Bacteria and Foraminifera: key players in a short-term deep-sea benthic response to phytodetritus

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ABSTRACT: The deep-sea floor has long been considered a ‘food desert’ but recent observations suggest that episodic inputs of relatively fresh organic matter (phytodetritus) occur and that benthic processing of this material may be rapid. Although the responses of the total community in terms of oxygen consumption and of some individual benthic groups have been identified, the quantitative role of the different groups in the short-term response remains largely unknown. We examined the short-term response in major benthic compartments in an in situ experiment in the NE Atlantic (2170 m water depth) using ¹³C-enriched diatoms as a tracer of labile carbon. Within 35 h, 6 mg C m⁻² was processed by the benthos, with the majority of the processed carbon recorded as respiration (45%). Among the fauna retained on a 300 µm sieve, Foraminifera were rapid consumers which, together with Bacteria, accounted for 50% of the processing. Therefore, although Bacteria dominate long-term carbon mineralization (as suggested by their general dominance in the benthic biomass), some faunal components, in this case Foraminifera, may play a central role in the rapid initial processing of fresh organic carbon in deep-sea sediments.

KEY WORDS: Deep-sea · Benthic-pelagic coupling · ¹³C labelling · Tracer · Respiration · Bacteria · PLFA · Foraminifera · Benthos

INTRODUCTION

The benthic processing of organic matter (OM) determines the amount and composition of organic carbon buried in marine sediments. A significant part of the carbon input to deep-sea sediments occurs in the form of short-lived deposition events of relatively fresh organic matter (Billett et al. 1983, Thiel et al. 1989) and a dynamic pelagic-benthic coupling may exist (Smith & Baldwin 1984, Graf 1989, Gooday & Turley 1990, Drazen et al. 1998). However, a benthic response is not always evident (Sayles et al. 1994, Ahrens et al. 1997, Pfannkuche et al. 1999, Smith & Kaufmann 1999) and may depend not only on the quality and quantity of the settling material (Sayles et al. 1994, Soetaert et al. 1996) but also on the timing of the observations (Pfannkuche et al. 1999). The benthic response in terms of sediment community oxygen consumption may be rapid and rather brief (Graf 1992) and therefore easily missed in field observations (Smith & Kaufmann 1999, Witte & Pfannkuche 2000). In terms of faunal response, the standing stock or biomass may not be a good indicator of direct processing of OM, as it reflects both the net outcome of individual OM uptake pathways and biological interactions (Pfannkuche et al. 1999). The use of tracers, however, allows a direct examination of deep-sea benthic responses. Pioneering studies revealed rapid assimilation of organic carbon by an undifferentiated micro-

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meiofaunal community (Cahet & Sibuet 1986, Cahet et al. 1990) and a rapid uptake of fresh phytodetritus by some individual benthic taxa (Blair et al. 1996, Levin et al. 1997, 1999). Similarly, the response of some individual groups has been examined in enrichment experiments with deep-sea sediments (e.g. Turley & Lochte 1990, Altenbach 1992, Linke 1992, Linke et al. 1995, Boetius & Lochte 1996, Relexans et al. 1996). Although several studies have demonstrated rapid deep-sea benthic response to fresh OM, the quantitative and relative role of the different benthic components in the short-term response remains largely unknown. For example, how does faunal uptake compare to that of the expected key player, Bacteria (Rowe & Deming 1985, Rowe et al. 1991, Relexans et al. 1996, Turley 2000)? In addition to the tracer recovered in benthic consumers, the algal carbon can also be immediately burnt for energy, with diatom carbon released as $^{13}$CO$_2$ (Levin et al. 1997); respiration therefore may be an important part of the benthic response. Recent studies have shown that the OM reaching the sediments may be very reactive (e.g. Soetaert et al. 1996, Drazen et al. 1998) and, as in shallow-water environments, deep-sea sediment communities may respond immediately to an input of fresh OM (Graf 1992, Levin et al. 1999).

