PREDATION OF MUDFLAT MEIO-MACROFAUNAL METAZOANS BY A CALCAREOUS FORAMINIFER, AMMONIA TEPIDA (CUSHMAN, 1926)

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ABSTRACT

Benthic foraminifera are heterotrophic protists that utilize different trophic mechanisms and nutritional resources. They exhibit a wide range of trophic behaviours: selective (grazing) and indiscriminate herbivory, symbiosis, carnivory, parasitism, uptake of dissolved organic matter, passive suspension feeding and, most commonly, deposit feeding. The benthic foraminifera Ammonia tepida, previously known as an herbivore, fed as a carnivore in laboratory experiments where mobile metazoans were provided. We observed predation on the three types of metazoa provided: nematodes, copepods, and a larval gastropod. This foraminifera used its pseudopodial network to entrap the invertebrates, which were then stripped of their soft internal tissues within 24 hours. Our experiments are the first to demonstrate that Ammonia tepida, despite its limited motility, is able to utilize larger mobile animals as a food source. The great abundance of small metazoans in most marine environments suggests that they are a food source for foraminifera. Further study of foraminiferal feeding strategies will enhance our understanding of their role in marine communities.

INTRODUCTION

Foraminifera are heterotrophic protists present in both pelagic and benthic marine environments. Benthic foraminifera inhabit all water depths and populate a variety of microhabitats on and in the substrate (Murray, 1991; Ellison, 1984; Chandler, 1989; Moodley and others, 1998, 2000, 2002, 2005; Murray and Alve, 2000). They utilize a diversity of trophic mechanisms and nutritional resources (Goldstein, 1999), and probably play an important role in food webs that influence the structure of the benthic community (Altenbach, 1992; Linke and others, 1995; Moodley and others; 2000; Fontanier and others, 2002; Nomaki and others, 2008; Suhr and others, 2008).

Trophic behaviours exhibited by foraminifera include selective (grazing) and indiscriminate herbivory (Jeeps, 1942; Sliter, 1965; Jones and Charnock, 1985; Lee and others, 1991; Moodley and others 2002; Witte and others, 2003; Nomaki and others, 2008), symbiosis (Lee and Anderson, 1991), parasitism (Cedhagen, 1994), uptake of dissolved organic matter (DeLaca and others, 1981; Lipps, 1983), passive suspension feeding (Lipps, 1983; Cedhagen, 1988; Lutze and Altenbach, 1988; Lutze and Thiel, 1989), and deposit feeding (Lipps 1983; Jones and Charnock, 1985). The majority of foraminiferal species are assumed to be omnivorous, feeding on organic detritus, bacteria, and algae (Lee, 1980; Lipps, 1983). Some foraminifera are known to feed on metazoans (Buchanan and Hedley, 1960; Bowser and others, 1986, 1992; Goldstein, 1999; Suhr and others, 2008), but most of them are not exclusively carnivorous and utilize carnivory in addition to at least one other trophic mechanism (Goldstein, 1999). Carnivory by planktonic foraminifera is well documented (e.g. Boltovskoy and Wright, 1976, Bé and others, 1977), but little is known about this behavior among benthic foraminifera (see review in Goldstein, 1999). The position of foraminifera in food webs remains conjectural despite direct observations on the diets of some species (Murray, 1963; Anderson and Lee, 1991; Lee and others, 1991; Bernhard and Bowser, 1992; Goldstein, 1999; Heinz and others, 2005; Pascal and others, 2008a). Foraminifera typically use their pseudopodia to gather and ingest food (Bowser and others, 1992). Planktonic foraminifera are known to prey upon copepods and other crustaceans (Anderson and Bé, 1976; Bé and others, 1977; Caron and Bé, 1984; Snider and others, 1984; Hemleben and others, 1988). Some larger benthic foraminifera also feed on metazoans. For example, Peneroplis pertusus (Forskál) feeds on copepods by ingesting the internal soft parts after which it discards the empty carapace (Winter, 1907 cited by Goldstein, 1999). The large agglutinated foraminifera Astrorhiza limnicola (Sandahl) feeds on crustaceans and echinoderms (Nyholm, 1956; Buchanan and Hedley, 1960); however, the same species has also been observed only as a suspension feeder (Cedhagen, 1988). Several authors (DeLaca, 1986; Bowser and others, 1992; Suhr and others, 2008) consider Astrammina rara (Rhumbler), another large agglutinated foraminifera, to be carnivorous. Relatively smaller benthic foraminifera such as Epithia and Pyrgo also have been observed capturing prey, but not feeding on them (Jepps, 1942; Suhr and others, 2008). Few studies suggest predation by small benthic foraminifera (Christiansen, 1971; Cedhagen, 1994; Hallock and Talge, 1994).

