- 1 Production of bioavailable and refractory dissolved organic matter by coastal heterotrophic
- 2 microbial populations
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Abstract

14	Production of dissolved organic matter (DOM) by heterotrophic microbial
15	communities isolated from Loch Creran (Scotland) was studied in time course incubations
16	in which cells were re-suspended in artificial seawater amended with variable proportions
17	of glucose, ammonium and phosphate. The incubation experiments demonstrated that
18	microheterotrophs released part of the substrate as new DOM, with a production efficiency
19	of 11 ± 1 % for DOC, 18 ± 2 % for DON and 17 ± 2 % for DOP. Estimating the impact of
20	this production in Loch Creran, showed that from $3 \pm 1 \%$ (DOC) to $72 \pm 16 \%$ (DOP) of
21	DOM could originate from the heterotrophic microbial community. The produced DOM
22	(PDOM) was both bioavailable (BDOM) and refractory (RDOM). Bioavailability as
23	assessed by the difference between the maximum and the end DOM concentration, was
24	generally higher than found in natural systems, with DOP (73 \pm 15 %, average \pm SD) more
25	bioavailable than DON (70 \pm 15 %), and DON than DOC (34 \pm 13 %).
26	The stoichiometry of PDOM was linked to both nutrient uptake and BDOM
27	ratios. Absorption and fluorescence of DOM increased significantly during the incubation
28	time, indicating that microheterotrophs were also a source of coloured DOM (CDOM) and
29	that they produce both bioavailable protein-like and refractory humic-like fluorophores.
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31	Key words: DOM, coloured DOM, bioavailable DOM, refractory DOM, stoichiometry,
32	microbial heterotrophy

1. Introduction

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River inputs of inorganic nutrients support high primary production rates in coastal marine ecosystems (Lalli & Parsons, 1997), which lead to the release of significant amounts of autochthonous dissolved organic matter (DOM) (Nagata, 2000). Although heterotrophic microbial communities are the major consumers of this DOM (Seitzinger & Sanders, 1997), considerably less attention has been paid to their ability to produce DOM. Some studies have suggested that microheterotrophs could be a significant but hardly quantified source of bioavailable and refractory DOM in marine systems (Ogawa et al., 2001; Kawasaki & Benner, 2007). Release of DOM by phytoplankton has been repeatedly observed in both laboratory and field studies, with exudation rates, stoichiometry and availability being related to nutrient levels (e.g. Obernosterer & Herndl, 1995). Some studies have suggested that bacteria are able to alter their C:N:P biomass ratios depending on the organic substrate ratios (e.g. Tezuka, 1990), while others find constant biomass ratios (Golman et al. 1987). However, the effect of changing C:N:P substrate ratios on the heterotrophic microbial production of DOM is largely unknown. Heterotrophic microbes have been shown to produce chromophoric DOM (CDOM) during mineralization processes (e.g. Rochelle-Newall & Fisher, 2002). CDOM absorbs light in the ultraviolet (UV) region declining to near–zero levels in the red region of the spectrum (Stedmon & Markager 2001). A part of the absorbed light is reemitted at longer wavelengths (FDOM), with two main DOM flourophores being identified: the protein- and humic-like (Coble et al. 1990; 1996). Protein-like fluorescence (FDOMt) has been

suggested as an indicator of freshly produced DOM, while humic–like flourophores (FDOMm) characterize older, more refractory DOM (Coble et al. 1996).

We hypothesised that marine heterotrophic microbial communities could be a quantitative important source of DOM in coastal marine waters, with the stoichiometry of the produced DOM depending on the substrate ratios. This was investigated by isolating coastal heterotrophic microbial populations from Loch Creran, Scotland and growing the community in 0.2 µm filtered artificial seawater, amended with carbon (glucose), inorganic nitrogen (ammonium) and phosphorus (phosphate) as growth media. Over the time course of the experiment dissolved organic carbon (DOC), nitrogen (DON), and phosphorus (DOP) concentrations, as well as DOM fluorescence were monitored as described below.

