Photosynthesis, respiration, and carbon turnover in sinking marine snow from surface waters of Southern California Bight: implications for the carbon cycle in the ocean

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ABSTRACT: Photosynthesis and respiration were measured in 1 to 6 mm large aggregates (marine snow) collected in the Southern Californian Bight, USA. The aggregates were freely sinking in a vertical flow system with an upward flow velocity which opposed the sinking velocity of individual aggregates during the measurements. The aggregates were net heterotrophic communities at light intensities <152 \pm 64 μ E m $^{-2}$ s $^{-1}$, and respiration comprised 75 \pm 21 % of gross photosynthesis at saturating light intensities >500 μ E m $^{-2}$ s $^{-1}$. Bacterial densities on aggregates were >2000-fold higher than in the surrounding water. Cytophaga was highly abundant in the aggregate-associated bacterial community as identified by in situ hybridization techniques. Both the respiration rate per aggregate volume and the bacterial densities decreased with increasing aggregate size. The respiration rates normalized to the number of bacteria in single aggregates were 7.4 to 70 fmol C cell $^{-1}$ d $^{-1}$. The aggregate community respired 433 to 984 ng C d $^{-1}$ per aggregate in darkness, which yielded a turnover time of 8 to 9 d for the total organic carbon in aggregates. Thus, marine snow is not only a vehicle for vertical flux of organic matter; the aggregates are also hotspots of microbial respiration which cause a fast and efficient respiratory turnover of particulate organic carbon in the sea.

KEY WORDS: Oxygen microelectrodes · Marine bacteria · Organic matter · Hydrolysis · Cylophaga

INTRODUCTION

Macroscopic organic aggregates larger than 0.5 mm (marine snow) comprise a significant fraction of the mass flux of organic matter in the ocean (Shanks & Trent 1980, Fowler & Knauer 1986, Alldredge & Silver 1988, Turley & Mackie 1994). Marine snow is, therefore, important for the export of particulate organic carbon (POC) from the euphotic zone to the sediments or the deep ocean, i.e. for the ocean's 'biological carbon pump'. The flux of POC, however, decreases with increasing depth in the ocean (Knauer et al. 1979,

Knauer & Martin 1981, Lee & Wakeham 1988). Marine snow is a microhabitat highly enriched in phytoplankton, bacteria, flagellates and detritus compared to the surrounding water, and the attached microorganisms have been considered to control the remineralization of the sinking fraction of POC (Caron et al. 1982, Alldredge & Silver 1988, Turley & Mackie 1994). However, on marine snow, the carbon demand of attached bacteria, estimated by their production rates, is often so small that months to years would be required for the bacteria to consume all the organic carbon of the aggregate after it is exported from the euphotic zone (Karl et al. 1988, Simon et al. 1990, Smith et al. 1992). This discrepancy between the observed rapid decrease in POC and the dense populations of microorganisms on marine snow and the surprisingly low bacterial carbon demand of the attached bacteria has been stated as a 'particle decomposition paradox' in the ocean (Karl et al. 1988).

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Particle dissolution through enzymatic hydrolysis of the aggregate carbon and particulate amino acids by the attached bacteria has been shown to be a significant and fast degradation process during sinking. The enzymatic hydrolysis and uptake of the hydrolysate by the aggregate-associated bacteria were not closely coupled since amino acids were released into the surrounding water (Smith et al. 1992). As a result, a large fraction of carbon and nutrients is processed and assimilated by free-living bacteria in the surrounding water rather than by the microbial community attached to the particles (Cho & Azam 1988, Smith et al. 1992, Grossart & Simon 1998).

Strong gradients in oxygen concentration and pH in marine snow, and even anoxic conditions in large fecal pellets, have been demonstrated by the use of microelectrodes (Alldredge & Cohen 1987). It has been hypothesized that diffusion limitation and the chemical microenvironment, e.g. anoxia and low pH, of sinking aggregates may limit bacterial growth of the attached community (Azam et al. 1993) as the growth rates of bacteria on marine snow seldom are higher than 1 division d⁻¹ (Alldredge & Gotschalk 1990, Simon et al. 1990).

The respiration rate of the attached community of microorganisms and the actual carbon turnover in sinking aggregates have remained unknown and the question as to why attached bacteria apparently do not utilize the aggregate carbon and nutrients more efficiently remains open. In the present study we have directly measured respiration rates in sinking marine snow to test if the turnover of particulate organic matter by the attached microorganisms is indeed as slow as estimated by the production measurements. Community photosynthesis, respiration, enzymatic hydrolysis, enumeration and identification of bacteria on marine snow were evaluated in relation to POC and particulate organic nitrogen (PON) content. This allowed us to determine the turnover time of the POC in the aggregates in relation to their estimated sedimentation time.

