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Tracing carbon flow from microphytobenthos to major bacterial groups in an intertidal marine sediment by using an in situ $^{13}$C pulse-chase method

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Abstract

Carbon flow from benthic diatoms to heterotrophic bacterial was traced in an intertidal sediment for 5 consecutive days. $^{13}$C-labeled bicarbonate was sprayed onto the sediment surface during low tide and $^{13}$C-label incorporation in major carbon pools, intermediate metabolites, and biomarkers were monitored. Phospholipid-derived fatty acid (PLFA) and ribosomal ribonucleic acid (rRNA) were used to identify the responsible members of the microbial community at class and family phylogenetic resolution. Diatoms were the predominant primary producers, and Gammaproteobacteria, Bacteroidetes, and Deltaproteobacteria were major heterotrophic bacterial groups. Both $^{13}$C-PLFA and $^{13}$C-rRNA data suggest a fast transfer of label from diatoms (60 nmol $^{13}$C g$^{-1}$ dry weight [dry wt]) to bacteria (7 nmol $^{13}$C g$^{-1}$ dry wt) during the first 24 h, which was probably due to the exudation of low-molecular-weight organic compounds by diatoms that could be directly utilized by bacteria. After this initial fast transfer, labeling of bacteria proceeded at a slower rate to 13 nmol $^{13}$C g$^{-1}$ dry wt on the third day of the experiment, which coincided with the degradation of carbohydrates in water-extractable extracellular polymeric substances (EPS) initially produced by the diatoms. Water-extractable EPS (primarily as glucose) was a major intermediate and its turnover explained 75% of the total carbohydrate processing in the sediment. Labeling in bacteria tracked labeling in the diatoms, suggesting a closely coupled system. The heterotrophic bacterial groups benefited equally from the organic matter released by the diatoms, suggesting limited specialization in this microbial food web.

Marine intertidal areas are highly productive ecosystems, and microphytobenthos contributes importantly to the total primary production of estuaries and other shallow-water coastal ecosystems (MacIntyre et al. 1996; Underwood and Kromkamp 1999). Benthic diatoms are typically the dominant microphytobenthos in marine intertidal sediments from temperate regions (Underwood and Paterson 2003), and they are known to exude large amounts of carbohydrates such as extracellular polymeric substances (EPS). Some, but not all, of the EPS is released as a consequence of the motility of the diatoms and their migration through the sediment. As a result of the EPS production, diatoms stabilize the sediment surface and avoid their re-suspension in the water column (Paterson and Black 1999). Hence, diatoms provide a major carbon source to the benthic food web, which includes heterotrophic bacteria in marine intertidal sediments (Smith and Underwood 1998; van Oevelen et al. 2006).

Diatoms exude different types of carbohydrates, which vary in structure and composition depending on the environmental conditions and the nutrient status (Smith and Underwood 2000; de Brouwer and Stal 2001). Low-molecular-weight compounds released by diatoms can be directly utilized by the heterotrophic microbial community (Sundh 1992), whereas high-molecular-weight compounds such as EPS first need to be hydrolyzed by extracellular enzymes produced by microorganisms (Fuchs et al. 1998; Hunter et al. 2006). The microbial community composition of intertidal sediments has been extensively studied (Rusch et al. 2003; Bühring et al. 2005; Hunter et al. 2006) as is the case for the EPS formation by benthic diatoms (Smith and Underwood 1998; Haynes et al. 2007). However, there are only a limited number of studies that link in situ carbon flow from diatoms directly to heterotrophic microbes and also identify in detail the responsible microorganisms (Taylor et al. 2013).

Middelburg et al. (2000) and Evrard et al. (2008) have reported rapid transfer of carbon from microphytobenthos to bacterial biomass by combining in situ $^{13}$C pulse-chase labeling and phospholipid-derived fatty acid (PLFA) biomarker analysis in order to differentiate between labeling in algae and heterotrophic bacteria. Previous studies suggested that EPS-derived carbohydrates were major intermediates in the transfer of organic matter between benthic diatoms and heterotrophic bacteria (Bellinger et al. 2009; Oakes et al. 2010; Taylor et al. 2013). Although it is possible to estimate the biomass based on the abundance of distinctive PLFAs (Middelburg et al. 2000), the phylogenetic resolution of these biomarkers is limited and the bacteria utilizing the organic matter produced by the benthic diatoms cannot be identified to sufficient detail (Boschker and Middelburg 2002). In general, 16S ribosomal ribonucleic acid (rRNA) provides a much higher phylogenetic resolution than is possible with

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PLFA, and it is widely used to identify microbial communities in natural environments. MacGregor et al. (2002) developed a method where 16S rRNA from defined phylogenetic groups was isolated by using specific oligonucleotide probes attached to paramagnetic beads for subsequent stable isotope analysis. Miyatake et al. (2009, 2013) further improved the stable isotope probing with magnetic bead capture hybridization (Mag-SIP) and demonstrated the utilization of organic substrates in marine sediments by certain microbial groups at the family level.

