Comparing taxonomic and morphological biodiversity of tintinnids (planktonic ciliates) of New Caledonia

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Abstract

Tintinnid ciliates are planktonic grazers of nanoplankton. They have a lorica (or shell) into which the ciliate cell can withdraw. The lorica provides information on both the identity and the ecology of the organism because characteristics of the lorica distinguish species and the diameter of the oral opening is related to the size of prey ingested. We examined the relationship between biodiversity estimates on the basis of classifying specimens as belonging to a species or a simple morphological group defined by lorica oral diameter (LOD) in a presumably species-rich area, a tropical lagoon. Two sites were sampled in the lagoon off Nouméa over an annual cycle. The tintinnid fauna was species-rich (76 species) and represented a relatively even distribution of LOD sizes compared to other tropical and temperate sites. Median LOD varied with the fraction of the chlorophyll concentration by >10 μm. Total chlorophyll concentration was related to tintinnid concentration and, in turn, weakly correlated with numbers of species and LOD size-classes. Numbers of species were closely related to numbers of LOD size-classes as were H’ of species and H’ (Shannon index) of LOD size-classes. Thus, metrics of a morphological characteristic, related to the ecology of the organisms, can be used to estimate species diversity.

With the exception of prokaryotes, organisms are distinguished from each other largely on the basis of morphology. Thus, biodiversity, generally taken to mean taxonomic diversity, should be related to morphological diversity in some manner. However, the relationship of taxonomic diversity (species, genera, family, etc.) and morphological diversity has received little attention with regard to planktonic organisms. Most studies have concerned benthic organisms and considered very large-scale, either temporal or spatial patterns. Among fossil taxa, morphological diversity, presumed to represent ecological diversity, appears to be poorly related to taxonomic diversity in terms of the appearance of high-level taxa (e.g., Foote 1997). Among extant mollusks, morphological diversity also appears only weakly related to taxonomic diversity across large geographic scales (e.g., McClain et al. 2004). Tintinnid taxonomic diversity has been related to morphological diversity and phytoplankton size-diversity across medium spatial scales (Dolan et al. 2002). However, to our knowledge, the relationship between morphological and taxonomic diversity has not been examined on medium temporal scales (weeks to months).

The general question we pose is: For a group of organisms (regardless of its definition as a taxonomic unit, a guild, a trophic level, etc.) is there a characteristic that can be directly related to taxonomic diversity in a given locality? If so, the identity of the characteristic can suggest the ecological mechanism governing taxonomic composition. For example, taxonomic diversity of a phytoplankton community may reflect physiological diversity in terms of nutrient demands or, alternatively, in terms of light-harvesting capabilities. Distinguishing between the two possibilities could allow the identification of either nutrient dynamics or the light field as a factor regulating taxonomic diversity of the phytoplankton.

In this report we summarize our investigation of morphological and taxonomic diversity in a group of planktonic herbivores, tintinnid ciliates. They are part of the microzooplankton, grazers on primarily autotrophic and heterotrophic nanoplankton. Occasionally, the feeding activity of tintinnids can dominate the consumption of phytoplankton (e.g., Karayanni et al. 2005), but they are generally a minority component of the microzooplankton compared to oligotrich ciliates or heterotrophic dinoflagellates. Nonetheless, tintinnids are much more abundant than foraminifera or radiolarians. Compared to foraminifera or radariolaria, their biogeography has received much less attention. Few generalities are known...
Biodiversity in tintinnids

Fig. 1. Examples of tintinnid morphology. Note that although all loricas are basically tube- or vase-shaped, different species show distinct lorica architecture. Examples are species common in samples from New Caledonia: (A) Epiploocyclus brandti, (B) Codonellopsis morchella, (C) Salpingella faurei, (D) Eutintinnus lusus-undae, (E) Steenstrupiella steenstrupii, (F) Tintinnopsis beroidea, (G) Protorhabdonella curta.