We examined the role of the major benthic groups in the short-term processing of fresh OM in the deep sea off NW Spain (42°37.59' N, 10°00.04' W, 2170 m water depth, bottom-water characteristics: 3.6°C and 256 µM O$_2$). The experiment was conducted on 13 to 16 May 1999 in benthic chambers mounted on a free-falling benthic lander (ALBEX). There was no visible evidence of natural phytodetrital deposits at the time of the experiment, but inputs of fresh material have been documented in the general vicinity (e.g. Pfannkuche 1993, Rice et al. 1994). Axenic $^{13}$C-enriched Thalassiosira pseudonana, a common diatom in the study area (Estrada 1984), was used as a tracer of labile carbon. Axenic diatoms do not directly mimic the natural OM input to the benthos, but the axenic state is necessary when bacterial assimilation is examined through $^{13}$C enrichment of bacterial biomarkers (see ‘Materials and methods’). Blair et al. (1996) and Levin et al. (1997, 1999) have shown $^{13}$C-enriched phytodetritus to be a powerful tracer in evaluating the potential of the deep-sea benthos to process labile carbon. Bacterial assimilation of the added carbon was assessed via carbon-isotope analysis of polar lipid-derived fatty acids (PLFAs) specific for Bacteria (Boschker et al. 1998, Middelburg et al. 2000, Moodley et al. 2000), respiration of the added carbon was assessed through $^{13}$C-enrichment of $\Sigma$CO$_2$, and uptake by benthic fauna through $\delta^{13}$C$_{org}$ measurements of their body tissue.

**MATERIALS AND METHODS**

An axenic clone of the diatom Thalassiosira pseudonana (CCMP 1013, Provasoli-Guillard National Center for Culture of Marine Phytoplankton, Bigelow Laboratory) was cultured in artificial seawater containing 30% $^{13}$C-enriched bicarbonate that produced algal carbon consisting of 20.3 ± 1.8% $^{13}$C. Diatoms were concentrated by centrifugation, washed several times to remove the $^{13}$C-bicarbonate, and freeze-dried. The axenic state of the diatom culture was verified microscopically, by separate PLFA analysis of the freeze-dried diatoms and by a 0 h sediment incubation experiment.

A detailed description of the design and working of the ALBEX lander as well as methods of sediment-community oxygen consumption (SCOC) measurements are given in Witbaard et al. (2000). One background SCOC measurement was done during the test deployment of the lander just prior to the uptake experiment. The $^{13}$C-uptake experiment was conducted during a single deployment, with the lander holding 3 benthic chambers; each chamber measured 12 × 12 × 20 cm and enclosed a water and sediment column of ~10 cm each; 1 chamber was used for background measurements (ALBEX 1) and the other 2 as experimental chambers (ALBEX 2 and ALBEX 3) which received a pulse of $^{13}$C-enriched diatoms. Diatoms (23.8 mg corresponding to 6.3 or 434 mg C m$^{-3}$) were mixed with 0.2 µm-filtered seawater and placed in a syringe. The syringe was placed upside down (to prevent clogging) and programmed to first extract water in order to re-suspend the diatoms before injection into the chamber. Excess water due to injection of diatom mix was expelled through an exit valve at the opposite end of the lid in order to limit the loss of diatoms during injection. The stirrers were switched on 1 h after diatom injection and the systems were preset to take water samples for CO$_2$ after 15 and 29 h. A final water sample was taken on deck (35 h after diatom injection). At the end of the experiment, the chambers were processed (within 2 h) in a temperature-controlled container (4°C). Two sub-cores (3.6 cm inner diameter) were taken: 1 for PLFA and bulk POC and 1 for meiofauna >63 µm. We chose a 63 µm sieve in preference to the 38 or 45 µm generally used in meiofauna studies in order to facilitate the isolation and hand-picking of the small organisms and a better removal of diatoms possibly attached to the outside of organisms. Three intervals of the remaining sediment (0–1, 1–3 and 3–5 cm) were sieved on a 300 µm sieve for the larger fauna and frozen in glass bottles. Megafauna were not present in our benthic chambers.

