Effect of phytoplankton bloom deposition on benthic bacterial communities in two contrasting sediments in the southern North Sea

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ABSTRACT: The response of benthic bacterial community composition, diversity, and biomass to phytoplankton deposition was investigated in 2 different sediment horizons at 2 contrasting sites in the southern North Sea. Differences in bacterial community composition between stations were observed. Seasonal differences in bacterial community composition were significant and were stronger in fine sediments, probably related to stronger fluctuations in food availability. Variation in community composition over the vertical sediment profile was different for both stations. In coarser sediment, the difference was mostly due to the absence of certain operational taxonomic units (OTUs) at the surface, while in fine sediment, 2 distinct communities were present. A RELATE test revealed that bacterial community composition was influenced by the amount of labile organic matter (estimated through chl a concentration in the sediment). Diversity in terms of OTU richness and Shannon-Weaver diversity index was higher in finer grained sediments. In coarser sediments, diversity at the surface layer was lower, which might be related to stronger hydrodynamic pressure at this station. These differences were not observed at the other station. Seasonal changes in diversity were not detected at either station. Bacterial biomass was slightly higher in finer sediments and was not correlated with either chl a or temperature. Seasonal differences in bacterial biomass followed those observed for community composition, while no vertical differences were detected.

KEY WORDS: Benthic bacteria · North Sea · Denaturing gradient gel electrophoresis · DGGE · Community composition · Diversity · Bacterial biomass
in both density and diversity, especially among the selective deposit-feeding nematodes (1A-nematodes; Wieser 1953, Vanaverbeke et al. 2004b) and deposit-feeding nematodes (1B-nematodes; Wieser 1953, M. Steyaert et al. unpubl.). This was partially explained by changes in bacterial diversity (Vanaverbeke et al. 2004b), since both 1A and 1B nematodes feed on bacteria (Wieser 1953), and nematodes can selectively feed on particular bacterial strains (Moens et al. 1999, De Mesel et al. 2004).

Bacteria have been reported to react fast on phytoplankton sedimentation in terms of biomass production (larger cells), cell division, and activity, which results in an increase in biomass, density, and productivity (Graf et al. 1982, Meyer-Reil 1983, Goedkoop & Johnson 1996, Boon et al. 1998). This response of the bacterial community to an input of organic matter seems to be influenced by the covariation of both food supply and temperature (Graf et al. 1982, Boon et al. 1998). Van Duyl & Kop (1994) indicated temperature and substrate availability for bacteria as the most important factors influencing bacterial production.

However, data concerning possible changes in benthic bacterial community composition and diversity after a mass input of organic matter (OM) are lacking. Applying denaturing gradient gel electrophoresis (DGGE) in microbial ecology allows for simultaneous analyses of multiple samples, which enables monitoring the complex dynamics that microbial communities may undergo by diel and seasonal fluctuations or after environmental perturbations (Muyzer 1999). This technique has been applied successfully to document changes in planktonic bacterial communities during phytoplankton bloom events (e.g. Van Hannen et al. 1999a, Riemann et al. 2000, Fandino et al. 2001, Riemann & Winding 2001, Van der Gucht et al. 2001, Muylaert et al. 2002, Rooney-Varga et al. 2005).

Here we aimed to examine the bacterial response to a pulsed food supply in terms of community composition, diversity, and biomass at 2 well studied contrasting sites on the BCS. The 2 study sites, Stns 115bis and 330, are biogeochemically different: 115bis is a depositional station, characterized by the presence of fine sediments and temperature (median grain size: 185 µm), with 4% of mud (M. Steyaert et al. unpubl.), while Stn 330 has permeable sediment containing medium sand (median grain size: 329 to 361 µm), devoid of mud (Vanaverbeke et al. 2004a,b). This results in strong differences in vertical profiles of chlorophyll (chl) a, which is a proxy for the availability of labile OM (Boon & Duineveld 1996). Especially in depositional stations, the differences in availability of labile OM between sediment depths during and after phytoplankton sedimentation can be striking (Graf 1992), creating very different biogeochemical conditions. After sedimentation of phyto-detritus, the sediment at Stn 115bis can become anoxic and remain so until mid autumn (M. Steyaert et al. unpubl.). As this affects metazoan meiobenthic vertical distribution patterns (Steyaert et al. 1999, M. Steyaert et al. unpubl.), it probably also has important consequences for bacterial life.

In this study, we therefore aimed to test whether there are significant differences in bacterial community composition and biomass (1) between the 2 stations and (2) at the surface and in a deeper sediment layer within a station, and (3) whether there are significant seasonal shifts related to the spring bloom deposition.

MATERIALS AND METHODS

Study area and sampling. Samples were taken from the BCS Stns 115bis, located close to the coast (51° 09.2' N, 02° 37.2' E; 13 m depth) and 330, located farther offshore (51° 26.0' N, 02° 48.5' E; 20 m depth; Fig. 1). The sampling sites were never located on slopes or tops of sand banks.