MATERIALS AND METHODS

Sampling of marine snow. During 6 to 28 June 1997 individual aggregates were collected by SCUBA in acid-cleaned glass bottles (120 ml, Wheaton, USA) at 0 to 15 m depth ca 2 km northeast of Two Harbors, Catalina Island, in the Southern Californian Bight, USA. The collection was done in the morning (08:00 to 10:00 h) and the aggregates remained in the glass bottles in a cooling box for <1 h until further analysis in the lab. The incident light intensity at the sea surface measured as downward irradiance varied from 200 µE

 $m^{-2}\,s^{-1}$ (08:00 h) to 2000 $\mu E\,m^{-2}\,s^{-1}$ (12:00 h). The depth at which the scalar irradiance was attenuated to 10 % of the downward irradiance at the sea surface was 27 m at 08:00 h.

Microelectrode measurements. Oxygen gradients were measured in sinking individual aggregates kept in suspension by an upward directed flow, which balanced their sinking rates as earlier described (Ploug & Jørgensen 1999). Ten liters of surface seawater was collected, filtered (GF/C) and used in the flow system. All measurements were done at in situ temperature (17°C), using a Clark-type microelectrode (Revsbech 1989) with a 4 µm wide sensing tip, a 90 % response time of 0.2 s, and a stirring sensitivity of <0.3%. The microelectrode was calibrated in air-saturated and in N₂-flushed seawater. The electrode signal was measured by a picoammeter connected to a computerbased data acquisition system with a data collection frequency of 510 Hz (LabVIEW 2, National Instruments). The position of the aggregate surface was determined by slowly advancing the microelectrode towards the aggregate until it visibly touched the upper surface as observed under a dissecting microscope. The gradients were measured after steady-state had been reached at the different light conditions or in darkness. The light source was a 150 W halogen fiberoptic lamp equipped with an infra-red cut-off filter (Schott, KI 1500). The light intensity was controlled by the use of neutral density filters (Oriel) and measured inside the chamber with an underwater scalar irradiance sensor (Biospherical Instruments QLS 100). The dimensions of the aggregates were measured under the dissecting microscope while they were in the flow system. Immediately after the microelectrode measurements, the aggregates were fixed in 4% paraformaldehyde solution and stored at 4°C for less than 2 wk until enumeration of attached bacteria.

Analysis of O_2 gradients. The measured oxygen gradients in the boundary layer were used to determine the oxygen fluxes and, thereby, the net photosynthesis and respiration rates in the aggregates (Ploug et al. 1997). The diffusion coefficient for oxygen is 1.88×10^{-5} cm² s⁻¹ (17°C) (Broecker & Peng 1974). The diffusion calculations were made for an ellipsoid with 3 half-axes a, b, and c, where a < b < c. The surface area of an ellipsoid is (Maas 1994):

$$A = 2\pi a c \left[\frac{c}{a} + \frac{b}{a} \sqrt{1 - \left(\frac{c}{a}\right)^2} E(\varphi, \kappa) + \frac{b}{a} \frac{c}{a} \frac{1}{\sqrt{1 - (c/a)^2}} F(\varphi, \kappa) \right]$$

where

$$\phi = \arcsin \sqrt{1 - \left(\frac{c}{a}\right)^2}; \qquad \kappa^2 = \frac{1 - (c/b)^2}{1 - (c/a)^2}$$

and $F(\phi, \kappa)$, and $E(\phi, \kappa)$ denote the (incomplete) elliptic integrals of the first and second kind, respectively, as

tabulated in Spiegel (1968). The volume of an ellipsoid is $\frac{4}{3}\pi abc$ (Spiegel 1968).

Gross photosynthesis was determined from the outward directed net fluxes of oxygen at saturating light intensity plus the inwards directed flux in darkness, i.e. assuming respiration during light to be equal to dark respiration. Photosynthesis and respiration rates were converted to carbon equivalents by using a conversion factor of $1.2 \text{ mol } O_2$ to 1 mol carbon.

The data for size-specific respiration rates and bacterial densities were fitted by an allometric function. The curve fitting was done with a non-linear regression Levenberg-Marquardt algorithm (Origon 3.0, Micro-Cal Software Inc.).

Enumeration of bacteria. Bacteria on marine snow and in the surrounding water were counted after DAPI staining by epifluorescence microscopy (Porter & Feig 1980). Two ml subsamples of surrounding water were filtered directly onto 0.2 µm nuclepore membranes. To remove bacteria from marine snow, single or pooled aggregates were treated with ultrasound in 2 mM Napyrophosphate (Velji & Albright 1986) prior to filtration.