The in-situ $^{13}$C pulse-chase method was used to trace the carbon flow in an intertidal benthic diatom mat for 5 d. In situ stable-isotope labeling approaches have the advantage over laboratory-based incubations that environmental conditions such as waves, tidal currents, sediment mixing, and pore-water flow are not disturbed (Middelburg et al. 2000). However, the inevitable loss of label that occurs in an open system could complicate quantitative interpretation of the data. We used 16S rRNA (Mag-SIP) and PLFA biomarkers for the identification of the major active microbial groups involved in the carbon transfer and to quantify their $^{13}$C-label incorporation rates. Furthermore, we used liquid chromatography combined with isotope ratio mass spectrometry (LC–IRMS) to trace $^{13}$C in intermediate metabolites such as the total carbohydrates in the sediment (bulk carbohydrates), water- and ethylenediamine-tetra-acetate (EDTA)–extractable carbohydrates, and short-chain organic acids (SCOA) including volatile fatty acids. Although other extraction schemes for carbohydrate extract (Underwood et al. 1995), we studied bulk carbohydrates as well as water- and EDTA-extractable carbohydrates, which are two fractions that are commonly used to characterize loosely and stronger bound extracellular carbohydrates, respectively, in benthic diatom mats (Underwood et al. 1995; de Brouwer and Stal 2001). A clone library derived from reverse-transcribed 16S rRNA was also constructed in order to determine the metabolically active microorganisms and relate this to the specific oligonucleotide probe set used in the Mag-SIP protocol.

In the present study, we followed the carbon flow through intermediates produced by the diatoms and identify water-extractable carbohydrates as a major intermediate. We also show that all members of the heterotrophic bacterial community that we took into account in our analyses appeared to be equally involved in the assimilation of organic matter produced by the diatoms, suggesting little specialization at the taxonomic level of this study.

Methods

Study site, in situ $^{13}$C labeling, and sampling—From 14 to 19 April 2009, an in situ $^{13}$C-labeling experiment was performed at an intertidal flat in the Zandkreek area (51°33′N, 3°53′E) of the Oosterschelde bay (The Netherlands). The sampling site was located 0.15 m below the mean tidal level, and the exposed period was ~6 h per tidal cycle (Fig. 1A). The sediment contained 6.8% silt (<63 μm particle size) and was covered by a diatom mat. Salinity and water temperature were 28.5° and 15°C, respectively, and were constant during the experimental period. The weather was mostly sunny with high light levels (photon irradiance up to 1500 μmol m⁻² s⁻¹; Fig. 1A) and an average mean air temperature of 16.2°C. Almost no rainfall occurred during the experimental period except for some minor drizzle on the third day.

During the first day of the experiment, shortly after the exposure of the site at low tide, two 50 × 50 cm stainless frames were inserted into the sediment to a depth of 8 cm in order to constrain the labeling and sampling area (the top rim of the frames was flush with the sediment surface). Initially, unlabeled control sediment and pore water ($n = 2$) were taken from just outside the frames as described below. The in situ labeling experiment was started by spraying the surface of the sediment within each frame with 250 mL of $[^{13}$C] sodium bicarbonate (99% $^{13}$C; Cambridge Isotope Laboratories). The label solution was at the ambient salinity and contained 1 g L⁻¹ of $[^{13}$C] sodium bicarbonate (equaled to 1 g $[^{13}$C] sodium bicarbonate m⁻²; Middelburg et al. 2000). The frames were divided in a 10 × 10 cm sampling grid. The first sampling of the labeled sediment was performed after 4 h at the end of the same exposed period (the pulse-labeling period), and the two frames were sampled in a time-course of 12 h, 1 d, 2 d, 3 d, and 5 d (the chase period).

The two frames were treated as duplicates ($n = 2$). At each sampling time, pore-water and sediment samples were collected and mixed from two randomly chosen positions within the sampling grid of each frame. Pore water (two times 1 mL) was sampled with porous polymer sippers (Rhizon Soil Moisture Sampler; Eijkelkamp Agrisearch Equipment) inserted into the upper 1.5 cm of the sediment. For $^{13}$C-DIC (dissolved inorganic carbon) analysis, 1 mL of mixed pore-water sample per frame was dispensed into air-tight headspace vials. The remainder of the pore-water sample was used for $^{13}$C-SCOA analysis. Subsequently, sediment was sampled by inserting a corer (inside diameter 10 cm) to a depth of 5 cm, and the top 1.5 cm of the sediment was collected. The sampling hole was filled with unlabeled sediment collected just outside the sampling frames and the corer was removed. The two sediment samples taken from each frame were homogenized, and divided in the field into eight samples of 20–25 g wet weight (wt) each for Mag-SIP analysis; one sample of 45 g wet wt for the measurement of total organic carbon (TOC), PLFA, and bulk carbohydrates; and a sample of 5 g wet wt for pigment analysis. Additionally, ~2 g wet wt of the homogenized sediment was directly processed for water-extractable carbohydrates by adding it to 4.5 mL Milli-Q water as described in de Brouwer and Stal (2001). The samples were immediately transferred on ice to the laboratory. In the laboratory, samples for water-extractable carbohydrates were shaken for 1 h at 30°C and the supernatant was collected after centrifugation at 4000 × g for 15 min and stored for carbohydrate analysis. The sediment pellet was re-extracted with 4.5 mL of 0.1 mol L⁻¹ EDTA by shaking for 4 h at room temperature. The supernatant was collected after centrifugation at 4000 × g for 15 min, and stored for EDTA-extractable carbohydrate analysis. Sediment samples for Mag-SIP and pigment analysis were stored at −80°C while the other samples were stored at −20°C.
To study dark fixation by chemoautotrophic and heterotrophic bacteria, two cores (7 cm internal diameter) were taken outside the frames and incubated in the dark for 4 h with the same amount of $^{13}$C-label (per m$^2$) added to the top of the sediment as in the field. The top 1.5 cm of these cores was sampled and analyzed for PLFA labeling.