Besides the fact that tintinnids supply a textbook example of the latitudinal species gradient (e.g., see fig. 3.14a in Gaston and Spicer 2003) and that high-latitude communities are distinct (e.g., Wasik 1998), tintinnids are ubiquitous in marine systems and a very specious group. The standard taxonomic monographs of Koifoid and Campbell (1929, 1939) catalogued more than 700 species. Identifications were made on the basis of characteristics of the lorica or shell into which the ciliate cell can withdraw. The lorica, formed of one or more layers, is composed primarily of proteinaceous material. Although the general shape of a lorica is a tube or vase, tintinnids can show very distinct lorica morphologies (Fig. 1). Like foraminifera and radiolarians, characteristics of gross morphology define species despite the fact that some genera are known to display plasticity in lorica morphology (e.g., Williams et al. 1994).

Tintinnids, in contrast to other groups of planktonic ciliates; for example, those known as oligotrichs, are a single order even among competing ciliate classification schemes. According to traditional ciliate taxonomy, and in agreement with the molecular data available to date (Agatha et al. 2005), they are a monophyletic group. Thus tintinnids are a coherent group ecologically as microzooplankters, morphologically as loricate ciliates, and phylogenetically as members of the order Tintinnida.

Not surprisingly, in tintinnids the diameter of the mouth end of the lorica, the lorica oral diameter (LOD), is related to the size of the food items ingested. For example, Heinboke (1978) found that the largest prey ingested were about half the LOD in longest dimension for a variety of tintinnid species. Studies of seasonal trends have documented shifts in overall community averages of LOD corresponding with changes in phytoplankton composition (e.g., Verity 1987). A more precise relationship was established by Dolan et al. (2002) who found that tintinnids removed prey at maximum rates that were 25% of LOD in size, based on an analysis of data from 11 species. Because lorica characteristics are related to both ecology and taxonomy, we hypothesized that morphological diversity should correspond with taxonomic diversity in this group over seasonal time scales.

Here we report on the relationship between morphological diversity, in terms of LOD diversity, and species diversity of tintinnids in a tropical lagoon system. We found that taxonomic diversity and diversity of LOD were tightly related, suggesting that temporal changes in prey-size diversity regulated temporal changes in tintinnid species diversity. We found then that a morphology-based biodiversity measure, based on a character of ecological importance, can provide evidence of the ecological mechanism regulating biodiversity.

Materials and methods

Study site and sampling—Sampling was conducted in the southwest lagoon of New Caledonia. The coral reef lagoon has an average depth of about 20 m and the water column is well-mixed. Southeast trade winds dominate hydrodynamics, giving average water residence times of about 10 d; chlorophyll a (Chl a) concentrations average about 0.5 μg L⁻¹ (Bujan et al. 2000). Samples were obtained from two stations in the lagoon, N12 (14-m depth) near the city of Nouméa and M33 (24-m depth) near the middle of the lagoon (Fig. 2) at 7- to 28-d intervals from April 2002 to March 2003. Niskin bottles were used to sample five depths in the water column at each station (3, 7.2, 11.5, 15.8, and 20 m for M33; 3, 4.7, 6.5, 8.2, and 10 m for N12). Samples were kept in Niskin bottles until pooled to produce a water column composite sample at the laboratory within 1.5 h after sample collection. The water column composite sample was subsampled for chlorophyll analysis and tintinnid ciliate determinations.

Sample processing—A 2-liter sample of the composite water column sample was fixed with Lugol’s (2% final concentration), sedimented in a graduated cylinder, and the concentrated sample was examined in a sedimentation chamber using an inverted microscope at ×200. Tintinnid species identifications were made at ×400 and made largely on the
basis of descriptions by Kofoid and Campbell (1929, 1939),
Campbell (1942), and Marshall (1969). Other works consulted
for species descriptions included Campbell (1942),
Hada (1938), Jorgensen (1924), and Rampi (1950).

