Chancres in bacterial community structure in seawater mesocosms differing in their nutrient status

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ABSTRACT: Quantitative and qualitative changes in bacterial communities from the Mediterranean Sea were analysed under eutrophication conditions simulated in batch mesocosms (addition of inorganic nutrients or phytoplanktonic lysate). A wide variety of methods including traditional microbial ecology techniques, molecular biology and flow cytometry were combined to determine abundances, production, cell size, activity, culturability, and genetic and taxonomic diversity. In all mesocosms, the increase in biomass was rapidly controlled by protozoan grazing. Morphological and physiological changes were observed during the growth phase of bacteria and under grazing pressure. The proportion of medium-size and culturable cells increased during the growth phase. Grazing eliminates preferentially active and medium-sized cells within communities regulating bacterial productivity. Small and large cells were produced as a consequence of grazing pressure, and the large active cells contributed to the remaining productivity after grazing. Although grazing had an effect on the genetic diversity of bacterial communities by eliminating some populations, other species were preserved. It seems that some species such as Alteromonas macleodii may have developed defence strategies to escape predation. We hypothesize that such species may escape grazing by producing small and/or large cells during their growing phase.

KEY WORDS: Mesocosm · Eutrophication · Bacteria · Biomass · Production · Diversity · Cell-specific activity · Grazing

INTRODUCTION

Heterotrophic bacteria are an abundant and important component of aquatic ecosystems. Their role in the functioning of biogeochemical cycles, energy flow and secondary production is now well established (Ducklow & Carlson 1992). Although top-down and bottom-up controls are widely known concepts in aquatic microbial ecology (Pace & Cole 1994), there is still disagreement as to whether the abundance and productivity of pelagic bacteria are determined mainly by predators or nutrients (Pernthaler et al. 1997). Further-
more, from a structural point of view, both predators and nutrients may be major forces shaping the genotypic and phenotypic composition of bacterial communities (Jürgens & Güde 1994).

Heterotrophic nanoflagellates and ciliates are usually the major consumers of bacteria in aquatic systems (Jürgens & Güde 1994). However, information about the strategies involved in the predator-prey interactions is scarce. It has been suggested that very small and large bacteria are partly or totally protected from protozoan grazing (Jürgens & Güde 1994). If filamentous and very small bacteria exhibit an ecologically significant defense strategy against protozoan grazing, it is not known to what extent their selection under grazing pressure affects the taxonomic structure and activity of natural communities and, thus, the ecological role of these communities. The activity level of filamentous cells remains unclear. On the one hand, Pernthaler et al. (1997) suggested that filamentous cells may be inactive; on the other hand, Gasol et al. (1995) suggested that the probability of a bacterium to be active was a linear function of its size. If we accept that the larger the cells, the higher their activity, the selection of large bacteria may result in the selection of very active and productive cells. Inversely, the selection of small cells may result in the selection of non-active cells.

There is a need to further understand the role of bottom-up and top-down controls on the physiological, taxonomic and functional diversity of natural communities and to understand how this diversity can be affected by changing environmental factors. Bottom-up control may play an important role in coastal eutrophized areas, depending on the nature and availability of inorganic and organic nutrients. Although these questions have long been unanswered due to methodological limitations, the recent development of new techniques including flow cytometry and molecular techniques may help to obtain new information.

The introduction of molecular biological techniques in microbial ecology has provided new approaches to the investigation of the structure of microbial communities (Ward et al. 1990, Muyzer et al. 1995). Phylogenetic analyses provide information on bacterioplankton population diversity, as well as nucleic acid probes, which can be used to establish the significance of cultured microorganisms and to study the temporal and spatial distributions of microbial taxa in marine ecosystems (Delong et al. 1993, Mulins et al. 1995).

The aim of this study was to analyse the effect of 2 simulated eutrophication conditions (addition of inorganic and organic nutrients) on changes in morphological, physiological and genetic diversity of bacterial communities. Thus, this study combines classical microbiological methods and techniques commonly used in microbial ecology with the recent cytometric and molecular biology tools.