**Analytical procedures**—Sediment samples for PLFA, TOC, pigment, and bulk carbohydrate analysis were freeze-dried. Lipids were extracted from 3 g of freeze-dried sediment using a modified Bligh and Dyer extraction (Boschker et al. 1999) from which the PLFA fraction was separated on a silica column. The PLFA fraction was converted to fatty-acid methyl esters and the carbon content and isotopic composition of these derivatives were measured by gas chromatography–combustion–isotope ratio mass spectrometry (Middelburg et al. 2000). The carbon content and isotopic composition of TOC was analyzed by using an elemental analyzer–isotope ratio mass spectrometer (EA–IRMS) after the removal of carbonate with hydrochloric acid (Boschker et al. 1999). For DIC analysis, pore-water samples were acidified by adding 0.1 mL of 19 mol L$^{-1}$ phosphoric acid (Miyajima et al. 1995), and headspace gas was injected into EA–IRMS in order to determine the concentration and isotopic composition of DIC. Pigments were extracted from freeze-dried sediment with acetone (90%, buffered with 5% ammonium acetate), and analyzed by reverse-phase high-performance liquid chromatography (HPLC; Dijkman and Kromkamp 2006). Carbon content and isotopic composition of SCOA were analyzed by LC–IRMS (Isolink interface and DELTA V Advantage IRMS; Thermo Fisher Scientific) equipped with an Aminex HPX-87H cation-exchange column (Bio-Rad). The eluent was 8 mmol L$^{-1}$ sulfuric acid at a flow rate of 0.4 mL min$^{-1}$ (Krumbock and Conrad 1991).

The monosaccharide composition of bulk, water-extractable, and EDTA-extractable carbohydrates was determined by LC–IRMS (Boschker et al. 2008). Freeze-dried sediment
and carbohydrate extracts were hydrolyzed with 1.1 mol L⁻¹ sulfuric acid for 1 h at 120 °C. The hydrolyzed carbohydrates were neutralized with strontium carbonate, and the precipitate was removed by centrifugation. Carbohydrate concentrations and isotope ratios were measured by LC–IRMS (see above) equipped with a CarboPac PA20 (Dionex Benelux). The eluent was 1 mmol L⁻¹ sodium hydroxide at a flow rate of 0.3 mL min⁻¹.

Concentration or ¹³C-labeling of compounds above was expressed as mole of carbon per gram of dry weight sediment (C or ¹³C g⁻¹ dry wt) in order to directly compare the size of each pool and to trace label transfer. Water content of the sediment was constant through the experiment at 0.32 ± 0.02 g per gram of sediment wet weight.

Mag-SIP analysis and clone library construction—A nested set of oligonucleotide probes was used for the Mag-SIP protocol (Table 1; Miyatake et al. 2013). The probes EUB338 (Amann et al. 1990) and DELTA495a (Loy et al. 2002) were used for most Bacteria and Deltaproteobacteria, respectively, even though they do not target all genera in these domains. The DELTA495a probe was used in combination with a competitor probe, cDELTA495a (Macalady et al. 2006), to avoid capture of Gammaproteobacteria, which have only one different base in the target region of DELTA495a. The family Desulfobacteraceae was covered with the probe Dbact653 (Miyatake et al. 2009) and the probe BG553 was used for Gammaproteobacteria (Miyatake et al. 2013). Both the Dbact653 and BG553 probes were used in combination with unlabeled helper probes complementary to the consensus sequence upstream and downstream of the probe target in order to increase yield (Fuchs et al. 2000; MacGregor et al. 2002). The probe CYA361 (Schönhuber et al. 1999) was used for Cyanobacteria and chloroplasts.

Hybridization of 16S rRNA with oligonucleotide probes and the isotope analysis of captured 16S rRNA were done as described in Miyatake et al. (2009). In short, total RNA was extracted from the sediment with the chloroform-phenol method and 20–40 μg total RNA was hybridized with biotin-labeled probes. The 16S rRNA-probe hybrids were captured with streptavidin-coated hydrophobic paramagnetic beads (Dynabeads MyOne Streptavidin T1; Invitrogen). Captured 16S rRNA was released from the beads by heat, and ~ 500 ng of this rRNA was used for stable isotope measurement by using flow injection analysis (FIA: Isolink interface with DELTA V Advantage IRMS; Thermo Fisher Scientific). Samples (in 50 μL) were directly injected into this FIA–IRMS operating in bulk injection mode. The carbon blank was also determined by performing the Mag-SIP protocol but without the RNA extract.

