Formation of transparent exopolymer particles, TEP, from dissolved precursor material

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ABSTRACT: Transparent exopolymer particles (TEP) form from polysaccharides released by many phytoplankton species, but this process by which dissolved organic matter becomes particulate is poorly understood. Here, the abiotic formation of TEP from precursors <0.2 μm and the minimum molecular weight (MW) of TEP-precursors were studied. In most samples TEP formed from material <0.2 μm (poly carbonate membrane filters, Whatman) when exposed to laminae shear in Couette (i.e. suspensions). This result was unexpected as no TEP formed from material <0.45 μm (polycap capsules). Whatman due to surface coagulation onto bubbles (Zhou et al. 1998, Limnol Oceanogr 43:1860–1871). Some TEP-precursors were able to pass through dialysis bags with a nominal pore size of 8 kDa (natural cellulose, Spectrum), although their MW is presumably 2 orders of magnitude larger, suggesting that TEP-precursors can be fibrillar. It is suggested that freshly released precursors are fibrillar and that the fibrillar precipitation form larger colloids and eventually TEP within hours to days after their release.

KEY WORDS: TEP · Particle formation · Exudation

INTRODUCTION

Dissolved organic carbon (DOC) increases during or shortly after blooms due to phytoplankton excretions (Kepkay et al. 1993, Niven et al. 1995). The fate of this newly generated carbon depends largely on how it is partitioned among the different carbon pools, because processes controlling the cycling of carbon in the ocean differ between DOC, colloidal organic carbon (COC) and particulate organic carbon (POC) (Gustafsson & Geschwend 1997). Solutes and colloids do not sink, but are transported passively with the ambient water, without substantial removal by settling. However, residence times of colloids in seawater are similar to those of small particles (Moran & Büsseler 1992, Huh & Frahl 1995) and, in contrast to solutes, colloids can coagulate and form aggregates (Wells & Goldberg 1993, Kepkay 1994) which eventually sediment or are grazed. Thus, in the cycling of carbon in the ocean is to be understood, mechanisms and rates of transformations between the different carbon pools must be understood.

The recent discovery of the high abundance (10^6 to 10^8·l⁻¹) of previously unrecognized particles called TEP (transparent exopolymer particles, Allerdige et al. 1993) further highlights the need to understand and quantify particle production in the ocean. TEP, which consist of acidic polysaccharides, are filterable on 0.4 μm filters and can be up to several 100 μm in length. TEP form from polysaccharides released by phytoplankton (Passow & Allerdige 1994, 1995a, Passow et al. 1994). The formation of TEP may potentially represent a pathway by which DOC released by phytoplankton is removed from the eutrophic zone, because, as an essential component of marine snow (Allerdige et al. 1993, Logan et al. 1995), TEP can sediment rapidly (Passow et al. 1994).

Transformations between the different fractions of organic carbon may be mediated by biotic or by abiotic processes. Biotic transformations include the formation of POC from DOC via bacterial utilization (Chrost & Faust 1983, Sundh 1989, Sel & Overbeck 1992, Alber & Valiela 1994) or via grazer feeding on colloidal mat-

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ter (Sherr 1988, Flood et al. 1992, Tranvik et al. 1993). Abiotically, particles can be generated from non-filterable material by adsorption to hard surfaces, like carbonate, clay or detrital matter (Otsuki & Wetzel 1973, Avnimelech et al. 1982, Jensen & Sondergaard 1982, Fattom & Shilo 1984, Bar-Or & Shilo 1987, Bender et al. 1994), or by surface coagulation onto rising bubbles (Baylor & Sutcliffe 1963, Riley 1963, Sutcliffe et al. 1983, Johnson 1975, Johnson & Cooke 1980). However, it is believed that colloids between 0.2 and 1.2 μm are needed for abiotic particle formation as no particulate matter is formed by bubbling material <0.2 μm (Johnson et al. 1986, Kepkay & Johnson 1989).

TEP can also form by 2 abiotic processes: adsorption onto surfaces (Passow & Wassmann 1994) and bubble scavenging (Mopper et al. 1993, Zhou et al. 1998). Precursors of TEP appear to also be >0.45 μm, as in previous studies very little TEP formed from substances <0.45 μm (Polycap capsules, Whatman) via bubble absorption (Zhou et al. 1998). However, TEP formed on particles (siliceous fibers) from mucus generated from Phaeocystis spp. colonies by pressure filtration through 0.2 μm (membrane) and enrichment onto 300 Da (Passow & Wassmann 1994).