The purpose of this study is to examine and document the trophic interactions between the smaller benthic foraminifera Ammonia tepida and several kinds of metazoans in laboratory-controlled experiments. Ammonia tepida is known as a deposit-feeder on algae (Lee, 1980; Stouff and others, 1999; Moodley and others, 2000) and bacteria (Goldstein and Corliss, 1994; Langezaal and others, 2005; Pascal and others, 2008a). Species of Ammonia have been used in numerous laboratory experiments because the genus is ubiquitous in inner-shelf, estuarine, and saltmarsh environments (Murray, 1991), where they tolerate wide ranges in temperature, salinity, and other physico-chemical parameters (e.g., Bradshaw, 1961; Schnitker, 1974; Walton and Sloan, 1990; Debenay and others, 1998). Being hardy and readily collected, Ammonia is ideal subjects for laboratory study of living specimens.

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METHODS

COLLECTION OF FORAMINIFERS AND METAZOANS

Foraminifera and invertebrates were sampled at low tide by scraping off the top centimeter of sediment from the upper-intertidal zone of Brouage mudflat, located on the French Atlantic coast about 20 km south of La Rochelle, 45°54'N, 1°7'W (Fig. 1). Time between collection and experiment was minimized. This environment supports a high density of living foraminifera, especially *Ammonia tepida* (Pascal and others, 2008b). The meiofaunal community of this mudflat is dominated by nematodes (95%), with subsidiary copepods (2%), both of which are present throughout the year (Rzeznik-Orignac and others, 2003). The deposit-feeding gastropod *Hydrobia ulvae* (common mudsnail) (Haubois and others 2004), as previously demonstrated by Rossignol and others (2007). After two days, most of the sediment had been ingested by the gastropods and excreted as fecal pellets, and the living benthic foraminifera that had eluded the gastropod were clean and readily visible for efficient picking with a very fine brush. Specimens of *A. tepida* were transferred to Petri dishes filled with 0.2 m of filtered seawater from the study area. Foraminifera were acclimated for a minimum of 24 hours at 18°C with a daily light-dark cycle before the feeding experiment commenced.

LIVING NEMATODES

Living nematodes were concentrated by first seawater-washing sediment samples through a 65-μm nylon mesh. Sediments >65 μm were distributed on a 0.5-cm-thick layer

LABORATORY PROCEDURES

Sediment samples were washed through a 50-μm sieve and the >50-μm fraction was distributed among several glass Petri dishes. Extraction of living *Ammonia tepida* was facilitated by exposing the sediment to 80 adult *Hydrobia ulvae* (common mudsnail) (Haubois and others 2004), as previously demonstrated by Rossignol and others (2007). After two days, most of the sediment had been ingested by the gastropods and excreted as fecal pellets, and the living benthic foraminifera that had eluded the gastropod were clean and readily visible for efficient picking with a very fine brush. Specimens of *A. tepida* were transferred to Petri dishes filled with 0.2 μm of filtered seawater from the study area. Foraminifera were acclimated for a minimum of 24 hours at 18°C with a daily light-dark cycle before the feeding experiment commenced.