In order to identify active members of the microbial community and determine the specificity of the oligonucleotide probes used in the Mag-SIP protocol, clone libraries derived from total RNA and captured 16S rRNA were constructed. The total RNA extracted from the sediment with the chloroform-phenol method (0 h sample) and captured 16S rRNA were reverse-transcribed with reverse primer DXR518 (Nogales et al. 1999). The reverse transcripts were amplified by polymerase chain reaction (PCR) using primers 27F-DXR518 (Mills et al. 2005; Martinez et al. 2006). The PCR products were ligated into pGEM T-easy vector (Promega) and transformed into Escherichia coli JM109 competent cells following the manufacturer’s instructions. Positive clones were re-amplified with M13 primers, and sequenced with the 27F primer on an ABI PRISM 3130 Genetic Analyzer (Applied Biosystems). Sequence chromatographs were manually checked using the Chromas Lite software (http://www.chromaslite.com.au/chroma_s_lite.html). Phylogenetic analysis was performed by using the fast aligner and treeing tools implemented in the ARB program package (Ludwig et al. 2004). Phylogenetic relationships were determined by inserting sequences from this study into an ARB tree composed of the Greengenes database (http://greengenes.lbl.gov/) to which additional Deltaproteobacteria clone sequences from marine environments had been added. Nucleotide sequences obtained in this study have been deposited in GenBank under accession numbers GU215079–GU215169.

Calculations of label incorporation—Stable carbon isotope ratios were expressed as δ¹³C values calibrated against the international standard Vienna Pee Dee Belemnite (VPDB). The delta notation is defined as

\[ \delta^{13}C_{\text{sample}}(\%o) = \left( \frac{R_{\text{sample}}}{R_{\text{std}}} - 1 \right) \times 1000 \]

where \( R_{\text{sample}} \) is the ratio of ¹³C in the sample and \( R_{\text{std}} \) is the ratio of the international standard VPDB (0.0111797).

Captured rRNA was measured by bulk injection and therefore a correction for the blank was necessary. Correction of δ¹³C RNA value for the blank is done as

\[ \delta^{13}C_{\text{rRNA}}(\%o) = \left[ \frac{\delta^{13}C_{\text{sample}} \times C_{\text{sample}}}{C_{\text{sample}} - C_{\text{blank}}} - \frac{\delta^{13}C_{\text{blank}} \times C_{\text{blank}}}{C_{\text{sample}} - C_{\text{blank}}} \right] \]

where \( C_{\text{sample}} \) is the amount of carbon in the sample, \( \delta^{13}C_{\text{blank}} \) is the \( \delta^{13}C \) value of the blank, and \( C_{\text{blank}} \) is the amount of carbon in the blank (Boschker 2004). ¹³C-label incorporation in rRNA is indicated as the increase of \( \delta^{13}C \) value between background (unlabeled sample) and labeled samples (\( \Delta\delta^{13}C; \%o \)).

The absolute amount of ¹³C atoms incorporated into different carbon pools over the background was expressed as excess ¹³C and calculated from \( \delta^{13}C_{\text{sample}} \) as

\[ \text{Excess}^{13}C(\text{mol}^{13}C \text{ g}^{-1} \text{ dry wt}) = \left[ \frac{\left( \frac{\delta^{13}C_{\text{sample}}}{1000 + 1} \right) \times R_{\text{st}}}{\left( \frac{\delta^{13}C_{\text{sample}}}{1000 + 1} \right) \times R_{\text{st}} + 1} - \left( \frac{\left( \frac{\delta^{13}C_{\text{background}}}{1000 + 1} \right) \times R_{\text{st}}}{\left( \frac{\delta^{13}C_{\text{background}}}{1000 + 1} \right) \times R_{\text{st}} + 1} \right) \right] \times C_{\text{sample}} \]

where \( \delta^{13}C_{\text{background}} \) is the \( \delta^{13}C \) value of the unlabeled sample and \( C_{\text{sample}} \) is the pool size in mole of carbon per gram of dry weight sediment (C g⁻¹ dry wt).
Excess $^{13}$C into bacterial biomass was estimated from the label in bacterial-biomarker PLFA as

$$\text{Excess}^{13}\text{C} - \text{bacterial biomass}(\text{mol}^{13}\text{C} \text{ g}^{-1}\text{dry wt}) = \sum \text{Excess}^{13}\text{C}_{\text{PLFA}_{\text{bact}}} / (0.056 \times 0.28)$$

where $^{13}\text{C}_{\text{PLFA}_{\text{bact}}}$ is $^{13}$C in bacterial-biomarker PLFA (i14:0, i15:0, a15:0, i16:0, and 18:1 $\omega 7c$). 0.056 represents the average PLFA content in bacteria in terms of carbon and 0.28 ± 0.04 is the average fraction of these bacterial-biomarkers PLFA among total PLFA in bacteria-dominated marine sediments (Middelburg et al. 2000). Excess $^{13}$C into algal biomass was calculated from the difference between excess $^{13}$C into all PLFA and excess $^{13}$C into bacterial-biomarker PLFA and also corrected for the typical PLFA content of diatoms:

$$\text{Excess}^{13}\text{C} - \text{algal biomass}(\text{mol}^{13}\text{C} \text{ g}^{-1}\text{dry wt}) = \left(\sum \text{Excess}^{13}\text{C}_{\text{PLFA}_{\text{all}}} - \sum \text{Excess}^{13}\text{C}_{\text{PLFA}_{\text{bact}}}\right) / 0.035$$

where $^{13}\text{C}_{\text{PLFA}_{\text{all}}}$ is $^{13}$C in all individual PLFA measured in this study and 0.035 represents the average PLFA content of diatoms (Middelburg et al. 2000). Algal and bacterial biomass was calculated as above in terms of carbon per gram of dry weight sediment using the biomarker PLFA concentrations instead of excess $^{13}$C.