In the laboratory, fauna sediment samples were thawed and all organisms handpicked. The large
fauna (retained on the 300 µm sieve) were isolated from different depth intervals but, due to high heterogeneity and low biomass, were combined for measurements. For this fauna, 4 groups were distinguished: (1) metazoan macrofaunal taxa (e.g. polychaetes, bivalves, crustaceans, and sipunculans); (2) metazoan meiofaunal taxa (taxa generally belonging to the meiobenthos, e.g. nematodes, copepods and ostracods); (3) Foraminifera (Protista) that can constitute a major part of the benthos, generally of the meiobenthic size class Foraminifera A (calcareous, agglutinated and soft-shelled); and (4) the group of less ‘familiar’ large agglutinated deep-sea Foraminifera, Foraminifera B (astrophizids and komokiaceans: Gooday et al. 1992).

The criterion for living Foraminifera was healthy-looking (as opposed to old, fossilised), cytoplasm-filled shells. In cases where this was not easily established, specimens were broken and examined for cytoplasm. However, for komokiacean Foraminifera, the living could not be distinguished from the dead. In the case of large agglutinated astrophizid Foraminifera, all specimens were broken; some specimens did not contain foraminiferal cytoplasm but were inhabited by polychaetes or isopods that were transferred to the metazoan macrofaunal taxa group. For the group of metazoan macrofaunal taxa and 1 genus of the Foraminifera B group (Rhizammina spp.), samples were first lyophilized and homogenised. For the other faunal groups, whole specimens were combined for measurements. Each sample was then treated with 5% HCl in silver boats and the excess acid was removed by drying overnight at 40°C.

Lipid extraction (from 3 g of lyophilized sediment per sediment interval) and carbon-isotopic analysis of PLFAs were done according to Boschker et al. (1999) and Middelburg et al. (2000). Details of the measurement of δ13C–ΣCO2 and δ13Corg of faunal compartments are given in Moodley et al. (2000).

Carbon isotopes are expressed in the delta notation (δ13C) relative to Vienna Pee Dee Belemnite (VPDB): 
\[ \delta^{13}C = \left( \frac{^{13}C/^{12}C}_{\text{sample}} / ^{13}C/^{12}C_{\text{VPDB}} - 1 \right) \times 1000 \]
Incorporation of 13C is reflected as excess (above background) 13C and is expressed in terms of specific uptake (i.e. δΔ13C = δ13Csample – δ13Cbackground) as well as total uptake (I). Specific uptake, as evident in Δδ13C (Δδ13C = 0 in the case of no uptake) is a clear indication of uptake of the 13C-enriched diatom, but remains a qualitative measure, as it is based on isotope ratio data and is influenced by the amount incorporated, the pool size, and the percentage of 13C in the diatoms (i.e. the intensity of the original labelling). Total uptake is a quantitative measure of 13C-uptake and is calculated as the product of excess 13C (E) and biomass of the consumers or concentration of CO2. Excess (E) 13C is the difference between the fraction (F) 13C of the background (Fbackground) and the samples (Fsample): 
\[ E = F_{\text{sample}} - F_{\text{background}} \text{ where } F = \frac{^{13}C/(^{13}C+^{12}C)}{R/(R+1)} \]
The carbon isotope ratio (R) was derived from the measured δ13C values as: 
\[ R = \delta^{13}C / 1000 + 1 \times R_{\text{VPDB}}, \text{ with } R_{\text{VPDB}} = 0.0112372 \]
For the Fbackground of CO2, we did not use the δ13C – ΣCO2 of the control chamber (average + 1.03‰) but that determined in a laboratory 35 h incubation with labelled diatoms using sterile sediment and 0.2 µm filtered seawater, which revealed a constant enrichment in 13CO2 equivalent to a δ13C – ΣCO2 of + 4.44‰. This reflects traces of 13C-bicarbonate in the added diatoms in spite of extensive rinsing with clean medium during diatom harvesting. The biomass of fauna is generally calculated from weight (wet or dry wt) or from size dimensions converted to carbon content. We calculated the organic carbon content (biomass) of the different faunal groups directly from the area counts given in the standard output of the isotope ratio mass spectrometer calibrated with sucrose (0 to 84 µgC, r² = 0.98). This method overcomes the need for separate biomass estimation or measurements and offers a direct and accurate calculation of carbon uptake and biomass in the same measurement. Bacterial data are based on the concentrations and 13C content of Bacteria-specific biomarkers (14:0 iso, 15:0 iso, 15:0 anteiso, 16:0 iso) assuming that bacterial biomarkers = 14% of total bacterial PLFAs (calculated from Rajendran et al. 1993, 1994, Guezennece & Fialamedioni 1996, Stewart et al. 1996, Boschker et al. 1998) and 0.056 g C PLFA g C⁻¹ biomass (Brinch-Iversen & King 1990), as described in Middelburg et al. (2000).