A ‘lumping’ approach to identification was employed: any
intermediate forms or slight variants were pooled with the
morphologically closest and most numerous species. For ex-
ample, as intermediate forms were observed, all varieties of
Dadayiella ganymedes (D. trachelium, D. bulbosa, and D.
acuta) were pooled; a single type was established for forms
ranging from Tintinnopsis parva to T. rapa, hence the des-
ignation in Table 1 of T. rapa-parva. Forms intermediate
between Climacocylis scalaria and C. scalaroides were ob-
served so, and C. scalaroides forms were designated C. sca-
laria. All empty lorica were ignored.

Taxonomic diversity was estimated for each sample as the
Shannon index (In-based) and species richness. Morpholog-
ical diversity was estimated by placing species into size-
classes of LOD. Each species was assigned the average di-

Dolan et al.

Table 1. Tintinnid species encountered in samples from New
Caledonia.*

<table>
<thead>
<tr>
<th>Acanthostomella c</th>
<th>Epiplocylis c</th>
<th>Rhabdonella w</th>
</tr>
</thead>
<tbody>
<tr>
<td>A. conicoides</td>
<td>E. acuta</td>
<td>R. amour</td>
</tr>
<tr>
<td>A. obtusa</td>
<td>E. acuminata</td>
<td>R. elegans</td>
</tr>
<tr>
<td>Amphorella c</td>
<td>E. brandti</td>
<td>R. polyclum</td>
</tr>
<tr>
<td>A. quadrilineata</td>
<td>E. undella</td>
<td>Salpingella c</td>
</tr>
<tr>
<td>Amphorelopsis c</td>
<td>Eutintinnus c</td>
<td>S. acuminata</td>
</tr>
<tr>
<td>A. tetragona</td>
<td>E. apertus</td>
<td>S. curta</td>
</tr>
<tr>
<td>Canthariella w</td>
<td>E. elegans</td>
<td>S. decurtata</td>
</tr>
<tr>
<td>C. pyrimidata</td>
<td>E. elongatus</td>
<td>S. faurei</td>
</tr>
<tr>
<td>Climacocylis w</td>
<td>E. fraknoi</td>
<td>Stenestrupiella c</td>
</tr>
<tr>
<td>C. scalaria</td>
<td>E. tubulosus</td>
<td>S. steenstrupi</td>
</tr>
<tr>
<td>Codonellopsis c</td>
<td>E. lusus undae</td>
<td>S. gracilis</td>
</tr>
<tr>
<td>C. inornata</td>
<td>E. macilentus</td>
<td>S. monacense</td>
</tr>
<tr>
<td>C. morchella</td>
<td>F. sp A</td>
<td>S. nivalis</td>
</tr>
<tr>
<td>C. orthocerca</td>
<td>F. azorica</td>
<td>T. sp A</td>
</tr>
<tr>
<td>Coxialla c</td>
<td>F. composita</td>
<td>Tintinnidium n</td>
</tr>
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<td>C. ampla</td>
<td>F. ehrenbergii</td>
<td>T. acuminata</td>
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<td>C. calyptra</td>
<td>Helicostomella n</td>
<td>T. beroidae</td>
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<td>C. lacinosa</td>
<td>H. fusiformis</td>
<td>T. brandti</td>
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<td>C. pelagica</td>
<td>H. edentata</td>
<td>T. campanula</td>
</tr>
<tr>
<td>C. pseudoannulata</td>
<td>Metacytis n</td>
<td>T. compressa</td>
</tr>
<tr>
<td>Craterella c</td>
<td>M. mediterranea</td>
<td>T. everta</td>
</tr>
<tr>
<td>C. tortulata</td>
<td>M. mereschkovskii</td>
<td>T. platensis</td>
</tr>
<tr>
<td>Dadayiella w</td>
<td>M. jorgensenii</td>
<td>T. radix</td>
</tr>
<tr>
<td>D. ganymedes</td>
<td>M. sp A</td>
<td>T. “rapa-parva”</td>
</tr>
<tr>
<td>Dictycysta c</td>
<td>M sp B</td>
<td>T. tocanthensis</td>
</tr>
<tr>
<td>D. duplex</td>
<td>Odontophorella w</td>
<td>T. turbo</td>
</tr>
<tr>
<td>D. lepida</td>
<td>O. sp. A</td>
<td>T. urnula</td>
</tr>
<tr>
<td>D. mira</td>
<td>Protorhabdonella c</td>
<td>Undella c</td>
</tr>
<tr>
<td>D. occidentalis</td>
<td>P. curta</td>
<td>U. clevei</td>
</tr>
<tr>
<td>D. pacifica</td>
<td>Prolectella w</td>
<td>U sp. A</td>
</tr>
<tr>
<td></td>
<td>F. perpusilla</td>
<td>Xystonella w</td>
</tr>
<tr>
<td></td>
<td></td>
<td>X. longicaudata</td>
</tr>
</tbody>
</table>