Bacteria and Cytophaga. Percentages of Bacteria and members of the Cytophaga-flexibacterium-bacteroides group (Cytophaga) were examined by *in situ* hybridization with 16S rRNA-targeted fluorescent oligonucleotide probes according to Weiss et al. (1996). Twenty µl aliquots of sonicated aggregates were pipetted onto gelatin-coated Teflon microslides (P. Marienfeld KG, Bad Mergentheim, Germany) and dried at 46°C. The samples were then fixed in 40 µl of fresh paraformaldehyde (4%) for 4 h at 4°C. Oligonucleotide probes specific for Bacteria (EUB, Amann et al. 1990) and Cytophaga of the CFB group (Amann et al. 1995) were used.

Hydrolytic enzyme activities. Hydrolytic enzyme activities of free-living and attached bacteria were measured with fluorogenic analogs of natural substrates (Hoppe 1993): L-leucine-methyl coumarinylamide (MCA, aminopeptidase), methyl umbelliferyl-aand β -D-glucoside (MUF, α - and β -glucosidase. Hydrolysis by free-living bacteria was measured by incubation of 5 ml subsamples with 50 µM fluorogenic substrates for 1 h at in situ temperature in the dark. Hydrolysis by aggregate-associated bacteria was measured on single aggregates in 5 ml seawater with 250 to 500 µM substrates. These concentrations assured maximum hydrolysis rates as determined by saturation kinetics. A control, heat-killed at 80°C for 20 min, was made for both free-living and aggregate-associated bacteria. Fluorescence was measured at 380/365 nm excitation and 440/455 nm emission for MUF / MCA substrates in a fluorometer (Hoefer TKO-100). The activity of aggregate-associated bacteria was calculated as the total activity minus that of free-living bacteria.

Dry weight and CHN analysis. For measurements of the dry weight and the CHN elemental composition, 10 individual aggregates were pooled and filtered onto precombusted and preweighed (W_0) glass microfiber filters (1 µm pore-size, Schleicher & Schüll, Germany). The filters were dried at 110°C for 1 h and weighed again (W_1). The mean dry weight of 1 aggregate was determined as ($W_1 - W_0$)/10. Filters were then fumed overnight with HCl (20% final conc.) to remove the inorganic carbon. The percentage of organic carbon and nitrogen was determined in a CHN analyzer (Perkin Elmer, USA). The amount of organic carbon or nitrogen per aggregate was calculated from the average dry weight and the average percentage of carbon or nitrogen.

Colonization/growth experiments. Individual aggregates were kept in the 120 ml sampling bottles at *in situ* temperature in a 12:12 h dark-light cycle for 1 to 3 d; bacterial numbers were counted after DAPI staining (see above), and the number and percentage of Bacteria and Cytophaga were determined after *in situ* hybridization (see above).

RESULTS

Characterization of aggregates

The mean equivalent spherical diameter (ESD) of the aggregates was 2.90 ± 1.28 mm. The ratio of the longest to the shortest axis was 1.23 ± 0.25 . Measurements of dry weight, POC and PON were made on 2 sampling dates (Table 1). The average dry weight of aggregates was 62.2 ± 44.4 and 91.1 ± 32.9 µg agg. on 27 and 28 June, respectively. The POC content was 3.87 ± 1.08 and 7.77 ± 1.07 µg C agg. on 27 and 28 June, respectively, and comprised 6.2 and 8.5% of the dry weight. The aggregates on 28 June were larger and had a higher C:N ratio (16.9:1) than those on 27 June (12.2:1). The phytoplankton of the aggregates was mainly composed of diatoms and cyanobacteria (Synechococcus sp.). The aggregates had an apparently high content of exopolymers and indefinable

Table 1. Dry weight, POC content, and C:N ratio in aggregates. Data show the mean ± SD values

	Dry weight (µg C agg. ⁻¹)	POC (µg C agg. ⁻¹)	C:N (w:w)
27 June	62.2 ± 44.4	3.87 ± 1.08	12.2
28 June	91.1 ± 32.9	7.77 ± 1.07	16.9

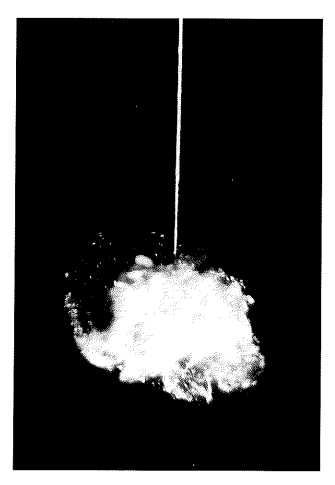


Fig. 1. A 3 mm large aggregate photographed through a dissection microscope is freely suspended by an upward flow velocity which balances its sinking velocity. An oxygen microelectrode is inserted from above

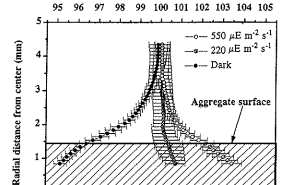
material. An example of the aggregates in the experimental setup is shown in Fig. 1.