### MATERIAL AND METHODS

**Experimental set-up.** The mesocosms consisted of 3 rectangular tanks of 401 capacity, 60 cm height, 60 cm width and 110 cm length. Each mesocosm was filled with 300 l of natural seawater collected from 1 m depth at a station (42° 31' N, 03° 11' E) located 2 miles (~3.2 km) off Banyuls-sur-Mer, France (Mediterranean Sea). Natural seawater was filtered through a 200 μm nylon mesh. This filtration removed large metazoans and detritus. Each mesocosm was temperature-controlled to fit the temperature of Mediterranean coastal waters at the time of the experiment by placing the mesocosms into larger tanks (30 cm height, 80 cm width, 140 cm length) continuously alimanted by natural seawater pumped from the shore. The temperature of the water was 17.7 to 19.6°C during the experiment. Two mesocosms (Mesocosms A and B) were illuminated by a fluorescent ramp consisting of 5 tubes (36 W, 120 cm) providing illumination on a 12:12 h light-dark period. Mesocosm C was non-illuminated. Photosynthetically active irradiance (PAR) was measured with a Li-cor quantum sensor. The PAR at the upper part of Mesocosms A and B was 200 μE m⁻² s⁻¹. The mesocosms were mixed by means of an immersed pump taking water from 2 opposite bottom corners and driving the water back below the surface.

**Mesocosm preparation.** The chemical compositions of seawater before and after the different treatments are reported in Table 1. Mesocosm A was used as a control to follow changes within the initial community without any nutrient addition. Mesocosm B was supplemented with inorganic nitrogen (NH₄, NO₃) and phosphorus (PO₄). Mesocosm C was supplemented with a mixture of 2 phytoplanktonic species: *Chaetoceros sp.* and *Porphyridium cruentum*. Each culture was first frozen and immediately autoclaved to facilitate cell lysis and then the introduction of molecular biological techniques, which can be used to establish the significance of cultured microorganisms and to study the temporal and spatial distributions of microbial taxa in marine ecosystems (Delong et al. 1993, Mulins et al. 1995).

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<table>
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<th>Nutrate</th>
<th>Ammonium</th>
<th>Phosphate</th>
<th>N.P</th>
<th>DOC</th>
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<td>0.13 (0.03)</td>
<td>0.03 (0.02)</td>
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<td>0.05 (0.03)</td>
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<tr>
<td>Mesocosm B</td>
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<td>1.98 (0.09)</td>
<td>0.66 (0.03)</td>
<td>6.7</td>
</tr>
<tr>
<td>Mesocosm C</td>
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<td>1.51 (0.05)</td>
<td>0.76 (0.05)</td>
<td>53.1</td>
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filtered onto 0.2 µM membrane filters (Millipore). Four liters of each culture were mixed and added to the mesocosm. Mesocosm C was supplemented after 113 h since it was not possible to process all mesocosms at the same time.

**Sampling strategy.** Five liters from each mesocosm were taken daily to analyse nutrients, phyto- and bacterioplankton counts, bacterial production and chlorophyll a. The temporal evolution of total bacterial counts was used to determine the sampling times at which all other parameters including the bacterial genetic diversity were also analysed. For these points, 10 l were taken and mixed. These sampling times were chosen corresponding to the starting time of each mesocosm, to the first significant increase in bacterial counts, and to the peak, and then after a significant decrease of total counts and at the end of experiments defined by the stabilisation of total counts.

**Nitrogen and phosphorus analysis.** For nitrate and phosphate determination, subsamples were collected in polyethylene bottles and directly frozen (−18°C) for later analysis. Nitrate and phosphate concentrations were determined with a Skalar autoanalyzer according to Treguer & Le Corre (1975). For ammonium analysis, subsamples were collected in 100 ml glass bottles and processed according to Koroleff's (1976) manual method. The reagents were added immediately after collection and measurements were performed 24 h later using a spectrophotometer (Spectronic 401). The optical density was also measured on each original sample in order to take into account the turbidity of the sample.

**Culturable counts.** Culturable bacteria were enumerated by plating 100 µl of serial dilutions of each water sample onto marine agar plates (MA Difco, Detroit, MI). In preliminary assays R2A medium with a seawater base was also used; however, marine agar was chosen for the assays since it yielded higher counts. After 7 d of incubation at room temperature (20 ± 3°C) colony forming units (CFU) were enumerated. Counts did not increase after prolonged incubation.