Sampling at both stations was conducted monthly from October 2002 to October 2003 from the RVs ‘Zeeleeuw’ or ‘Belgica.’ Sampling took place on the same days at both stations. In December 2002, it was not possible to sample station 330 due to bad weather and sea conditions.

The water column was sampled at 3 m depth and 1 m above the sea floor for phytoplankton analysis using 10 l Niskin bottles. To obtain pigment samples, 500 ml of water from each depth were filtered onto GF/F glass microfiber filters (i.d. 4.7 cm) using a vacuum pump. Three replicated samples were obtained. Water temperature was recorded simultaneously. The samples were kept in the dark and preserved at –20°C on board and stored at –80°C in the laboratory.

Sediment was sampled using a Reineck box corer (surface area 180 cm²) or another box corer with a greater surface area (February, April, and October 2003). The box corer was deployed 3 times at each sampling station. In February, only 2 box cores were obtained for station 330.

From each box corer, 2 perspex cores (i.d. 3.6 cm) were taken: 1 for pigment analysis and another for bacterial counts and biomass analysis. These cores were sliced in 1 cm slices down to 10 cm. Pigment samples were preserved at –20°C on board and stored at –80°C in the laboratory. Bacterial counts and biomass samples were preserved in a 4% formaldehyde-tap water solution, thoroughly shaken, and stored in the fridge until processed further.

During February, April, and October 2003, additional cores for bacterial community composition
analysis (i.d. 6 cm) were taken. The bacterial cores were carefully closed in order to retain the overlaying water and preserved at 4°C until further processing in the laboratory. Additional deployments of the Reineck box corer or other box corer were done to obtain samples for bacterial biomass assessment.

**Laboratory treatment of samples.** The pigments sediment samples were weighed and pigment contents (chl a, phaeophytin, and phaeophorbid) were analyzed by HPLC (Gilson) following Wright & Jeffrey (1997). The ratio of phaeopigments to the sum of chl a + phaeopigments (PAP ratio) was calculated as an indication of the freshness of the material deposited on the sediment (Boon et al. 1998).

Samples for bacteria counting were prepared following Starink et al. (1994). Milli-Q water was added to the bacterial sample to a volume of 9.5 ml, and 0.5 ml Na2P2O7 (0.2 M) was added to reach a final concentration of Na2P2O7 of 0.01 M. This mixture was sonicated 5 times for 30 s using a Soniprep 150 (10 W). Between sonication pulses, the sample rested on ice for 30 s. Before staining, samples were diluted 10 to 1000 × using Tris. A subsample was filtered onto a 0.2 µm polycarbonate filter and stained with Sybrgold (Molecular Probes). Filters were then mounted on a slide. Counting of bacterial cells was performed using a Leica confocal microscope connected to QWIN software. From every image, 16 scans were made 0.2 µm vertically apart from each other. In this way, a 3.2 µm thick image was analyzed. From each scan, all particles >0.2 µm were counted and allocated to a size class. Per slide, 50 to 100 images were analyzed. The volume of the bacteria was calculated as a sphere volume \(4 \times \pi \times \frac{r^3}{3}\) with a radius estimation of half the average of lower (L) and upper (U) diameters, and was converted to carbon content (bacterial biomass) by using a conversion factor of 310 fg of C µm⁻³ (Fry 1990). Biomass was therefore calculated as: \[\{4 \times 3.141593 \times \frac{[(L + U)/ 4]^3}{3}\} \times 310.\] Data from Stn 115bis were only available from January 2003 onwards.

In a flow bench, 2 subsamples were taken from each of the sediment cores using a sterile syringe from which the tip was removed. From each of the 2 subsamples, the layers between 0 to 1 and 4 to 5 cm (1 ml each) were preserved at –80°C until further processing.

Although in most occasions 3 replicates per station and sediment layers were obtained, methodological problems prevented complete sampling for station 115bis in February (2 replicates from the 0 to 1 cm layer), in April (1 replicate from the 4 to 5 cm layer), and in October (2 replicates from the 4 to 5 cm layer).