Finally, the uptake of total added algal carbon (12C + 13C) was calculated as the quotient of total uptake (l) and the fractional abundance of 13C in the algae (0.203). All error terms presented in the text are standard errors.

RESULTS AND DISCUSSION

Approximately 229 ± 74 mg C m⁻² of labelled diatom carbon was recovered in the upper 5 cm of the sediment column as bulk POC (calculated from the excess PO13C and POC content) indicating that ~40% of the diatoms was lost during deployment or injection. The amount of recovered diatom carbon represented 0.7 ± 0.1% of the natural particulate organic carbon in the 0 to 1 cm interval of the sediment (33.5 ± 5.4 g C m⁻²).

Background sediment community oxygen consumption measured during the test deployment (0.53 mmol O₂ m⁻² d⁻¹) was similar to that measured in the control (ALBEX 1) benthic chamber (0.48 mmolO₂ m⁻² d⁻¹; Table 1). The SCOC measured in the experimental chambers was, on average, twice that of the back-
ground value (Table 1), but because of limited replication this difference has low significance. However, independent of significant changes in SCOC, 13C-enrichment in \( \Sigma \)CO2 is a sensitive tool for detecting respiration of the pre-labelled added carbon (see also Blair et al. 1996). In our \textit{in situ} experiment, a rapid benthic response was recorded in the respiration of the added carbon, as reflected in a linear increase in excess \(^{13}\)CO\(_2\) during the time period measured (Fig. 1). Quantitatively, within 35 h, 6.3 \pm 0.6 mg m\(^{-2}\) of the added carbon was processed by the benthic compartments examined, i.e. Bacteria in the upper 5 cm, fauna of the upper 5 cm retained on a 300 \(\mu\)m sieve, and total community respiration. The role of the smaller fauna (meiobenthos retained on a 63 \(\mu\)m sieve) will be discussed separately because 1 replicate (ALBEX 2) was lost during processing. The major signal of benthic processing was recorded as CO\(_2\) (45.6 \pm 2.6\%).

Respiration of the added carbon represented 16.4 \pm 2.6\% of total respiration (calculated from sediment oxygen consumption using a respiratory quotient of 1; Table 1). This supports previous evidence that benthic processing of labile carbon may be rapid in deep-sea sediments under well-oxygenated conditions (e.g. Graf 1992, Blair et al. 1996, Levin et al. 1997, 1999). Bacterial assimilation accounted for 21.7 \pm 0.6\% of the processed carbon. Among the fauna retained on a 300 \(\mu\)m sieve, metazoan meiofaunal taxa took up <0.1\%, metazoan macrofaunal taxa ingested 3.5 \pm 1.2\%, and 29.1 \pm 2.0\% of the processed carbon was recovered in Foraminifera (Table 1). The ultimate long-term fate of OM escaping burial is respiration to CO\(_2\) (e.g. Rowe et al. 1991). It is evident that respiration may be a primary sink of labile OM also in the very short term (hours to days). Similarly, in shallow marine environments, CO\(_2\) was found to be a major sink of organic matter (e.g. Olofsson et al. 1999, Moodley et al. 2000). A transient fate of OM reaching the seafloor is benthic biomass, which in the deep sea is primarily dominated by Bacteria (Rowe & Deming 1985, Rowe et al. 1991, Relexans et al. 1996, Turley 2000). Bacteria account for >95\% of the biomass standing stock at our study site (Table 1), and accordingly Bacteria are key players in processing the added carbon (Fig. 2). It is important to note that incorporation of phytodetritus-derived \(^{13}\)C in bacterial PLFA represents assimilation only; another part of bacterial uptake has been respired and is probably the main contributor to total community respiration (Table 1).

