REMAINS OF THE PROTOZOAN STICHOLONCHE ZANCLEA IN THE FAECAL PELLETS OF Paracalanus quasimodo, Parvocalanus crassirostris, Temora stylifera AND Temora turbinata (COPEPODA, CALANOIDA) IN BRAZILIAN COASTAL WATERS

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Descriptors: Copepods, Faecal pellets, Feeding, Protozoa, Brazil.
Descritores: Copépodos, Pelotas fecais, Alimentação, Protozoário, Brasil.

Studies of copepod feeding have identified these organisms as key species in marine ecosystems, because they have a pivotal position in food webs: they control primary production through grazing, and are also a crucial link between the microbial food web and higher trophic levels, especially because they are able to predate on protozoa (STOECKER; CAPUZZO, 1990). The ingestion of protozoans has attracted particular attention, not only because it extends our understanding of the manifold energy pathways that structure aquatic food webs, but also because it involves an important behavioral aspect, i.e., what strategies are used to locate and capture preferred prey (ÁZEMAR et al., 2007). Copepods of the genera Paracalanus and Temora are believed to devote most of their time to filter feeding (VAN DUREN; VIDELER, 1995). The feeding habits of the copepods Paracalanus quasimodo, Temora stylifera and T. turbinata have shown that these species are omnivorous, but primarily opportunistic herbivores and that carnivory is infrequent (TURNER, 1984a, 1984b).

Paracalanus quasimodo, Parvocalanus crassirostris, Temora stylifera and T. turbinata are abundant in Brazilian coastal waters, and are assumed to play a fundamental role in the control of primary production. Scanning electron microscopy analysis of the faecal pellets of these copepods was performed to assess the main food item ingested during the occurrence of different water masses in the São Sebastião Channel (SSC), a coastal ecosystem located on the northeast coast of São Paulo State. Sampling was carried out every three months in the SSC, from January 1996 to July 1997 and also in July 1998 and January 1999, at two sampling points (map available in Eskinazi-Sant’Anna; BJÖRNBÄR, 2006). Surface (0.5 m) water samples (250 ml) for phytoplankton and protozooplankton analyses were taken with a Van Dorn bottle (5 L) and preserved in Lugol’s solution. Phytoplankton was enumerated using the inverted microscope method and carbon content determined using the volume:carbon conversions.

Mesozooplankton was collected during 2-min surface tows with a 150-mm mesh net. Adult females of the copepods Parvocalanus crassirostris, Paracalanus quasimodo, Temora stylifera and Temora turbinata were sorted under a WILD M8 stereomicroscope, isolated within 20 min of collection, and transferred to Petri dishes containing collected surface sea water, and then left for 30 to 120 min to produce faecal pellets. The expelled faecal pellets (38 from P. crassirostris, 78 from P. quasimodo, 37 from T. stylifera and 21 from T. turbinata) were individually removed by pipette and placed in petri dishes containing a mixture of filtered sea water (Nuclepore cellulose acetate membrane filters 20 mm) and 10 mm screened surface sea water. The pellets were then left for 24 h at 24°C (± 2°C) for microbial stripping of their peritrophic membranes. All pellets were subsequently preserved in 4% formalin-seawater-glutaraldehyde solution. Ashore, preserved pellets were individually removed by pipette, transferred to Nuclepore filters, and then left to adhere. Filters with attached pellets were washed in distilled water for salt elimination, dehydrated in a graded ethanol series (50, 60, 70, 80, 90, 95, and 100%), left to dry at room temperature, and then coated with gold. The faecal pellets were examined with a Zeiss DSM-950 Scanning Electron Microscope (SEM).

Analysis of the faecal pellets of Paracalanus quasimodo, Parvocalanus crassirostris, Temora stylifera and Temora turbinata indicated the presence of the protozoan S. zanclea fragments and remains of diatoms, mainly Chaetoceros sp., Thalassiosira frauenfeldii, Skeletonema costatum and Thalassionema nitzschioides. Identification of S. zanclea fragments was only possible under SEM, which revealed their characteristic ornamentation,
consisting of a series of fringes along the spine (Fig. 2A and B). Fragments of *S. zanclea* were abundant in the faecal pellets of *P. quasimodo*. In some pellets, these fragments were the only food item found, which reinforces the indications of selective capture of this protozoan (Fig. 1). Besides *S. zanclea* remains, fragments of diatoms (*Chaetoceros* sp., *Skeletonema costatum*, *Thalassiosira nitzschioides* and *Thalassiothrix frauenfeldii*) were also observed (Fig. 2E). Analysis of *P. crassirostris* faecal pellets showed, besides phytoplankton items, the presence of remains of *S. zanclea* (Fig. 2C), suggestive of selective feeding, because of the large size of the protozoan (> 200 µm) and the small size of *P. crassirostris* individuals (< 0.6 mm). Unidentified centric and pennate nanoplanktonic cells (< 20 mm) were observed in the faecal pellets of all the species, especially *P. crassirostris*, illustrating the importance of these copepods to the energy pathway in the nano-trophic food web compartment (Fig. 2D).