**DNA extraction.** Total DNA was extracted from about 1.5 g of sediment following the procedure of Demba Diallo (2003). Sediment (1.5 g) was mixed with 1.5 ml Na2HPO4 (0.1 M) and washed by shaking for 30 min at room temperature. After centrifugation at 7000 × g (10 min at 4°C), the supernatant was removed. The pellet was resuspended in 500 µl of lysis buffer (2% cetyltrimethylammonium bromide [CTAB], 0.15 M NaCl, 0.1 M Na2EDTA [pH 8], 1% polyvinyl polypyrrollidone [PVPP]), and 7.5 mg lysozyme (Sigma) were added. Samples were incubated overnight at 37°C. We added 25 µl of proteinase K (20 mg ml⁻¹, Boehringer), and the tubes were incubated at 50°C for 40 min. The temperature was increased to 65°C for 20 min, and 300 µl extraction buffer (0.2 M NaCl, 0.1 M Tris-HCl pH 8, 2% SDS) were added. The mixture was then incubated at 65°C for another 10 min. After addition of 350 µl of 5 M NaCl, the samples were cooled on ice for 15 min. The supernatant was collected after centrifugation (7000 × g, 10 min, 4°C) and transferred into 2 ml centrifuge tubes. To precipitate the crude DNA, 75 µl of 5 M potassium acetate (Kac) and 250 µl 40% polyethylene glycol 8000 (PEG) were added, and the mixture was incubated at –80°C for 1 h. The pellet, obtained by centrifugation (13 000 × g for 15 min at 4°C), was resuspended in 900 µl 2× CTAB (2% CTAB, 1.4 M NaCl, 0.1 M Na2EDTA) and incubated for 15 min at 68°C. After addition of 900 µl of chloroform, the solu-
tion was gently mixed and centrifuged at 13 000 × g for 10 min at room temperature. The DNA was precipitated by addition of 1 ml of isopropanol and incubated for at least 15 min at 20°C. The pellet, obtained by centrifugation (13 000 × g for 15 min at 4°C), was dissolved in 450 µl 2.5 M ammonium acetate (NH₄OAc), and subsequently the DNA was precipitated by the addition of 1000 µl 95% ethanol and then incubated for at least 15 min at –20°C. The pellet of DNA was obtained by centrifugation at 13 000 × g for 15 min at 4°C and resuspended in 200 µl sterile water (Sigma). For each sediment sample, we performed 2 independent 1.5 g sediment preparations, of which the purified DNA samples were pooled and stored in a single vial. A 100 µl aliquot of the crude extract was further purified using the Wizard® DNA CleanUp kit (Promega).

**PCR amplification of 16S rRNA.** In total, 6 µl (ca. 100 ng) of the purified DNA were amplified in a Genius temperature cycler. For each sample, 2 PCRs were performed. The PCR mixture contained 6 µl of template DNA, 0.5 µM of each of the appropriate primers, 200 µM of each deoxynucleoside triphosphate, 5 µl of 10× PCR buffer (100 mM Tris-HCl pH 9), 500 mM KCl, 15 mM MgCl₂, 20 ng of bovine serum albumin, and 2.5 U of Taq DNA polymerase (AmpliTaq Perkin Elmer). Each mixture was adjusted to a final volume of 50 µl with sterile water (Sigma).

The primers were: F357GC (5’-CGCCCGCGCGCCG- CCGCGCCGCCGCCGCCGCCGCCGCCG-3’) and R518 (5’-ATTACCGCG GCCCGCGCCGCCCCCCTA CGGGAGGCAGCAG-3’). Both primers, which were designed by Muyzer et al. (1993), are used to amplify the 16S rDNA region corresponding to positions 341 to 534 in *Escherichia coli*. Primer F357GC, which contains a GC-rich clamp, is specific for most bacteria, and R518 is specific for most *Bacteria, Archaea*, and *Eucarya* (Van Hannen et al. 1999b). In order to improve the specificity of the amplification and to reduce the formation of spurious by-products, a ‘touchdown’ PCR (Don et al. 1991, Muyzer et al. 1993) was performed starting with 5 min at 94°C, followed by 20 cycles of denaturation at 94°C for 1 min, annealing at 65°C (the temperature was decreased by 0.5°C every cycle until the touchdown temperature of 56°C was reached) for 1 min, and primer extension at 72°C for 1 min. Ten additional cycles were carried out at an annealing temperature of 55°C. The tubes were then incubated for 10 min at 72°C. The presence of PCR products and their concentration were determined by analyzing 5 µl of PCR product on a 2% agarose gel. A molecular weight marker (Smartladder-Eurogentec SA) was included.

**Denaturing gradient gel electrophoresis (DGGE) analysis.** The DGGE technique was carried out using the D-Code System (Bio-Rad Laboratories). The PCR products were loaded onto 8% (w/v) polyacrylamide gels of 1 mm thickness, in 1× TAE buffer (20 mM Tris-acetate with pH 7.4, 10 mM acetate, 0.5 mM disodium EDTA). The denaturing gradient contained 35% to 70% denaturants (100% denaturant corresponded to 7 M urea and 40% [vol/vol] deionized formamide). The total lane intensity was normalized between the samples at 400 ng of DNA. Electrophoresis was performed at a constant voltage of 75 V for 16 h and at a constant temperature of 60°C. After electrophoresis, the gels were stained for 1 h in 1× TAE containing ethidium bromide (0.5 mg l⁻¹; Muyzer et al. 1993). The bands were visualized on a UV transillumination table equipped with a digital CCD camera.