Photosynthesis and respiration

The oxygen gradients in the diffusive boundary layer (DBL) at the aggregate-water interface and 0.5 mm below the upper surface of the aggregates were measured in darkness, at the compensating light intensity, and at the light intensity at which photosynthesis was saturated. The gradients were small but significant as shown by the small standard deviations of the mean concentrations at the different light conditions. At the aggregate surface, the $\rm O_2$ concentration ranged from 96% in darkness to 102% of air saturation measured at 550 $\rm \mu E~m^{-2}~s^{-1}$ (Fig. 2). At the compensating light intensity all the $\rm O_2$ produced by photosynthesis is consumed by the heterotrophic processes within the community, and there is no net exchange of oxy-

gen between the aggregate and the surrounding water. The compensating light intensity was 220 uE m⁻² s⁻¹ in the aggregate shown in Fig. 2. The aggregates were very sticky on the electrode and the distribution of O2 could, therefore, not be determined through whole aggregates without pushing the aggregates with the electrode during the measurements. Oxygen concentrations inside aggregates in the dark were always >90% of air saturation (data not shown). The net oxygen exchange between the aggregate community and the surrounding water occurs through the DBL. The gradients through the DBL at the aggregate-water interface are, therefore, determined by the net photosynthesis and respiration rates of the whole aggregate community. The net oxygen fluxes of sinking aggregates have been shown not to differ significantly between the upstream and downstream side (Ploug et al. 1997, Ploug & Jørgensen 1999). The gradients in the DBL of only the upper (i.e. downstream) side were, therefore, used to determine net photosynthesis and respiration rates of the whole aggregate community.

Net photosynthesis ranged from 0 to 340 ng C agg. $^{-1}$ d $^{-1}$ at saturating light intensity, and the dark respiration ranged from 80 to 3600 ng C agg. $^{-1}$ d $^{-1}$ in aggregates of different sizes (Fig. 3). The gross photosynthesis calculated as net photosynthesis + dark respiration ranged from 110 to 3600 ng C agg. $^{-1}$ d $^{-1}$. The average compensating light intensity was 152 ± 61 μ E m $^{-2}$ s $^{-1}$, and dark respiration comprised on average 75 ± 21% of gross photosynthesis at saturating light intensities of >500 μ E m $^{-2}$ s $^{-1}$. In some aggregates the dark respiration was equal to the gross photosynthesis also at saturating light intensities, meaning that net photosynthesis was zero. The autotrophic and heterotrophic



O2 (in % of air saturation)

Fig. 2. Oxygen gradients in the diffusive boundary layer (DBL) of the aggregate shown in Fig. 1 measured in darkness and at 2 different light intensities. Data points represent the mean \pm SD of 3 gradients Note the expanded O_2 scale

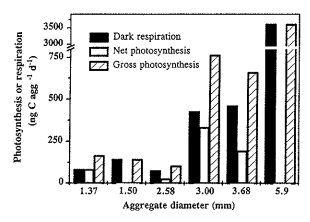


Fig. 3. Gross photosynthesis, net photosynthesis, and respiration rates in individual aggregates as determined from the oxygen fluxes in dark and at saturating light intensities (see text). Absence of photosynthesis columns means zero rate. Each bar represents the mean of 3 gradient measurements

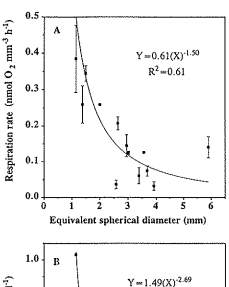
processes were therefore tightly coupled through carbon and oxygen being recycled within the aggregate. Marine snow of the surface waters were, thus, net heterophic communities not only in the dark, but also when the light intensity was below $152 \pm 61 \,\mu\text{E} \,\text{m}^{-2} \,\text{s}^{-1}$.

Community respiration rates and bacterial numbers in aggregates

The respiration rate per aggregate volume decreased with increasing aggregate size (Fig. 4A), although the total respiration rate increased with increasing aggregate size (Fig. 3). Volume-specific respiration rates could be described by: $0.61 \times ESD^{-1.50}$ nmol O_2 mm⁻³ h⁻¹, where ESD is in mm. R^2 was 0.61.

