Ecology of amorphous aggregations (marine snow) in the Northern Adriatic Sea. II. Microbial density and activity in marine snow and its implication to overall pelagic processes

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ABSTRACT: In the Northern Adriatic Sea, marine snow dry mass varied between 3.4 and 9.1 mg l⁻¹ during summer. In situ measurements of O₂-fluxes mediated by marine snow revealed a gross primary production (GPP) ranging from 3 to 23 µg O₂ mg⁻¹ (marine snow ash-free dry wt, AFDW) h⁻¹. O₂-consumption rates ranged from 1.35 to 3.7 µg O₂ mg⁻¹ (marine snow AFDW) h⁻¹ and were ca 5 times the mean consumption obtained under laboratory conditions. Based on these O₂-flux measurements, we estimate that more than 90 % of the pelagic GPP is bound to marine snow aggregations and ca 70 % of the mineralization activity is mediated by marine snow associated microbes during summer. Total carbohydrate (TCHO) content in marine snow was found to be enriched by a factor of 304 as compared to the surrounding water (F = 0.82 TCHO ml⁻¹). Batch cultures with marine snow were designed to investigate the influence of marine snow on the free-living microbial community. In the chambers containing marine snow, growth rates (v) of the bacterial community were 3 times the rates obtained for unenriched media (containing 1 µm filtered seawater only). Based on the batch culture experiment, it is concluded that marine snow enhances not only microbial growth on/in the organic matrix of the aggregations but also favors microbial activity of the surrounding water. The degree of development of the microheterotrophic food chain was quantified using the ratio (nanoflagellate biomass/bacterial biomass) × 100. In the incubation media containing marine snow, ratios of >100 were maintained most of the time, indicating that additional food sources other than free-living bacteria were exploited by the nanoflagellate population. It is suggested that more sensitive sampling methods should be used to resolve such nutrient patches which cause heterogeneity of the pelagic environment.

INTRODUCTION

Marine amorphous aggregations ('marine snow') are a ubiquitous component of pelagic marine waters (Fowler & Knauer 1986). The chemical and biological components of these aggregations may vary according to their origin. Estuaries are a site of particle formation. Terrigenous humic matter transported by rivers into estuaries rapidly flocculates with increasing salinity (Jackson 1975, Sholkovitz 1976). These terrigenous humic substances have been shown to constitute about 60 to 80 % of the dissolved organic carbon (DOC) in rivers (Reuter & Perdue 1977). The flocculation of humics at the freshwater-seawater boundary has been attributed to adsorption/(co)precipitation of humic substances by iron (III) oxyhydroxides (Krom & Sholkovitz 1977, Poutanen & Morris 1983). Organic coating of negatively charged particles (of µm size) facilitates aggregation with other particles (McCave 1984, Kranck et al. unpubl.).

If water-current velocities are sufficiently high to prevent particle settlement, these aggregates are transported offshore (Eisma 1986). This means that the organic fraction of the aggregates increases with distance from the estuary. Amorphous aggregates of estuarine origin, however, never reach as high an organic content as that usually detected in marine snow of biological origin (Amy et al. 1987). An important role in the formation of amorphous aggregations may be performed by bacteria, which convert dissolved...
While marine snow particles of estuarine origin contain little biological material, aggregations of biological origin harbor a rich microbial community of living phytoplankton, Protozoa and bacteria (Silver et al. 1978, 1984, Caron et al. 1982, Beers et al. 1986). Usually, these aggregates range from 0.5 mm to several cm in diameter.

In the Northern Adriatic Sea these particles tend to aggregate to a size of up to 20 cm in diameter during calm weather conditions. Typically, 'stringers' are formed at the initial stage of aggregation, while the more advanced stage of succession is characterized by large coagulations (Eisma 1986, Herndl & Peduzzi 1988). Their large dry mass - up to 10 g (marine snow dry wt) m⁻³ - in the waters of the Gulf of Trieste during summer and the high densities of both autotrophic and heterotrophic organisms embedded in the organic matrix, led to the hypothesis that these aggregations may contribute significantly to the metabolism of the pelagic phase (Herndl & Peduzzi 1988).

The present study evaluates the contribution of marine snow to overall metabolism of the pelagic phase; it also assesses the extent to which decomposition of the organic matrix of marine snow enhances planktonic production of microheterotrophs.

