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quantification using $^{15}$N enriched bacteria

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Abstract

The fate of benthic bacterial biomass in benthic food webs is a topic of major importance but poorly described. This paper describes an alternative method for evaluation of bacterial grazing rate by meiofauna and macrofauna using bacteria pre-enriched with stable isotopes. Natural bacteria from the sediment of an intertidal mudflat were cultured in a liquid medium enriched with $^{15}$NH$_4$Cl. Cultured bacteria contained 2.9 % of $^{15}$N and were enriched sufficiently to be used as tracers during grazing experiments. Cultured-bacteria presented a biovolume (0.21 $\mu$m$^3$) and a percentage of actively respiring bacteria (10 %) similar to those found in natural communities. The number of Operational Taxon Units found in cultures fluctuated between 56 and 75 % of that found in natural sediment. Despite this change in community composition, the bacterial consortium used for grazing experiments exhibited characteristics of size, activity and diversity more representative of the natural community than usually noticed in many other grazing studies. The bacterial ingestion rates of three different grazers were in the range of literature values resulting from other methods: 1149 ngC ind$^{-1}$ h$^{-1}$ for the mud snail *Hydrobia ulvae*, 0.027 ngC ind$^{-1}$ h$^{-1}$ for the nematode community, and 0.067 ngC ind$^{-1}$ h$^{-1}$ for the foraminifera *Ammonia tepida*. The alternative method described in this paper overcomes some past limitations and it presents interesting advantages such as short time incubation and *in situ* potential utilisation.

KeyWords

Bacteria, grazing, sediment, stable isotope, tracer

Introduction

Development of improved methods for measuring bacterial abundance and production have radically changed the perception of the role of bacteria in pelagic marine ecosystems.
Bacteria are known to play a major role in organic matter degradation and regeneration of nutrients. Moreover the “microbial loop” model (e.g. Azam et al. 1983) considers bacteria as a “link” more than a “sink”, increasing the ratio of primary production available for higher trophic levels. Therefore, bacteria appear to play a major role in pelagic foods web models (e.g. Vézina and Savenkoff 1999).

Bacterial abundance in marine soft sediments is relatively constant, around $10^9$ cells.ml$^{-1}$ porewater (Schmidt et al. 1998), being a thousand times more abundant than in pelagic systems. Moreover, high rates of production have been measured in aquatic sediments (e.g. van Duyl and Kop 1990). These findings have driven a debate on the fate of bacteria in benthic food webs. Due to technical limitations, studies dealing with benthic bacterivory are not as developed as pelagic ones (Kemp 1990).

Pelagic bacteria are mainly grazed by protozoa (e.g. Sherr and Sherr 1994) and a similar pattern was expected in benthic systems (van Duyl and Kop 1990; Bak et al. 1991; Hondeveld et al. 1994). Nevertheless, numerous authors consider the bacterial grazing by benthic ciliates and flagellates as insignificant (Alongi 1986; Kemp 1988; Epstein and Shiaris 1992; Epstein 1997). Depending on the studies, meiofauna grazing is considered either as (i) high enough to structure microbial communities (Montagna 1984b), (ii) using 3% of bacterial production (van Oevelen et al. 2006a), or (iii) negligible (Epstein and Shiaris 1992). Data on macrofaunal grazing rates are not less variable than on meiofaunal ones. In a synthesis, Kemp (1990) asserted that bacteria density is not high enough to play a major role in macrobenthos diet. These contrasting conclusions probably reflect the use of different methods. In conclusion, it appears that drawing a general view of the role played by microfauna, meiofauna and macrofauna in bacterial grazing is presently difficult (Kemp 1990).

Most of the benthic studies on trophic process employ tracers. Labels can be added directly to sediment. In such a situation, bacteria incorporate labels and are simultaneously grazed by
predators (Montagna 1995; van Oevelen et al. 2006a; van Oevelen et al. 2006b). This technique minimizes disturbance of the spatial distribution and metabolism of grazers and bacteria (Carman et al. 1989). Nevertheless, only a small part of the bacterial assemblages takes up detectable quantities of labels (Carman 1990b). Moreover, the main drawback to this method is that a large part (up to 83% in Montagna & Bauer (1988)) of the total labels uptake may be attributable to processes other than grazing. Grazers may become labelled by absorption and adsorption of dissolved organic matter (DOM) (Montagna 1984a) or by uptake of labels by non-prey microorganisms associated with grazers (e.g. epicuticular or gut microorganisms) (Carman 1990a).