Results

Incorporation of $^{13}$C in major carbon pools and intermediate metabolites—Concentrations of DIC and TOC in the sediment were 1.2 ± 0.1 $\mu$mol C g$^{-1}$ dry wt and 309 ± 20 $\mu$mol C g$^{-1}$ dry wt, respectively, throughout the experimental period. Label recovered in the DIC after 4 h (25 mmol $^{13}$C g$^{-1}$ dry wt; Fig. 1B) was already substantially lower than expected from the total amount of DIC-label that was initially added (600 mmol $^{13}$C g$^{-1}$ dry wt), which was the result of photosynthesis by diatoms and probably also exchange with the atmosphere in this open system. DIC $^{13}$C-labeling further decreased sharply during the first day of the experiment followed by more gradual decrease until the end of the experiment when almost no $^{13}$C label remained (Fig. 1B). Label incorporation in TOC increased sharply during the first low tide and remained approximately constant at 200 nmol $^{13}$C g$^{-1}$ dry wt during the first 3 d of the experiment, except for a small peak after 12 h. Finally, the label in TOC decreased to 120 nmol $^{13}$C g$^{-1}$ dry wt on day 5 (Fig. 1C). These data suggest a rapid incorporation of the $^{13}$C-DIC label by the benthic primary producers during the pulse-labeling period followed by minimal losses for the remainder of the experiment until day 5.

The bulk carbohydrates content was 27.1 ± 1.7 $\mu$mol C g$^{-1}$ dry wt throughout the experimental period and showed little variation (Fig. 2A). Glucose and galactose were the two major components and their contributions remained constant (~30% and 15%, respectively). Water-extractable carbohydrates accounted for ~10% of the bulk carbohydrates and varied substantially during the experiment (Fig. 2B). The concentration of water-extractable carbohydrates was lower for the 12 h and day 3 samplings. The concentration of EDTA-extractable carbohydrates was ~2–3 times lower than the water-extractable carbohydrates and remained approximately constant during the experiment except on the day 3 when values decreased by a factor of 2–3 (Fig. 2C). Glucose was the main carbohydrate in both the water-extractable EPS and EDTA-extractable fractions explaining between 30% and 70% of the total carbohydrates.

The limited turnover of the $^{13}$C-label in the TOC pool during the experimental period was not seen in the bulk and water-extractable carbohydrates. Excess $^{13}$C in bulk, water-extractable, and EDTA-extractable carbohydrates were highest between 4 h and 12 h after labeling and decreased with time (Fig. 2D–F). More than 90% of $^{13}$C labeling was recovered in glucose in the three analyzed pools. Initially, $^{13}$C content in the water-extractable carbohydrates explained about half of the $^{13}$C label of the bulk carbohydrates. Labeling of the EDTA-extractable carbohydrates was substantially lower (one order of magnitude) than of the water-extractable carbohydrates. The label decreased faster in the water-extractable carbohydrates than in the bulk and EDTA-extractable carbohydrates, and had
almost disappeared at the end of the experiment. Label incorporation in carbohydrates other than glucose was much lower and remained more-or-less constant. Hence, the decrease in $^{13}\text{C}$-label incorporation in bulk carbohydrates between 4 h and day 5 was mainly due (by about 75%) to the decrease of $^{13}\text{C}$ label in glucose from water-extractable carbohydrates and to a much lesser extent by the EDTA-extractable carbohydrates.

SCOA in the pore water was the most dynamic pool of metabolites analyzed during this experiment. SCOA detected in this study were formate, acetate, oxalate, malate, lactate, and succinate. Lactate and succinate could not be separated sufficiently by our HPLC protocol and are therefore reported together. The propionate peak overlapped with an unknown peak. Significant changes in the concentrations of lactate-succinate and acetate were observed. Other SCOA detected in this study had low and constant concentrations during the experiment (20–30 nmol C g$^{-1}$ dry wt) and showed limited labeling; hence, only lactate+succinate and acetate are depicted in Fig. 3. Concentrations of lactate+succinate and acetate were similar at the start of the experiment, but lactate+succinate was higher than acetate after addition of the tracer and increased again on day 3. The concentration of...
acetate was highest at 0 h followed by sharp decrease at 4 h, then gradually increased to day 5. Pore-water concentrations of acetate (range 100–1200 nmol C mL$^{-1}$) and lactate + succinate (range 750–2300 nmol C mL$^{-1}$) were high. $^{13}$C-label incorporation was only detected for lactate + succinate and acetate, and was higher in lactate + succinate than in acetate (Fig. 3B). Excess $^{13}$C in lactate + succinate and acetate was highest at 4 h, and then significantly decreased at 12 h followed by gradual increase before diminishing at day 5.