**MATERIAL AND METHODS**

**Study locations and sampling procedure.** Investigations were conducted in the Gulf of Trieste (Northern Adriatic Sea) between April and October 1987. Marine snow abundance was determined at irregular time intervals in various depth layers using the procedure described in Herndl & Peduzzi (1988). Briefly, a translucent tube (50 cm in length, 5 cm in diameter) was slowly moved horizontally through a specified depth layer of the water column. Within 30 min the contents of each tube were filtered through a precombusted Whatman glass-fiber filter (GF/F) and rinsed with distilled water to obtain the dry wt (12 h at 70 °C) of marine snow. Zooplankters not attached to the organic matrix of marine snow were carefully removed under a dissecting microscope by means of forceps prior to drying. After dry wt determination, ash-free dry wt (AFDW) was determined after combustion at 450 °C for 6 h.

For chemical and biological analyses, marine snow and surrounding water were collected using syringes of various size (10 to 930 ml). Although a great deal of effort was exerted to minimize the volume of ambient water during marine snow sampling, we unavoidably sucked a varying volume of water into the 'marine snow'-syringes. Therefore, we used 2 methods for analysis of marine snow: (1) We examined marine snow-seawater slurries sampled by syringes; (2) marine snow-seawater slurry was poured in 5 ml glass jars, and marine snow allowed to settle for 10 min prior to collection with Pasteur pipettes. Samples for enumeration of organisms were preserved as described below as soon as they were brought aboard. Samples of ambient water were taken and treated in the same way as marine snow.

**In situ and laboratory O₂-flux measurements.** In situ measurements of O₂-flux in marine snow and ambient water were performed around noon for 4 h. Marine snow-seawater slurries and seawater were sampled at the same depth and subsequently incubated using 930 ml syringes with an opening of 1 cm in diameter. Syringe contents from a given depth were poured into translucent and dark glass jars, respectively. Care was taken to avoid enclosure of air bubbles. Light and dark bottles containing marine snow-seawater slurries and seawater, respectively, were deployed in duplicate at 3, 6, 9, and 12 m depth. The O₂-contents of the incubation media were measured at the beginning and the end of the deployment by a battery-driven respiration set equipped with polarographic oxygen sensors (Yellow Springs Instrument; Svoboda & Ott 1983).

At the end of the incubation, the marine snow-seawater slurries were filtered onto precombusted and preweighed glass fiber filters (Whatman GF/F) for dry wt determination, ash-free dry wt (AFDW) and chemical analyses. In addition to in situ O₂-flux measurements, laboratory experiments were performed for comparison using the continuous O₂-recording system and incubation chambers with a volume of 23 ml (Svoboda & Ott 1983, Peduzzi 1987). Incubation chambers were filled with 0.45 μm Nuclepore-filtered seawater and marine snow particles were carefully transferred from storage vials to incubation chambers. Marine snow was stored in dark at in situ water temperature for ca 1 h prior to the O₂-flux measurements. Between 1.0 and 7.3 mg (marine snow dry wt) were incubated in the incubation chambers and the O₂-flux recorded for 2 to 4 h.

**Organic matter (DOM) into a particulate form, and aggregate particles with their slime (Paerl 1975, Biddanda 1985, 1988, Biddanda & Pomeroy 1988).**
O₂-concentration of the incubation media never deviated by more than 30 % of the initial concentration.

**Batch culture experiments.** To evaluate the influence of marine snow on microbial dynamics of the surrounding water, incubation experiments were performed at 20 °C in the dark at the Stazione Biologia Marina at Trieste-Aurisina (Italy). Incubations were performed in triplicate; additionally, one jar containing 800 ml of 1 μm Nuclepore-filtered seawater only served as a control. Two hundred ml of a freshly collected marine snow-seawater slurry (corresponding to 30 to 35 mg marine snow dry wt) were added to each of the 3 jars containing 1 μm filtered seawater. Incubated marine snow was kept in suspension by gently aerating the incubation media with 0.45 μm filtered air. Water samples were taken at various time intervals for bacteria and nanoflagellate enumeration and carbohydrate (CHO) analysis. Additionally, marine snow particles were removed occasionally from the media for particulate CHO-determination.