To reduce this bias, microbial prey can be prelabelled with fluorescent products, or isotopes either stable or radioactive. Fluorescent Labelled Bacteria (FLB) with monodispersed FLB or whole-sediment staining methods are used mainly to assess grazing activity of small predators like flagellates and ciliates (Novitsky 1990; Epstein and Shiaris 1992; Hondeveld et al. 1992; Starink et al. 1994; Hamels et al. 2001). meiofauna studies using FLB are seldom because FLB detection is difficult and time consuming in large sized grazers. Consequently, only a small number of specimens can be examined preventing detection of inter individual variations in grazing rate. Therefore, grazers like nematodes are able to discharge various digestive enzyme to realise extracorporeal hydrolyse of food (Riemann and Helmke 2002). In such a case, pre-digested FLB ingested are impossible to detect in grazers. Nevertheless bacterivory levels by foraminifera (Langezaal et al. 2005) and nematodes (Epstein 1997) were assessed using FLB.

Bacterivory assessment by the way of adding radioactive or stable isotope to sediment was performed on meiofauna and macrofauna (e.g. Montagna 1984b; Sundback et al. 1996; van Oevelen et al. 2006a; van Oevelen et al. 2006b).
Bacterivory assessment using prelabelled bacteria was performed with radioactive isotopes (Rieper 1978; Carman and Thistle 1985). To our knowledge, the use of stable isotopes on prelabelled bacteria has never been performed until present. Compared to radioactive isotopes, bacteria enriched with stable isotopes are more convenient to use, since they can be used in situ without negative environmental effects and legal restrictions. This method will help investigators who are limited by radioactive material prohibition.

The aim of this paper is to describe a method using $^{15}$N stable isotope to prelabel bacteria in the view to assess bacterivory of large size benthic organisms (meiofauna and macrofauna). Experiments were performed mainly to assess the validity of this method, taking in consideration size, diversity, and activity of the prelabeled bacteria in order to be close to natural population parameters. The method was applied to 3 grazers from an intertidal mudflat in order to appreciate its potential generalization: one mollusc Hydrobia ulvae, a nematode community and the foraminifera Ammonia tepida.

**Material and methods**

*Bacterial culture*

Superficial sediment (1 cm) was collected from the Brouage intertidal mudflat located in the eastern part of Marennnes-Oléron Bay (45°55N, 1°06W) on the Atlantic Coast of France. One ml of this sediment was added to 20 ml of bacterial liquid culture medium and kept in the dark at 13°C during 24 hours. The liquid bacterial culture medium was composed of: peptone 3 g.l$^{-1}$ (BioRad), yeast extract 1 g.l$^{-1}$ (BioRad), $^{15}$NH$_4$Cl 1 g.l$^{-1}$ (99 % $^{15}$N-enriched NH$_4$Cl CortecNet); sodium glycerophosphate 0.025 g.l$^{-1}$ and sequestren Fe 6 g.l$^{-1}$. It was completed with 0.2 μm filtered distilled water (500 ml) and 0.2 μm filtered sea water (500 ml) at pH 7.4. The first culture was subcultured during 24 hours under the same incubation conditions in the view to reach approximately $2 \times 10^9$ cells.ml$^{-1}$. Bacteria were rinsed (i.e. separated from
culture medium) by the means of 3 centrifugations (3500 g, 10 min, 20°C) in 0.2 µm filtered
sea water, then frozen in liquid nitrogen and kept frozen (-80°C) until grazing experiments.
The bacteria δ¹⁵N was measured on an Eurovector Elemental Analyser coupled with an
Isotope Mass Ratio Spectrometer (Isoprime, Micromass). Nitrogen isotope composition is
expressed in the delta notation (δ¹⁵N) relative to air N₂: δ¹⁵N = [(¹⁵N/¹⁴N)sample / [(¹⁵N/¹⁴N)reference]-1] × 1000. Rinsing efficiencies were tested using bacteria cultured in the
medium previously described with non-enriched NH₄Cl. These bacteria were killed by
formalin (2 %), placed in the ¹⁵N-enriched culture medium previously described, harvested by
the means of 3 centrifugations (3500 g, 10 min, 20°C) before isotope ratio measurement.