The number of bacteria attached to the aggregates was on average 1.16×10^6 agg.⁻¹, or ca 2000-fold higher per aggregate volume than in the surrounding water. The number of bacteria per aggregate volume decreased nearly 2 orders of magnitude with increasing aggregate size according to bacterial density = 1.49 $\times 10^9 \times ESD^{-2.69}$ bac. ml⁻¹; R² = 0.98 (Fig. 4B). The bacterial densities decreased relatively more with increasing aggregate diameter than the volume-specific respiration rates. Community respiration rates per bacterial cell were lower in small aggregates than in large aggregates (Fig. 5A). The large aggregates harbored less bacteria per aggregate volume, and community respiration rates per bacterial cell decreased with increasing bacterial density as described by: 10.4 × (bac. ml⁻¹)^{-0.54} fmol O_2 bac.⁻¹ d⁻¹; $R^2 = 0.66$ (Fig. 5B).

The high respiration rates per bacterial cell indicate that aggregate-associated bacteria were catabolically highly active. The bacterial number in aggregates



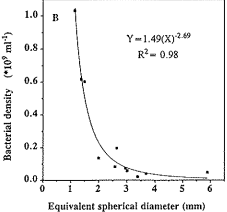


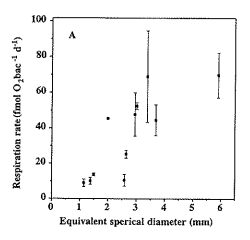
Fig. 4. (A) Respiration rates in the dark and (B) bacterial densities normalized to aggregate volume for different aggregate sizes. Each data point represents the mean \pm SD of the mean for 3 series of measurements

fixed immediately after collection was 2.4-fold lower than of the number fixed after the microelectrode measurements or after 1 to 3 d of incubation in the glass sampling bottles without stirring (Table 2). A high percentage of all aggregate-attached bacteria belonged to the Bacteria domain (92 to 99.2%), of which 8.9 to 73.7% belonged to Cytophaga. The total number of bacteria and the percentage of Bacteria as well as of the Cytophaga group increased with time after collection. The high detection level by the 16 S rRNA oligonucleotide probes as well as the increase in total number with time indicates that bacteria on marine snow were highly active. The increase in bacterial numbers with time, however, can partly be due to cellular growth of the aggregate-attached bacteria as well as increased colonization by bacteria from the surrounding water.

Enzymatic hydrolysis rates. The potential hydrolysis rates by α - and β -glucosidase, aminopeptidase, phosphatase, and chitinase were measured at high sub-

Table 2. Bacteria number per aggregate and community composition versus age. Data show the mean \pm SD values. Data range is	١	CO	. C	ì	1 (i c	d c	d (d (i c	C	or	m	n	າເ	ur.	ì	ty	y	C	OD	aj)(25	si	ti	oı	ì	V	6	rs	sι	ıs	a	ge	€.	Γ)a	ta	S	h	OΜ	' t	h€	n	ıea	ın	±S	SD) v	al	ue	S.	D	ata	ŧΪĉ	ınς	je:	is	
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Age at fixation	DAPI-stained	Bacteria	Cytophaga
	bacteria (× 10 ⁶)	(% of DAPI)	(% of Bacteria)
Freshly collected aggregates (n = 27 ; each sample = 10 pooled agg.)	0.48 ± 0.16	96.3	22.9
	(0.13-0.79)	(92.0–98.5)	(8.90–62.7)
Fixed after O_2 measurements within the day of collection (n = 15)	1.16 ± 1.13 $(0.37-5.7)$	97.0 (95.7–99.0)	28.4 (13.7–71.9)
Fixed after 1 d incubation without stirring (n = 7)	1.17 ± 0.27	98.2	32.2
	(0.77–1.36)	(94.5-99.2)	(13.0-50.6)
Fixed after 3 d incubation without stirring $(n = 4)$	1.00 ± 0.49 $(0.67-1.73)$	96.7 (95.3–98.5)	33.22 (17.3–73.7)



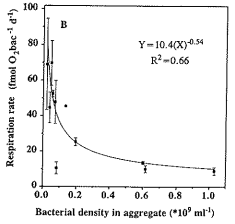


Fig. 5. (A) Respiration rates per bacterial cell relative to the aggregate size, and (B) the bacterial density. Each data point represents the mean \pm SD for 3 series of measurements

Table 3. Potential hydrolysis rates in aggregates. Data show the mean \pm SD values. Data range is shown in parentheses

Enzyme activity	Rate (fmol bac. $^{-1}$ h $^{-1}$)
21-22 Jun	
α-Glucosidase	$5.23 \pm 5.0 (0.83 - 13.5)$
β-Glucosidase	$2.43 \pm 1.44 (0.75 - 4.2)$
Phosphatase	$1.22 \pm 1.37 \ (0.04 - 3.20)$
Chitinase	$2.43 \pm 1.44 \ (2.37 - 6.93)$
Aminopeptidase	$21.1 \pm 7.44 \ (12.9 - 36.0)$
27-28 Jun	
Aminopeptidase	187 ± 120 (69.5-368)

peptidase activity was 21.1 fmol bac.⁻¹ h⁻¹ on 21-22 June and increased to 187 fmol bac.⁻¹ h⁻¹ on 27-28 June.