**Microbial and chemical analysis.** For microbial analysis, all surrounding water samples, marine snow particles, and slurries were fixed with concentrated, unbuffered formalin to a final concentration of 2 % (v/v). Bacteria and heterotrophic flagellates were enumerated using epifluorescence microscopy and acridine orange direct counting (Hobbie et al. 1977). Flagellates lacking distinct red fluorescent chloroplasts were assigned as heterotrophs. Dimensions of organisms enumerated by epifluorescence microscopy (magnification: 1250 ×) were measured using a calibrated ocular micrometer. Volumes of these organisms were calculated assuming a spherical shape for heterotrophic coccoid bacteria, a cylindrical shape for rods, and an ellipsoid shape for microflagellates. For microflagellates, the cell-volume to cell-carbon conversion factor (220 fg C μm⁻³) was used (Borsheim & Bratbak 1987). The carbon content of bacteria was calculated using the conversion factor 380 fg C μm⁻³ cell volume (Lee & Fuhrman 1987).

The enrichment factor (EF) was calculated for each category of microorganism (n = number of organisms): EF = n ml⁻¹ (marine snow)/n ml⁻¹ (surrounding water). Marine snow volume was measured by means of graded pipettes.

Water samples for dissolved CHO analysis were filtered through precombusted glass fiber filters (Whatman GF/F) within 30 min after sampling and stored frozen (−20 °C) until analysis. For particulate CHO, marine snow aggregations were filtered onto glass fiber filters (which were treated as described above). CHO-contents were measured spectrophotometrically according to the methods of Burney & Sieburth (1977) and Johnson & Sieburth (1977). In order to present the data on CHO in terms of carbon we assumed that hexoses, containing 72 g C mol⁻¹ (MW = 180, carbon content 40 % (w/w), dominated the carbohydrates.

**RESULTS**

**Marine snow abundance and organic content**

Marine snow dry mass varied between 3.4 and 9.1 mg l⁻¹ and increased steadily with depth (Fig. 1). The organic content of marine snow, however, decreased with depth. While at 3 m depth 80 % of marine snow dry wt was organic material this percentage dropped to 60 % at 20 m depth (Fig. 1).

**Heterotrophic microorganisms and CHO-content of marine snow and surrounding water**

The microheterotrophic community in marine snow, as well as in the surrounding water, was dominated by heterotrophic bacteria, choanoflagellates and monads.
In marine snow the microheterotrophic density was always higher than that of the ambient water (EF > 1) (Table 1) and varied over 2 orders of magnitude, depending on the sampling method (see 'Material and Methods'). Microbial densities 1 order of magnitude higher were obtained when marine snow was allowed to settle in the storage vials prior to collection for analysis (Table 1). Despite this methodological variation introduced, microbial density varied considerably, even between consecutive days. No group in the microbial community examined had accumulated to an extent significantly greater than that of any other (Kruskal-Wallis, \( p > 0.2 \)). Maximum EF-values were obtained in marine snow kept in a 500 ml storage vial in the dark at in situ temperature (18 °C) for 4 d (Table 1).

Microbial density in marine snow exhibited an inconsistent pattern between various depth layers during most of the sampling dates. After wind-induced mixing of the water column on 24 June, however, an exponential increase in bacterial density with depth was detected in marine snow on 25 and 30 June (Fig. 2a). During this period, monads were the most abundant eukaryotes in marine snow in the upper layers of the water column while in deeper horizons choanoflagellates were the most prominent eukaryotes (Fig. 2b).

No significant differences in the biovolume of marine snow bacteria and free-living bacteria, respectively.

Table 1. Mean microbial density (± SD) ml\(^{-1}\) marine snow and enrichment factors (EFs) averaged over the water column during consecutive days