*Cultured bacteria size*
Size of bacteria from original sediment and cultures were measured. For sediment samples,
particle-associated bacteria were detached by pyrophosphate (0.01M) and sonication. Bacteria
were stained using 4,6-diamidino-2-phenylindole dihydrochloride (DAPI) (2500 µg.L⁻¹) and
filtered onto 0.2 µm Nucleopore black filters (Porter and Feig 1980).
Length (L) and width (2r) of each bacteria was determined by a computer-assisted image
analysis (AxioVision Release 4.3) with an epifluorescence microscope (Axioskop 2 mot plus
– Zeiss) equipped with a charge-coupled device camera (AxioCam MRc5 – Zeiss). Bacterial
biovolumes (V) were calculated for cultured bacteria (N = 1981) and natural bacteria (N =
1806) as follows: $V = \pi r^2.(L-2/3 r)$ (Fuhrman 1981).

*Cultured bacteria activity*
Frozen aliquots of cultured bacteria were thawed and immediately incubated with 5-cyano-
2,3-ditoyl tetrazolium chloride (CTC) (final concentration of 5 mM). After 2, 3 and 5 hours of
incubation, experiments were stopped with formalin (2%) and stored at 4°C. Bacterial
samples were processed as described above for the DAPI staining in order to simultaneously
count total cells (UV excitation) and active cells (green excitation) on same slide.
Cultured bacteria diversity

The bacterial diversity of original sediment and culture aliquots was assessed. The DNA was extracted using an Ultraclean Soil DNA Kit (MO BIO, Oxyme) for sediment samples and a QIAamp DNA Mini Kit (Qiagen) for cultures. Bacterial 16S rRNA gene fragments of about 520 bp (the V6-V8 regions of 16r DNA (Gelsomino et al. 1999)) were amplified by PCR using primers 968fGC (5’-AACGCGAAGAACCTAC-3’[with GC clamp 5’]) and 1401r (5’-CGGTGTGTACAAGGCC-3’).

PCR products (300 ng) were loaded onto polyacrilamide gel (8% w/v, 7M urea) in TAE 1 X buffer. Electrophoresis was processed under a constant voltage of 68 V, during 17 h, with a thermal gradient from 66 to 69.7°C increasing at the rate of 0.2°C h⁻¹ (Dcode™ System: Biorad). The gel was stained with 0.5 μg.ml⁻¹ Gelstar (BMA) in 1.25× TAE buffer during 30 min and checked through a UV transilluminator system (Versa Doc (Bio-Rad)) equipped with a camera. Temperature Gradient Gel Electrophoresis (TGGE) banding patterns were automatically calculated by the Bionumerix software (Applied Biomaths, Koutrai, Belgium) using the Dice coefficient (DC), without band weighing by both the complete linkage and unweighted pair group method with arithmetic mean (UPGMA) algorithms (threshold of 1%).

Grazing experiments

The first centimetre of sediment was collected from a square meter patch during ebb tide from the Brouage intertidal mudflat (France) on March 13, 2006. It was sieved on 500 μm, 200 μm and 50 μm in order to concentrate respectively H. ulvae, A. tepida and nematodes. Choice of these organisms was driven by their high natural abundance in the study area. Each type of grazer was placed in individual microcosms. Seventeen handpicked specimens of H. ulvae were placed in polypropylene Petri dishes (diameter 9 cm). For the foraminifera and nematode experiments, 1 ml of the fraction remaining on the 200 μm and on the 50 μm mesh sieves respectively were placed in 100 ml Pyrex beakers. Each experiment was carried out in
triplicate, along with triplicate controls. Control samples were frozen (-80°C) during 12 hours in order to kill grazers before thawing.

Sediment that passed through the 50 μm mesh was mixed with $^{15}$N enriched bacteria. Abundance of sediment and cultured bacteria were counted using the methods previously described. This slurry containing $1.05 \times 10^9$ bacteria ml$^{-1}$ and $^{15}$N enriched bacteria were twice as abundant as natural ones. Seventeen ml of this slurry were placed in *H. ulvae* microcosms and 4 ml were placed in nematode and foraminifera microcosms. Grazing incubations were run in the dark at 20°C. Incubations were stopped by freezing the microcosms at -80°C.

*H. ulvae* were separated by hand from their shell and all specimens of each microcosm were pooled and homogenised using a Potter-Evelhjem. Nematodes were extracted from sediment using ludox (Heip et al. 1985). Approximately 700 nematodes were randomly handpicked from each sample. Foraminifera were stained with rose Bengal in order to identify living specimens. As Rose Bengal is an organic compound, it could affect isotopic composition but control experiments were also stained in order to take this bias into account.