Among the fauna, a surprisingly large fraction of the processed carbon was recovered in Foraminifera (Table 1). The ultimate fate of OM escaping burial is respiration to CO\(_2\) (e.g. Rowe et al. 1991). It is evident that respiration may be a primary sink of labile OM also in the very short term (hours to days). Similarly, in shallow marine environments, CO\(_2\) was found to be a major sink of organic matter (e.g. Olofsson et al. 1999, Moodley et al. 2000). A transient fate of OM reaching the seafloor is benthic biomass, which in the deep sea is primarily dominated by Bacteria (Rowe & Deming 1985, Rowe et al. 1991, Relexans et al. 1996, Turley 2000). Bacteria account for >95\% of the biomass standing stock at our study site (Table 1), and accordingly Bacteria are key players in processing the added carbon (Fig. 2). It is important to note that incorporation of phytodetritus-derived \(^{13}\)C in bacterial PLFA represents assimilation only; another part of bacterial uptake has been respired and is probably the main contributor to total community respiration (Table 1).

Among the fauna, a surprisingly large fraction of the processed carbon was recovered in Foraminifera (Fig. 2). The highest \(^{13}\)C-enrichment was also found in Foraminifera (Table 1). With uptake experiments, there is always the possibility of artefacts due to adhesion of

**Table 1. Weighted average \(^{13}\)C signatures (‰), biomass and carbon uptake (mg C m\(^{-2}\)) of different benthic components in the control benthic chamber (ALBEX 1) and 2 experimental benthic chambers (ALBEX 2 and ALBEX 3). Values are for 0 to 5 cm sediment depth, except for meiofauna >63 \(\mu\)m (0 to 1 cm interval). SCOC: sediment-community oxygen consumption; PLFAs: polar lipid-derived fatty acids**

<table>
<thead>
<tr>
<th>Parameter</th>
<th>ALBEX 1</th>
<th>ALBEX 2</th>
<th>ALBEX 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>SCOC (mmol O(_2) m(^{-2}) d(^{-1}))</td>
<td>0.48</td>
<td>0.74</td>
<td>1.37</td>
</tr>
<tr>
<td>Respiration of added carbon(^a) (mg C m(^{-2}) 35 h(^{-1}))</td>
<td>–</td>
<td>2.47</td>
<td>3.31</td>
</tr>
<tr>
<td>(^{13})C</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bacterial biomarkers(^b)</td>
<td>–24.4</td>
<td>–11.5</td>
<td>–13.9</td>
</tr>
<tr>
<td>Foraminifera A(^c)</td>
<td>–21.4</td>
<td>+2.952</td>
<td>+3.993</td>
</tr>
<tr>
<td>Foraminifera B(^d)</td>
<td>–22.1</td>
<td>+518</td>
<td>+100</td>
</tr>
<tr>
<td>Metazoan meiofauna(^e)</td>
<td>–19.0</td>
<td>–12.63</td>
<td>–8.6</td>
</tr>
<tr>
<td>Macrofauna(^f)</td>
<td>–20.5</td>
<td>+121</td>
<td>+53.5</td>
</tr>
<tr>
<td>Meiofauna &gt;63 (\mu)m(^g)</td>
<td>–19.7</td>
<td>Lost</td>
<td>+3.254</td>
</tr>
</tbody>
</table>