* Genera noted as c = cosmopolitan, w = warm, or n = neritic, following
the classification of Pierce and Turner 1993.

Blanca Estuary (Barria de Cao 1992). Data from two tropical
systems were included: the coastal Caribbean waters of Ja-
mica (Gilron et al. 1991) and tropical estuarine and man-
grove waters on the southeastern coast of India (Godhantari-
aman 2002). Systems were compared in terms of numbers
of species and numbers of lorica size-classes. Each reported
species was assigned the average dimensions reported by
Kofoid and Campbell (1929, 1939), Campbell (1942), and
Marshall (1969) unless data were provided in the report (e.g.,
Verity 1987). Consequently, unidentified species were ig-
nored and such species (generally about 10% of the species
numbers) may have occupied unique LOD size-classes in the
assembly. For each system LOD frequency distribution was
plotted.

Chlorophyll concentrations were estimated in three frac-
tions for each sample: whole water, water filtered through a
10-µm Nuclepore filter, and water filtered through a 2-µm
Nuclepore filter. For each fraction, replicate samples of
300–500 mL were filtered using Whatman GF/F filter. Ma-
terial retained on the filters was extracted using methanol,
and Chl a concentrations were determined fluorometrically
(Holm-Hansen et al. 1965). The size-fractionated Chl a con-

Fig. 2. Map of station locations in New Caledonia.
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Fig. 3. Frequency distribution of LOD found in the pool of tintinnid species from the lagoon of New Caledonia compared with species pools reported in seasonal studies of tropical and temperate systems. Data for the coastal waters of the Bay of Bengal from Godhantaraman (2004), Jamaica from Gilron et al. (1991), Gulf of Naples from Modigh and Casaldo (2002), Narragansett Bay from Verity (1987), Damariscotta Estuary from Sanders (1987), Buzzards Bay from Pierce and Turner (1994), Nervion Estuary from Urrutxurtu (2004), Bahia Blanca Estuary from Barria de Cao (1992), and Hiroshima Bay from Kamiyama and Tsunjino (1996).

Fig. 4. Temporal changes in the concentrations of Chl $a$ and tintinnids between April 2002 and March 2003. (A) Data from the nearshore Sta. N12 and (B) data from the midlagoon Sta. M33. Statistical relationships are given in Table 2.

centrations (<2 $\mu$m, 2–10 $\mu$m, and >10 $\mu$m) were transformed into percentages of total chlorophyll and then used to estimate an index of the size-diversity of chlorophyll (Shannon index, ln-based). Statistical relationships between diversity and concentration estimates were examined using nonparametric Spearman rank correlation.

Results

General characteristics of New Caledonia tintinnids—A total of 76 species in 27 genera were found (Table 1). A complete listing of the abundance by date of each species, along with the lorica dimensions used for each species, is available on request from the corresponding author. The taxonomic richness found represents about 13% of the species and 50% of the genera in a ‘global’ tintinnid data set composed of the species cataloged in the monographs of Kofoid and Campbell (1929, 1939), Campbell (1942), and Marshall (1969). Following the distributional classification of genera proposed by Pierce and Turner (1993), most of the New Caledonia species (43) belong to cosmopolitan genera with the remainder species of either neritic (25) or warm-water genera (9). Thus, the tintinnid fauna of New Caledonia is not dominated taxonomically by tropical forms.