**Flow cytometry analysis of bacteria.** Total counts of SYTO 13-stained bacteria were performed with a FACSCalibur flow cytometer (Becton Dickinson, San Jose, CA) equipped with an air-cooled argon laser (488 nm, 15 mW). Living samples were stained with SYTO 13 (Molecular probes, Eugene, OR) at 5 µM for 30 min. One µM of fluorescent beads (Polysciences, Warrington, PA) was systematically added to samples in order to normalise cell fluorescence emissions and scatters. According to their fluorescence (FL1) and right angle light scatter (RALS, related to cell size) signatures, different bacterial populations could be discriminated and enumerated. Bacterial enumerations by flow cytometry performed as described above were found to be in close concordance with bacterial enumerations by epifluorescence microscopy after 4,6-diamidino-2-phenylindol (DAPI) staining (del Giorgio et al. 1996a).

In order to calibrate the flow cytometric scatter measurements, bacterial cell lengths and biovolumes were estimated from epifluorescence examination of DAPI stained cells of fixed samples (n = 17) (Troussellier et al. 1995). The relationship between RALS and cell volumes showed high and significant correlation (r = 0.869, p ≤ 0.0001). The same correlation was obtained when using cell lengths rather than biovolumes (Troussellier et al. 1999). RALS values were not transformed as equivalent spherical diameters. Several bacterial cell populations were discriminated from RALS, and a green fluorescence (FL1) cytogram was obtained for each sample. Cells with high RALS values were related to long rods observed by epifluorescence microscopy. From one sample to another, these populations were not exactly located at the same place on cytograms. Thus, in order to objectively group the different populations observed in all analysed samples, a clustering method (Ward’s algorithm; Ward 1963) was applied to all the identified populations. Each population was characterised by its RALS and FL1 mean values. Clustering was carried out with the ‘R package’ (Legendre & Vaudor 1991). Three main clusters with an increasing estimated mean size (cell length) (C1 ≤ 1.0 µm, 1.0 µm < C2 ≤ 2.5 µm, and C3 > 2.5 µm) were discriminated in each sample.

**Microscopic counts of protozoa.** The abundance of protozoa was determined by epifluorescence microscopy (Leitz, Laborlux D) after DAPI staining. Twenty ml of glutaraldehyde-preserved (0.5% final concentration) water samples were stained with DAPI (0.1 µg l−1, final concentration) for 15 min. Filters were mounted on microscopic slides and stored at +4°C until examination. Stained protists were collected by filtration on a 0.8 µm pore size black filter (Nucleopore). Pico-sized (0.2 to 2 µm in diameter) and nano-sized (2 to 20 µm in diameter) micro-organisms were identified, counted and measured at a magnification of 1250× and 625×, while micro-sized (20 to 200 µm in diameter) micro-organisms were analysed at a magnification of 125×. A minimum of 100 organisms per filter was counted. Autotrophic species were distinguished from heterotrophs by the red autofluorescence of chlorophyll a observed under blue light excitation. Dinoflagellates, flagellates and ciliates were enumerated distinctly. Data presented in this paper concern only the heterotrophic protozoa.

**Respiring cell counts.** 5-cyano-2,3-dimethyl tetrazolium chloride reduction (CTC; Polyscience Europe) was used to identify active respiring cells by the method of Rodriguez et al. (1992). CTC was added (5 mM final concentration) to 2 ml samples. CTC incubations were
performed in the dark at room temperature during 8 h. The fraction of respiring cells was determined as previously described (Joux et al. 1997).

**Bacterial activity and production.** Four 10 ml subsamples were incubated in the presence of 4 different concentrations (6.5, 13, 26 and 39 nM) of tritiated thymidine (Amersham 84 Ci mmol⁻¹) for 1 to 2 h in the dark at the temperature of the mesocosms. After incubation, cold trichloroacetic acid (TCA) was added (5% final concentration) and the samples were filtered through 0.2 μm pore size cellulose acetate membrane. Radioactivity associated with the filters was estimated by liquid scintillation. Incorporation rates expressed in pmol 1⁻¹ h⁻¹ at the different thymidine concentrations were calculated and plotted against the added thymidine concentrations. The maximum incorporation rates were estimated by best fitting a hyperbolic function to the experimental data using a software based on the least squares criterion (Servais 1995). Data presented in the paper are maximum incorporation rates.

Bacterial production (P) expressed in cells produced per unit of volume and time was calculated by multiplying the maximum incorporation rate by a conversion factor equal to 0.5 x 10¹⁸ cells produced per mole of thymidine incorporated into cold TCA insoluble material (Servais 1992).