**Biomass (C uptake)**

<table>
<thead>
<tr>
<th></th>
<th>ALBEX 1</th>
<th>ALBEX 2</th>
<th>ALBEX 3</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bacteria</td>
<td>1.360</td>
<td>1.700 (1.21)</td>
<td>1.570 (1.53)</td>
</tr>
<tr>
<td>Foraminifera A(^c)</td>
<td>9.9</td>
<td>8.2 (1.29)</td>
<td>8.4 (1.74)</td>
</tr>
<tr>
<td>Foraminifera B(^d)</td>
<td>9.4</td>
<td>13.9 (0.49)</td>
<td>15.9 (0.11)</td>
</tr>
<tr>
<td>Metazoan meiofauna(^e)</td>
<td>5.2</td>
<td>7.6 (&lt;0.01)</td>
<td>6.4 (&lt;0.01)</td>
</tr>
<tr>
<td>Macrofauna(^f)</td>
<td>44.1</td>
<td>35.8 (0.27)</td>
<td>39.1 (0.16)</td>
</tr>
<tr>
<td>Meiofauna &gt;63 (\mu)m(^g)</td>
<td>11.8</td>
<td>Lost</td>
<td>22.5 (3.84)</td>
</tr>
</tbody>
</table>

\(^a\)Based on excess \(^{13}\)CO\(_2\) and CO\(_2\) concentration at the end of the incubation  
\(^b\)Depth average of PLFAs 14:0 iso, 15:0 iso, 15:0 anteiso and 16:0 iso  
\(^c\)Group of ‘familiar’ Foraminifera  
\(^d\)Group of less ‘familiar’ large agglutinated Foraminifera  
\(^e\)Metazoan meiofaunal taxa  
\(^f\)Bulk measurement of macrofaunal taxa retained on 300 \(\mu\)m sieve  
\(^g\)Metazoan meiofaunal taxa plus Foraminifera but excluding group of less ‘familiar’ large agglutinated Foraminifera

**Fig. 1.** \(^{13}\)C-enrichment of total inorganic carbon (\(\Delta\delta\Sigma^{13}\)CO\(_2\)) versus time. Values from experimental chambers, ALBEX 2 (○) and ALBEX 3 (●)
the added algal material onto the exterior of organisms (Levin et al. 1999). However, a 6 h on-deck incubation revealed limited $^{13}$C-enrichment in both the group of ‘familiar’ Foraminifera as well as the group of large agglutinated Foraminifera (average $\Delta^{13}$C +15.4‰) compared to an average $\Delta^{13}$C of +3494 and +331‰ respectively for the 2 groups after 35 h in situ. This large difference clearly indicates that the high $^{13}$C-enrichment recorded in the Foraminifera after 35 h in situ is not an artefact of adhesion but reflects active uptake. Additionally, for a group of Foraminifera in which maximum adhesion may be expected (Rhizammina spp., due their coarse agglutinated branching shells), $^{13}$C-enrichment was very limited (an average $\Delta^{13}$C of +12.3‰). This demonstrates that our sieving and cleaning procedures were efficient. Additionally, other studies have also demonstrated that benthic organisms, including Foraminifera, do indeed ingest and assimilate labelled diatoms very rapidly (e.g. Blair et al. 1996, Levin et al. 1997, 1999, Herman et al. 2000, Middelburg et al. 2000, Moodley et al. 2000). Foraminiferal ingestion of algae under high pressure and low temperature has also been documented (Turley et al. 1993).