<table>
<thead>
<tr>
<th>Date</th>
<th>Bacteria ( N \times 10^6) ml(^{-1})</th>
<th>EF</th>
<th>Choanoflagellates ( N \times 10^3) ml(^{-1})</th>
<th>EF</th>
<th>Monads ( N \times 10^5) ml(^{-1})</th>
<th>EF</th>
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</thead>
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<tr>
<td>19 Apr*</td>
<td>55.9</td>
<td>93</td>
<td>17.5</td>
<td>253</td>
<td>4.0</td>
<td>67</td>
</tr>
<tr>
<td>( n = 3 )</td>
<td>(28.2)</td>
<td></td>
<td>(7.0)</td>
<td></td>
<td>(1.4)</td>
<td></td>
</tr>
<tr>
<td>23 Jun</td>
<td>15.2</td>
<td>30</td>
<td>12.9</td>
<td>161</td>
<td>8.4</td>
<td>105</td>
</tr>
<tr>
<td>( n = 11 )</td>
<td>(3.4)</td>
<td></td>
<td>(3.4)</td>
<td></td>
<td>(3.0)</td>
<td></td>
</tr>
<tr>
<td>25 Jun</td>
<td>2.5</td>
<td>2</td>
<td>0.3</td>
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<td>0.3</td>
<td>7</td>
</tr>
<tr>
<td>( n = 4 )</td>
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<td>(0.1)</td>
<td></td>
<td>(0.1)</td>
<td></td>
</tr>
<tr>
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<td>7</td>
<td>1.8</td>
<td>27</td>
<td>0.4</td>
<td>8</td>
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<td>(0.04)</td>
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<tr>
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<td>(0.1)</td>
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<td>(0.02)</td>
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<tr>
<td>30 Jun</td>
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<td>1</td>
</tr>
<tr>
<td>( n = 4 )</td>
<td>(0.7)</td>
<td></td>
<td>(0.02)</td>
<td></td>
<td>(0.1)</td>
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<tr>
<td>2 Jul*</td>
<td>110.4</td>
<td>133</td>
<td>20.7</td>
<td>207</td>
<td>11.6</td>
<td>290</td>
</tr>
<tr>
<td>( n = 3 )</td>
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<td></td>
<td>(3.9)</td>
<td></td>
<td>(5.0)</td>
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<tr>
<td>3 Jul</td>
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<tr>
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<td>(0.03)</td>
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</tr>
<tr>
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<tr>
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<td>(2.6)</td>
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<td>(0.7)</td>
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</tr>
<tr>
<td>Marine snow</td>
<td></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>aged 4 d**</td>
<td>204.4</td>
<td>232</td>
<td>46.6</td>
<td>259</td>
<td>14.2</td>
<td>129</td>
</tr>
<tr>
<td>( n = 3 )</td>
<td>(35.6)</td>
<td></td>
<td>(5.6)</td>
<td></td>
<td>(6.3)</td>
<td></td>
</tr>
</tbody>
</table>

* Marine snow was allowed to settle in storage vials for 10 min prior to sampling for analysis
** Marine snow was incubated in 1 µm Nuclepore-filtered seawater in dark at 20 °C for 4 d and kept in suspension by a gentle stream of 0.45 µm filtered air
were found between sampling dates. Therefore, the volumes of marine snow bacteria were pooled regardless of sampling date as well as free-living bacteria. Mean bacterial biovolume for the 2 categories (i.e. rods and cocci) were higher in marine snow than in the surrounding water. For rod-shaped bacteria, mean volume in marine snow was 0.25 μm³ (SD = 0.05, n = 350), and therefore significantly higher than that of ambient water (X = 0.185 μm³, SD = 0.038, n = 325, ANOVA, p < 0.005). The corresponding volumes for coccoid bacteria were 0.067 μm³ (SD = 0.009, n = 175) in marine snow and 0.042 μm³ (SD = 0.009, n = 185, ANOVA, p < 0.001). Both groups of flagellates, however, did not differ significantly in volume between marine snow and ambient water (choanoflagellate mean volume: 11.43μm³; SD = 1.9; n = 270; mean monad volume: 2.46μm³; SD = 0.83, n = 236).

Total carbohydrate content (TCHO) of marine snow averaged 251 pg TCHO ml⁻¹ (marine snow) (SD = 43.8, n = 10) with no discernible pattern between depth layers; however, TCHO in marine snow was found to be enriched by a factor of 304 as compared to the surrounding water (X = 0.82 μg TCHO ml⁻¹).