Experiments presented in this paper were conducted to determine if TEP can form from non-filterable precursors at natural concentrations in the absence of particles or bubbles and if TEP can form from precursors <0.4 μm in size. Another goal was to determine the minimum size of precursors from which TEP can form abiotically. Finally, experiments were conducted with samples representing different plankton assemblages under various growth conditions to assess if characteristics of abiotic TEP formation varied as a function of the phytoplankton releasing the precursors.

METHODS

The abiotic formation of TEP from precursors <0.2 μm (polycarbonate membrane, Poretics) was investigated by exposing precursors to laminar shear in Couette flocculators (flocculator experiments). The minimum size of precursors able to form TEP was studied in dialysis bag experiments. A total of 14 flocculator and 11 dialysis bag experiments were conducted using water from various sampling sites reflecting different ecological conditions. Between 1 and 4 dialysis bag experiments accompanied most flocculator experiments. Table 1 gives an overview of the experiments conducted.

Flocculator experiments. Flocculator experiments were primarily designed to study the abiotic formation of TEP from material <0.2 μm in the absence of bubbles or particles. TEP are, by definition, particles retained by 0.4 μm filters (polycarbonate membrane, Poretics), whereas TEP-precursors include all substances which pass through 0.4 μm filters, but will potentially form TEP. In flocculator experiments the formation of TEP was studied from precursors <0.2 μm (polycarbonate membrane, Poretics), to be conservative. A second goal was to use TEP production as an estimate of relative amounts of precursor concentration. Precursor concentration can currently not be measured directly, as TEP-precursors appear to be a chemically complex group of acidic polysaccharides which can form particles (TEP). The formation of TEP under constant conditions should represent a relative estimate of precursor concentration.

A small (volume: 160 ml) Couette flocculator was used for the 5 experiments conducted in June 1995 and a larger Couette flocculator (volume: 1600 ml) was used for experiments conducted during the mesocosm study in November 1995 and the study with Phaeocystis spp. in April 1996. A Couette flocculator, which consists of a fixed inner and a rotating outer cylinder, provides a quantifiable 2-dimensional laminar shear in the annular space between the cylinders (Duvaren 1968). Expts F1 to F5 (Table 1) were rotated at a shear of 152 s⁻¹, whereas the larger flocculator generated a shear of 24.9 s⁻¹. In all experiments samples were pre-filtered through 0.2 μm filters (polycarbonate membrane, Poretics) and incubated in the flocculator for 24 h. When pre-filtering water for flocculator experiments, care was taken to avoid collecting TEP in the filtrate by keeping the filtration pressure below 150 mm Hg and by not letting the filter become dry. The concentration of TEP was determined initially in the 0.2 μm filtered water and again after 24 h of incubation. Flocculators and flasks were rinsed well with ethanol, and then 3 times with distilled water to avoid bacterial contamination of 0.2 μm filtered seawater. Production of TEP in flocculators was calculated by subtracting the initial concentration of TEP in flocculators from the final concentration.

Dialysis bag experiments. Dialysis bag experiments were primarily designed to estimate the minimum molecular weight (MW) of TEP-precursors. For each experiment seawater was incubated within dialysis bags of varying MW cut-off, and the formation of TEP monitored both within bags and in the water surrounding the bags. It was hypothesized that TEP-precursors generated by the assemblage of organisms growing within the dialysis bags would potentially diffuse through bags and that TEP would form abiotically in the pre-filtered, autoclaved seawater (PASW) surrounding the bag. Total production of TEP, which should reflect ambient precursor concentration + newly released precursors, was calculated by adding the amount of TEP formed in PASW and within bags at the end of the experiment and subtracting the initial concentrations.
Table 1. Flocculator and dialysis bag experiments conducted at each of the sampling dates (MW: molecular weight cut-off of dialysis bags, n: number of replicate dialysis bag experiments)