Carbon turnover by microbial respiration on marine snow

The mean respiration rates were 0.433 ± 0.215 and $0.984 \pm 1.78 \,\mu g \, C \, agg.^{-1} \, d^{-1}$, respectively, on 27 and 28 June. The respiration rates were calculated as the mean of aggregates of random sizes, which explains the high standard deviation of the mean. The carbon turnover time due to respiration alone estimated from the ratio of the POC content and the mean respiration rates was 7.9 and 8.9 d, respectively, on 27 and 28 June. Settling velocities of the aggregates were very variable (10 to 65 m d⁻¹). The average settling velocity was relatively low 31 ± 18 m d⁻¹, presumably due to their high mucus content. As the water column was approximately 200 m deep, the average minimum residence time was 7 d. Assuming that the respiration rate was independent of time and quality of the carbon pool, 80% of the POC content could potentially be respired before the aggregates reached the sea floor.

Carbon to support the respiration in aggregates may, however, also be scavenged from the surrounding water by sinking aggregates in the ocean.

DISCUSSION

Photosynthesis and respiration

Photosynthesis, bacterial production, and enzymatic hydrolysis have been studied intensively on marine snow (Alldredge et al. 1986, Alldredge & Gotschalk 1990, Simon et al. 1990, Smith et al. 1992, Turley & Mackie 1994); however respiration rates have not previously been quantified. Photosynthesis has been measured by ¹⁴C-techniques in disrupted marine snow. The rates were found to vary widely between 8.1 and 2170 ng C agg. ⁻¹ d⁻¹ (Alldredge & Gotschalk 1990, Simon et al. 1990).

In the present study, net and gross photosynthesis were calculated from the $\rm O_2$ fluxes to or from intact sinking aggregates during darkness and in light. The respiration in light cannot be detected by the steady-state fluxes; however it can be detected from the dynamic changes of oxygen concentrations during shifts between light and dark (Revsbech et al. 1981). Respiration during photosynthesis has been shown to be up to 5-fold higher than dark respiration in diatom aggregates, presumably due to photorespiration in the algae and/or excretion of easily degradable carbohydrates which may stimulate bacterial respiration (Ploug & Jørgensen 1999). The gross photosynthesis may, therefore, be underestimated, as the respiration in light probably was higher than the dark respiration.

Net photosynthesis at light saturation ranged from 0 to 340 ng C agg. -1 d-1. Gross photosynthesis was much higher and ranged from 110 to 3600 ng C agg.⁻¹ d⁻¹, as most of the oxygen produced during photosynthesis was respired within the aggregates. The aggregates were net heterotrophic at light intensities up to 152 \pm 64 μE m⁻² s⁻¹. Compensating light intensities in colonies and aggregates of the planktonic microalgae, Phaeocystis sp. have previously been found to be much lower (7 to 10 μ E m⁻² s⁻¹) in early stages during spring blooms in the North Sea and in the Barents Sea, and to increase at later stages of blooms (80 µE m⁻² s⁻¹), presumably due to increased bacterial colonization during the bloom. Dark respiration in such colonies and aggregates only comprised 12% of the gross photosynthesis measured from the steady-state fluxes at the early stages of the blooms (Ploug et al. in press). In the present study, the heterotrophic processes were much more important in the aggregate communities, as reflected in the relatively high light intensity at which photosynthesis could balance respiration within the

community and the high dark respiration which comprised 75 \pm 21% of gross photosynthesis at light saturation.

Dark respiration and oxygen microenvironments

Steep gradients of oxygen and pH have previously been demonstrated in marine aggregates and fecal pellets placed in a small cone made of a plankton net (Alldredge & Cohen 1987). In a recent study it was shown that sinking of aggregates greatly facilitates diffusion to the aggregates and results in 9-fold higher O_2 concentration inside the aggregates as compared to aggregates sitting on a solid surface (Ploug & Jørgensen 1999). The flow past the sinking aggregates is therefore important for its chemical microenvironment and, thus, for the growth conditions of the microbial community. Measured oxygen concentrations in darkness inside the aggregates of the present study were ≥90% of air saturation (225 µM), at which the organisms cannot be diffusion limited by oxygen since the half-saturation constant, K_S , for oxygen respiration is on the order of 1 µM in heterotrophic microorganisms (Focht & Verstraete 1977). If the aggregates had occurred in an oxygen minimum zone with a bulk water concentration of 10% of air saturation they may have been anoxic.