**O₂-flux mediated by marine snow and surrounding water**

In situ measurements of O₂-fluxes mediated by marine snow revealed a gross primary production (GPP) ranging from 3 to 23 μg O₂ mg⁻¹ (marine snow AFDW) h⁻¹ (Fig. 3). Highest marine snow associated O₂ production rates were obtained in the 6 m layer, lowest values for the 9 m horizon. Although PAR levels at 12 m depth were only about half the levels of the 9 m horizon, GPP in marine snow was higher in the deeper layer. O₂ consumption rates of marine aggregates varied within a more narrow range throughout the water column. Mean O₂ consumption averaged over depths amounted to 1.35 μg O₂ mg⁻¹ (marine snow AFDW) h⁻¹ (SD = 0.2, n = 16) on 27 June; for 28 June a mean consumption of 3.7 μg O₂ mg⁻¹ (marine snow AFDW) h⁻¹ (SD = 1.6, n = 16) was calculated. These consumption rates were 2.5 and 7 times, respectively, the mean consumption obtained under laboratory conditions. In laboratory incubations, mean O₂ consumption was 0.54μg O₂ mg⁻¹ (marine snow AFDW) h⁻¹ (SD = 0.19, n = 9).

**Influence of decomposing marine snow on free-living microbes**

In the incubation flasks containing marine snow, free-living rod-shaped bacteria increased rapidly in number until 20 h after starting the incubation, while cocci did not exhibit such a pronounced reaction during the initial incubation phase. For the initial period (0 to 13 h) of the incubation, rods increased from 6.1 to 33.9 × 10⁵ cells ml⁻¹ and cocci from 7.0 to 16.9 × 10⁵ cells ml⁻¹, corresponding to a mean generation time (g) for rods of 5.4 h and 10.5 h for cocci. During this period, rods increased in the control jar (containing 1 μm fil-
tered seawater only) from $3.5 \times 10^5$ cells ml$^{-1}$ ($g = 13.9\ h$) while cocci increased from $4.4 \times 10^6$ cells ml$^{-1}$ ($g = 6\ h$). The observed increase in bacterial biomass in the marine snow flasks was attributed to rods (Fig. 4a). Thereafter, rod-shaped bacterial density dropped off rapidly to $3 \times 10^5$ cells ml$^{-1}$ (SD = 1.5) 53 h after starting the experiment. In the subsequent period, total bacteria fluctuated in a very narrow range (Fig. 4a) with densities 2 to 3 times the bacterial abundance in the control. Biovolume of free-living rods incubated in the marine snow jars increased from $0.377\ \mu m^3$ (SD = 0.112, $n = 178$) at the beginning of the incubation to $0.459\ \mu m^3$ (SD = 0.108, $n = 156$) (ANOVA, $p < 0.01$) 4 h after starting the incubation. After 34 h, rod-shaped bacteria, however, decreased again in volume approaching their initial size (mean = $0.361\ \mu m^3$, SD = 0.19, $n = 285$). No such changes in biovolume were noticed in coccoid bacteria or in the bacterial community of the control.

The corresponding development of the 2 dominant nanoflagellate categories (choanoflagellates and monads) are shown in Fig. 4b. In the marine snow chambers choanoflagellates increased continuously from $<10^3$ cells ml$^{-1}$ at the beginning to $23 \times 10^3$ cells ml$^{-1}$ after 53 h when they reached their peak density. Concurrent with the decline in choanoflagellate abundance, monads appeared, reaching densities up to $10^6$ cells ml$^{-1}$. Monad biomass remained relatively stable from 72 h after starting the experiment to the end. In terms of carbon, monads maintained a biomass of 300 to 400 $\mu g\ C\ l^{-1}$ in the marine snow chambers during the second half of the incubation, which is about twice the bacterial biomass (Fig. 4b). In the control, maximum choanoflagellate biomass (ca 100 h after the beginning of the incubation) exceeded the peak biomass of choanoflagellates in the marine snow jars containing marine snow. Monads in the control maintained a biomass only one third of that of the marine snow incubations. No significant changes were found in the biovolumes of either category of nanoflagellates during the course of the incubation in either the control or in the marine snow jars. Mean choanoflagellate volume was $17.8\ \mu m^3$ (SD = 1.4, $n = 75$), average monad volume was $0.84\ \mu m^3$ (SD = 0.21, $n = 78$).