Faecal pellets of *Temora stylifera* contained a wide variety of food remains, including diatoms, dinoflagellates, silicoflagellates, radiolarians and amorphous material, in addition to *S. zanclea* remains (Fig. 2F and G). Remains of *S. zanclea* were also common in the faecal pellets of *T. turbinata* (Fig. 2G and H).

All the copepods examined fed in situ upon *S. zanclea*. In their faecal pellets, the proportion of *S. zanclea* spines was in some cases higher than the proportion of diatom frustules. Eskinazi-Sant’anna (2006) also registered the importance of *S. zanclea* as food item for other mesozooplanktonic copepods (*Centropages velificatus* and *Paracalanus parvus*) and the macrozooplanktonic Euphausiacea *Euphausia* sp., reinforcing the idea of *S. zanclea* being a central trophic link in the food web of SSC.
There is little published information on predation on protozoans under natural conditions. Evidence of the ingestion of protozoans comes, mostly, from predation experiments done in situ or in the laboratory (SANCHEZ, 2011), through the use of biomarkers (LI et al., 1996), filming (VERTY; PAFFENHOFER, 1996), or through identification of hard parts such as loricae and skeletal material in gut contents or in faecal pellets (CONOVER, 1982). In tropical coastal waters, the trophic model linking copepods and primary production is the classical representation of the regional marine food web. In fact, tropical copepods are considered “small-sized” species (in general, < 2 mm) and the occurrence of predation on large-sized components of the protozoan communities is not considered a regular trophic pathway, since the algal resource is abundant year-round and the size spectrum of the phytoplankton is diverse and appropriate for filter-feeding copepod species (~20 and 100 µm). Because phytoplankton is available in the SSC (phytoplankton density and biomass and protozooplankton abundance data are available in Eskinazi-Sant’Anna; Björnberg, 2006), questions that arise are: why is S. zanclea predated on by copepods, including small copepods such as Parvocalanus crassirostris and at high frequency? Also, what would be the competitive advantage for the active capture of a species that is relatively large (>200 mm) and rare in the environment? Copepods can switch their feeding modes depending on algal resources available, requirements for micronutrients and competition pressure. The search for additional food sources can also occur when phytoplankton biomass falls to very low levels, or when drastic changes in phytoplankton composition occur (COWLES et al., 1988).

The SSC is dominated by Coastal Water (CW: T > 20°C; S > 36 ‰), but during spring and summer a distinct flow of nutrient-rich, high-salinity, low-temperature water (South Atlantic Central Water, SACW: T < 20°C; S > 35 ‰) can be detected in the deepest layers. The CW is superficial, with oligotrophic characteristics, marked by the supremacy of flagellates (GIANESELLA-GALVÃO et al., 1999) and enriched seasonally by intrusion of the SACW. Diatoms need high nutrient concentrations, and for this reason are scarce in oligotrophic waters. Because diatoms are the main food item ingested by calanoid copepods (KÖSTER et al., 2011), the quantitative dominance of flagellates in the SSC waters, may be the determining factor for incorporation of heterotrophic food sources. Utilization of varied food sources may also be associated with specific micronutrient requirements, which would help maintain the nutritional status of individuals. In addition, a large-sized species may be selected because it is easily detectable and offers an important energy source. Protozoans are richer in proteins and lipids than diatoms, which may be why they are targets of such intense predation.

Parvocalanus crassirostris, Paracalanus quasi modo, Temora stylifera and Temora turbinata are perennial species in the SSC. The apparent ability to switch between two feeding modes might be selectively advantageous in environments with great variations in prey size and abundance, which is typical of coastal waters. In addition, the results underline the role of protozooplankton to structure food webs in oligotrophic tropical coastal waters. The next step to comprehend this trophic link between calanoid copepods and the protozoan S. zanclea is to carry out experimental situations considering different types of food particles (diatoms) versus S. zanclea abundance. These experiments could be fundamental to understand why this protozoan is the target of such intense predation in Brazilian coastal waters.

ACKNOWLEDGEMENTS

The author is grateful to the Center of Marine Biology (CEBIMar) of the University of São Paulo, and the Center of Electronic Microscopy (CEMEL) of the Federal University of Minas Gerais for the use of their facilities. I would also thank Dr. Tagea Kristina Simon Björnberg for her advice and Dr. Janet Reid for the revision of the English text. Financial support was provided by FAPESP (Fundação de Amparo à Pesquisa do Estado de São Paulo).

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(Manuscript received 06 August 2012; revised 16 February 2013; accepted 05 March 2013)