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2. Material and methods

68 2.1. Incubation experiments

- Samples for the microbial inoculum were obtained during winter and spring (16

 January and 20 March 2007) from a depth of 5 m in the Scottish fjord Loch Creran. These
 sampling dates were chosen to investigate the production of DOM by microbial
 heterotrophic populations collected under different biological and hydrological conditions
 (See Lønborg et al. 2009).
- The artificial seawater was prepared with water treated in a Milli-Q ultraviolet (UV) plus purification unit, which resulted in low carbon and nutrient content. NaCl, KCl,
- 76 NaHCO₃, Na₂SO₄, MgCl₂·6H₂O, CaCl₂·2H₂O, KBr, H₃BO₃, NaF, SrCl₂·6H₂O,
- Na₂SiO₃·6H₂O, FeCl₃·6H₂O, MnSO₄·6H₂O, ZnSO₄·9H₂O and CoSO₄·9H₂O modified from
- Harrison et al. 1980, were added to the Milli–Q water to reach salinity levels equal to the

microbial inoculum, 28.2 (16 January) and 28.3 (20 March) repectively. NaCl, Na₂SO₄, and KCl were combusted (450°C for 4 hours) before use to remove organic carbon. Glucose (C₆H₁₂O₆), ammonium (NH₄Cl) and phosphate (KH₂PO₄) were added as substrate and pH was adjusted to 8.0 with either HCl or NaOH. The microbial inoculum was prepared by gravity filtering the seawater from Loch Creran two times through precombusted GF/F filters. The GF/F filtrate was thereafter inoculated into the media in a 1: 50 dilution. A total of 16 incubations were conducted in duplicate in two litre amber glass bottles and incubated at a constant temperature of 14°C in the dark. The DOC concentration of the artificial seawater, 20 umol L⁻¹, originated from the added salts and the inoculum it was considered refractory and was therefore subtracted from all samples. If we consider that the 20 µmol L⁻¹ is 100% bioavailable and that the microbes would convert 11% (as found in this study) of this DOC into new DOC, this would lead to an underestimation of the microbial produced pool of $\sim 2\mu \text{mol L}^{-1}$ and 1.5 $\mu \text{mol L}^{-1}$ for the refractory pool (considering a bioavailability of 33%). Dissolved inorganic nitrogen (DIN) and phosphorus (DIP) concentrations of the artificial seawater were below the detection limit. Glucose, ammonium and phosphate concentrations added to the medium ranged from 163 to 867 µmol L⁻¹ of C, 21.4 to 157.8 μ mol L⁻¹ of N, and 1.3 to 13.6 μ mol L⁻¹ of P (Table 1). C:N ratios of the added substrate varied between 2 and 17, C:P ratios from 32 to 311 and 5 to 57 for N:P. These ratios were chosen to vary widely around the average C:N:P ratio of bacterial biomass (50:10:1) (Fagerbakke et al. 1996). Since dissolved organic nitrogen (DON) and phosphorus (DOP) concentrations were below the detection limit at the beginning of the experiments, it has been assumed that the DON and DOP produced during the experiment was of microbial origin, and was named as

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produced DOM (PDOM). Sub–samples were collected at days 0, 3, 11, 30, 50 and 100 after the beginning of the experiments to follow: dissolved organic matter absorbance (CDOM) and fluorescence (FDOM), glucose, DOC, DIN, DIP, total dissolved nitrogen (TDN) and phosphorus (TDP), total organic carbon (TOC), total nitrogen (TN) and phosphorus (TP). Samples for the dissolved phase were filtered through prewashed (> 1 litre of sterile filtered Milli–Q water) 0.2 μ m polycarbonate membranes. Samples for CDOM analysis were stored in amber glass bottles at 4°C in the dark. Glucose and FDOM samples (20 mL) were collected in pre–combusted (450°C for 6 hours) glass ampoules and stored frozen (–20°C) in the dark until analyzed. The effect of freezing the FDOM samples was measured by taking samples over time with no major impact found. Samples (10 ml) for TOC, DOC, TN and TDN analysis were collected in pre–combusted (450°C for 6 hours) glass ampoules and preserved by adding 10 μ L 85 % H₂PO₄. Samples for DIN and DIP were stored in acid washed polyethylene bottles and kept frozen (–20°C). TDP and TP samples were collected in glass bottles and frozen (–20°C) until analysis. All glassware used was acid–washed (2 μ mol L⁻¹ HCl) for 24 hours and washed three times with Milli–Q water before use.

2.3. Sample analysis

The absorption of CDOM was measured in four replicates in a Thermo Nicolet Evolution 300 Turn spectrophotometer using Milli–Q water as a blank. Before analysis samples and Milli–Q water were warmed to room temperature. The absorption was measured in a 1 cm quartz cuvette at 375 nm. The absorption coefficients (α_{λ}) were calculated as follows:

$$\alpha_{\lambda} = \frac{2.303 \cdot A_{\lambda}}{L}$$

Where A_{λ} is the optical density and the denominator L is the cell path-length in metres 125 (Stedmon & Markager, 2001). 126 FDOM was measured in duplicate in a Perkin Elmer LS 55 luminescence 127 spectrophotometer. The instrument was equipped with a xenon discharge lamp, using a 1 128 cm quartz fluorescence cell. Milli-Q water was used as a reference, and subtracted the 129 FDOM measurements. Discrete excitation/emission (Ex/Em) pair measurements were 130 performed at peaks T (aromatic amino acids, average Ex/Em, 280/350 nm; FDOMt) and M 131 132 (marine humic substances, average Ex/Em 320/410 nm; FDOMm), according to Nieto-Cid 133 et al. (2006). Following Nieto Cid et al. (2006), fluorescence units were expressed in ppb equivalents of trypthophan (ppb Trp) for FDOMt and ppb equivalents of quinine sulphate 134 (ppb QS) for FDOMm. 135 Glucose was measured in triplicate at days 0, 3, 11 and 30, using the enzyme assay 136 described by Hicks & Carey (1968). Concentrations were determined using a four-point 137 standard curve following the subtraction of a Milli-Q blank. DOC, TOC, TDN and TN 138 were measured in at least triplicate, using a nitrogen-specific Antek 7020 nitric oxide 139 chemiluminescence detector, coupled in series with the carbon–specific infrared gas 140 analyser of a Shimadzu TOC-5000 organic carbon analyser. Three to five replicate 141 injections of 150 µl were performed per sample. Concentrations were determined by 142 143 subtracting a Milli–Q blank and dividing by the slope of a daily standard curve. DON concentrations were obtained by subtracting DIN from TDN (DON = TDN - DIN), with 144 the standard error calculated as $SE_{DON}^2 = SE_{TDN}^2 + SE_{DIN}^2$. DIN (ammonium and 145 146 nitrate+nitrite) was measured in four replicates with a Lachat (quickchem 500) auto analyzer using standard protocols (Hansen & Grasshoff 1983). Triplicate samples were 147 analyzed for DIP by the standard molybdenum blue technique using a 5 cm cuvette 148

- (Koroleff, 1983). TDP and TP were measured in triplicate by oxidation to soluble reactive
- phosphorous with the addition of sulphuric acid and persulphate (Koroleff, 1983),
- following autoclaving at 100°C for 90 min. The standard error for DOP was estimated as:
- $SE_{DOP}^2 = SE_{TDP}^2 + SE_{SRP}^2$. POM concentrations were calculated as the difference between
- TOC and DOC for POC, TN and TDN for PON and TP and TDP for POP. The
- 154 corresponding standard errors where calculated as $SE_{POC}^2 = SE_{TOC}^2 + SE_{DOC}^2$, $SE_{PON}^2 = SE_{DOC}^2$
- $SE_{TN}^2 + SE_{TDN}^2$ and $SE_{POP}^2 = SE_{TP}^2 + SE_{TDP}^2$, respectively.
- Regression model II was used to examine the linear relationship between pairs of variables
- 157 (Sokal & Rohlf 1995). In the cases where the intercept was not significantly different from
- zero, the intercept was set to zero and a new slope was calculated. The confidence level was
- set at 95% with the statistical analysis conducted in Statistica 6.0.

3. Results

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3.1. Production of DON and DOP by heterotrophic microbial populations

TN (Fig. 1b) and TP (Fig. 1c) remained approximately constant throughout the 162 incubations, TOC decreased rapidly during the first phase of all the cultures (initial 30 163 days) and thereafter more slowly (Fig. 1a). Particulate organic matter (POM) reached 164 maximum values at day 10 with values between 12 and 138 μ mol L⁻¹ for POC (Fig. 1a), 165 4.7 to 49.3 µmol L⁻¹ for PON (Fig. 1b) and 0.6 to 4.4 µmol L⁻¹ for POP (Fig. 1c). DIN and 166 DIP concentrations were not significantly different from TN and TP at the beginning of the 167 incubations. The time profile of nutrients concentrations was V-shaped, with a pronounced 168 decrease to a minimum during the first and an increase during the second phase of the 169 cultures (Figs 1b, c). Nitrate+nitrite concentrations were undetectable at the beginning of 170 the incubations, but increased from day 50 onwards in all experiments, concomitantly with 171