**Incorporation of ³H-leucine.** Four 10 ml subsamples were incubated in the presence of 4 different concentrations of leucine (2 nM of tritiated leucine, Amersham 151 Ci mmol⁻¹, with 0, 25, 50, 75 nM non-radioactive leucine) for 1 to 2 h in the dark at the temperature of the mesocosms (Servais 1995). After incubation, cold TCA was added (5% final concentration) and subsamples were heated at 85°C for 30 min after acidification following the procedure proposed by Kirchman et al. (1985) to specifically measure the incorporation into proteins. After cooling, the samples were filtered through 0.2 μm pore size cellulose acetate membrane. Maximum incorporation rates into proteins were calculated as for thymidine incorporation. Data presented in the paper are maximum incorporation rates into proteins.

**Growth rate.** Growth rates (μ, h⁻¹) were calculated using the following model: specific growth rate μ = ln(1 + P/N), where P is cellular production (cells 1⁻¹ h⁻¹) and N is bacterial abundance (cells 1⁻¹) (Lovejoy et al. 1996). Doubling time (h) was calculated as (ln2)/μ.

**Denaturing gradient gel electrophoresis (DGGE) analysis.** Water samples were taken from the mesocosm and approximately 2.5 x 10⁸ bacterial cells were collected by filtration (Millipore GVWP 02500, 0.22 μm, 25 mm O). Filters were frozen in liquid nitrogen immediately. DNA and rRNA were extracted and purified from filters as described by Teske et al. (1996). The extracted genomic DNA was used as target DNA in the polymerase chain reaction (PCR; Saiki et al. 1988) to amplify fragments of bacterial 16S rRNA genes suitable for subsequent DGGE analysis using primer combinations 341F-GC/907RC and 341F-GC/907RM (Muyzer et al. 1995). At the 5’ end of the forward primer 341F-GC (sequence 5’-CCT AGC GGA GCC AGC AG-3’) a 40 bp GC-rich sequence (GC-clamp) was attached (sequence: 5’-CGC CCG CCC CCG CCG CCC GTC CCG CCC G-3’). The 2 reverse primers used differed in 1 nucleotide position, at which 907RM has a degeneracy at a position with some variation in the bacterial target site (primer 907RC: 5’-CCG TCA ATT CCT TTG AGT TT-3’ and primer 907RM sequence: 5’-CCG TCA ATT CMT TTG AGT TT-3’). PCR products obtained with the 2 primer sets were used for independent DGGE analyses. DGGE was performed essentially as described by Muyzer et al. (1998) using linear denaturing gradients of 10 to 70% or 20 to 80% urea and formamide. After DGGE, electrophoresis gels were stained in an ethidium-bromide solution (0.5 mg 1⁻¹) and photographed with a Polaroid system.

For cluster analysis of banding patterns the gel images were analysed for similarities of banding patterns by scoring absence or presence of apparent bands with 0 and 1, respectively. In this way a binary matrix was created which was subsequently analysed with the UPGMA option in PAUP* using the correction of Nei & Li (1979).

**Molecular identification.** Preparation of genomic DNA from pure cultures isolated from mesocosms and amplification of 16S rDNA by PCR was done as described by Rainey et al. (1996).

**Phylogenetic analyses.** Sequences were manually aligned with sequences published previously. These were stored in the DSMZ internal database consisting of more than 5000 16S rDNA sequence entries, including those from the Ribosomal Database Project (Maidak et al. 1997) and EMBL. Similarity values were transformed into phylogenetic distance values that compensate for multiple substitutions at any given site in the sequence (Jukes & Cantor 1969). The neighbor joining method contained in the PHYLIP package (Felsenstein 1993) was used in the construction of phylogenetic dendrograms. All analyses were done on a SUN SparcII workstation.

**RESULTS**

**Nutrients**

The initial seawater was characterised by low concentrations in phosphate, but the N:P ratio was close to the Redfield ratio (Table 1). The phosphate concentra-
tions fell close to the detection limit after 144 h in Mesocosm A (Fig. 1A). Ammonium concentrations were maintained constant at a low concentration. In Mesocosm B, phosphorus was consumed rapidly during the first 5 d and then stabilised but was never exhausted (Fig. 1B). Inversely, nitrogen fell close to the detection limit after 72 h. Both phosphorous and nitrogen (nitrate and ammonium forms) concentrations were high after addition of the phytoplankton lysate in Mesocosm C and were never exhausted (Fig. 1C).