Compared to other systems, the tropical tintinnid fauna of New Caledonia appears rich in both numbers of species and size-classes of LOD (Fig. 3). Comparing communities found over an annual cycle, tropical systems do not necessarily show higher species or morphological diversity. Whereas the tintinnid community found in coastal waters of India appears to be more diverse than those of the temperate communities of the Atlantic and Pacific (47 vs. 20–32 species), Jamaican waters harbored relatively few species (18), representing only 9 size-classes of LOD. Thus, the diversity found New Caledonia waters can not be attributed to latitude alone. In contrast to the disparity found among tropical systems, temperate coastal communities appear remarkably consistent. Excluding the northwestern Mediterranean, which is arguably not a typical temperate site, the systems from the northwestern Pacific to three coasts of the Atlantic ranged narrowly in numbers of species (19–30) and numbers of LOD size-classes (9–13).

Temporal trends—At Sta. N12, chlorophyll averaged about 1 $\mu$g L$^{-1}$, and tintinnids averaged 250 cell L$^{-1}$. The midlagoon station showed lower average concentrations of both chlorophyll (0.25 $\mu$g L$^{-1}$) and tintinnids (50 cells L$^{-1}$). Tintinnid and total chlorophyll concentrations covaried loosely at both the nearshore Sta. N12 and the midlagoon Sta. M33 (Fig. 4). Chlorophyll and tintinnid concentrations were highest during the austral autumn and lowest in the austral spring period when water temperatures are lowest (21–22$^\circ$C). Pooling data from the two stations, chlorophyll and tintinnid concentrations were significantly correlated (Table 2).

A morphological characteristic of the tintinnid community can be related to a chlorophyll size parameter: the percentage greater than 10 $\mu$m. Shifts in the proportion of chlorophyll greater than 10 $\mu$m, which did not appear to follow any seasonal rhythm, were generally followed by shifts in the
Table 2. Spearman rank correlation coefficients (nonparametric) of pooled data (n = 64). *

<table>
<thead>
<tr>
<th></th>
<th>Chl a</th>
<th>Chl size H'</th>
<th>Tin conc.</th>
<th>Median LOD</th>
<th>No. of tin spp.</th>
<th>Tin spp. H'</th>
<th>No. of tin LODs</th>
<th>tin LOD H'</th>
</tr>
</thead>
<tbody>
<tr>
<td>Chl a &gt;10 µm</td>
<td>0.785***</td>
<td>0.352***</td>
<td>0.543***</td>
<td>0.364**</td>
<td>0.410***</td>
<td>0.214</td>
<td>0.400***</td>
<td>0.103</td>
</tr>
<tr>
<td>% Chl a &gt;10 µm</td>
<td>0.432***</td>
<td>0.396***</td>
<td>0.193</td>
<td>0.111</td>
<td>−0.046</td>
<td>−0.195</td>
<td>0.035</td>
<td>−0.172</td>
</tr>
<tr>
<td>Chl size H'</td>
<td>0.111</td>
<td>−0.046</td>
<td>0.195</td>
<td>0.762***</td>
<td>0.814***</td>
<td>0.448***</td>
<td>0.746***</td>
<td>0.273*</td>
</tr>
<tr>
<td>Tin concentration</td>
<td>0.350</td>
<td>0.118</td>
<td>0.212</td>
<td>0.608***</td>
<td>0.833***</td>
<td>0.834***</td>
<td>0.570***</td>
<td>0.596***</td>
</tr>
<tr>
<td>Median LOD</td>
<td>0.079</td>
<td>0.814***</td>
<td>0.448***</td>
<td>0.746***</td>
<td>0.834***</td>
<td>0.834***</td>
<td>0.570***</td>
<td>0.596***</td>
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<tr>
<td>No. of tin spp.</td>
<td>0.118</td>
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<td>0.834***</td>
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<tr>
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<td>0.834***</td>
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<td>tin LOD H'</td>
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<td>0.834***</td>
<td>0.834***</td>
<td>0.570***</td>
<td>0.596***</td>
</tr>
</tbody>
</table>