The total biomass of Foraminifera is negligible compared to that of Bacteria (Table 1), but uptake of added carbon is more in the range of Bacteria than other faunal groups. Apart from a possible preference against fresh OM, the differential short-term uptake among the faunal groups (Fig. 2) reflects primarily their efficiency in acquiring food particles. Foraminifera possess granuloreticulate pseudopodia that can form long, extensive networks (Travis & Bowser 1991) that constitute an efficient system for trapping, gathering and accumulating food particles (Bowser et al. 1992, Gooday et al. 1996; present authors, pers. obs.). Foraminifera are evidently important consumers of fresh organic matter but their relatively low biomass suggests that they are probably more dependent on fresh material, presumably of limited input, than are Bacteria. Additionally, Foraminifera may be important links in the benthic food chain in the deep sea. In 1 of the experimental chambers, the gut of a scaphopod was found packed with calcareous Foraminifera, and there are numerous reports of predation on Foraminifera by a wide variety of organisms (e.g. Ahrens et al. 1997, Gudmundsson et al. 2000), including scaphopods (Davies 1987 in Gooday et al. 1992). The high $^{13}$C-enrichment found in Foraminifera, especially in the ‘familiar’ group (average $\Delta^{13}$C +3,421‰), suggests that they are important packaging agents of fresh OM, and that subsequent predation results in efficient trophic transfer. In contrast, Bacteria had relatively low $^{13}$C-enrichment (average $\Delta^{13}$C +11.7‰), indicating either that the bacterial community is slow-growing or that only a fraction of the bacterial community is involved in the short-term response. However, as noted above, a major part of community respiration may be attributable to bacterial processing of the added carbon. On the basis of bacterial assimilation (A) and total community respiration (R), bacterial assimilation efficiency (A/[A+R]) is >0.32, consistent with values reported for oxic shallow marine and lake environments (del Giorgio & Cole 1998) as well as for other deep-sea sediment bacterial assemblages (Boetius et al. 2000).

Among the fauna retained on a 300 µm sieve, the uptake by the macrofaunal taxa was relatively low and that of metazoan meiofaunal taxa was extremely limited (Fig. 2). However, the role of meiofauna is by definition underestimated when using a 300 µm sieve. We examined surface sediments (0 to 1 cm) using a sieve with a smaller mesh size (63 µm) excluding the group of large agglutinated Foraminifera. The meiofauna assemblage retained on the 63 µm sieve consisted primarily of Foraminifera and a strong enrichment in $^{13}$Corg was evident (Table 1). The numbers of nematodes may have been larger if a finer sieve had been used, but this is not expected to have major consequences for biomass or uptake trends. Other studies have recorded limited short-term uptake of diatoms by nematodes (Levin et al. 1999, Olafsson et al. 1999, Middelburg et al. 2000). It is unfortunate that the number of metazoan meiofauna was too low to measure separately. When the uptake in the smaller size fraction is taken into account, then the amount of added carbon processed in ALBEX 3 (the meiobenthos sample of ALBEX 2 was lost during processing) increases from 6.9 to 9.0 mg, with 36.9% recovered as CO₂, 17.1% assimilated by Bacteria, 42.9% taken up by the meiofauna retained on a 63 µm sieve (primarily Foraminifera), 1.7% ingested by the macrofauna and 1.2% taken up by the group of less ‘familiar’ large agglutinated Foraminifera retained on a 300 µm sieve. Bacteria (assimilation and respiration) and Fora-

Fig. 2. Proportional division of biomass and processed carbon among the different benthic compartments. Fauna are those retained on a 300 µm sieve and values are average of the 2 experimental chambers.
minifera were evidently the key players in this short-
term benthic response to an input of fresh material. 
This indicates that small-sized organisms, in particular
Bacteria, play a dominant role in carbon-processing 
(Pfannkuche 1993). Although Foraminifera were clearly
important competitors of Bacteria at this deep-sea site,
other faunal groups may out-compete Foraminifera
under different environmental settings. In a conti-
nental slope setting (850 m depth) supporting a rich
benthic community, some species of deposit-feeding
polychaetes exhibited relatively strong 13C-enrich-
ment, indicating rapid ingestion of algal carbon, but at
a relatively organic carbon-poor site, phytodetritus up-
take by large agglutinated Foraminifera was more pro-
nounced (Levin et al. 1999). Earlier studies have also
highlighted that Foraminifera may be rapid and impor-
tant consumers of phytodetritus (Meyer-Reil & Koster 
kuch 1993, Graf & Linke 1992, Linke et al. 1995,
Drazen et al. 1998). How this quantitatively compares
to bacterial assimilation and uptake by other faunal
groups in the different environmental settings remains
to be established. Although this study demonstrates
quantitatively that Foraminifera play an important role
in benthic carbon cycling, their relative role can be in-
fluenced by several factors: e.g. the history and quality
of the food supply and the size (and structure) of the
benthic community. The high dominance of bacterial
biomass suggests that Bacteria dominate long-term
carbon mineralisation and are probably, in contrast to
faunal components, less dependent on highly reactive
carbon. Specific studies are required to elucidate
whether Bacteria are sinks or links of organic carbon.