**Microbial community composition**—Throughout the 5 d of the experiment, the sediment was covered by a diatom mat. Chlorophyll $a$ content was $17.2 \pm 0.9 \mu g \cdot g^{-1} \cdot dry \ wt$, and the content of fucoxanthin (a marker for diatoms [Wright and Jeffrey 1987; Dijkman and Kromkamp 2006]) was $6.1 \pm 0.3 \mu g \cdot g^{-1} \cdot dry \ wt$ throughout the experiment, indicating a high proportion of diatom biomass. The 16S rRNA clone library derived from total RNA reflects the metabolically active community and confirmed the dominance of benthic diatoms (44% of the sequences belonged to chloroplasts of benthic diatoms), whereas cyanobacterial clones were fewer (6%; Fig. 4). Moreover, the content of specific biomarkers PLFA for diatoms was much higher ($86.5 \pm 9.3$ and $495.5 \pm 110.4 \text{ nmol C g}^{-1} \cdot dry \ wt$ for 16 PUFA and 20:5$\omega$3, respectively) than for Cyanobacteria ($51.8 \pm 10.1$ and $30.9 \pm 5.5 \text{ nmol C g}^{-1} \cdot dry \ wt$ for 18 PUFA and 18:2$\omega$6c, respectively), and hence, Cyanobacteria contributed much less to the microphytobenthos biomass than diatoms (Fig. 5). The 16S rRNA clone library showed that Gammaproteobacteria (21% of the sequences), Bacteroidetes (8%), and Deltaproteobacteria (7%) were the major groups in the bacterial community in addition to the Cyanobacteria (Fig. 4). EUB338 was used as a general bacterial probe in this study and it covered most of the Bacteria detected in the clone library and only missed the Planctomycetes. The combination of the other specific probes covered 77% of the 16S rRNA targeted by the EUB338 probe but did not include the Bacteroidetes. The specificity of each oligonucleotide probe used with Mag-SIP was $\geq 90\%$.

**Incorporation of $^{13}$C into biomarkers**—Label ($^{13}$C) incorporation in PLFA biomarkers and in 16S rRNA of
specific bacterial groups was measured in order to trace the carbon transfer from diatoms to heterotrophic bacteria. The total amount of RNA extracted per dry weight unit of sediment was constant throughout the experiment (10.9 ± 1.5 μg RNA g⁻¹ dry wt), as was the case with PLFA (3.22 ± 0.35 μmol C g⁻¹ dry wt). The PLFA content in diatoms and bacteria is relatively constant and conversion factors are known that can be used to infer the actual amount of label in their biomass (Middelburg et al. 2000; Flärdh et al. 1992; Kerkhof and Kemp 1999). Therefore, 13C incorporation in 16S rRNA was expressed as Δδ¹³C, which is a measure of specific ¹³C-label incorporation into 16S rRNA of the target group relative to the unlabeled 16S rRNA of this group.

Based on PLFA data, ¹³C labeling in diatom biomass rapidly increased during the first 12 h and subsequently decreased somewhat until day 1, which again was followed by a secondary increase until day 3 (Fig. 6B). The ¹³C labeling in bacterial biomass also increased rapid during the first 12 h followed by a slower increase until day 3 and a decrease at day 5 (Fig. 6B). Note that hardly any ¹³C-DIC label remained in the sediment after the first day (Fig. 1). This suggests that the increase in label incorporation in the diatoms and bacteria after day 1 can be attributed to the consumption of the ¹³C-labeled organic pool produced during the first 4 h of the experiment. This second increase in ¹³C-label incorporation in diatoms and bacteria (Fig. 6B) coincided with the decrease in ¹³C-labeled glucose originating from the water-extractable carbohydrates (Fig. 2E). There was no detectable label incorporation in PLFA in dark incubations (data not shown), indicating that dark fixation by chemoheterotrophic bacteria and anaplerotic carbon fixation by heterotrophs were not important. Hence, the PLFA data suggest a rapid transfer of organic carbon from diatoms to bacteria in the first 12 h, followed by a slower secondary utilization by both diatoms and bacteria of primarily ¹³C-labeled glucose from the water-extractable carbohydrates.

Label incorporation in 16S rRNA that was captured with group-specific oligonucleotide probes also showed two maxima, the first on day 1 and the second on day 3 (Fig. 7A). Unlabeled controls of 16S rRNA possessed δ¹³C values between −15‰ and −20‰, which is within the typical range for marine benthic diatoms and bacteria (Coffin et al. 1990; Boschker and Middelburg 2002). The Δδ¹³C value of diatom and Cyanobacteria 16S rRNA represents primarily the label incorporation by diatoms.

Fig. 5. Average concentration of individual PLFA in unlabeled sediment (0 h, n = 2). Error bars indicate the range in the duplicate data.

Fig. 6. (A) Average concentration of biomass carbon and (B) excess ¹³C in diatoms and in bacteria calculated from concentrations of biomarker PLFA and ¹³C label incorporated into PLFA, respectively (n = 2). The bacterial-biomarker PLFA used are i14:0, i15:0, a15:0, i16:0, and 18:1ω7c. Biomass carbon and ¹³C-label incorporation of diatoms were calculated from the difference between total PLFA and bacterial PLFA. Error bars show the range in the duplicate data.
because the Cyanobacteria biomass was low. Remarkably, $\Delta^{13}C$ ratios of all captured 16S rRNA fractions were similar and showed the same trend during the experiment. For instance, the difference in the $\Delta^{13}C$ pattern of the total bacteria (EUB338-captured 16S rRNA) and of the family Desulfobacteraceae (Dbact653-captured 16S rRNA), which comprised only 3% of the clone library, was small. Only Deltaproteobacteria 16S rRNA showed a delay in accumulating label during the second part of the experiment. On day 3, the label in Deltaproteobacteria 16S rRNA was less than in the other captured 16S rRNAs and reached a maximum only on day 5 at the end of the experiment.