<table>
<thead>
<tr>
<th>Sample description</th>
<th>Collection date (mo/d/yr)</th>
<th>Flocculator experiments</th>
<th>Dialysis bag experiments</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Expt no.</td>
<td>Shear (s⁻¹)</td>
</tr>
<tr>
<td>Diatom bloom in Santa Barbara Channel</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diatom bloom at peak</td>
<td>6/17/95</td>
<td>F1</td>
<td>15.2</td>
</tr>
<tr>
<td>Diatom bloom at peak</td>
<td>6/17/95</td>
<td></td>
<td></td>
</tr>
<tr>
<td>fraction &gt;5 μm</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diatom bloom, cloudy</td>
<td>6/18/95</td>
<td>F2</td>
<td>15.2</td>
</tr>
<tr>
<td>Diatom bloom, flocced</td>
<td>6/19/95</td>
<td>F3</td>
<td>15.2</td>
</tr>
<tr>
<td>Diatom bloom grown in a mesocosm</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lag-phase</td>
<td>10/28/95</td>
<td>F8</td>
<td>24.9</td>
</tr>
<tr>
<td>Growth phase</td>
<td>10/29/95</td>
<td>F9</td>
<td>24.9</td>
</tr>
<tr>
<td>Growth phase</td>
<td>10/30/95</td>
<td>F10</td>
<td>24.9</td>
</tr>
<tr>
<td>Late growth phase</td>
<td>11/1/95</td>
<td>F11</td>
<td>24.9</td>
</tr>
<tr>
<td>Senescence</td>
<td>11/2/95</td>
<td>F12</td>
<td>24.9</td>
</tr>
<tr>
<td>Senescence</td>
<td>11/3/95</td>
<td>F13</td>
<td>24.9</td>
</tr>
<tr>
<td>Non-diatom systems</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Growing Phaeocystis spp.</td>
<td>4/18/95 or 4/25/96*</td>
<td>F15</td>
<td>24.9</td>
</tr>
<tr>
<td>Growing Phaeocystis spp.</td>
<td>4/18/96</td>
<td></td>
<td></td>
</tr>
<tr>
<td>fraction &lt; 200 μm</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Growing Phaeocystis spp.</td>
<td>4/19/96</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Growing Phaeocystis spp.</td>
<td>4/19/96</td>
<td></td>
<td></td>
</tr>
<tr>
<td>fraction &lt; 200 μm</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Phaeocystis spp. aged 5 d</td>
<td>4/18/96</td>
<td>F16</td>
<td>24.9</td>
</tr>
<tr>
<td>Phaeocystis spp. aged 5 d</td>
<td>4/18/96</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Deep water (chl a-poor)</td>
<td>7/12/96</td>
<td>F4</td>
<td>15.2</td>
</tr>
<tr>
<td>Nanoflagellate bloom</td>
<td>6/21/95</td>
<td>F5</td>
<td>15.2</td>
</tr>
</tbody>
</table>

*D8 and D9 were conducted with growing Phaeocystis spp. collected on 18 April 1996, whereas F15 was conducted with growing Phaeocystis spp. collected on 25 April 1996

For each dialysis bag experiment, between 9 and 27 clear 280 ml bottles were rotated on rolling tables at in situ temperature and low light (20 to 60 \( \mu \text{E m}^{-2} \text{s}^{-1} \)) for 44 to 72 h (Table 1). Each bottle contained a dialysis bag (Spectra/Por 7 from Spectrum, made of natural cellulose) filled with 50 to 100 ml of fresh, raw seawater. Dialysis bags with a MW cut-off between 2 and 50 KDa were used (Table 1). Dialysis bags were suspended in pre-filtered (0.2 μm), autoclaved seawater (PASW) collected from below 100 m. The sterile PASW contained no TEP and little or no TEP-precursors. The continuous rotation on rolling tables ensured mixing during the experiment and facilitated formation of TEP as rolling tables generate turbulent shear (Jackson 1994). Concentrations of TEP were determined initially and at the end both within the bags and in the PASW surrounding the bags. Each experiment consisted of 3 to 4 replicate bottles plus 2 to 4 control bottles. Empty dialysis bags suspended in PASW were used as controls. All containers were either autoclaved or washed with ethanol and rinsed 3 times with sterile water to avoid contamination and growth of bacteria in the water surrounding the dialysis bags.

Additional dialysis bag experiments (D1b, D8b, D9b) were conducted to test which size fraction of the phytoplankton community was responsible for the production of TEP-precursors. In experiment D1b, particles <5 μm were removed from the sample (particles retained on 5 μm mesh were resuspended in 0.2 μm filtered seawater), incubating predominately diatoms. In experiments D8b and D9b Phaeocystis spp. colonies were excluded by filtration through a 200 μm mesh (Table 1). Single Phaeocystis spp. cells, which were not excluded, did not form colonies during the experiment.