the corresponding decrease of ammonium (Fig. 1d). The total uptake of DIN (UDIN) and 172 DIP (UDIP) (Table 1) was calculated as UDIN = DIN₀- LDIN and UDIP = DIP₀ - LDIP, 173 where DIN₀ and DIP₀ are the initial nutrient concentrations and LDIN, LDIP are the 174 minimum DIN and DIP concentrations obtained at the end of the first phase of the cultures. 175 DOM maximum was found at day 30 of the incubations (Fig. 2), associated with complete 176 depletion of the added glucose (data not shown). The produced DOM (PDOM) reached 177 concentrations of 38 to 81 µmol L⁻¹ for PDOC, 2.2 to 9.1 µmol L⁻¹ for PDON, and 0.24 to 178 0.78 µmol L⁻¹ for PDOP (Table 1). Calculating the bioavailability (BDOM) as the 179 180 difference between PDOM and the refractory DOM (RDOM) at the end of the incubations, it resulted that 33 \pm 12 % (average \pm SD) of PDOC, 70 \pm 16 % of PDON and 79 \pm 14 % of 181 PDOP were bioavailable. The PDOM C:N ratios varied between 6 and 16, C:P ratios from 182 20 to 224 and 5 to 21 for N:P. Resulting C:N:P ratios varied between 20: 5: 1 and 225: 14: 183 1 (average 103: 11: 1). The stoichiometry of PDOM was linearly related with the C: N: P 184 ratios of UDOC, UDIN and UDIP (Fig 3a, c, e). 185

3.3. Production of chromophoric DOM

Table 3a shows how a375 increased from being undetectable at the beginning of the incubations reaching an average value of $0.47 \pm 0.26 \,\mathrm{m}^{-1}$ at the end. In parallel to the production of CDOM, FDOM also increased during the incubations above background levels in all cases, with a mean (\pm SE) of 12.9 ± 5.6 ppb Trp for FDOMt and 6.4 ± 3.8 ppb QS, for FDOMm (Table 3b, c). The amounts of FDOMt and FDOMm produced increased with substrate uptake and produced DOM (Table 4).

4. Discussion

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Single–substrate approaches as used in this study are an obvious simplification of reality, and will not fully represent the natural microbial growth substrate. These experiments should therefore be seen as a first trial to quantify heterotrophic microbial production of DOM.

Two contrasting phases have been identified during all the incubations (Fig. 1): i) an initial net-anabolic phase, when the added glucose was still detectable, as characterised by the net consumption of the glucose, DIN and DIP and a net production of DOM; and ii) a final net-catabolic phase, where glucose was undetectable, characterised by the net production of DIN and DIP and the net consumption of the DOM produced during the anabolic phase of the cultures. Goldman and Dennet (1991) also observed in their cultures that ammonium was taken up as long as a readily assimilable carbon source such as glucose was present, whilst nutrient regeneration was evident only after glucose was completely utilized.

3.1 Production of DON and DOP by heterotrophic microbial populations

POM and inorganic nutrients concentrations showed opposite patterns, POM increased during the net-anabolic phase of the cultures, reaching maximum values at day 10 (Fig. 1a-c) and subsequently decreased. The increase in nitrate+nitrite following the decrease in ammonium concentrations (Fig. 1d), suggested, that ammonium was consumed i) as a substrate during the net-anabolic phase of the cultures (Kirchman, 1994); and ii) as an energy source during the net-catabolic phase of the cultures on basis of the complete oxidation of ammonium to nitrate, i.e. nitrification (Wada & Hattori, 1991).

In the net-anabolic phase, the use of glucose and inorganic nutrients by the cultured community of microheterotrophs was accompanied by increases in DON and DOP

concentrations (Fig 1a, c). We hypothesise that bacteria take up the dissolved substrate, converting it into organic matter which is subsequently released as DOM by direct extra cellular release and/or by protist grazing and viral lysis (Caron et al., 1985; Riemann & Middelboe, 2002; Kawasaki & Benner, 2007). A DOM maximum was found at day 30 of the incubations (Fig. 2), associated with complete depletion of the added glucose (data not shown). The efficiency of DOM production was found using the slope of the linear regressions between UDOC (glucose) and PDOC ($11 \pm 1 \%$), UDIN and PDON ($18 \pm 3 \%$), and UDIP and PDOP ($17 \pm 3\%$) (Table 2), which was comparable findings in similar experiments (Kawasaki & Benner, 2007). In the net-catabolic phase of the incubations, DOM concentrations decreased until the end of the experiments (Fig. 1; Fig. 2). The calculated bioavailability (BDOM) suggest that PDOC and PDON were more bioavailable than found in natural marine waters (9–30 %, DOC; 30-40 %, DON; Hopkinson et al., 2002; Lønborg & Søndergaard, 2008), while PDOP bioavailability was comparable with natural systems (70–80%; Nausch & Nausch, 2006). Thereby PDOP was more bioavailable than PDON, and PDON more than PDOC consistently with findings in natural marine systems (Hopkinson et al., 2002). The slopes of the linear regressions between BDOM and PDOM indicate that most of the variations in PDOM were due to the bioavailable fraction (Table 2). The significant origin intercepts of those regressions together with the significant DOM end levels (Table 1, Fig. 2) showed that part of the PDOM persisted at the end of the incubations, demonstrating that the microbial communities also produce refractory DOM (RDOM) in agreement with previous studies (Ogawa et al., 2001; Kawasaki & Benner, 2007). Estimating the environmental impact of the microbial DOM production, we can use published DOC and nutrient data from Loch Creran (Lønborg et al. 2009), together with average efficiency of DOC (11 \pm 1