Oxygen gradients are proportional to the community respiration rates in the aggregates when K_S is much lower than the oxygen concentration inside the aggregates (Ploug et al. 1997). Anoxia inside the aggregates of the present study with air-saturated water would, therefore, have required ≥10-fold higher cell-specific respiration rates, or 10-fold higher community densities. The bacterial numbers in those aggregates which we analyzed with microelectrodes ranged between 0.37×10^6 and 5.7×10^6 . Other studies have also shown that bacteria numbers in aggregates larger than 3 mm usually range between 0.41×10^6 and 1.7×10^6 in the Southern California Bight (Alldredge et al. 1986, Simon et al. 1990). Such bacterial densities with similar respiration rates as estimated in the present study would be 10-fold too low to create anoxic conditions inside the aggregates. Higher bacterial numbers have been observed in aggregates from the NE Atlantic, but bacterial numbers in the range of 107 to 108 in aggregates are rare unless the aggregates are very large (Alldredge et al. 1986, Alldredge & Gotschalk 1990). This higher limit of bacterial densities in aggregates may be determined by diffusion limitation of bacterial growth, available surface area, and probably grazing.

Flagellates have been shown to occur in high numbers on aggregates (Caron et al. 1982, Turley & Mackie 1994). Their contribution to the community respiration

is not known since it is difficult to measure separately. The average bacterial biomass on aggregates was 6.5 nmol C, assuming a bacterial carbon content of 5 fmol C bac. (Simon et al. 1990). If 50% of the bacterial biomass was grazed per day by flagellates growing with a 50% growth efficiency and a respiratory quotient of 1.2 mol O_2 : mol CO_2 , the respiration rate by flagellates would be $[(0.5 \times 0.5 \times 6.5 \text{ nmol } CO_2 \times 1.2 \text{ mol } O_2 \text{ mol } CO_2^{-1})/24 \text{ h}] = 0.081 \text{ nmol } O_2 \text{ h}^{-1}$. The average community respiration rate of the measured aggregates was $2.7 \pm 4.2 \text{ nmol } O_2 \text{ h}^{-1}$. The flagellates would, thus, contribute with 3% to the total community respiration.

Dark respiration and carbon turnover

The respiration rates normalized to bacterial number ranged from 1.5- to 12-fold the bacterial carbon biomass (7.4 to 58 fmol C bac.⁻¹ d⁻¹) assuming an average cell-specific carbon content of 5 fmol C bac.-1. The bacterial carbon content on marine snow, however, can vary 4-fold (Simon et al. 1990). Respiration rates of free-living bacteria have been found in the same range in seawater samples highly enriched with glucose (Stoderegger & Herndl 1998). The cell-specific bacterial production rate has also been shown to increase with increasing aggregate size and decreasing bacterial density in marine aggregates. At low densities, bacteria may compete less for available resources and therefore grow faster (Alldredge & Gotschalk 1990). The high respiration rates measured in the present study suggest high growth rates or a low growth yield of the attached bacteria. The doubling time of bacteria attached to aggregates has been shown to range between 0.5 and 10 d (Alldredge & Gottschalk 1990, Simon et al. 1990, Grossart & Simon 1998). Unfortunately, we could not measure bacterial growth rates in this study.

Recent studies have shown that, whereas bacterial production is high and enzymatic hydrolysis is low in newly formed aggregates, bacterial production decreases and hydrolysis increases in aggregates after ca 3 d (Unanue et al. 1998). The potential hydrolysis rates and the C:N ratio were high on the aggregates, which indicate that the aggregates were not newly formed (Table 3). Cytophaga were highly abundant on the aggregates, and the relative abundance of this group was shown to increase with time when aggregates were allowed to age in the lab (Table 2). Similar results were also found in a limnic study (Schweitzer 1998). Bacteria of the cytophaga group have previously been shown to dominate the bacterial community on marine snow, and they are known to be motile and can hydrolyze a variety of more refractory polymeric compounds, especially chitin and other polysaccharides (DeLong et al. 1993, Rath et al. 1998). The short turnover time of carbon by respiration may be a consequence of the cost of producing exoenzymes, which may lead to a low growth yield of the attached bacteria. Bacterial growth efficiencies have been shown to be 8 to 39% for free-living bacteria degrading high molecular weight DOC, and 16 to 66% when they degrade low molecular weight DOC (Amon & Benner 1996).