As standards, we used a mixture of DNA from 9 clones (Van Der Gucht et al. 2001). On each gel, 3 standard lanes were analyzed in parallel to the samples, to facilitate comparison between gels. The banding patterns were then analyzed using BioNumerics 5.1 (Applied Maths BVBA). By measuring an optical density profile through each lane (corresponding to a single sample), this software identifies the band positions and calculates the contribution of the intensity of each band to the total intensity of the lane. This procedure yields a matrix with the relative intensity of each band in all samples.

**Bacterial biodiversity.** Ecological diversity is considered a function of the number of different classes (richness) and the relative distribution of individuals among these classes (evenness; Washington 1984). By treating each band as an individual ‘operational taxonomic unit’ (OTU), the richness (number of OTUs present) and the Shannon-Weaver diversity index (H'; Shannon & Weaver 1963) were calculated using PRIMER 5. As rare populations (less than 1% of the total community) might not be represented in a DGGE gel (Muyzer et al. 1993), the bacterial community richness (total number of OTUs detected) and diversity (Shannon-Weaver diversity index) calculated here only refer to the dominant bacteria populations and should be interpreted as an indicator of the minimum diversity of the bacterial community.

**Statistical analysis.** All correlation tests were done using Spearman rank R. STATISTICA 6 software was used, and a confidence level of 0.05 was considered in all test procedures.

The analysis of the banding profiles of the DGGE gels was done by multivariate tests using PRIMER 5 software. All tests were carried out on log (x+1) transformed data. The Bray-Curtis index was used as a similarity coefficient. A multi-dimensional scaling (MDS) analysis places samples in a multi-dimensional space, based on the similarities between them. In 2D ordination plots, similar samples are therefore placed together (Clarke 1993). The relative intensity of each band was therefore taken into account, which appears
to be more appropriate than using only presence/absence of the bands (Muylaert et al. 2002).

One-way analysis of similarity (ANOSIM) was used to test for statistical differences between the community composition of the 2 stations (115bis and 330). For each station, a 2-way crossed ANOSIM was used to test for statistical differences between the community composition at different depths (0 to 1 cm and 4 to 5 cm) and the sampling months February, April, and October 2003 (Clarke 1993). Whenever significant differences were found, pairwise tests were done.

The relationship between chl $a$ and the community structure was assessed by calculating rank correlations between similarity matrices derived from chl $a$ concentration in the sediment and the banding profiles (RELATE procedure) using the software package PRIMER 5. The same RELATE test was used between similarity matrices derived from bacterial counts per size class and bacterial biomass per size class.

One-way and 2-way analyses of variance (ANOVA) were performed to test for statistical differences in the Shannon-Weaver diversity index and OTU richness and in the bacterial counts and biomass for the months of February, April, and October 2003. Homogeneity of variances was tested using Hartley $F_{\text{max}}$, Cochran $C$, and Bartlett $\chi^2$, and data were transformed whenever necessary. Whenever significant differences were found, a posteriori Tukey HSD tests were performed. One-way ANOVA was used to test for significant differences between the 2 stations. A 2-way ANOVA was used to test for the effect of depth, time, and their interaction for bacterial counts and biomass at both stations and for the Shannon-Weaver diversity index and OTU richness at Stn 330. At Stn 115bis, due to the lack of replication in April in the 4 to 5 cm layer, a 2-way ANOVA could not be performed on the Shannon-Weaver diversity index and OTU richness. One-way ANOVA was then used to test for significant differences between the sampling dates in the 0 to 1 cm layer; a $t$-test was used to test for significant differences between the sampling dates February and October for the 4 to 5 cm layer, and a 1-way ANOVA was performed to test for statistical differences between the 2 sampling depths. STATISTICA 6 software was used, and a significance level of 0.05 was considered in all test procedures.

RESULTS

Environmental variables

At both stations, the surface and bottom water showed comparable concentrations of chl $a$, indicating a well-mixed water column (Fig. 2). At Stn 115bis, the chl $a$ concentration in the water was on most occasions higher than at Stn 330. Chl $a$ concentrations in the water at both stations started rising in February, peaked in April (48 mg m$^{-3}$ at Stn 115bis and 32 mg m$^{-3}$ at Stn 330), and decreased afterwards (Fig. 2). Smaller peaks were observed in July for both stations and only at Stn 330 in September, reaching values no higher than 17 mg m$^{-3}$.

At the sediment surface, chl $a$ concentrations were considerably higher at Stn 115bis than at Stn 330 (Fig. 3) throughout the sampling period and followed the patterns observed in the water column, with a peak in April. At Stn 330, other smaller peaks were observed in August and October 2003, corresponding to the deposition of the late summer and autumn blooms. For both stations, chl $a$ in the bottom water was correlated with the chl $a$ at the sediment surface (115bis: $R = 0.73$, $p < 0.05$; 330: $R = 0.62$, $p < 0.05$).

At Stn 115bis, the PAP ratio was relatively low in winter, started increasing in April, and remained relatively stable until it reached the highest value in October 2003. At Stn 330, the PAP ratio varied more abruptly, showing a similar value in winter, decreasing abruptly in April, and peaking 2 mo later. From June onwards, the PAP ratio decreased gradually (Fig. 3).