For each sample, 150 specimens of *A. tepida* were picked individually and cleaned of any adhering particles.

$\delta^{15}$N of grazers was determined using the technique described above. Incorporation of $^{15}$N is defined as excess (above background) $^{15}$N and is expressed in terms of specific uptake ($I$) (gN ind$^{-1}$). $I$ was calculated as the product of excess $^{15}$N ($E$) and biomass of N per grazer. $E$ is the difference between the fraction $^{15}$N in the background ($F_{\text{background}}$) and in the sample ($F_{\text{sample}}$): $E = F_{\text{sample}} - F_{\text{background}}$, where $F = ^{15}$N / ($^{15}$N + $^{14}$N) = $R / (R + 2)$ and $R$ is the nitrogen isotope ratio. For the $F_{\text{background}}$, we used control values measured with killed grazers (frozen). $R$ was derived from the measured $\delta^{15}$N values as: $R = ((\delta^{15}N/1000)+1) \times R_{\text{airN2}}$ with $R_{\text{airN2}} = 7.35293 \times 10^{-3}$ (Mariotti 1982). The uptake of bacteria (gC ind$^{-1}$ h$^{-1}$) was calculated as
Uptake = \((I \times (\% \text{C}_{\text{enriched bacteria}} / \% \text{N}_{\text{enriched bacteria}})) / (F_{\text{enriched bacteria}} \times \text{incubation time})\). This uptake was multiplied by the ratio between the abundance of total and enriched bacteria, determined from DAPI counts. Uptake \((gC_{\text{bacteria}}/h/gC_{\text{grazer}})\) were obtained by dividing uptake of bacteria \((gC/\text{ind}/h)\) by grazer mean weight \((gC/\text{ind})\).

### Results

**Characteristics of enriched bacteria**

Bacteria cultured in a liquid medium with a 18 mM \(^{15}\text{NH}_4\text{Cl}\) subsequently centrifuged to remove unincorporated label were found to contain \(2.88 \pm 0.03 \% \text{^{15}N}\). Bacteria killed by formaldehyde before being placed in the same culture medium and centrifuged, contained \(0.028 \% \text{^{15}N}\). Cultured bacteria were enriched enough to allow their detection in the three studied grazers (Tab. 1). Bacterial abundance was not affected by the liquid nitrogen freezing process.

Cell volume of cultured bacteria \((0.21 \mu m^3 \pm 0.26)\) was not significantly (bilateral unpairwise student test; \(p = 0.07\) ) different from cell volume of natural bacteria \((0.23 \mu m^3 \pm 0.62)\).

The ratio between active and non active bacteria increased significantly with time in cultured bacteria (Analyse of variance, \(p < 0.01\) ) and evolved from 9.9 to 12.9 \% during the first five hours after thawing.

The number of Operational Taxon Units (OTU) found in cultures fluctuated between 56 and 75 \% of that found in natural sediment (Fig. 2). The resulting dendrogram of TGGE patterns for cultured and natural sediment samples displayed two clusters. These clusters, of similar community composition, were defined by 49 \% pattern similarity. Subculturing does not seem to affect community composition to a great extent (75 \% of similarity) and did not change bacterial diversity. Freezing process induced a decline of 25 \% in the diversity of bacteria and slightly affected the bacterial community composition (84 \% of similarity).
**Bacterial ingestion rates**

After grazing experiments with pre-labelled bacteria, frozen grazers (control) were systematically less $^{15}$N enriched than living ones for the three grazers types under study (Tab. 1). $^{15}$N concentration increased linearly in grazers according to incubation time (Fig. 3). This linearity pointed that ingestion rates were constant during incubation period studied: 2 hours for *H. ulvae* and 5 hours for the nematode community and *A. tepida*. Raw data used for ingestion rates calculations are presented in Tab. 1. The mud snail *H. ulvae* grazed 1149 (± 0.285) ngC ind$^{-1}$ h$^{-1}$, each nematode in the community grazed 0.027 (± 0.005) ngC ind$^{-1}$ h$^{-1}$ and the foraminifera *A. tepida* grazed 0.067 (± 0.013) ngC ind$^{-1}$ h$^{-1}$.