A recent study of respiration rates and carbon turnover in lake snow in Lake Kinneret, Israel, showed respiration rates of 0.65 to 1.04 nmol O₂ agg.⁻¹ h⁻¹ and a POC turnover time of 14 to 50 d, as the respiration rates were lower and the POC content was higher in the aggregates of that study (Parparov et al. 1998). The POC content in the aggregates of the present study was in the same range but the dry weight was in the high range of that which has been previously found in marine aggregates (Alldredge 1979, 1998). The POC content of aggregates has been shown to comprise 8 ± 1% of the dry weight during summer in the Gulf of California and 15 to 19% of the dry weight during winter in the Santa Barbara Channel (Alldredge 1979). The POC content as a percentage of dry weight is generally lower and the C:N ratio is higher in miscellaneous aggregates as compared to other types of aggregates of recognizable organic matter, e.g. diatoms, larvacean or fecal pellets (Alldredge 1998). The low POC content relative to the dry weight and the high C:N ratio of the aggregates in the present study indicates that the aggregates were mainly dominated by inorganic matter and more refractory organic matter, presumably due to high hydrolysis and respiration rates.

Implications for the vertical flux of carbon

The respiration rates per aggregate volume decreased with increasing aggregate size (Fig. 4A). Also the dry weight, the POC and the PON content per aggregate volume showed a similar decrease with increasing aggregate size (Alldredge 1998). The common size-dependent distributions of dry weight, POC, and PON in aggregates may all be related to a general fractal geometry of the aggregates, because porosity increases with increasing aggregate size as the aggregates are formed by coagulation of smaller particles (Logan & Wilkinson 1990, Mari & Kiørboe 1996, Mari & Burd 1998).

The size-dependent respiration rate as measured in the present study versus the size-dependent POC content of aggregates as determined by Alldredge (1998) is shown in Fig. 6A. The carbon turnover time as a function of aggregate size was calculated from the POC content

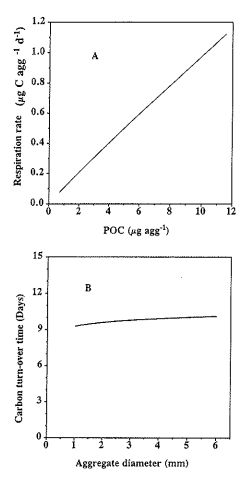


Fig. 6. (A) Respiration rates and (B) carbon turnover time in aggregates of different sizes, calculated using the size specific POC content in aggregates as determined by Alldredge (1998)

(Alldredge 1998) divided by the measured respiration rate as a function of aggregate size from the present study (Fig. 6B). Due to the similar size dependency of respiration and POC content in marine aggregates, the respiration rates appeared to be proportional to the POC content, which suggests a tight coupling between POC and respiration in aggregates. As a consequence, the carbon turnover time was rather independent of aggregate size. The turnover time only increased from 9.3 to 10.1 d for 1 to 6 mm large aggregates (Fig. 6B). Our estimates of the carbon turnover time of 8 to 9 d was, thus, close to that predicted in Fig. 6.

If the carbon turnover time is the same for all aggregate sizes as indicated in the present study, the degree of carbon remineralization in aggregates is determined by their residence time in the water column, which also depends on aggregation, scavenging, and disintegration processes. Large aggregates are less common than small ones in surface waters, because they rapidly sink out of the surface waters or disintegrate and produce smaller aggregates due to turbulence (Alldredge

et al. 1990, Jackson et al. 1997). The residence time of aggregates in the water column and the carbon fluxes are further dependent on grazing and defecation by zooplankton (Lampitt et al. 1993, Green & Dagg 1997) and fishes (Grossart et al. 1998). Since small aggregates are more common in the ocean than larger ones, and small aggregates have much higher respiration rates and POC content per volume compared to large aggregates, they may contribute more to the carbon remineralization in surface waters than previously presumed.

The flux of POC and PON decreases with increasing depth in the ocean, the highest losses occurring in the upper ocean (<150 m) followed by a more moderate decrease in the deeper parts of the water column (Knauer et al. 1979, Knauer & Martin 1981, Lee & Wakeham 1988). In the present study, we have estimated that 80% of the aggregate carbon would be respired within the aggregate during sedimentation through a water column of 200 m, as these aggregates were relatively slowly sinking. As a consequence of the short turnover time of the aggregate carbon, the respiration in aggregates will become carbon limited and decrease during sedimentation when degradable carbon sources to support respiration can no longer be produced within the aggregates as exudates and hydrolysis products or be scavenged from the surrounding water.

The present study directly demonstrates that sinking aggregates are not only a sink of CO_2 fixed in the euphotic zone but they can also be a source of CO_2 to the surrounding waters at relatively high light intensities up to 150 $\mu E \ m^{-2} \ s^{-1}$ and during sedimentation out of the euphotic zone. The carbon turnover in sinking aggregates due to respiration, only, can be so fast that most of the aggregate carbon will be respired in the surface waters and only a minor fraction of the aggregate carbon will reach the sea floor.

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