In order to compare the label incorporation in 16S rRNA and PLFA, the $\Delta^{13}C$ values of several representative biomarker PLFAs for microphytobenthos (16 PUFA, 18 PUFA, 18:2ω6c, and 20:5ω3) and biomarker PLFAs for heterotrophic bacteria (i14:0, i15:0, a15:0, i16:0, and 18:1ω7c) are depicted in Fig. 7B and 7C. Biomarker PLFA $\Delta^{13}C$ values for diatoms (16 PUFA and 20:5ω3) and Cyanobacteria (18 PUFA and 18:2ω6c) showed different trends (Fig. 7B). Labeling in biomarker PLFA for Cyanobacteria initially increased sharply followed by a gradual decrease until the end of the experiment. In contrast, diatom biomarker PLFA labeling showed a fast increase until 12 h followed by a more gradual increase until day 3, and it then decreased slightly. Although $\Delta^{13}C$ values of biomarker PLFA for Cyanobacteria were higher than for diatoms, excess $^{13}C$ in diatom PLFA was much higher because diatoms were the dominant primary producers at the study site. Representative biomarker PLFA for bacteria showed similar labeling patterns as biomarker PLFA for diatoms (16 PUFA and 20:5ω3) with a fast increase during the first day of the experiment followed by a slower labeling until a maximum on day 3 (Fig. 7C). Furthermore, the highest $\Delta^{13}C$ values in the different captured 16S rRNA and biomarker PLFAs were at a similar range (50–100%) except for cyanobacterial biomarker PLFAs, which was, however, only a minor part of the microbial community. To summarize, the combined results of the biomarker labeling suggest that carbon fixed by diatoms was rapidly and evenly utilized by the heterotrophic bacterial community.

Discussion

In this study, the transfer of carbon from benthic diatoms to the heterotrophic bacterial community was traced by following the $^{13}C$-labeling dynamics in different biologically derived compounds and major groups within the microbial community. We demonstrated that Mag-SIP can be used to trace carbon flow in a benthic diatom mat and we measured the $^{13}C$-label incorporation in heterotrophic bacteria that were identified to the family level. Other in situ $^{13}C$-labeling studies used PLFA (Middelburg et al. 2000; Evrard et al. 2008; Oakes et al. 2012) and D-alanine (Veuger et al. 2006) as biomarkers, but these have a phylogenetic resolution that does not exceed the level of domains or kingdoms. In addition, we also measured $^{13}C$ labeling in bulk, water-extractable, and EDTA-extractable carbohydrates as well as SCOA in pore water using recently developed LC–IRMS methods to determine which intermediates were important in the transfer of organic matter from diatoms to heterotrophic bacteria. Although the relationship between diatom carbohydrates exudates and the heterotrophic bacterial community have been studied in intertidal sediments (Hanlon et al. 2006; Haynes et al. 2007; Bellinger et al. 2009; Oakes et al. 2012; Taylor et al. 2013), we included SCOA as important intermediates in the anaerobic degradation of diatom material (McKew et al. 2013). The results were used to construct a conceptual model of the transfer of carbon in the diatom mat (Fig. 8A) and are summarized in a $^{13}C$-budget (Fig. 8B). We propose two main events during which organic carbon was
transferred from benthic diatoms to heterotrophic bacteria, which operated on different time scales.

The $^{13}$C-label incorporation in PLFA and 16S rRNA biomarkers suggested that there was a fast transfer of organic substrates from diatoms to heterotrophic bacteria already during the first day of the experiment (Figs. 6, 7, 8). This initial fast transfer was probably the result of the utilization of low-molecular-weight (< 800 Da) organic compounds exuded by diatoms that could be directly utilized by heterotrophic bacteria. These low-molecular-weight exudates may include a wide range of different compounds including monomeric carbohydrates, amino acids and organic acids, some of which may be difficult to track with the currently available $^{13}$C-methods due to their fast turn-over and typical low concentrations. However, the fast transfer of label coincided with a sharp peak in SCOA labeling (Fig. 3), which indicated that some low-molecular-weight exudates produced by the diatoms (such as glucose) were quickly fermented by anaerobic bacteria. Fermentation products such as acetate and lactate are important substrates for anaerobic bacterial communities (Jørgensen 2006; McKew et al. 2013). During photosynthesis, carbon assimilated by the diatoms is in part stored as intracellular glucan that consists of 90% glucose (Underwood et al. 2004) and excess carbon is exuded from the cells as water-extractable carbohydrates (Smith and Underwood 2000; de Brouwer and Stal 2001). At daytime in intertidal sediments, up to 80% of this water-extractable carbohydrate can be mono- and oligo-saccharides (Underwood et al. 1995; Underwood and Smith 1998; Smith and Underwood 2000), which are readily available for heterotrophic bacteria and may explain the initial fast transfer.

Between 12 h and 24 h, the label recovered in water-extractable carbohydrates explained 70% of the label in the
bulky carbohydrates (Fig. 2D, E) and between 20% and 25% of the label in TOC (Fig. 1C, 6B). Among the carbohydrates, most of the $^{13}\text{C}$ label was recovered in glucose (> 90%), which was highest at 4 h after labeling and subsequently decreased sharply until it had disappeared almost completely by day 3 (Fig. 2D–F), explaining most of the total loss of labeled glucose in the diatom mat. Glucose is the first compound synthesized by phototrophic organisms and it was therefore not surprising that it had the highest label, confirming observations by Bellinger et al. (2009) and to a lesser extent by Oakes et al. (2010). Water-extractable EPS carbohydrates were therefore major intermediates released by the diatoms as detected in this study.