**Discussion**

**Discussion on methodology**

◆ Success of enrichment

Rinsing efficiency was tested by placing non enriched killed bacteria in enriched medium and by separating them from this medium by centrifugations. Those bacteria were poorly enriched in $^{15}$N, showing that the bacterial rinsing centrifugation process was efficient. Thus, $^{15}$N enrichment of bacteria was due to a bacterial assimilation and not to culture medium remaining between bacterial cells. This high bacterial rinsing efficiency is essential, since some grazers are able to consume directly DOM (Montagna and Bauer 1988) from the culture medium.

There is one disadvantage in using $^{15}$N instead of $^{13}$C enriched bacteria. As grazers contain more C than N, more biomass is required for isotopic measurements. However, the use of $^{15}$N avoids the decalcification step required by $^{13}$C and bias associated with this decalcification (Jacob et al. 2005). Moreover, grazing experiences are based on the assumption that isotopic composition of bacteria remains constant during the incubation period. The isotopic
composition of $^{13}$C enriched bacteria will vary quickly due principally to respiration loss and
to a lower degree to production of DOM (Ogawa et al. 2001; Kawasaki and Benner 2006).
The use of $^{15}$N permits to limit this respiration loss bias so isotopic composition of bacteria
remains more stable during incubation.

♦ Size of cultured bacteria

Discrimination of prey by grazers on the basis of size can influence the estimate of total
bacterivory. Bacterial selection according to size has been well documented in planktonic
protozoa (Pérez-Uz 1996; Hahn and Höfle 1999). Most protists graze preferentially on
medium-sized bacterial cells, grazing being less efficient with smaller and larger cells (review
in Hahn and Höfle 2001). The soil nematode, *Caenorhabditis elegans* feeds on bacteria
suspended in liquid and smaller bacteria are better food sources than larger ones for this
species (Avery and Shtonda 2003). Since, in our study, cultured and natural bacteria presented
a similar average size, it can be inferred that there is only a small bias if any due to cell size
selection by grazers.

♦ Activity of cultured bacteria

Few data are available on CTC activities of natural benthic bacteria for comparison with
our results. In superficial sediments from intertidal mudflats, 4 to 20% of benthic bacteria
were found to be active (van Duyl et al. 1999). Proctor and Souza (2001) found 9 to 10 %
active cells in river sediments and 25 % in intertidal sediments in the Gulf of Mexico.

Enriched bacteria with activity levels different from those of the natural community may
induce a bias if grazers select bacteria according to prey activity. Nematodes can discriminate
bacteria exhibiting different physiological or nutritional states (Grewal and Wright 1992).
*Pellioditis marina* do not assimilate heat-killed bacteria even though it feeds on live cells at
high rates (Moens 1999). In contrast, *Diplolaimelloides meyli* is more attracted by killed than
by live bacteria (Moens et al. 1999a). The foraminifera *Ammonia beccarii* collects dead and
living stained bacteria without discrimination (Langezaal et al. 2005).

Many existing methods to quantify bacterivory use bacterial communities with activity
levels different from those of the natural communities. For instance, labels directly added to
the sediment are only incorporated by the active fraction of the bacterial community: 100 %
of labelled bacteria are active. In contrast, prelabelled bacteria such as standard FLB are
generally dead (heat-killed): 0 % of labelled bacteria are active. With the method described in
our study, 10 % of labelled bacteria are active after thawing of frozen cultures (Fig. 1). This
activity is included in range found in natural environments (van Duyl et al. 1999; Proctor and
Souza 2001; Haglund et al. 2002). With the present method, grazers have the opportunity to
pick up active or inactive bacteria according to their preference like in the natural
environment.

- Diversity in the cultured bacteria

Subculturing of bacteria produces a final culture free of sediment. Freezing allows storage
of aliquots that may be enriched under standardised conditions at any time. The freezing step
induces small variations in the diversity of the bacterial community that must be nevertheless
kept in mind when using this method.

Grazers may be highly selective of prey species. To our knowledge, selection of bacteria
has never been observed for macrofauna but demonstrated for nematodes (Moens et al.
1999a) and foraminifera (Lee et al. 1966; Bernhard and Bowser 1992; Langezaal et al. 2005).