Apparent mass transfer coefficients, which can be viewed as the inverse of the resistance against precursors passing through bags, were estimated for each experiment because they allow direct comparisons.
between experiments, even when experimental conditions and type of bag vary. Theoretically, the amount of TEP formed in PASW depended on the diffusion of precursors through bags. Besides being a function of the MW of precursors in relation to the MW cut-off of bags, the rate of diffusion depended upon the duration of the experiment, the surface area of the bag and the concentration gradient of the precursors. Mass transfer coefficients normalize for these latter variables. However, as I could not measure precursor concentrations directly, the production of TEP was used as a relative estimate of ambient precursor concentration. Apparent mass transfer coefficients were thus estimated from TEP concentrations using the following relationship (Cussler 1984): the flux through dialysis bags \( F_{\text{TEP}} \) (\( \mu g \) xan. eq. m\(^{-2}\) s\(^{-1}\)) equals the concentration gradient times the apparent mass transfer coefficient (k: m s\(^{-1}\)). Since TEP concentration is used as a measure for the precursor concentration:

\[
F_{\text{TEP}} = k \cdot (\text{TEP} - \text{TEP}_{\text{PASW}})
\]

TEP, and TEP\( _{PASW} \) (\( \mu g \) xan. eq. m\(^{-2}\)) represent the concentration of TEP inside and outside of bags respectively at the end of the experiment.

**Sample types. Diatom blooms:** Some flocculator (F1 to F3) and dialysis bag (D1 and D2) experiments were conducted with water collected at the chlorophyll (chl) a maximum (at 10 to 15 m) during the flocculation of a diatom bloom in the Santa Barbara Channel (SBC) in summer 1995. Flocculator (F8 to F14) and dialysis bag (D4 to D7) experiments were also conducted with samples collected during the growth and decline of a diatom bloom grown in a 1400 l mesocosm in fall 1995. To initiate the bloom, raw seawater from a depth of 15 m was inoculated with 50 l of surface water from the SBC and nutrients were added (NO\(_3\) + NO\(_2\) = 60 \( \mu M\), SiO\(_2\) = 50 \( \mu M\), PO\(_4\) = 5 \( \mu M\)). The temperature in the mesocosm was held constant at 12\(^\circ\)C, a 14:10 h light:dark regime was imposed and the mesocosm was mixed by a propeller rotating slowly near the bottom (for a detailed description of the mesocosm see Aldredge et al. 1995).

**Non-diatom phytoplankton:** For comparison, experiments were also conducted with samples stemming from blooms dominated by Phaeocystis spp. or nanoflagellates. Expts F15, D8 and D9 were run with water collected during the growth phase of a bloom dominated by colonial Phaeocystis pouchetti which developed in Balsfjord Svarnines, Norway (69\(^\circ\) 21.8' N, 19\(^\circ\) 06.5' E) in April 1996. An aliquot of the water sampled on 18 April was aged at in situ temperature (2\(^\circ\)C) in the dark for 5 d and then used in lieu of a senescent Phaeocystis spp. population from the field for Expts F16, D10 and D11. Expts F5 and D3 were conducted with samples from a summer bloom of autotrophic nanoflagellates collected on 21 June in the SBC. One flocculator experiment (F4) was conducted with water from below the euphotic zone (150 m).

Chemical analysis. TEP was determined following the colorimetric method (Passow and Allerdge 1995b). Briefly, TEP were filtered onto 0.4 \( \mu m\) filters, stained with an Alcian Blue solution and soaked in 80% sulfuric acid. The dye redissolves in sulfuric acid and its concentration may be determined spectrophotometrically at 787 nm. Blanks (empty, but stained filters) were subtracted and absorption calibrated with gum xanthan. TEP concentrations are expressed as gum xanthan equivalent. This unit does not reflect the absolute weight of TEP and potentially the equivalent amounts of TEP generated under different conditions could result in varying degrees of staining. Standard deviations between the 4 replicate measurements conducted for each sample were usually below 5%, in a few cases between 5 and 10%. Chl a was determined in sample water using 2 replicate 20 ml samples filtered onto GF/F filters and analyzed following standard fluorometric methods (Parsons et al. 1984) using a Turner Model 110 fluorometer.

**RESULTS**

**Diatom blooms**

**Flocculator experiments**

Initial concentrations of TEP in 0.2 \( \mu m\) pre-filtered seawater were always near the detection limit of the method (<30 \( \mu g\) xan. eq. l\(^{-1}\)), but after subjecting the samples to shear generated in a flocculator for 24 h TEP was always present. Production of TEP during the diatom bloom in the SBC was highest on the first sampling day, 17 June (F1). About half as much TEP was generated in the sample collected 1 d later (F2) and correspondingly little TEP the following day (F3) (Fig. 1a). During the growth period of the mesocosm bloom (F9 to F13), TEP production in flocculators was relatively constant, and lower than during the lag phase (F8, Fig. 1b). During experiments conducted at the end of the mesocosm bloom (F14), however, TEP production in the flocculators increased appreciably.