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**Fig. 4.** Batch cultures with marine snow. (a) Development of free-living bacterial biomass in incubation media; triangles: rod-shaped bacteria, circles: cocci. (b) Development of flagellate biomass; triangles: monads, circles: choanoflagellates. (c) Course of dissolved MCHO concentration. Closed symbols indicate mean (SD) of 3 replicate incubations; open symbols stand for control to which no marine snow was added.
Dissolved monosaccharide (DMCHO) concentrations in both marine snow chambers and control revealed similar trends (Fig. 4c). During the first 4 h of incubation, however, DMCHO-concentrations in the marine snow jars were about 3 times lower than the value of the control. DMCHO increased during the first 24 h and subsequently declined rapidly to concentrations below 200 µg MCHO-C l⁻¹. The following small peak in DMCHO reached only about one third the height of the initial peak; thereafter, DMCHO concentrations remained well below 100 µg MCHO-C l⁻¹ in both control and marine snow chambers during the latter stage of the incubation.

**DISCUSSION**

Abundance of marine snow in the water column of the Northern Adriatic Sea was shown to range from 3 to 9 mg (marine snow dry wt) l⁻¹ in 1987, and is therefore similar to the mean value of 5 mg (marine snow dry wt) l⁻¹ reported for the preceding year (Herndl & Peduzzi 1988). Unlike the previous study, the lack of a pronounced thermocline led to a continuous increase in marine snow dry mass with depth (Fig. 1). This increase in dry mass is compensated by decreasing organic matter content with depth (Fig. 1). Based on the regression given in Fig. 1, at 3 m depth mean marine snow dry mass amounts to 5.3 mg l⁻¹ with an average organic matter content of 80 % or 4.2 mg l⁻¹. At 20 m depth (ca 4 m above bottom) the 6.7 mg marine snow dry wt contains only 61 % organic matter or 4.1 mg l⁻¹. The combined effect of microbial decomposition of the organic matrix of marine snow during sedimentation and the attachment of resuspended inorganics may be responsible for this alteration.

Considerable variations in microbial density of marine snow particles were detected between consecutive days (Table 1). Stable weather conditions with low current velocities favor the development of a rich microbial community on/in marine snow while wind induced currents lead – at least partly – to a renewal of the water body, as indicated by changes in temperature and salinity (data not shown). Such an event was observed on 24 June, from 29 to 30 June and in the evening of 2 July. As demonstrated in Table 1, on each consecutive day marine snow microbes exhibited EF values between 1.3 and 4. These values are considerably lower than the EF value obtained during more stable weather conditions. The rapid increase of the EF during the period following such an event may indicate rapid microbial colonization of recently developed marine snow. It should be noted, however, that all EF values of marine snow are conservative estimates since the samples were always diluted with surrounding water. Generally, ca one order of magnitude higher EF values were obtained if the marine snow was allowed to settle in the storage vials prior to analysis, indicating the dilution effect of surrounding water (Table 1). The settling period for marine snow of 10 min seems to be too short to account for significant microbial growth.

The degree of development of the microheterotrophic food chain was quantified using the ratio (heterotrophic nanoflagellate biomass/bacterial biomass) × 100. Fig. 5 presents the development of the microheterotrophic food chain during the course of marine snow incubation; for comparison, indices are also given for marine snow particles and the surrounding water. In the incubation media containing marine snow, ratios of > 100 were maintained most of the time, suggesting that additional food sources other than free-living bacteria were exploited by the nanoflagellate population. The indices of both the control medium and the surrounding water are similar to values reported for eutrophic or stratified waters of various regions (Sorokin 1977, Linley et al. 1983, Davoll & Silver 1986). Extremely high ratios of > 200 were also recorded for marine snow in April and at the end of June (Fig. 5). In general, enforced renewal of the water body by unstable weather conditions leads to lower ratios in marine snow.

As stated by Smetacek (1985), nutrient-depleted diatom blooms rapidly release cellular material into the surrounding water leading to flake formation. These flakes may aggregate further and reach dimensions of up to a few cm. In the Northern Adriatic Sea, marine snow rapidly increases in abundance during April which coincides roughly with the decay of the diatom bloom and the change from a 'new' to a 'regenerating' system (a concept first introduced by Dugdale & Goering 1967). Marine snow (of biological origin) may therefore be considered as a typical sign of the regenerating phase in the Northern Adriatic Sea. While new systems are based on material resources transported by the environment, in regenerating communities the material resource is not necessarily lost from the system (Smetacek & Pollehne 1986). In the latter, the bulk of primary production is based on short-term recycling, implying that the microbial loop plays an important role in system respiration. All these criteria are fulfilled by marine snow aggregations.