At Stn 115bis, chl $a$ concentration in the sediment was always highest at the surface (Fig. 4). This profile was most evident in April 2003 while deposition was occurring; the PAP ratio showed the opposite trend and increased with depth. At Stn 330, no clear vertical gradient could be found for either chl $a$ concentration or PAP ratio.

Water temperature was lowest in December (4.9°C), remained low until March, and increased from April onwards. Maximum values were observed in August (22°C), after which temperature decreased gradually.

![Fig. 2. Chl $a$ concentration in the surface and bottom water (mg m$^{-3}$) at Stns 115bis and 330 from October 2002 to October 2003. Error bars represent SE](image-url)
Bacterial biomass

Bacterial counts varied between 1.9 (± 0.3 SE) and 14.1 (± 0.9 SE) × 10^12 m⁻² at Stn 115bis and between 2.0 (± 0.9 SE) and 10.6 (± 4.6 SE) × 10^12 m⁻² at Stn 330 (Fig. 5). Bacterial counts were strongly correlated with bacterial biomass (R = 0.94; p < 0.001). Moreover, when taking into account the bacterial counts and biomass distributed per size class, a RELATE test revealed a strong relationship between both variables (ρ = 0.90; p < 0.01). Both tests indicated that bacterial counts and bacterial biomass varied in the same way. Therefore, the results are focused on bacterial biomass, assuming bacterial counts varied in the same way.

The bacterial biomass ranged from 0.2 (± 0.1 SE) to 2.8 (± 0.8 SE) gC m⁻² at Stn 115bis and from 0.2 (± 0.1 SE) to 1.9 (± 0.8 SE) gC m⁻² at Stn 330, showing different seasonal patterns when comparing both stations (Fig. 5). Bacterial biomass at Stn 115bis was much more variable than at Stn 330. Bacterial biomass was low in January and February 2003. Peak values were reached in March 2003, with intermediate values in April and May 2003. A progressive decrease toward initial values was then observed. Bacterial biomass at Stn 330 was lower compared to Stn 115bis, fluctuating around 1 gC m⁻² from January to July 2003. Prior to and after that period, bacterial biomass values were considerably lower. Biomass at 4 to 5 cm resembled that in the upper layer in all seasons at both stations (Fig. 5). Biomass showed no correlation with either chl a or temperature (p > 0.05) at either station.

Analyzing bacterial biomass for the sampling dates of February, April, and October 2003, no statistical differences were observed between the 2 stations (F = 0.243, df = 1, p > 0.05). At Stn 115bis, significant differences were found between sampling dates (F = 5.24, df = 2, p < 0.05) but not between layers (F = 0.08, df = 1, p > 0.05), nor for the interaction term (F = 0.35, df = 2, p > 0.05). Post hoc comparisons showed that April significantly differed from February and October. At Stn 330, significant differences were found between sampling dates (F = 5.99, df = 2, p < 0.05) but not between layers (F = 0.56, df = 1, p > 0.05) nor for the interaction term (F = 0.77, df = 2, p > 0.05). Post hoc comparisons showed that April and October were significantly different.

Bacterial community composition

Each band on the DGGE represents one OTU (Fig. 6). In this study, 52 different OTUs were identified. Only 25 of these were detected at both stations; 21 were found only at Stn 115bis and 6 only at Stn 330. Twenty-one of the 52 OTUs were detected during the 3 sampling campaigns, while 15 were restricted to 1 sampling period. Sixteen OTUs were detected in only 1 of the 2 layers. Five OTUs were only detected once. A 1-way ANOSIM considering all the samples showed significant differences between the 2 sampling stations (Global R = 0.721; p < 0.05).

MDS carried out for both stations separately revealed seasonal and vertical differences in bacterial
Fig. 4. Vertical profiles of chl a concentration (mg m⁻²) and ratio of phaeopigments to chl a + phaeopigments (PAP ratio) in the sediment at Stns 115bis and 330 in February, April, and October 2003. Error bars represent SE. Note different scales on top x-axes.
community composition (Fig. 6). When considering Stn 115bis, a 2-way crossed ANOSIM showed significant differences between the sampling months (Global R = 0.991, p < 0.05) and sediment depths (Global R = 0.788, p < 0.05). When considering pairwise tests (Table 1), all sampling months were significantly different from each other (p < 0.05). At Stn 330, the 2-way crossed ANOSIM again showed significant differences between sampling months (Global R = 0.884, p < 0.05) and depth layers (Global R = 0.726, p < 0.05). Pairwise tests (Table 1) revealed that communities in October were significantly different (p < 0.05) from those in February and April, while no differences were found between the 2 latter months (p > 0.05).

The bacterial community composition at both stations at all time and depths was significantly influenced by the chl a concentration in the sediment as shown by the significant RELATE test (ρ = 0.26, p < 0.05).