**Dialysis bag experiments**

TEP could only form in PASW of dialysis bag experiments if precursors were (1) present within bags and (2) small enough to diffuse through bags. TEP was generated abundantly within dialysis bags of all exper-
Passow: Abiotic TEP formation

Fig. 1. Results of flocculator experiments. (a) diatom bloom in the Santa Barbara Channel, (b) diatom bloom in a mesocosm and (c) non-diatom systems. In each experiment concentration of TEP was measured in 0.2 μm filtered seawater initially (t = 0) and after 24 h incubation in a flocculator (flocculator 24 h). Error bars are standard deviations of replicate TEP measurements (analytical replicates). Note change of scale.

Experiments conducted with diatoms, showing that precursors were always present in the water during diatom blooms. TEP were not generated in PASW of controls, implying that (1) PASW was free of TEP and their precursors, (2) the bags did not generate any TEP-like artifacts and (3) no TEP-generating organisms (bacteria for example) were present. The absence of TEP in PASW of controls confirms that TEP in PASW of experiments must have formed abiotically from precursors passing through the bags.

TEP was formed in PASW during experiments with the diatom bloom from the SBC (D1 and D2, Fig. 2a,b) indicating that precursors passed through the 50 kDa bags. The amount of TEP formed in PASW was lower during Expt D2 sampled 1 d after Expt D1, but only a small amount of TEP formed within bags and the apparent transfer coefficient in Expt D2 was an order of magnitude higher compared to that in Expt D1 (Table 2). The higher apparent transfer coefficient in D2 suggests that precursors passed through bags more easily compared to those in Expt D1. Results of Expt D1b, where only the size fraction >5 μm was incubated, were the same as those of Expt D1a, where the whole water sample was incubated, indicating that the size fraction >5 μm, which consisted predominantly of diatoms, was primarily responsible for the release of TEP-precursors.

In experiments conducted at the beginning of the mesocosm diatom bloom (D4), the concentration of TEP in PASW was not significantly different from controls, suggesting that precursors did not pass through 8 kDa bags (Fig. 2c). However, the formation of TEP in PASW later in the bloom (D5 and D6) implies that TEP-precursors passed through 8 kDa bags at this time (Fig. 2d), which is confirmed by the increase of the apparent transfer coefficients by an order of magnitude (Table 2). During the experiment conducted at the end of the mesocosm bloom (D7), the median concentration of TEP in PASW was only marginally higher than that of controls (Fig. 2e), resulting in a very small apparent transfer coefficient (Table 2). The small apparent transfer coefficient suggests that precursors did not pass through 8 kDa bags at the end of the bloom.

Non-diatom blooms

Flocculator experiments

After subjecting the samples to shear some TEP had formed in the experiments with the growing Phaeocystis spp. (F15), but TEP production in the experiments conducted with the aging Phaeocystis spp. (F16) was negligible (Fig. 1c). TEP formed in the sample collected during a nanoflagellate bloom (F5, Fig. 1c), but almost no TEP was generated in the water collected from below the euphotic zone (F4). These results suggest that concentrations of TEP-precursors <0.2 μm in the deep water sample and in the sample of the aging Phaeocystis spp. were insufficient to form TEP. Precursor concentrations in water of the growing Phaeocystis spp. and the nanoflagellate bloom, however, were high enough for TEP to form abundantly.

Dialysis bag experiments

No TEP formed within bags of experiments conducted with the growing Phaeocystis spp. (D8 and D9) (Table 2), indicating that precursor concentrations
were low in samples of the growing *Phaeocystis* spp. Consequently no TEP appeared in the PASW of Expts D8 and D9 (Fig. 2i). Expts D8a and D8b as well as Expts D9a and D9b were similar, emphasizing that neither *Phaeocystis* spp. colonies (fraction >200 μm, D8a and D9a) nor organisms in the smaller size fraction (D8b and D9b) released significant amounts of TEP precursors during the experiments. Also, precursor concentrations at the time of sampling were low. Although TEP were formed abundantly within bags during experiments with aged *Phaeocystis* spp. colonies (D10 and D11, Table 2), TEP was negligible in PASW whether the 25 kDa (D10) (Fig. 2g) or the 2 kDa (D11) dialysis bags were used, implying that precursors were >25 kDa in samples with aged *Phaeocystis* spp. Small apparent transfer coefficients confirm that precursors generated by the aging *Phaeocystis* spp. could not pass through the bags (Table 2).