**Bacterial and protistan abundances**

Initial bacterial numbers were about $5 \times 10^5$ cells ml$^{-1}$ in all mesocosms (Fig. 2). Although bacterial abundances increased during the first few days in all mesocosms, this increase was more important in Mesocosm B ($5 \times 10^6$ cells ml$^{-1}$ after 120 h). The peaks of abundances were observed after 136, 120 and 122 h in Mesocosms A, B and C, respectively. The increase in total bacterial counts was followed by a rapid increase in protozoa counts with a delay of approximately 24 h. The increase in the number of protozoa resulted in a rapid and important decrease of bacterial counts. Then, the grazing pressure was reduced with decreasing bacterial counts. After grazing, the total numbers of bacteria were close to the initial counts.

**Bacterial activity**

The rate of $^{3}$H-thymidine incorporation into DNA was used to estimate bacterial cell replication (Fuhrman & Azam 1982) and that of $^{3}$H-leucine into bacterial proteins in order to estimate the protein synthesis rate (Kirchman et al. 1985). Although both methods are usually well correlated (Servais 1992), they give complementary information on the bacterial metabolism as one measures cell multiplication and the other one biomass synthesis (Shiah & Ducklow 1997).
Maximum incorporation rates of thymidine and leucine presented the same trends, although slight differences were observed in some situations (Fig. 3). Bacterial production as determined by the rates of leucine incorporation into protein and thymidine incorporation into DNA were low in the initial seawater (respectively 100 pmol leucine l⁻¹ h⁻¹ of and 2 pmol thymidine l⁻¹ h⁻¹). The incorporation rates increased slightly during the first 3 d in the control (Mesocosm A) and then stabilised. In Mesocosm B, both incorporation rates increased exponentially until Day 2 and then high activity values were maintained until Day 6 when they fell rapidly within 24 h to values which remained higher than those measured at the beginning of the experiment. Thereafter, leucine and thymidine incorporation rates remained low. In Mesocosm C, leucine and thymidine incorporation rates increased strongly within 48 h after nutrient addition (at 113 h) and then decreased very rapidly. Maximum values reached on Day 6 were higher than those obtained in other mesocosms. However, these high values decreased rapidly compared to those reported in Mesocosm B, which were lower but remained high and almost constant during the last 4 to 5 d.

**Culturability and activity of bacterial cells**

The fraction of culturable cells was low in the initial seawater (less than 0.1%) and increased during the growing phase in all mesocosms (Fig. 4). In the control, the culturability remained low up to 144 h, was maximum (30%) at 185 h, and then stabilized between 20 and 30%. The number of actively respiring bacteria increased more rapidly and was higher than culturable counts during the growing phase of bacteria (from 0 to 144 h). Active cell counts decreased sharply up to 185 h. In Mesocosm B, both active and culturable cells
increased up to 113 h. The fraction of respiring cells decreased rapidly and remained low after 185 h. Conversely, an important fraction of cells (18 to 35%) retained culturability during the grazing phase. In Mesocosm C, there was a concomitant increase in both active and culturable cells followed by a rapid decrease within 24 h. Thereafter, both fractions remained constant and equal to approximately 20%. In all mesocosms, although total cell counts were almost similar at time 0 and after grazing, there were important changes in the community structure.

### Bacterial cell biovolume

In the initial seawater, C1 cells contributed about 50–60% to total bacterial biovolume (Fig. 5). In all mesocosms the contribution of C1 cells to total biovolume decreased continuously during the growing phase and was inversely related to an increasing contribution of C2 cells. This inverse trend was more rapid and pronounced in Mesocosm B. In Mesocosm C the trend was similar to that in Mesocosm A during the first slight increase of total bacterial counts and then was amplified within 24 h after nutrient addition, corresponding to the rapid shift in total counts. Thereafter, there was a rapid decrease in the contribution of C2 cells which was concomitant with the decrease in total bacterial counts under grazing pressure. There was a pronounced inverse trend of increasing contribution of C1 and C3 cells and decreasing contribution of C2 cells to total biovolume.