Bacterial biodiversity

Both OTU richness and the Shannon-Weaver diversity index were relatively higher at Stn 115bis than at Stn 330 (Fig. 7). At Stn 115bis, OTU richness and Shannon-Weaver diversity index were similar for both sediment layers and for all seasons, with lower values detected only in the 4 to 5 cm layer in April. The OTU richness and Shannon-Weaver diversity index of the communities at Stn 330 were lower in the surface layer than in the 4 to 5 cm layer and showed minor changes with time. The sample from 4 to 5 cm in April at Stn 115bis was removed from the statistical analysis, since it had no replication. A 1-way ANOVA performed on the Shannon-Weaver diversity index showed significant differences between the 2 stations (F = 15.20, df = 1, p < 0.001). At Stn 115bis, no significant differences were found between sampling dates for the 0 to 1 cm layer (F =
0.035, df = 2, p > 0.05); a t-test showed no significant differences between February and October for the 4 to 5 cm layer (t = 0.071, p > 0.05), and a 1-way ANOVA showed no significant differences between the 2 depth layers (F = 0.129, df = 1, p > 0.05). At Stn 330, a 2-way ANOVA showed significant differences between the depth layers (F = 14.310, df = 1, p < 0.05) but not between sampling months (F = 0.540, df = 2, p > 0.05), nor when combining both effects (F = 1.312, df = 2, p > 0.05). The same analyses performed on the OTU richness showed the same significant differences as shown by the bacterial Shannon-Weaver diversity index.

DISCUSSION

Environmental variables

As described previously (Vanaverbeke et al. 2004a,b, M. Steyaert et al. unpubl.) a strong phytoplankton bloom occurred in spring as indicated by the seasonal pattern of the concentration of chl a in the water column. The higher chl a concentration in the water column at Stn 115bis during the spring phytoplankton bloom was to be expected, since primary production is higher closer to the coast (Joint & Pomroy 1993).

Patterns of chl a concentrations in the sediment also showed a strong seasonal signal as already described (Vanaverbeke et al. 2004a,b, M. Steyaert et al. unpubl.), but in contrast to the water column values, differed considerably between the stations. Chl a concentration at the sediment surface at Stn 115bis was about 10× higher than at Stn 330. This, together with the absence of clear vertical profiles in the sediment, corroborated the idea that Stn 330 has more permeable sediment, where stronger bottom water currents prevent the deposition of sedimenting phytodetritus (Precht & Huettel 2004) and induce subsurface chl a peaks (Ehrenhauss & Huettel 2004, Ehrenhauss et al. 2004a). At Stn 115bis, vertical gradients were very obvious, especially during the deposition of the phytoplankton bloom, corroborating that 115bis is a deposition station.

The pattern of the PAP ratios showed little seasonal variation at Stn 115bis compared to Stn 330 (Fig. 3). The comparatively higher PAP ratio at Stn 330 following bloom senescence indicate a rapid degradation of organic matter, which is typical of permeable coastal sands (Ehrenhauss et al. 2004b). Advective transport allows fast removal of decomposition products (Huettel et al. 1998) resulting in an up and down fluctuation of the PAP ratio at Stn 330. At Stn 115bis, there was a carbon build-up (phytodetritus derived) in the sediment during spring and subsequent degradation and mineralization in late summer as previously reported in the North Sea for this type of sediment (Boon & Duineveld 1998, P. Provoost et al. unpubl.). Therefore, the PAP ratio did not show major fluctuations but slowly increased toward late summer and autumn.

Table 1. Pairwise test for bacterial community composition in the different sampling months for both stations. *p < 0.05

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<th>Stn 115bis</th>
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<th>Stn 330</th>
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<tr>
<td></td>
<td>R statistic</td>
<td>Signif.</td>
<td>R statistic</td>
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<tr>
<td>Feb vs. Apr</td>
<td>0.958</td>
<td>0.010*</td>
<td>0.500</td>
</tr>
<tr>
<td>Feb vs. Oct</td>
<td>1.000</td>
<td>0.025*</td>
<td>1.000</td>
</tr>
<tr>
<td>Apr vs. Oct</td>
<td>1.000</td>
<td>0.033*</td>
<td>0.944</td>
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Fig. 7. Shannon-Weaver index (H') and number of OTUs identified in the bacterial community from Stns 115bis and 330 in the sampling campaigns in February 2003, April 2003, and October 2003, in the 0 to 1 cm and 4 to 5 cm sediment layers. Error bars represent SE.
Bacterial biomass