Significant amounts of TEP formed in PASW in the experiments with the nanoflagellate bloom, but because some of the controls were high, this result is ambiguous (Fig. 2h).

**Minimum size of precursors**

Actual MW cut-off of ultra-filters may differ from the nominal value given by the vendor and depend,
Table 2. Results of dialysis bag experiments (μg xanthan g⁻¹ = μg xanthan equivalent l⁻¹). n.v: no value

<table>
<thead>
<tr>
<th>Expt</th>
<th>Description</th>
<th>MW of bags (kDa)</th>
<th>Chl a within bags (μg l⁻¹)</th>
<th>TEP within bags t lig (μg xanthan g⁻¹)</th>
<th>TEP in PASW, t lig (μg xanthan g⁻¹)</th>
<th>Transfer coefficient, k (10⁻³ m s⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diatom bloom in Santa Barbara Channel</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>D1a Bloom peak</td>
<td></td>
<td>30</td>
<td>3.2</td>
<td>122</td>
<td>6309</td>
<td>958</td>
</tr>
<tr>
<td>D2 Cloning</td>
<td></td>
<td>50</td>
<td>1.9</td>
<td>245</td>
<td>405</td>
<td>347</td>
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<td>Diatom bloom grown in a mesocosm</td>
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<tr>
<td>D4 Lag-phase</td>
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<td>8</td>
<td>1.7</td>
<td>298</td>
<td>5941</td>
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<tr>
<td>D5 Growth phase</td>
<td></td>
<td>8</td>
<td>13.9</td>
<td>516</td>
<td>4643</td>
<td>2135</td>
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<td>D6 Growth phase</td>
<td></td>
<td>8</td>
<td>37.2</td>
<td>935</td>
<td>4794</td>
<td>1810</td>
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<tr>
<td>D7 Senescence</td>
<td></td>
<td>8</td>
<td>220.0</td>
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<td>D8a Growing P. th.</td>
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<td>25</td>
<td>2.8</td>
<td>770</td>
<td>833</td>
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<tr>
<td>D9a Growing P. th.</td>
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<td>2</td>
<td>2.8</td>
<td>770</td>
<td>843</td>
<td>78</td>
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<tr>
<td>D10 P. th. aged 5 d</td>
<td></td>
<td>25</td>
<td>nv</td>
<td>560</td>
<td>2488</td>
<td>115</td>
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<tr>
<td>D11 P. th. aged 5 d</td>
<td></td>
<td>2</td>
<td>nv</td>
<td>560</td>
<td>2177</td>
<td>115</td>
</tr>
<tr>
<td>D3 Nanoflagellates</td>
<td></td>
<td>8</td>
<td>2.0</td>
<td>140</td>
<td>871</td>
<td>902</td>
</tr>
</tbody>
</table>

¹Transfer coefficient of Expt 1b: 5.5 × 10⁻³ m s⁻¹  
²Transfer coefficients calculated from samples taken at different time steps during time-series studies were similar, indicating that apparent transfer efficiency remained the same during several days of the experiments.  
| k of Expt D2: at 48 h = 187 × 10⁻³ m s⁻¹; of D2: at 72 h = 282 × 10⁻³ m s⁻¹; of D2: at 96 h = 252 × 10⁻³ m s⁻¹ |
production in floculator experiments decreased even further (Fig. 3b).

The mesocosm bloom, which was also dominated by Chaetoceros spp., entered its exponential growth after 29 October, when both primary production and chl a concentration increased rapidly (Fig. 3c) until nutrients became limiting. Nutrients were depleted on 2 November (author's unpubl. results) and the bloom entered senescence, which was also reflected in a drastic increase of pheopigment concentration (12 to 146 μg l⁻¹) between 2 and 3 November. TEP concentrations, which had increased exponentially during the growth phase, dropped off at senescence (Fig. 3c). TEP production in floculator experiments decreased at the onset of growth, remained low during growth and increased at senescence, whereas production of TEP in
dialysis bag experiments remained constant at the beginning of the growth period and increased slightly at senescence (Fig. 3d). No large diatom aggregates formed to terminate the bloom, but during late senescence a few small micro-aggregates consisting of detrital matter and bacteria were observed.