After the initial fast transfer of organic matter from diatoms to heterotrophic bacteria, label incorporation into both diatoms and heterotrophic bacteria proceeded more slowly from day 1 until day 3 (Figs. 6, 7, 8). Almost no label was left in the DIC between day 1 and 3 and, hence, the second label incorporation must have been due to organic carbon utilization that was initially produced by the diatoms. This second peak of label incorporation in bacteria coincided with the disappearance of label in water-extractable carbohydrates (Fig. 2E) and also with the second increase of label in SCOAs (Fig. 3B). It was therefore concluded that the origin of this second labeling between day 1 and 3 was based on the consumption of the EPS fraction of water-extractable carbohydrates that was exuded during the first day after labeling and was partially fermented under anaerobic conditions. Approximately 15% or more of the diatom exudates in the light is EPS (Smith and Underwood 2000; Underwood and Paterson 2003). The chemical properties of EPS make them more recalcitrant to degradation in contrast to the low-molecular-weight exudates. Bacteria can utilize high-molecular-weight compounds such as EPS only after extracellular enzymatic hydrolysis to low-molecular-weight compounds (Fuchs et al. 1998; Hunter et al. 2006), which would explain the slower labeling rate of bacterial PLFA and 16S rRNA between day 1 and 3 (Fig. 7A,C). Surprisingly, we also detected this secondary labeling in the diatoms, which may be attributed to the heterotrophic metabolism of these organisms because hardly any DIC $^{13}$C label was left. This second labeling coincided with the decrease of labeling in carbohydrates, which suggested that the diatoms used EPS as an external carbon storage as has been shown for pure cultures of diatoms (Staats et al. 2000; de Brouwer and Stal 2001) and for slurry incubation to which diatom-derived EPS was added (Taylor et al. 2013). Miyatake et al. (2013) recently showed that benthic diatoms in coastal sediment incorporated a variety of organic substrates, and active heterotrophic growth of diatom cultures on glucose is well-known (Lewin and Hellebust 1976; Admiraal and Pelletier 1979). Alternatively, it is also possible that diatoms gradually re-incorporated intracellular storage glucans into cell materials. This was not specifically investigated in our study but it is well-known that diatoms utilize these reserve carbohydrates during the dark (Smith and Underwood 2000). For instance, Smith and Underwood (2000) reported that pure cultures of diatoms produced EPS in the dark from intracellular glucan during up to 3 d, which agrees with the second peak of label incorporation of diatoms that we observed.

We used both PLFA analyses and Mag-SIP to determine the labeling dynamics of major microbial groups in the diatom mat. Labeling levels and timing were mostly similar, although labeling of 16S rRNA seemed to be more dynamic than that of PLFA (Fig. 7). The highest $\Delta^{13}\text{C}$ values of diatom PLFA and bacterial PLFA were in the same range at 50–100% in this study. This is consistent with the study of Bellinger et al. (2009), but disagrees with those of Middelburg et al. (2000) and Evrard et al. (2008), who reported an approximately five-fold higher labeling in terms of $\Delta^{13}\text{C}$ values in diatom PLFA than in bacterial PLFA. Another difference is that Middelburg et al. (2000) and Evrard et al. (2008) reported a much faster decay of total labeling stocks. A possible reason for these differences is that our experiment and that of Bellinger et al. (2009) were done early in spring when thick diatom mats are present, whereas the other studies were done later in the season. In our study, macrofauna had not yet fully developed and therefore the grazing pressure was low. Hence, the diatom mat was at a climax with high biomass and limited growth, and a close coupling between the diatoms and the heterotrophic bacterial community.

Given the importance of water-extractable carbohydrates as an intermediate in diatom mats, we expected that specialized bacteria would be involved in the coupling between diatoms and bacteria (Taylor et al. 2013). However, our results suggest that all major groups within the heterotrophic bacterial community equally utilized the diatom-derived organic matter because the labeling of 16S rRNA captured by all different oligonucleotide probes was more or less the same throughout the experiment (Fig. 7). Only the labeling of the *Deltaproteobacteria* 16S rRNA was somewhat slower compared with other groups, but this was only the case between day 1 and 5. The *Deltaproteobacteria* in the diatom mat are mostly related to sulfate-reducing bacteria and the slower labeling could be due to the fact that their substrates had first to be produced by fermentation under anaerobic conditions by other members of the bacterial community (Miyatake et al. 2013). The even labeling between different bacterial groups may be explained by a combination of diatom primary production dominating heterotrophic carbon cycling in this sediment and a wide variety of exudates produced by diatoms (although our results indicate that glucose in water-extractable EPS was a major intermediate). Taylor et al. (2013) reported high label incorporation by diatoms, *Alpha*, and *Gammaproteobacteria* from isolated water-extractable EPS with RNA-SIP, but also showed the even labeling of general bacterial PLFA with PLFA-SIP. Miyatake et al. (2013) also reported the even utilization of glucose (and other organic substrates) by most major groups in the microbial community of the sediment surface layer (0–2 cm). This also hints at a limited specialization of the bacterial community in these intertidal sediments; however, we cannot exclude that there may still be specialized bacteria within the relatively broad phylogenetic groups that were targeted in our study (Taylor et al. 2013). We conclude, therefore, that the heterotrophic bacterial community relies to a similar extent on the organic
carbon produced by the diatoms, resulting in a closely coupled microbial food web.

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