Bacterial counts make no distinction between dead, active, or inactive bacteria. Coastal marine sediments contain a large fraction of dead or inactive bacteria (Luna et al. 2002), and the percentage of active bacteria may change rapidly (Créach et al. 2003). In deep Mediterranean sediments, the number of active bacteria depends on the organic substrate in the sediment derived from the photic layer (Luna et al. 2004). However, according to the same study, no relationship was observed between total bacterial counts and the content of organic substrates in the sediment. In our study, bacterial biomass, calculated from bacterial counts, also had no correlation with either temperature or chl a (as an indicator of the amount of labile OM) at either station. Both temperature and food availability seem to be key factors influencing the response of benthic bacterial communities to an input of organic matter (Graf et al. 1982, Boon et al. 1998), and a relationship between bacterial community composition and chl a concentration in the sediment was detected. Bacterial biomass and/or counts alone might not be a reliable indicator of changes in bacterial activity, although coupled with other data, like DGGE banding profiles, they can help achieve a better understanding of changes in the bacterial community. A significantly higher biomass was detected in April at Stn 115bis (compared to February and October), which can be coupled with the seasonal shifts in bacterial community composition observed at this station (discussed below, in ‘Bacterial community composition and diversity’). Also, a significantly lower biomass detected in October at Stn 330 might be the cause of this community composition being significantly different from that in February and April (discussed below, in ‘Bacterial community composition and diversity’).

Methodological considerations

As with any PCR and DNA based techniques, DGGE of PCR-amplified 16S rDNA (Muyzer et al. 1993) has its own methodological limitations, including the inhibition of PCR amplification by co-extracted contaminants, deferential amplification, or formation of artefactual PCR products or contaminating DNA, and 16S rRNA sequence variations. Co-migration of DNA, single bands representing more than 1 bacterial strain or slightly different rRNA gene sequences resulting in multiple bands are problems that can occur in DGGE gels (Nübel et al. 1996, Palys et al. 1997, Vallaeys et al. 1997). Nevertheless, PCR-mediated analysis of 16S rRNA is a powerful tool for the determination of microbial diversity of environmental ecosystems (von Wintzingerode et al. 1997), and DGGE techniques are quite effective when characterizing bacterial community structure, particularly for monitoring changes in occurrence and/or relative frequency of the different bacterial populations (Fromin et al. 2002). Since samples were treated in the same way during the whole study, the method allows for a proper comparison of the results beyond a mere qualitative approach (Fromin et al. 2002). The Shannon-Weaver index is the most common diversity index generally used by ecologists (Washington 1984) and has also been applied to DGGE fingerprints to estimate bacterial diversity (e.g. Nübel et al. 1999, Boon et al. 2002, Dilly et al. 2004, Haack et al. 2004, Gafan et al. 2005, Xia et al. 2005, Lagacé et al. 2006), even though in a DGGE gel, populations representing less than 1% of the total community might not be represented (Muyzer et al. 1993).

Bacterial community composition and diversity

Our results showed that bacterial community composition was significantly different at both stations. Only 6 of the 52 OTUs were exclusively detected at Stn 330, while 21 OTUs were found exclusively at Stn 115bis (data not shown). OTU richness and Shannon-Weaver diversity index were mostly higher at Stn 115bis than at Stn 330, especially at the sediment surface. This may be a reflection of higher food availability at Stn 115bis, since substrate availability affects the bacterial community composition (Van Hannen et al. 1999a, Muylaert et al. 2002) and a high concentration of potentially available substrate can sustain a higher bacterial diversity (Luna et al. 2004). Even though we did not detect the whole community by means of DGGE, the results on the dominant OTU (relative density higher than 1%) seem to indicate a relatively more diverse community at Stn 115bis.

In bacterioplankton, both free-living bacteria and particle-associated bacteria can occur as 2 distinct communities (Fandino et al. 2001, Riemann & Winding 2001, Rooney-Varga et al. 2005), although this should not be generalized (Sapp et al. 2007). If this were true for the sediment as well, we would expect that the establishment of free-living bacteria populations would become more difficult at Stn 330 due to the high bottom water currents and the permeability of the sediment. This would be more evident at the surface of the sediment, where our results indicate the lowest bacterial diversity for Stn 330. At Stn 115bis with no such currents, both free-living and attached bacteria could coexist as they can in the plankton.

Another factor that could be responsible for different community composition at both stations is the possible coexistence of aerobic and anaerobic bacteria at Stn 115bis. At this station, the sediment becomes reduced...
after a spring bloom (M. Steyaert et al. unpubl.), while at Stn 330, sediments at the depths studied are well aerated (Vanaverbeke et al. 2004a,b).

Significant vertical differences in bacterial community composition were observed at both stations, which might have different causes at each station. Only 1 OTU was found exclusively at the surface layer at Stn 330. This OTU was also found exclusively at the surface layer at Stn 115bis. All other OTU encountered at the surface were also found in the 4 to 5 cm layer, generally with relatively higher densities. Twelve OTUs were found exclusively in the 4 to 5 cm layer at Stn 330, half of which were also observed at the sediment surface at Stn 115bis (data not shown). The differences in the community composition between the 2 layers at Stn 330 were a consequence of the non-detection of certain bacterial populations at the surface of the sediment. Bacterial populations in the upper layers of Stn 330 might not be able to cope with the high hydrodynamic stress (advective currents through the sediment) prevailing there (Precht & Huettel 2004), which results in both lower Shannon-Weaver diversity index and number of OTUs, and different bacterial community composition compared to deeper layers.