### Taxon affiliation of selected isolates

Of the 430 strains isolated from Mesocosms A, B and C, 36 strains were picked to represent the variation of pigments and colony forms. The majority of the analysed organisms belong to the alpha and gamma subclass of *Proteobacteria* (36 and 44%, respectively), while the number of Gram-positive bacteria was much smaller (16%). The majority of strains clustered close to the described species, e.g. *Sulfitobacter pontiacus* (99.0 to 99.7% similarity) and *Roseobacter algicola* (97.0 to 100% similarity) in the alpha subclass of *Proteobacteria*, and members of *Pseudoalteromonas haloplanktis* (99.4 to 100% similarity), *Alteromonas macleodii* (95.5% similarity), *Marinobacter hydrocarbonoclasticus* (95.6 to 96% similarity) and *Vibrio splendidus* (95.3 to 97.6% similarity) in the gamma subclass of *Proteobacteria* (see Table 2).

### Genetic fingerprints from mesocosm water samples and identification of dominant isolates

The genetic diversity of bacterial communities was also studied at different sampling points of each mesocosm by means of DGGE. According to the DGGE profiles (Fig. 6), there were marked changes in the number and position of bands suggesting temporal modifications of the community structure. The number of apparent bands slightly decreased during the increase of bacterial biomass and then increased again after grazing. Furthermore, important changes in the intensity of the different bands suggested changes in the relative abundance of the different 16S rDNA defined populations. Cluster analysis of the banding patterns was performed; the resulting dendrogram is shown in Fig. 7. The samples fell into 2 clusters that represented the status before and after the peak of grazing activity in each mesocosm. Furthermore, consecutive sampling times of mesocosms clustered together (e.g. A065 and A113, B065 and B113, C144 and C185). Also, the patterns of B281 and C281 were similar.
DISCUSSION

The 3 mesocosm experiments presented herein were done to study the effect of bottom-up and top-down controls on the structure of bacterial communities as affected by 2 laboratory-simulated eutrophication conditions. Although these results did not determine if similar processes occur within natural ecosystems and did not take into account the potential role of viruses, the combination of methods used in this study have been useful for formulating some new hypotheses and suggesting new experimental approaches in aquatic microbial ecology.

The bacterial production was probably limited in Mesocosms A and B by phosphate and nitrogen, respectively. Similar P limitation has already been demonstrated for Mediterranean bacterial communities (Thingstad & Rassoulzadegan 1995). In all situations, including the control, we observed a more or less important increase of total bacterial counts rapidly controlled by grazing. In all cases, the abundances before nutrient supply and after grazing were almost similar. However, the production, physiological state, genetic diversity and size distribution of the communities were subjected to important changes.

Active cells and production

The fraction of active cells is often based on determinations of the activity of the electron transport system by tetrazolium salts such as INT and CTC (Zimmermann et al. 1978, Rodriguez et al.
Changes in bacterial communities

Although this method has limitations (Thom et al. 1993, Lovejoy et al. 1996, Ullrich et al. 1996) it has been shown that the number of actively respiring cells determined by CTC reduction is generally well correlated with bacterial production estimates [thymidine (TdR) and leucine (Leu) incorporation rates] and that the active cell counts vary in an ecologically interpretable manner (del Giorgio & Scarborrough 1995).

The high correlation between the number of active bacteria and the global incorporation rates suggests that at a given time only a small fraction of the bacteria within natural communities participates in nutrient cycling (Table 3). Better correlations could probably be obtained by quantifying respiration, since the respiration rate of a single cell may vary depending on its metabolic activity and, thus, on its productivity. However, if we assume that these detectable respiring cells are those which participate in cell production, average growth rates can be estimated more accurately than with total cell counts.

Generation times estimated from total bacterial counts varied from 2.4 to 346 h. Inversely, the specific growth rates calculated for each sampling time for active bacteria ranged from 0.93 to 0.11 h⁻¹ and correspond to doubling times of 44 min to 6.02 h (Table 4). These doubling times were in good agreement with those reported in the euphotic zone of winter waters in the Gulf of St. Laurent by Lovejoy et al. (1996). The very short generation times reported in this study can be explained by an underestimation of active cells due to the low sensitivity of the CTC detection by microscopic methods, by the selection of species which cannot reduce tetrazolium salts, or by inappropriate application of a single conversion factor to transform thymidine incorporation rates into cell production. Although these limitations may contribute to these low generation times, strong modifications were observed in each mesocosm. The shorter generation times were observed after the decrease of protozoa counts, suggesting that bacteria which escaped grazing were very active cells. The mean generation time calculated after the decrease of total and active cell counts (at times 185 h in Mesocosm A and 161 h in Mesocosms B and C) was short and varied between 44 and 59 min when calculated based on active cells.