*Tint = tintinn ids. *p < 0.05, **p < 0.005, ***p < 0.0005.

median size of tintinnid LOD (Fig. 5). The percentage of chlorophyll greater than 10 µm was, on average, much higher in the nearshore station (about 40%) compared with that of the midlagoon station (about 20%). These differences corresponded to differences in median LOD values between the two stations, as the nearshore community averaged about 40 µm, and the midlagoon community averaged around 30 µm. Pooling data from both stations, the percentage of chlorophyll >10 µm was significantly correlated with the median LOD of the tintinnid community (Table 2).

Temporal changes in metrics of tintinnid diversity closely tracked one another. The numbers of species oscillated with the numbers of LOD size-classes found (Fig. 6). Both were highest during the austral autumn and lowest in the austral spring when both chlorophyll and tintinnid concentrations were minimal. An especially close association was found between H' values of species and LOD diversity (Fig. 7). Not surprisingly then, diversity metrics were strongly correlated with each other and to a lesser degree with tintinnid concentration (Table 2).

Scatter plots showed that taxonomic and morphological diversity measures were not related to each other in the same fashion, corresponding to the differences observed in temporal patterns. For the two closest relationships, numbers of species and LOD size classes displayed a curvilinear relationship, while H’ of species and H’ of LOD size-classes showed a near linear relationship (Fig. 8).

**Discussion**

Tintinnid abundance in the lagoon was similar to that of other systems, given the concentration of Chl a. The average concentrations of 10^1–10^2 tintinnid cells L^-1 corresponded with Chl a concentrations ranging from 0.2 to 2 µg L^-1, which is similar to reports for the southwestern Atlantic, (Thompson and Alder 2005), the Mediterranean (Dolan 2000), and Indian Ocean (Modigh et al. 2003). Although the abundance of tintinnids appeared predictable given chlorophyll concentration, the community composition was not.

Taxonomically, the assemblage did not appear to be a predictable assemblage simply from the location of New Caledonia. Species of tropical or warm-water genera were not dominant, but rather, species of cosmopolitan or neritic genera were. *Tintinnopsis* and *Eutintinnus* species dominated samples from the nearshore and midlagoon stations, respectively (Table 1). Species of these genera are common in both temperate and tropical coastal systems, ranging from both

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**Fig. 5.** Temporal changes in the relative importance of the larger fraction of chlorophyll greater than 10 µm and the median LOD of the tintinnid community. (A) Data from the nearshore Sta. N12 and (B) data from the midlagoon Sta. M33. Statistical relationships are given in Table 2.

**Fig. 6.** Temporal changes in the numbers of species and numbers of LOD grouped in 4-µm size-classes. (A) Data from the nearshore Sta. N12 and (B) data from the midlagoon Sta. M33. Statistical relationships are given in Table 2.
sides of the North Atlantic (Sanders 1987; Verity 1987; Urrutxurtu 2004), to Jamaica (Gilron et al. 1991), India (Gohhantaraman 2002), and the northwestern Mediterranean Sea (Cariou et al. 1999; Modigh and Castalado 2002). An explanation might be that the relatively shallow nature of a lagoon environment favors species of these genera, which are known to form benthic cysts.

In taxonomic terms there did not appear to be a well-defined tintinnid assemblage; that is, a tintinnid ‘fingerprint’ (i.e., Modigh et al. 2003) for the lagoon of New Caledonia. However, comparing the frequency distributions of LOD among various systems (Fig. 3), the New Caledonia community appears distinct in that most of the LOD size-classes between 10 and 100 µm were occupied.