The ability of the deep-sea benthic community to
respond rapidly is clear from the processing of 6 to
9 mg C m⁻² within 35 h. This, however, still represents
a small fraction of the available carbon. Even if we
double the total faunal uptake to take into account
other protists (e.g. flagellates and ciliates) and smaller
meiofauna in the deeper layers, a maximum of ~13 mg
(6%) of the added carbon would have been processed.
In contrast, in a vastly different marine environment,
for an intertidal sediment community incubated at
16°C, respiration alone accounted for the processing of
~61 mg (12%) of the added diatom carbon within 35 h
(authors’ unpubl. data). This suggests that the prevail-
ing deep-sea conditions of low temperatures in combi-
nation with low biomass concentrations and composi-
tion limits and slows down the recycling of organic
matter, at least in the short term.

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LITERATURE CITED
Ahrens MJ, Graf G, Altenbach AV (1997) Spatial and tempo-
ral distribution patterns of benthic Foraminifera in the
Northeast Water Polynya, Greenland. J Mar Syst 10:
445–465
Altenbach AV (1992) Short-term processes and patterns in the
foraminiferal response to organic flux rates. Mar Micro-
 paleontol 19:119–129
Billett DSM, Lampitt RS, Mantoura RFC (1983) Seasonal
sedimentation of phytoplankton to the deep-sea benthos.
Nature 302:520–522
Blair NE, Levin LA, DeMaster DJ, Plaia G (1996) The short-
term fate of fresh algal carbon in continental slope sedi-
ments. Limnol Oceanogr 41:1208–1219
hydrobiontic potentials and growth of bacteria in deep-sea
sediments of the deep Arabian Sea in relation to vertical
Boschker HTS, Nold SC, Wellsbury P, Bos D, de Graaf W, Pel
R, Parkes RJ, Cappenberg TE (1998) Direct linking of mi-
crobial populations to specific biogeochemical processes
Boschker HTS, de Brouwer JFC, Cappenberg TE (1999) The
contribution of macrophyte derived organic matter in
microbial biomass in salt marsh sediments: stable carbon-
isotope analysis of microbial biomarkers. Limnol Oce-
anogr 44:309–319
Bowser SS, Alexander SP, Stockton WL, DeLaca TE (1992)
Extracellular matrix augments mechanical properties of
pseudopodia in the carnivorous foraminifera Astram-
Brinch-Iversen J, King GM (1990) Effects of substrate concen-
tration, growth rate, and oxygen availability on relation-
ships among bacterial carbon, nitrogen and phospholipid
phosphorus content. FEMS Microbiol Ecol 74:345–356
Cahet G, Sibuet M (1986) Activité biologique en domaine pro-
fond: transformations biochimiques in situ de composés
organiques marqués au carbone-14 à l’interface eau-
sédiment par 2000 m de profondeur dans le Golfe de
Cahet G, Daumas R, Sibuet M (1990) In situ experimentation
at the water/sediment interface in the deep sea. 2. Bio-
transformation of dissolved organic substrates by the
microbial communities at 2000 m depth in the Bay of Bis-
cay. Prog Oceanogr 24:169–178
del Giorgio PA, Cole JJ (1998) Bacterial growth efficiency in
Drazen JC, Baldwin RJ, Smith KL Jr (1998) Sediment commu-
nity response to a varying food supply at an abyssal station
Estrada M (1984) Phytoplankton distribution and composi-
tion off the coast of Galicia (northwest of Spain). J Plankton
Res 6:417–434
Gooday AJ, Turley CM (1990) Responses of benthic organ-
isms to inputs of organic material to the ocean floor. Phil


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