The water column – considered a regenerating system during summer – is modified due to unstable weather conditions which enforce the renewal of the water body. These events shift the pelagic phase toward a new system with a diminished nanoflagellate population, resulting in flagellate/bacteria ratios typical for new systems. Due to the high growth rates of nanoflagellates in waters where marine snow is abundant (discussed below), the former stage of the system
is established again rapidly. As pointed out by Smetacek & Pollehne (1986), mucus release (of phyto- and zooplankton) may favor the development of short-circuit nutrient cycles and small-scale nutrient patches in the pelagic environment. Such a mucus-based system with a high microbial activity is the interstitial space between coral branches (Schiller & Herndl in press).

Incubation experiments were designed to assess the microbial response of the surrounding water to marine snow. In the present experiment marine snow dry mass $1^{-1}$ was ca 4 to 10 times higher than recorded for the Northern Adriatic Sea (Herndl & Peduzzi 1988, this study). The initial, pronounced increase in bacterial biomass of free-living rods in the incubation media containing marine snow does not simply reflect detachment of marine snow bacteria since the 200 ml marine snow-seawater slurry added contained only about 3.6 x 10$^7$ bacteria or 4.5% of the bacterial population of the incubation media (800 ml). In terms of carbon, 5.1 µg C (or 5%) of marine snow bacteria were added to 96 µg C of free-living bacteria. Hence we obtained a ca 5 fold increase in bacterial biomass within 20 h. It seems unlikely that detached marine snow bacteria contribute significantly to the free-living bacterial population (Fig. 4a). The peak in bacterial biomass is caused almost exclusively by the rod-shaped bacterial population. This may indicate that rods took up nutrients more actively than cocci. This pattern is consistent with observations made in other environments. It is generally believed that rods are characteristic for waters of high nutritive value. Kogure et al. (1987) demonstrated that large bacteria grow much faster than small cells (<0.6 µm) and that the former are the main contributors to the turnover of organic matter. In the experiment presented in Fig. 4a, coccoid bacteria reached their maximum biomass 14 h after the biomass peak in rods. At that time (34 h after starting the incubation) rod-shaped bacterial biomass had already diminished – most likely due to enhanced nanoflagellate predation. Rods started to decrease when the nanoflagellate population entered exponential growth.

Based on these incubation experiments, parameters describing growth can be derived (Table 2). Bacteria and nanoflagellate growth rates ($\mu$) und growth yields ($Y$) obtained in our incubation experiments are among the highest reported in the literature. However, the growth yields for nanoflagellates derived from our incubations are most likely overestimates since the bacterial population was probably still growing during the observed decrease in biomass. Therefore, the total consumption of bacteria is higher than the net removal of bacterial biomass. Even taking these overestimates into account, marine snow enhances microbial growth and leads to ca 3 times higher growth rates than incubations with unenriched media (Table 2). In our experiment, the control media were sampled in the same parcel of water in which the marine snow was sampled. Since Herndl & Peduzzi (1988) detected a net release of dissolved organic carbon from marine snow into the surrounding water, we suggest that the control medium was at least partly enriched by marine snow derived from the incubation media.
organic nutrients. This assumption is supported by a similar incubation experiment in which decomposition of coral mucus was investigated (Herndl & Velimirov 1986); a mean bacterial growth rate in the control jar of 0.038 h⁻¹ was obtained using similar experimental conditions. Therefore, we conclude that marine snow enhances not only microbial growth on/in the organic matrix of the particles but also favors microbial activity of the surrounding water.

The reason for such favorable conditions for microbial life is the concentration of easily utilisable organic matter in marine snow. TCHO-concentrations in marine snow were 300 times higher than in the same volume of surrounding water. In the batch culture, the available dissolved MCHO was utilized within 100 h (Fig. 4c). At the end of the incubation, MCHO concentrations were close to the detection limit of the method used.