The temperate communities of tintinnids, ranging across three coasts of the Atlantic and the North Pacific, were quite similar to each other in both the low number (10–13) and wide dispersion (20–120 µm) of LOD size-classes. Most of the temperate communities were dominated by two LOD size-classes; one around 20 µm and another around 40 µm. The two peaks in species-richness are likely related to seasonal changes in the size-spectrum of the phytoplankton community. Many temperate systems are characterized by the occurrence of two temporally and compositionally distinct peaks of phytoplankton concentration.

Field studies in distinct systems, both temperate and tropical, have shown that the tintinnid community characteristic of average LOD changes seasonally with the size spectrum of prey (Middlebrook et al. 1987; Verity 1987; Gilron et al. 1991). Laboratory studies of tintinnids established that LOD determines the maximum size of food particle ingested (Heinbokel 1978), and for ciliates in general with oral cilia, the minimum size of prey is likely related to the distance between the membranelles of the oral cilia (Fenchel 1987). Reviewing data on 11 tintinnid species, Dolan et al. (2002) found that LOD was about 4 times the size of prey items fed on most efficiently, in terms of clearance rate. Given these relationships, it is not surprising that in the lagoon of Nouméa, the most common LOD size-class was 20–24 µm, as most of the chlorophyll stock was <10 µm in size. Nor is it surprising that a measure of community morphology, the community median LOD, varied with the proportion of chlorophyll >10 µm in size (Fig. 5). Verity (1987) previously noted a similar relationship for Narragansett Bay tintinnids. Such observations support the idea put forward by Admiraal and Venekamp (1986) that community LOD is controlled by the size spectrum of phytoplankton prey items.

The surprising finding in our data was not that species diversity was related to LOD diversity, but rather to the strength of the relationship. The number of species found was reflected in the number of LOD size-classes present in a curvilinear manner (Fig. 8). When species abundance was low (<10 species), nearly each species represented a unique size-class; at higher species abundance, a size-class was occupied by up to five species. Biodiversity measured as H’ values, indicating both the number of different species and their relative abundance, were nearly equivalent to LOD H’ values (Figs. 7 and 8). This reflects the fact that a given LOD size-class, when composed of several species, was often dominated by a single species.

The close match of diversity metrics based on LOD or species suggests that in tintinnids, LOD data can be employed as proxy measure of diversity. We employed 4-µm size-classes as the finest distinction possible and reasonable. LOD appears to be about four times the size of prey most efficiently filtered (Dolan et al. 2002), thus 4-µm classes would differ in “preferred prey size” by 1 µm; intuitively,
Fig. 9. Scatterplots of species richness versus richness of LODs using groupings of 2-, 4-, and 8-μm size-classes. See Discussion for details.

Fig. 10. Scatterplots of LOD size-class richness and species richness of the tintinnid community encountered throughout annual cycles across different systems. Tropical systems are coastal waters of New Caledonia, Bay of Bengal, and Jamaica. Temperate systems are Buzzards Bay, Hiroshima Bay, Narragansett Bay, Damariscotta River Estuary, Nervion River Estuary, Bahia Blanca Estuary, and the Gulf of Naples. Data sources given in the legend to Fig. 3. 

an absolute lower limit of size distinction. Second, specimens were examined at ×200 or ×400; therefore, LOD estimates made in assigning species were then ±2 μm magnitude.

In a previous study, the relationship of morphological diversity and taxonomic diversity was examined using standard deviations (SDs) of average community lorica dimensions (Dolan 2000). This approach showed that SD was an indicator of taxonomic diversity, especially SD of lorica lengths, but was weakly correlated comparing communities across a large spatial scale (Dolan 2000). In a subsequent study, diversity of LOD size-classes was employed and shown to be closely related to both taxonomic diversity and the size diversity of phytoplankton across a large spatial scale (Dolan et al. 2002). In contrast to the data presented here, LOD size-classes were variable, set as equal to the number of species encountered, to maximize the possibility of distinguishing species via LOD (Dolan et al. 2002). The method we used here does not require knowledge of the number of species present; each sample is analyzed in a similar manner.

