum gustoće u rujnu (35 ind. m⁻³). Acartia tonsa je bila dominantna u uzvodnom estuariju s maksimumom gustoće u prosincu (2372 ind. m⁻³) i listopadu (1056 ind. m⁻³). U nizvodnom estuariju ova je vrsta pokazala veću gustoću u rujnu (52 ind. m⁻³). Dvije vrste se u estuariju javljaju istovremeno, ali su prostorno izrazito odijeljene.

Ključne riječi: estuarij, kopepodi, Acartia, ekologija