Nematodes used to be considered as generalist feeders, but they were recently shown to be
selective feeders exhibiting various preferences for algal and microbial prey. Their
reproduction rates differ according to the ingested strain of bacteria (Venette and Ferris 1998;
Blanc et al. 2006). Moens *et al.* (1999a) show that monhysterid nematodes are able to select
bacterial strain. Selection can be due to the bacterial size: filamentous bacteria escape uptake
by nematodes with small buccal cavities (Blanc et al. 2006). Moens et al. (1999a) consider
that the chemotaxic responses of nematodes to their bacterial prey may be due more to
chemical cues produced by the bacteria than to bacterial cell-wall structure that determine
their palatability. Nematodes are also able to significantly modify the composition of a
bacterial community by their species-specific bacterial food preferences (De Mesel et al.
2004).

The foraminifera *A. beccarii* distinguishes food and non-food particles during collection
(Langezaal et al. 2005). Two allogromiidae species (Foraminifera) have been shown to be non
selective grazers, actively harvesting bacterial biofilm from 3 different inocula (Bernhard and
Bowser 1992). Lee et al. (1966) found that most species of bacteria do not serve as food for
foraminifera whereas selected species of bacteria are consumed in large quantity.

Each bacterial species presents characteristics such as cell surface, nutritional quality or
chemical cues which may influence bacterial grazer behaviour. These differences have not
been evaluated between cultured versus natural bacteria in the present study. However,
estimation of total community composition and diversity gives us an approximate idea of
these differences. This molecular approach has the advantage to target dominant community
members. The cultured community presents 49 % of similarity with the natural bacterial
community. Although cultivation of natural bacteria induces a shift in community
composition (Fig. 2), this bacterial consortium seems more representative of the natural
community than that of many other grazing studies. The majority of experiments that use FLB
are done with monospecific bacteria or with a really limited number of bacterial species. Even
if natural and cultured community are not strictly identical, the probability for grazers to find
and ingest their preferred bacterial species is higher in the supplied bacterial consortium than
with monospecific bacteria.
Characteristics of grazing experiment

All various methods developed and applied to measure bacterivory in natural communities possess methodological shortcomings that make interpretation of the resulting data problematic. The method presented in this study, using \(^{15}\text{N}\) pre-enriched bacteria also presents bias. Sieving the sediment changes the bacterial availability for predators, bacteria being not attached to particle as in natural situation. The best way to minimize this artefact is to add the label directly to sediment in order to label bacteria while they are being grazed. This method is problematic as a high fraction of label found in grazers is due to processes other than grazing as underlined before. This requires control of incubations with a prokaryote activity inhibitor and the effectiveness of this inhibitor has to be tested for each grazer. The pre-enriched bacteria technique does not require the use of such inhibitors and only necessitate one control to determine adsorption of enriched bacteria on grazers.

Nematodes (Gerlach 1978), foraminifera from the genera *Ammonia* (Chandler 1989) and *H. ulvae* secrete mucus. During experiments with pre-enriched bacteria, controls must be performed to determine abundance of enriched bacteria stuck in the mucus secreted by grazers. Stuck bacteria modify the isotopic composition of grazers and controls are required to evaluate this bias due to non-grazing processes. In this study, freeze-killed grazer controls were used to determine this adsorption assuming that mucus post-freezing and mucus never frozen absorb bacteria at the same rate.

During grazing experiments, prey egestion from grazers may occur when chemical preservatives are used and this leads to underestimating grazing rates. Nematodes can egest a significant part of their gut contents when killed with formaldehyde (Moens et al. 1999b). In this study, grazers were frozen at -80°C to reduce this bias.

Prior to grazing experiments, enriched bacteria were frozen in liquid nitrogen to prevent spontaneous and enzymatic degradation and maintain their viability. Bacteria are commonly
cryopreserved during long-term storage in liquid nitrogen. Some agents like glycerol and methanol can be used to enhance cryopreservation, but in the case of grazing experiments, they can be toxic to grazers. Low-temperature storage enables standardised cultures, helping to ensure reproducible results in a series of experiments. This storage is really useful when monitoring over long periods is considered.

The method used allows short incubations that limit bias due to recycling. Bacterial ingestion is detectable after 2 hours of incubation for 3 grazers (Fig. 3). During incubation, labelled bacteria may be first ingested by grazers that are themselves preyed by studied grazers. In such a situation, it is impossible to determine the part of label present in studied grazers that is provided respectively by bacteria and first grazers. Even if a short incubation time does not prevent this type of bias, it reduces it substantially.