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of coarse sand (1 mm grain size) over a 20-μm nylon mesh placed above a seawater bath and exposed to light for two days. Following this negative phototropism method described by Rzeznik-Orignac and others (2004), living nematodes migrated from the sand, through the nylon mesh, and into the seawater. The living nematodes were then cleaned of organic matter, transferred to 0.2-μm filtered seawater from the study area, and acclimated prior to the feeding experiment, as described above for A. tepida.

Ten live A. tepida and 10 live nematodes or copepods were placed in 0.2-μm-filtered seawater on the same 5-cm diameter Petri dish. The control dish similarly received 10 live nematodes or copepods but no foraminifera. Foraminiferal vitality was verified by observing pseudopodial activity, while nematode and copepod vitality was verified by their mobility. All feeding experiments were carried out at room temperature (18°C). Nematodes were kept in the dark, whereas copepods had a 24-hour light-dark cycle. Each part of the experiment was repeated three to five times. The behaviors of foraminifera, nematodes, and copepods were observed under a stereomicroscope regularly over the 24 hours. Photographs were taken with a LEICA DM IRB inverted stereomicroscope (×400 maximum magnification) equipped with an Olympus DP-70 digital camera, and processed with Visilog software.

An additional series of experiments was carried out where each dish had one juvenile Hydrobia ulvae gastropod and five A. tepida, under the same light-dark cycle that was applied in the copepod experiments. Observations were made with an Olympus SZX-12 stereomicroscope equipped with a Olympus E-330 digital camera.

**SPECIMEN PREPARATION FOR ELECTRON MICROSCOPY**

The various steps of this preparation method were performed with small microcentrifuge (Eppendorf-type) tubes that were conducive to the preservation of the fine, fragile pseudopodia. When nematodes were captured by foraminifera, samples were fixed in a solution of 2.5% glutaraldehyde diluted in 0.45-μm filtered seawater for 24 hours. To avoid desiccation and consequent loss of detail, the foraminiferal specimens with prey were washed with 0.2 M cacodylate buffer (pH 7.4). Post-fixation was accomplished using 2% OsO₄ diluted in the cacodylate buffer, followed by three rinses with 0.45-μm-filtered seawater and dehydration in a 50–75–95% series of ethanol baths, plus two baths of 100% ethanol. Specimens were then immersed in HMDS (hexamethyldisilazane) for 10 minutes, after which they air-dried. Dried specimens were mounted on SEM stubs covered with carbon-conductive adhesive tape then double-coated with carbon. Observations and imaging were obtained with a Jeol 6301F SEM at the SCIAM (Service Commun d’Imagerie et d’Analyses Microscopiques) of the University of Angers.

**RESULTS**

**PREDATOR-PREY RELATIONSHIP BETWEEN FORAMINIFERA AND NEMATODES**

Acclimated specimens of Ammonia tepida extruded a dense network of pseudopodia that extended onto the dorsal and ventral surfaces of the test. The foraminifera generally oriented themselves perpendicular-to-oblique relative to the bottom of the Petri dish, and used their pseudopodial network to attach at two main points on the glass bottom (Fig. 2a).

Nematodes encountering the networks were immediately entrapped (Fig. 2b), adhering to the pseudopodia or to the cyst on the ventral side of the tests (Figs. 2f–i). When stuck in the foraminiferal pseudopodia, the nematodes struggled but rarely escaped. Some nematodes were captured by two foraminifera (Figs. 2f, 2g). About 18 hours after initial contact, the foraminifera began to empty the nematode of its soft tissues (Fig. 2e). Within six hours, only an empty cuticula remained (Fig. 2c, 2d). Each cast-off cuticula was characterized by a hole that had been created by the predatory foraminifera (Figs. 2d). The predation of nematodes by A. tepida occurred every time these two meiofaunal organisms were observed in contact. The control nematodes that were not in placed into association with foraminifera remained alive and unscathed after 24 hours.