Automated or semiautomated systems of plankton analysis capable of producing data on tintinnid LODs potentially can be employed to estimate diversity. Sophisticated systems capable of resolving species within a tintinnid genus have been described (Culverhouse et al. 1994; Williams et al. 1994). However, our analysis suggests that relatively crude data on LOD sizes may suffice. Clearly, the image capture and analysis systems must be capable of resolving differences of about 2 μm and distinguishing empty from occupied lorica. It should be noted that results are dependant on the resolution of size-classes. As an example, Fig. 9 shows the relationship between numbers of species and LOD size-classes using different magnitudes of size-classes. Grouping in 8-μm intervals underestimates species abundance compared with 4-μm grouping. Using a finer grouping of 2-μm intervals yields estimates closer to our estimates of actual species richness. However, if the data employed were individual LOD measurements (rather than assigned LOD values), the 2-μm grouping may yield more LOD groups than species, because individual variability within a species exists.

The relationship between total species richness in various systems and LOD size-class richness in those systems is shown in Fig. 10. In this crude cross-system comparison,
morphological diversity is also closely related to taxonomic diversity. The abundances of LOD size-classes found appear to be one-half to one-quarter of the number species described as occurring in the system. This suggests that, most species are not the sole occupants of a given size-class. Over an annual cycle, two to four species can be found for each LOD size class.

In our study, each species was assigned a name on the basis of lorica morphology, including a reliance on lorica dimensions, and each species was assigned an average LOD on the basis of a description of the species. Thus, no allowance was made for plasticity in a given species. This is a possible source of error. Species exhibiting multiple morphologies in a sample would inflate diversity estimates. Species exhibiting distinct morphologies in distinct samples are less likely to have influenced diversity metrics. We have no data to permit an assessment of either possibility. However, species of genera known to show considerable plasticity of lorica morphology such as *Favella* (Laval-Peuto 1983), *Cymatocylis* (Wasik 1998), and *Psychocylis* (Davis 1981) were rare or absent in our samples. Species of the genus best known to show seasonal shifts in lorica dimensions, *Tintinnopsis* (Gold and Morales 1976), varied in length but little in LOD in our samples, with the exception of *T. campanula*, a minor component of the tintinnid community. LOD appears to be a conservative characteristic (Laval-Peuto and Brownlee 1986).

Morphological diversity studied here is a relatively simple analysis, focusing on a single character, LOD, compared to investigations of multicellular organisms. Morphological diversity in fossil or living taxa has been defined as an estimate derived from a consideration of a multivariate morphospace (e.g., Roy et al. 2004). In protistan and metazoan taxa, compared to gastropods, for example, perhaps both the strength and weakness of investigating morphological diversity is the relative simplicity of their morphologies. Thus, the limited number of characters available increases the opportunity to find and focus on a character or characters related to their ecological niche. However, it should be recalled that characters should not be restricted to morphology, but rather, extended to include physiology.

Tintinnids in the tropical Pacific lagoon of New Caledonia are a species-rich assemblage; 76 species were distinguished. The community showed temporal changes in diversity and morphology (median LOD), which were only weakly related to changes in trophic resources measured as Chl a. However, metrics of morphological diversity based on LOD were correlated with metrics of taxonomic diversity. Because LOD is related to the size of prey exploited in tintinnids, correlation of taxonomic and LOD diversity supports the hypothesis that the taxonomic diversity of tintinnids in the lagoon reflects a trophic diversity. Although the relationship of primary producer diversity generating consumer diversity is intuitively appealing, this has rarely been shown (Turner and Hawkins 2004). Furthermore, metrics of LOD diversity in tintinnids appear to be a good proxy measure of taxonomic diversity both within and among systems.

References


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