Active cells and size distribution

The high growth rate of bacteria after grazing may be related to significant changes in the cell size distribution. In Mesocosms A and B, there was an increasing fraction of large and small cells with a high and low contribution to apparent total biovolume, respectively. The high growth rate reported in Mesocosm A at 185 h was concomitant with the presence of very large cells representing more than 80% of the total biovolume. In Mesocosm C, the selection of small and large cells was also observed but was less pronounced, probably because the physiological state of the community was more active at the onset of the nutrient supply and resulted in a very rapid growth of bacterial cells and control by protozoa (see below). In all cases, the question whether large cells have a higher contribution than small cells to the whole community production remains unclear. However, the high growth rates reported in Mesocosm A in the presence of large cells suggest that these cells were very productive. A positive relationship between cell size and activity has already been suggested in the literature (Bird & Kalff 1993, Gasol et al. 1995).

Table 3. Matrix of coefficients of determination (r²) for total bacterial abundance (Tot), abundance of active cells (Act), leucine (Leu) and thymidine (TdR) incorporation rates. Data are log10-transformed, n = 32, p < 0.001 for all values

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<th>Act</th>
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combined with leucine incorporation may be one way to confirmed by the diversity of bands on the DGGE pat-
trates. The use of flow cytometry and cell sorting com-
gate the relationship between cell size and growth

the initial culturable community was characterized
and, thus, on the functioning of natural communities.

These results are in good agree-
ment with these, with respect to grazing, advanta-
geous morphological properties or if they express these characteristic features only under strong grazing pressure. If most of these filamentous forms express these features only under grazing pressure, their selec-
tion may have little effect on the taxonomic structure
ities. Gasol et al. (1995) suggested that protozoa prefer-
etentially crop the active portion of the bacterial com-
munity if they select their prey according to size, consi-
tering that activity is related to cell size. Recently,

Table 4. Bacterial growth rates (h⁻¹) calculated from cellular
production and abundances of active cells (or total cell
counts) at different sampling times of each mesocosm (A, B
and C). nd: not determined

<table>
<thead>
<tr>
<th>Time (h)</th>
<th>A</th>
<th>B</th>
<th>C</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>0.164 (0.002)</td>
<td>0.336 (0.004)</td>
<td>0.408 (0.002)</td>
</tr>
<tr>
<td>42</td>
<td>0.316 (0.033)</td>
<td>0.598 (0.156)</td>
<td>nd</td>
</tr>
<tr>
<td>65</td>
<td>0.674 (0.061)</td>
<td>0.439 (0.098)</td>
<td>nd</td>
</tr>
<tr>
<td>89</td>
<td>0.127 (0.014)</td>
<td>0.285 (0.074)</td>
<td>nd</td>
</tr>
<tr>
<td>113</td>
<td>0.115 (0.016)</td>
<td>0.245 (0.073)</td>
<td>0.256 (0.039)</td>
</tr>
<tr>
<td>122</td>
<td>nd</td>
<td>nd</td>
<td>0.605 (0.166)</td>
</tr>
<tr>
<td>137</td>
<td>nd</td>
<td>nd</td>
<td>0.439 (0.280)</td>
</tr>
<tr>
<td>144</td>
<td>nd</td>
<td>nd</td>
<td>0.307 (0.157)</td>
</tr>
<tr>
<td>161</td>
<td>0.171 (0.023)</td>
<td>0.850 (0.087)</td>
<td>0.697 (0.124)</td>
</tr>
<tr>
<td>185</td>
<td>0.933 (0.054)</td>
<td>nd</td>
<td>0.720 (0.126)</td>
</tr>
</tbody>
</table>

Grazing and defense mechanisms

Little is known about the effect of size selection on both taxonomic diversity and productivity of communities. Gasol et al. (1995) suggested that protozoa preferentially crop the active portion of the bacterial community if they select their prey according to size, considering that activity is related to cell size. Recently, Pernthaler et al. (1996, 1997) suggested that 'bacteria can adopt at least two distinct strategies as a reaction to intense flagellate predation: to outgrow predation pressure or to develop inedible, inactive filaments'. *Vibrio* species should be able to form chains composed of several cells as a defense mechanism against protistan attacks (Jürgens & Güde 1994, Jürgens et al. 1997, Hahn & Hofle 1998).