At Stn 115bis, 10 OTUs were found exclusively at the surface and 8 OTUs only in the 4 to 5 cm layer (data not shown), indicating that at this station different communities might have developed at each layer. If the sample from the 4 to 5 cm layer in April at Stn 115bis was omitted, no significant differences were observed in the Shannon-Weaver diversity index and OTU richness between layers, even though community composition was different. At this sampling occasion, it was only possible to obtain 1 replicate for the 4 to 5 cm layer. The DNA extraction and consequent PCR was not as successful as for other samples. Since bacterial biomass was not lower, this might be related to chemical inhibition. Because there was no replication, results referring to the 4 to 5 cm layer at Stn 115bis in April should be interpreted with care.

Steep vertical gradients in chl a concentration and PAP ratio were registered at station 115bis. Moreover, deep sediment layers at this station showed strong negative redox potential values and a build-up of NH₄⁺ (M. Steyaert et al. unpubl.). Vertical differences in the benthic bacterial community have previously been reported (Luna et al. 2004) in Mediterranean sediments with oxygen depletion problems in which the vertical differences in the bacterial community were closely related to redox potential changes (Uraakawa et al. 2000).

The statistical analysis of the DGGE fingerprints also showed that bacterial community composition at both stations changed with time. In the planktonic system, shifts in plankton species composition are responsible for changes in the attached bacterial community composition (Rooney-Varga et al. 2005). Carbon-rich mucilage sedimentation, as secreted by Phaeocystis, can be a post-bloom food source (Riebesell et al. 1995). Experimental degradation of such compounds in agar from different sources and under different oxygen conditions presented very different microbial communities with only a few overlapping species (Janse et al. 2000). Changes in organic matter quality and quantity can induce shifts in bacterial community structure (Luna et al. 2004). Therefore, seasonal changes in food availability and oxygenation are likely to produce distinct bacterial communities. This was supported by the significant RELATE test observed between the bacterial community composition and the concentration of chl a in the sediment.

Temporal differences in bacterial composition were stronger at Stn 115bis, where all sampling months were significantly different from each other. Also, the percentage of OTUs that were present in the 3 sampling campaigns was lower at Stn 115bis than at Stn 330 (data not shown), indicating higher shifting of the bacterial populations at this station. Moreover, bacterial biomass was higher in April, implying shifts in bacterial biomass from February to April and again from April to October. Stronger variations in chl a concentration in the sediment at Stn 115bis, probably also related to shifts in bacterial biomass, would imply higher temporal differences in the bacterial community composition at Stn 115bis.

Besides food availability, the oxygenation of the sediment can also change more drastically at Stn 115bis (M. Steyaert et al. unpubl.), which is not the case for Stn 330 where the redox potential remains positive throughout the studied sediment depths (Vanaverbeke et al. 2004a,b).

At Stn 330, the bacterial community composition in October differed from the other 2 sampling campaigns, which was also the case for bacterial biomass. Temporal variability was also present at this station, albeit not strongly.

Even though the bacterial community composition changed with time, such differences were not detected in the Shannon-Weaver diversity index and OTU richness at Stn 330, nor at Stn 115bis when the sample from the 4 to 5 cm layer in April was omitted. At each station, we observed that with time OTUs were disappearing, new ones were emerging, and others were always present but their density changed. The regular seasonal environmental changes might be responsible for maintenance of diversity, since environmental fluctuations can provide temporal niche opportunities, allowing species to coexist (Chesson & Huntly 1997). This does not mean that the same bacteria were not always present, but changes in dominance might push certain bacteria above the detection limit (Hedrick et al. 2000).
CONCLUSIONS

We have shown that bacterial community composition differed when comparing fine and coarser marine sediments, and our results indicated that diversity in general tends to be higher in finer sediments. Bacterial community composition also differed vertically within the sediment, but not in the same way for both sediment types. In the fine sediment, 2 relatively distinct communities developed at the surface and deeper in the sediment. In coarser sediments, vertical differences were related mostly to the non-detection of certain bacteria populations at the surface of the sediment where hydrodynamic stress is stronger. Seasonal patterns in food availability played a key role in the bacterial community composition, which changed more drastically within fine sediment, where the input of organic matter was stronger.

These changes in bacterial community may also be of importance when considering higher trophic levels. Changes in nematode community composition after bloom sedimentation toward feeding types that feed on bacteria has been observed at these 2 stations (Vanaverbeke et al. 2004b, M. Steyaert et al. unpubl.). Vanaverbeke et al. (2004b) hypothesized that an increase in nematode species richness could have been caused by an increase in bacterial diversity. Indeed, a shift in bacterial community composition was observed.

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