Demonstration of applicability

Data from literature to compare with our values are scarce. First, these predator species have not been systematically studied. Secondly, herbivory is more commonly studied than bacterivory. Thirdly, the manner in which to report grazing rates depends on the aim of the study. Studies dealing with carbon flow generally report values on a biomass basis (e.g. ngC ind\(^{-1}\) h\(^{-1}\)). When the aim of the study is the impact of grazers on microbial community, grazing rates are generally reported as rate constants (e.g. h\(^{-1}\)) (Montagna 1995).

While meiofaunal grazers are adapted to pick out specific microbial particles, macrofaunal deposit feeders process large volumes of sediment. Mud snails of the genus *Hydrobia* assimilate epipelagic diatoms and attached bacteria (Newell 1965; Kofoed 1975; Lopez and Levinton 1978; Jensen and Siegismund 1980) contained in the ingested sediment. To our knowledge bacterial ingestion rates have never been determined but data is available concerning algal ingestion rates. The bacterial ingestion rate found in our study is in the same
range as algal ingestion rates found by Forbes and Lopez (1989), Blanchard et al. (2000) and Habois et al., (2005) (Tab. 2).

In literature, grazing rates of nematodes are strongly variable with a range of fluctuations of more than two orders of magnitude (Tab. 2). Thus, comparison of our data with literature is difficult. Those discrepancies may arise from a lot of reasons such as the use of different techniques or the experimental conditions. When grazing experiments are performed in monoxenical conditions, nematodes are in an environment constituted by water (or agar) and bacteria. Nematodes would probably present higher grazing rates in such conditions than during grazing experiments where bacterial food is mixed with minerals and refractory organic matter and therefore is less available. However, when our results are compared to values resulting from experiments using nematodes from mudflat grazing on labelled bacteria mixed with sediment, it appeared that they are in the range of grazing rates found by Epstein & Shiariis (1992) but more than ten times lower than those found by Montagna (1984b).

Algal ingestion rates by *A. tepida* are higher (Moodley et al. 2000) than bacterial ingestion rates found in our study (Tab. 2). Langezaal et al. (2005) used FLB in simplified microcosms with one specimen of *A. beccarii* in a reduced volume of water. Their bacterial grazing rate is lower than ours. This may be linked to the bacterial concentration used in microcosms (1.4 *10³* cell ml⁻¹), which is considerably lower than benthic bacterial abundance in the natural environment (c.a. *10⁹* cell ml⁻¹) and in the present study.

**Conclusion**

The fate of benthic bacterial biomass is a topic of major importance in microbial ecology and in food web studies. All various methods developed and applied to measure bacterivory in natural communities possess artefacts and difficulties that make interpretation of the resulting data problematic. Our experimental approach is not an exception and also presents shortcomings. These bias are due principally to sediment manipulation. Labelled bacteria are
not available to bacterivores in the same manner as bacteria in unmanipulated sediment.

However, grazing experiments with $^{15}$N pre-enriched bacteria also present several advantages: (i) they can be performed *in situ* without environmental consequences, (ii) they do not require long incubations, so bias due to recycling is minimized, (iii) they require quite simple control tests with freezing of enriched prey, (iv) they can be performed at different times under standardised conditions, (v) they can be extended to other types of sediment or soil and (vi) they can be used in double-labelling experiments with $^{13}$C enriched algae, in order to simultaneously measure bacterial and algal ingestion rates.
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### Tables

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<td>δ¹⁵N living grazers</td>
<td>7068.2</td>
<td>95.0</td>
<td>20.6</td>
<td>20.0</td>
</tr>
<tr>
<td>δ¹⁵N dead grazers</td>
<td>10.7</td>
<td>11.1</td>
<td></td>
<td>16.7</td>
</tr>
<tr>
<td>Incubation time (h)</td>
<td>2</td>
<td>5</td>
<td>5</td>
<td></td>
</tr>
</tbody>
</table>

Ratio (enriched/non enriched bacteria) = 1.5

<table>
<thead>
<tr>
<th>Ingestion rate (ngC/ind/h)</th>
<th>1149.16</th>
<th>0.03</th>
<th>0.07</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ingestion rate (10⁻³ gC_{bacteria} / gC_{grazer}/h)</td>
<td>6.43</td>
<td>0.23</td>
<td>0.06</td>
</tr>
</tbody>
</table>