We have shown that during the initial phase of increasing production there was a shift in cell sizes from the small size class to the middle size class and then this fraction of new cells was selectively removed by protistan grazers. Similarly, the fraction of *active cells* was rapidly consumed during the grazing phase, suggesting that the proportion of metabolically active cells was strongly top-down controlled. The effect of grazing on the productivity of the whole bacterial communities in the 3 mesocosms confirms that it affects preferentially the fraction of active cells. Selective grazing allows the persistence of an important pool of large and small cells. These results are in good agreement with other reports (Gasol et al. 1995, del Giorgio et al. 1996b). Although the inactivity of small cells is not demonstrated, the persistence of large cells may be responsible for the maintenance of a significant level of bacterial production. However, further investigations should be done using these methods to better investigate the relationship between cell size and growth rates. The use of flow cytometry and cell sorting combined with leucine incorporation may be one way to overcome methodological limitations. The potential role of viruses was not considered in this study, but one should keep in mind that viruses may also shape bacterial communities as recently suggested in freshwaters in relation with bacterial size classes (Weinbauer & Hofle 1998).

Changes in genetic diversity

The initial culturable community was characterized by an important diversity of bacterial species and was confirmed by the diversity of bands on the DGGE pat-
terns. The cluster analysis of DGGE banding patterns shows that samples are divided into 2 main clusters: those samples taken before the peak in grazing activity, and those obtained thereafter. This supports the notion that grazing was an important factor, controlling not only the density of the bacterial community but also its genetic diversity. The dominant isolates belonged to Alteromonas, Roseobacter, Rhodobacter, Pseudoalteromonas and Sulfitobacter species and were also present in the late sampling points after grazing. These bacterial genera have already been found in a wide variety of marine ecosystems (Giovannoni et al. 1990, Gonzalez & Moran 1997, Suzuki et al. 1997). The results obtained from clone libraries in Mesocosms A and B suggest that bacteria related to A. macleodii are dominant after grazing (Pukall et al. 1999). Our results suggest that some species are maintained while others were not detectable after grazing. Species which were present in the initial water and were also present after grazing, such as A. macleodii, may have developed defense strategies to escape predation. A size polymorphism has already been shown for P. haloplanktis and elongated multigenomic cells were observed under starvation conditions (Lebaron & Joux 1994). This size polymorphism could explain why some species escape to predation. Although this polymorphism was not demonstrated in the presence of grazers, it may result in an indirect survival advantage for this species, but it might also be induced in the presence of grazers.

The key species reported here which seem to escape grazing may be considered as a first step in understanding the effect of grazing on the taxonomic structure of natural communities and the importance of these defense mechanisms towards bacterial groups. Their identification will allow the definition of taxonomic probes for the further investigation of these questions.

CONCLUSIONS

Our results suggest that on the one hand nutrients and grazing play a key role in shaping the morphologic, genotypic and phenotypic composition of bacterial communities by regulating abundances and, on the other hand, grazing preserves a minimum background of bacterial genetic diversity since it is density-dependent and also because some marine species seem to be able to develop defense mechanisms such as rapid growth and induction of a size polymorphism. Although the role of viruses was not investigated in this study due to methodological limitations, it should also be considered in order to better understand these regulations. Furthermore, as grazing may affect the production of cells with the highest growth rates, it may eliminate preferentially the most active cells within natural communities and regulate bacterial productivity if we assume that size and activity are positively correlated. It may explain why the fraction of active cells as determined by the CTC method, although this method suffers from limitations, is often not more than 30 to 40% within marine communities. Because the increase in actively respiring cells does not correlate well with the increase of total bacterial counts, it is possible that the rapid inactivation of new cells may be one way to preserve the diversity of natural communities, since these inactivated cells may return to the background of grazing resistant bacteria and non-active cells.

Further experiments coupling flow cytometry cell sorting based on the relative cell size of bacteria and genetic analysis of these cells will be very useful for validating some of the hypotheses presented herein. These investigations are actually in progress in our group and within the ELOISE-CHABADA project supported by the European Community. It is also important to understand to what extent similar behavior can be observed in the natural environment and how often and how long such perturbed food web structures exist.

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