In marine snow, the high concentration of microbial biomass with autotrophic and heterotrophic compartments may considerably influence overall pelagic metabolic processes during summer. Compared to the in situ mean O₂-consumption in dark chambers of 2.5 µg mg⁻¹ (marine snow AFDW) h⁻¹, O₂-consumption in the laboratory was 2 to 7 times lower. In the laboratory marine snow was incubated in 0.45 µm Nuclepore filtered seawater; thus organisms not associated with marine snow were excluded. In situ, however, planktonic organisms from the surrounding water were unavoidably enclosed during sampling of marine snow and consequently interfered with the in situ measurements.

In conjunction with marine snow dry mass data, these measurements enable us to estimate roughly the contribution of marine snow to overall water column processes. Mean midday GPP of the water column amounted to 15 mg C m⁻³ h⁻¹, water column respiration to 5.6 mg C m⁻³ h⁻¹. Mean marine snow content over the water column averaged 4.1 mg (marine snow AFDW) m⁻³ during investigation. Average GPP of marine snow over the water column meant 3.54 µg C mg⁻¹ (marine snow AFDW) h⁻¹, or 14.5 mg C m⁻³ h⁻¹. In situ mean respiration rate was 0.94 µg C mg⁻¹ (marine snow AFDW) h⁻¹, or 3.8 mg C m⁻³ h⁻¹. Although these values are only crude estimations and do not allow extrapolation of the GPP of the system, they indicate that more than 90 % of the pelagic GPP is bound to marine snow aggregations and ca 70 % of the mineralization activity is mediated by marine-snow-associated microbes. The high contribution of marine snow to pelagic GPP during summer may be explained by its autotrophic community, which exhibits EF values of up to 1000 (Herndl & Peduzzi 1988). Knauer et al. (1982) found that up to 58 % of the nearshore primary production may occur in association with marine snow, and Aldridge & Cohen (1987), using O₂-microelectrodes, clearly documented the intensive O₂-production and consumption of individual marine snow particles. Considering the high abundance and the large size of marine snow in the waters of the Northern Adriatic Sea during summer, the calculated contribution of these aggregations to overall pelagic metabolism seems reasonable.

In summary, these marine snow aggregates seem to be vital signs of regeneration systems in which the microbial loop is an important pathway in the pelagic food web (Azam et al. 1983, Smetacek & Pollehne 1986). These fragile aggregations escape conventional sampling methods. They form nutrient patches and

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**Table 2.** Comparison of parameters of growth and feeding in heterotrophic bacteria and nanoflagellates obtained from marine snow batch cultures with data given in literature. Gross grow rate \( \mu = (\ln X_1 - \ln X_0)/t \), where \( X_1 \) and \( X_0 \) = density of organisms at end and beginning of time interval \( t \) (in h); \( Y_N \) = (eventual number of flagellates/initial number of bacteria) × 100; \( Y_B \) = (flagellate C-biomass/bacterial C-biomass) × 100; – not determined.

<table>
<thead>
<tr>
<th>Source</th>
<th>Bacteria ( \mu ) (h⁻¹)</th>
<th>Nanoflagellates</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>( \mu ) (h⁻¹)</td>
<td>( Y_N ) (%)</td>
</tr>
<tr>
<td>This study</td>
<td></td>
<td></td>
</tr>
<tr>
<td>marine snow</td>
<td>0.104</td>
<td>30.4</td>
</tr>
<tr>
<td>control</td>
<td>0.067</td>
<td>7.3</td>
</tr>
<tr>
<td>Andersen &amp; Sorensen (1986)</td>
<td>0.035</td>
<td>–</td>
</tr>
<tr>
<td>Bratbak (1987)</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Fenchel (1986)</td>
<td>–</td>
<td>37</td>
</tr>
<tr>
<td>Herndl &amp; Velimirov (1986)</td>
<td>coral mucus</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>control</td>
<td></td>
</tr>
<tr>
<td></td>
<td>0.077</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>0.038</td>
<td>–</td>
</tr>
</tbody>
</table>

\( a \) Value for choanoflagellates

\( b \) Value for monads
stimulate microbial life not only within the aggregations but also in the surrounding water. More intensive and sensitive sampling methods capable of detecting such nutrient patches will make it possible to resolve the heterogeneity of the pelagic environment, in which the function of mucus as a binding agent has not yet received adequate attention.

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