Table 1. Calculation of ingestions rates of three different grazers
<table>
<thead>
<tr>
<th>Grazers</th>
<th>Grazing rate (10^-3 gC_{bacteria} h^{-1} gC_{grazer})</th>
<th>Grazing rate (ngC h^{-1} ind^{-1})</th>
<th>Labelling method</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gastropoda</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>Hydrobia ulvae</td>
<td>6.43</td>
<td>1149.16</td>
<td>Stable isotope pre-enriched bacteria</td>
<td>Present study</td>
</tr>
<tr>
<td>Hydrobia ulvae</td>
<td>40-2080</td>
<td></td>
<td>Radioactive prelabelled algae</td>
<td>(Haubois et al. 2005)</td>
</tr>
<tr>
<td>Hydrobia ulvae</td>
<td>896-1064</td>
<td></td>
<td>Radioactive prelabelled algae</td>
<td>(Blanchard et al. 2000)</td>
</tr>
<tr>
<td>Hydrobia truncata</td>
<td>506-2873</td>
<td></td>
<td>Radioactive prelabelled algae</td>
<td>(Forbes and Lopez 1989)</td>
</tr>
<tr>
<td>Nematode</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Mudflat Nematode community</td>
<td>0.23</td>
<td>0.03</td>
<td>Stable isotope pre-enriched bacteria</td>
<td>Present study</td>
</tr>
<tr>
<td>Subtidal Nematode community</td>
<td>2.59-3.66</td>
<td></td>
<td>In situ radioactive labelled bacteria</td>
<td>(Montagna 1984b)</td>
</tr>
<tr>
<td>Subtidal Nematode community</td>
<td>0.01</td>
<td></td>
<td>In situ radioactive labelled bacteria</td>
<td>(Montagna et al. 1995)</td>
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<tr>
<td>Plectus palustris</td>
<td>10.54*</td>
<td></td>
<td>Radioactive prelabelled bacteria</td>
<td>(Duncan et al. 1974)</td>
</tr>
<tr>
<td>Monhystera disjuncta</td>
<td>0.15-0.49 *</td>
<td></td>
<td>Radioactive prelabelled bacteria</td>
<td>(Herman and Vranken 1988)</td>
</tr>
<tr>
<td>Diplolaimelloides meyli</td>
<td>11-17</td>
<td></td>
<td>Radioactive prelabelled bacteria</td>
<td>(Moens and Vinx 2000)</td>
</tr>
<tr>
<td>Pellioditis marina</td>
<td>55-60</td>
<td></td>
<td>Radioactive prelabelled bacteria</td>
<td>(Moens and Vinx 2000)</td>
</tr>
<tr>
<td>Mudflat Nematode community</td>
<td>0.02*</td>
<td></td>
<td>Fluorescent prelabelled bacteria</td>
<td>(Epstein and Shiaris 1992)</td>
</tr>
<tr>
<td>Foraminifera</td>
<td></td>
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<td></td>
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</tr>
<tr>
<td>Ammonia tepida</td>
<td>0.06</td>
<td>0.07</td>
<td>Stable isotope pre-enriched bacteria</td>
<td>Present study</td>
</tr>
<tr>
<td>Ammonia tepida</td>
<td>2.18</td>
<td>2.18</td>
<td>Stable isotope pre-enriched algae</td>
<td>(Moodley et al. 2000)</td>
</tr>
<tr>
<td>Ammonia beccarii</td>
<td>0.5 \times 10^{-4} - 1.6 \times 10^{-7} *</td>
<td></td>
<td>Fluorescent prelabelled bacteria</td>
<td>(Langezaal et al. 2005)</td>
</tr>
</tbody>
</table>

Table 2. Ingestion rate of bacteria observed in this study and compared with data from the literature concerning bacterivory and herbivory. *

ingestion rate converted with a bacterial biomass of 35 fgC cell^{-1} (Theil-Nielsen and Søndergaard 1998).
Figure captions

Fig 1. Evolution of the percentage of CTC + cells related to total bacteria after thawing cultured bacteria. Bars indicate standard deviation (n = 3).

Fig 2. TGGE analyses of natural sediment, first culture, subculture from the first culture and subculture from the first frozen culture. The right panel shows the relating band similarity (%) of bacterial communities.

Fig 3. Evolution of isotopic composition of three types of living (●) and dead (○) grazers placed in contact with $^{15}$N enriched bacteria. Bars indicate standard deviation (n = 3).
Figure 1.
Fig 2.
Fig 3.