No measurable TEP formed during dialysis experiments conducted with material collected on 18 April (D8 and D9), when the colony density of the growing Phaeocystis spp. bloom reached about 25 colonies ml⁻¹ (chl a: 2.8 μg l⁻¹). This contradicts results of the respective floculator experiment F15, where TEP was formed. Expt F15 was conducted on 25 April, when the colony density had reached about 50 colonies ml⁻¹. Samples for both experiments were, however, taken a week apart and apparently reflected differences (physiological or environmental) that were not observed. TEP was generated within bags of dialysis experiments with senescent Phaeocystis spp. colonies, although no TEP formed in the respective floculator experiment, suggesting that TEP-precursors must have been >0.2 μm in samples of the aging Phaeocystis spp. colonies. TEP may have been formed directly from pieces of disintegrating colonies (Passow & Wassmann 1994, Hong et al. 1997) rather than from dissolved precursors.

**DISCUSSION**

Results clearly indicate that TEP can form abiotically from dissolved precursors small enough to pass through 0.2 μm filters (poly carbonate membrane). The production of TEP from material <0.2 μm was surprising, as experiments on the formation of TEP by bubble scavenging had suggested that TEP-precursors were larger than 0.45 μm (Polycap capsules) (Zhou et al. 1998). Results presented here show not only that TEP can form from material <0.2 μm, but that this happened in the absence of bubbles or particles. Instead, precursors were subjected to laminar shear in floculator experiments and turbulent shear on rolling tables (Jackson 1994). Using shear to generate POC from DOC is a new and, it appears, very efficient method to generate TEP and presumably other organic particles. The highest concentration of TEP generated in floculator experiments from material <0.2 μm was about 2 orders of magnitude larger (>2000 μg xan. eq. l⁻¹) than the highest concentration of TEP generated by bubbling material <1.0 mm (30 μg xan. eq. l⁻¹, normalized to 15.3 l of water bubbled at optimized extraction efficiency: Zhou et al. 1998). However, a direct comparison between both methods is impossible, as scavenging efficiency by bubbling depends on bubble concentration and size distribution which can not be compared directly to shear rate.
TEP consist of surface active, acidic polysaccharides enriched in covalently bound sulfate (Passow & Allerdige 1995b, Zhou et al. 1998). Polysaccharides are large molecules with a MW of several 100s to 1000s of kDa. Microbe-produced polysaccharides appear similar to TEP under the microscope, such as alginate or xanthan (Passow & Allerdige 1995b), for example, are polydisperse with molecular weights of 200 to 2000 and 100 to 2500 kDa, respectively (Buffle et al. 1998). TEP-precursors were thus initially not expected to pass through 50 kDa bags. However, in some dialysis bag experiments TEP was formed from precursors passing through dialysis bags with a nominal pore size of 6 to 8 kDa. Presumably, this large discrepancy cannot be solely explained by artifacts (fouling or aging of membranes) or by the assumption that the nominal MW cut-off of dialysis bags as specified by the producers is too low (see Büsseler et al. 1996, Gustafsson et al. 1996). However, polymers which are freshly released by diatoms exist as long fibrils with a diameter of about 2 to 10 kDa (Leppard 1995, Leppard et al. 1997). It appears that because of their fibrillar nature these polymers can pass through an ultra-filter with a pore size of about 10 kDa, although their MW is 2 orders of magnitude higher (Santschi et al. 1998).

The abiotic formation of large colloids or particles from small fibrillar polymers involves their assembly to nanogels, which then form larger colloids by annealing and aggregation (Chin et al. 1998). Fibrillar polymers may also aggregate with inorganic colloids by bridging, forming loose aggregations (Buffle & Leppard 1995). Thus, it may be assumed that freshly released fibrillar TEP-precursors form larger colloids and eventually TEP as they age. If, because of their fibrillar nature, freshly released TEP-precursors are able to pass through 8 kDa filters, whereas larger colloids formed from these fibrillar precursors cannot, the fraction of precursors able to pass through 8 kDa filters is primarily a function of the age of precursors and of the conditions influencing polymer assembly and aggregation.