**PREDATOR-PREY RELATIONSHIP BETWEEN FORAMINIFERA AND OTHER METAZOA ORGANISMS**

As with the nematodes, copepods also fell prey to A. tepida whenever the two made contact. Despite vigorous attempts to escape, copepods could not free themselves from the pseudopodial mesh. Often, two foraminifera trapped the same copepod (Fig. 3a). After ingesting the soft tissues of the prey, the empty carapace was discarded.

One observation was made of A. tepida feeding on a juvenile gastropod. The foraminifera attached to the aperture of a living Hydrobia and, about 20 hours later, the snail shell was empty (Fig. 3b, 3c).

**DISCUSSION**

The carnivorous behavior of Ammonia tepida in the laboratory has also been observed on meio-macrofauna. It appears that this carnivorous behaviour of A. tepida is not specific for the specimens of our study area. A similar carnivorous behaviour has been observed in A. tepida collected from the Japanese coast (H. Nomaki, personal communication, 2008) indicating that carnivory in A. tepida is a general feeding strategy under laboratory conditions. But is this feeding strategy used by A. tepida in its natural environment?

In mudflat environments, due to the abundance of nematodes (on average, 42 times more abundant than foraminifera) and foraminifera, especially in the surface sediments of mudflats (Pascal and others, 2008b), contact between A. tepida and small metazoans is likely to occur very frequently. In the Brouage mudflat, for example, nematode abundances range 800–4,050 × 10³ m⁻² (mean of 2,100 × 10³ m⁻²) with maximum abundances in winter and spring (Rzeznik-Orignac and others, 2003), while foraminifera range 20–170 × 10³ m⁻² (mean of 50 × 10³ foraminifera m⁻²) with their maximum in winter (Pascal and others, 2008b). By preying on meiofauna, A. tepida may be considered to be on a trophic level similar to metazoan consumers, but it is probably not alone among the benthic foraminifera. Microscope imaging with fatty-
Acid biomarker analysis strongly suggests that the diet of *Astrammina rara* (Rhumbler) includes polychaetes, crustaceans, molluscs, and echinoderms (Suhr and others, 2008). Chandler (1989) reported on *Ammonia beccarii* thought to possibly be an amensal relationship with the copepod *Amphiascoidea limicola* (Brady). The presumption is that this is an indirect result of having consumed most of the microflora, leaving the less-nutritious detritus for the copepod. We suggest that the lower copepod densities observed by Chandler (1989) could also be due to foraminiferal predation or copepod avoidance of sediments supporting high densities of *Ammonia*. However, our results suggest that *A. tepida* are able to directly feed on large and actively moving grazers. *Ammonia* species have typically been considered herbivores, with *Ammonia tepida* feeding upon algae and bacteria (Goldstein and Corliss, 1994; Moodley and others, 2000; Langezaal and others, 2005; Pascal and others, 2008a). Moodley and others (2000) reported *Ammonia* sp. exhibiting rapid uptake of freshly deposited algal carbon. It is now evident that some of those foraminifera are likely to be omnivorous. *Ammonia*, the most common benthic foraminiferal genus, ubiquitous in inner shelf, estuarine, and saltmarsh environments, may be able to employ various feeding strategies according to the most available food sources. Thus, *A. tepida*’s position in the benthic food web is complex, occupying both primary and secondary consumer positions. Thus, they may have a greater impact on benthic community structure than previously suspected. Because of their limited motility, active hunting for prey is probably not common among carnivorous foraminifera. Whereas *A. tepida* spreads its pseudopodial network more extensively in this feeding mode than when grazing, the trap appears to be intentionally set to entangle larger, mobile prey that unwarily wander into it. Additional experiments are needed to confirm if *A. tepida* switches feeding modes according to the most available food sources.