Theoretical considerations suggest that in aquatic systems free macromolecules disappear quickly (within hours to days), whereas particles in the size range of TEP are more stable (Buffle & Leppard 1995). As TEP-precursors are highly surface active (Mopper et al. 1995), the formation of larger non-fibrillar colloids and particles should be rapid. Gelation theory suggests that after large colloids and particles are removed by filtration a new equilibrium between polymers and large colloids will be reached within hours to days (Chin et al. 1998). The fact that comparably little TEP is generated by bubbling water with low algal activity (pre-bloom or post-bloom) compared with bubbling water with high algal activity (Zhou et al. 1998) suggests that turn-over times of TEP-precursors in situ are also in the range of hours to days. If freshly released precursors aggregate up the size scale to form larger colloids and ultimately TEP within hours to days, the precursor concentration in the water would be expected to mirror the ecological growth situation of phytoplankton fairly closely, without much lag-time. High concentrations of TEP and their precursors should thus be associated with phytoplankton blooms releasing these substances.

Results of experiments presented here are consistent with such an interpretation. By combining data from flocculator and dialysis bag experiments, changes in the release rate of precursors were estimated and compared to apparent transfer coefficients. Increases in the release rate should occur simultaneously with increases in apparent transfer coefficients, because a larger fraction of freshly released precursors implies a larger fraction of precursors able to pass through bags, which results in a larger apparent transfer coefficient. A comparison between production of TEP in flocculator and dialysis bag experiments allowed estimates of relative changes in the release rate of precursors, because TEP production in flocculator experiments depended on the ambient concentration of precursors <0.2 μM, whereas TEP production in dialysis experiments (in PASW + within bags) depended on the concentration plus the release of precursors.

The steeper slope of the decrease in total TEP production during dialysis experiments compared to flocculator experiments during the early aggregation of the diatom bloom in SBC (Fig. 3b) indicates that release rate of precursors decreased during the aggregation phase. As predicted, the apparent transfer coefficients decreased simultaneously. During the growth phase of the mesocosm bloom, the large relative decrease between the production of TEP in dialysis and flocculator experiments compared to the lag-phase or senescence suggests that during growth the release rate was higher (Fig. 3d), which was also mirrored by the apparent transfer coefficients.

Precursor concentration, as estimated from TEP production in flocculator experiments, also varied over the course of the blooms. Precursor concentrations which were high during the peak of the diatom bloom in the SBC decreased as the bloom began aggregating (Fig. 3b), suggesting rapid turnover of precursors to TEP. Ambient concentration of TEP, however, did not increase simultaneously. The decrease in chl a concentration and the observation that aggregates were sinking imply that an appreciable amount of TEP sedimented, probably as marine snow, since TEP are an essential part of diatom aggregates (Allerdige et al. 1993, Logan et al. 1995, Passow & Allerdige 1995a). During the aggregation phase of this bloom, loss of TEP due to sinking must have balanced the production of TEP from precursors.
Although the release rate of precursors increased at the onset of growth of the mesocosm bloom, and remained high, no accumulation of precursors was observed. The ambient precursor concentration decreased and remained low (Fig. 3d). The simultaneous exponential increase of TEP during growth (Fig. 3c) suggests the rapid formation of TEP from freshly released precursors. Low precursor concentration and a rapid turnover to TEP in the mesocosm experiment may have been facilitated by the mixing of the mesocosm (70 cm propeller stirred at 2.6 rpm, energy dissipation rate = 0.0137 cm$^2$ s$^{-3}$).

Results of low precursor concentration in deep water fit observations that fibrillar colloids, which are very abundant in productive surface water, are much rarer in deep water (Sanitschi et al. 1998). The low concentration of precursors in deeper waters also confirms that TEP-precursors are generated primarily by phytoplankton and that the turnover times of precursors to TEP are rapid. Although, phytoplankton release a wide variety of organic substances (Decho 1990), whose chemical composition varies between species (Allen et al. 1972, Myklestad 1974, Painter 1983) and physiological stage (Myklestad et al. 1972, 1989, Myklestad 1995), TEP-precursors appear more homogeneous. The monomer composition of TEP remained fairly constant, despite changes in algae composition, and all TEP analyzed at date (sampled from 4 different sites and ecological situations) has consisted of strongly sulfated polysaccharides enriched in desoxy-sugars and galactose (Mopper et al. 1995, Zhou et al. 1998). The high concentration of covalently bonded sulfates may explain why TEP themselves are frequently found in deeper waters (Passow & Alledgerde 1994), as preliminary data suggest that hydrolysis rates of polysaccharides high in sulfates are much slower compared to other polysaccharides (Arnosti 1999). TEP and their precursors appear to be a chemically distinct group of polysaccharides whose production and standing stocks are uncoupled from bulk carbohydrates (Mopper et al. 1995).

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