Three significant questions arise from our results:

1. **How do prey remain attached to foraminiferal pseudopodia?**

   Goldstein (1999) stated that the pseudopodia of carnivorous foraminifera are specifically designed to catch prey.
**FIGURE 3.** *Ammonia tepida* predation on copepods and gastropod larvae (a–c: photomicroscopic images; d: electron micrograph) a Two foraminifera (denoted by stars) feeding on a copepod; note the part of the carapace visible on the right is now empty; b Benthic juvenile gastropod *Hydrobia ulvae* caught alive by three foraminiferans (denoted by stars); c After 20 hours, the gastropod shell is empty; note the cyst-covered foraminiferan; d Oblique umbilical view of *A. tepida*; e detailed view of imaged.
Foraminifera secrete an adhesive (Buchanan and Hedley, 1960), possibly in their Golgi vesicles (Goldstein, 1999), which is then transported to the peripheral cytoplasm and released by expulsion in the vicinity of prey (Anderson and Lee, 1991; Bowser and others, 1992). This substance is very sticky and thought to be non-toxic (Langer and Bell, 1995). We observed that the pseudopodia and ventral cyst of A. tepida are capable of retaining very active nematodes and copepods. Thus, A. tepida is able to produce an adhesive substance that can be used for capturing meio-macrofauna. However, we cannot confirm that this adhesive substance is produced only for capturing food. In the case of feeding on bacterial biofilm, Bernhard and Bowser (1992), using time-lapse microscopy, revealing that biofilm parcels are transported extracellularly toward the foraminiferal cell body by pseudopodia, an observation which further implicates pseudopodial function in foraminiferal trophic mechanisms.

2. How do foraminifera immobilize prey?

It is not clear from our observations whether foraminifera sedate or kill their prey prior to ingestion. It is conceivable that foraminifera use narcotic agents or extracellular enzymes to immobilize or kill their prey, as Anderson and Bé (1976) suggested for planktonic foraminifera. In either case, the digestion of prey by foraminifera usually occurs via extruded pseudopodia (Buchanan and Hedley, 1960), and that process will eventually terminate any active resistance. Extensions of the pseudopodia push the prey against the foraminiferal cell body, an observation which further implicates pseudopodial function in foraminiferal trophic mechanisms.

3. How do foraminifera penetrate the cuticula of nematodes?

Austin and others (2005) propose that foraminifera such as Haynesina Banner and Culver use their test ornamentation to mechanically break diatom frustules. Anomonia tepida also present pustules around the aperture of the test (Fig. 3d, e) and this ornamentation may be used in penetrating cuticula. It has been suggested that secretions by pelagic foraminifera aid in digesting prey (Snider and others, 1984). Spindler and others (1984) claim that pelagic foraminiferal pseudopodia are not physically capable of boring through crustacean carapaces, and therefore some substance must be secreted to dissolve the carapace at the point of entry. Benthic foraminifera such as A. tepida may also use chemical digestion. Bowser and others (1985) observed pseudeopodial activities in Allogromia, and reported that pseudopodia tore small pieces from a gel, a behavior termed skyllocytosis. The authors suggested that skyllocytosis may also be used by carnivorous foraminifera to obtain prey tissues. Skyllocytosis is an alternative hypothesis for how A. tepida penetrates the nematode cuticula.

CONCLUSIONS

Although previous studies reported Anomonia tepida as an herbivore that feeds on algae and bacteria, the species behaved as a carnivore in our laboratory experiments, intentionally orienting itself and extending its sticky pseudopodial network to capture metazoa that wandered too close. We observed A. tepida preying upon nematodes, copepods, and a gastropod larva that were placed near them. Thus, the species is probably omnivorous and a secondary consumer in its natural environment, possibly switching feeding modes to take advantage of the food that is most readily available. Further experiments are needed to determine whether A. tepida has a trophic preference when multiple food types are simultaneously available. We also need to quantify in situ carbon fluxes between nematodes, copepods, juvenile Hydrobia, and this foraminifera, and then integrate these fluxes in a food web as we continue unravelling the complexities of the carbon cycle in coastal mudflats.

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