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- %), DON (18 \pm 3 %), and DOP (17 \pm 3 %) production found in this study. These 241 calculation showed that the heterotrophic microbial community on average could produce 5 242 \pm 2 μ M DOC, 0.7 \pm 0.5 μ M DON and 0.08 \pm 0.04 μ M DOP, corresponding to 3 \pm 1 %, 8 \pm 243 7 % and 72 \pm 16 % of the measured DOC, DON and DOP in Loch Creran. 244 The average PDOM C:N:P stoichiometry (103: 11: 1), was generally richer in 245 carbon than typically found for bacterial biomass (50: 10: 1; Goldman et al., 1987; 246 Fagerbakke et al., 1996), and more N and P rich than for both algae produced (170: 6.5: 1; 247 248 Conan et al., 2007) and marine bulk DOM (300: 22: 1; Benner, 2002). The stoichiometry of 249 PDOM was linearly related with the C: N: P ratios of UDOC, UDIN and UDIP (Fig 3a, c, e). The slopes of the linear regressions show that PDOM had C:N and C:P ratios which 250 were 63 ± 13 % and 68 ± 14 % lower than the substrate, while the N:P ratios (105 ± 15 %) 251 were similar, indicating that the stoichiometry of PDOM is linked to the substrate ratios. 252 The fact that the C:N and C:P ratios of PDOM were 30–40 % lower than the corresponding 253 substrate ratios can be attributed to the bacterial respiration of glucose, which previously 254 has been found to represent 30–60 % of the total consumption (Bianchi et al., 1998). The 255 stoichiometry of BDOM was similar to ranges reported for bacterial C: N: P ratios 256 (Goldman et al., 1987; Fagerbakke et al., 1996). The slopes of the correlation between 257 PDOM and BDOM stoichiometries (Fig. 3b, d, f) indicate that BDOM had C:N and C:P 258
- 261 *4.2. Production of chromophoric DOM*

were not significantly different from PDOM.

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Recent studies suggest that marine bacteria produce CDOM and FDOM in coastal ecosystems depending on the chemical composition of the substrate (e.g. Rochelle–Newall

ratios that were 53 ± 19 % and 44 ± 8 % lower respectively, while N:P ratios (92 ± 25 %)

& Fisher, 2002; Yamashita & Tanoue, 2008). Table 3a shows how a375 increased both during the net-anabolic and net-catabolic phase from being undetectable at the beginning of the incubations reaching higher values (average - $0.47 \pm 0.26 \text{ m}^{-1}$) at the end. This increase indicates microbial production of coloured high molecular weight aromatic DOM (Pages & Gadel, 1990). In parallel to the production of CDOM, FDOM also increased during both the net-anabolic and net-catabolic phase above background levels in all cases (Table 3b, c), in agreement with previous studies (Kramer & Herndl, 2004; Yamashita & Tanoue, 2004). The amounts of FDOMt and FDOMm produced increased with substrate uptake and produced DOM (Table 4). Although protein-like fluorophores have been suggested to represent freshly produced DOM (Yamashita & Tanoue, 2004), our study found links between FDOMt and DOM bioavailability, but also the production of refractory FDOMt. It firstly indicates that protein-like fluorophores not only provide a measure of labile material; and secondly suggests that biological processes could be a source of refractory FDOMt. FDOMm was related linearly with both BDOC and RDOC, confirming it as a product of microbial degradation processes (Yamashita & Tanoue, 2004; 2008; Nieto-Cid et al., 2006).

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Figure legends 379 Figure. 1. Examples of time evolution (Exp. 1 shown in table 1), of a) total (TOC) 380 particulate (POC) and dissolved organic carbon (DOC), b) total nitrogen (TN), particulate 381 organic nitrogen (PON), dissolved inorganic (DIN) and organic (DON) nitrogen; c) total 382 phosphorus (TP), particulate organic phosphorus (POP), dissolved inorganic (DIP) and 383 organic (DOP) phosphorus; d) nitrate/nitrite (NO₃-/NO₂-) and ammonium (NH₄+). Error 384 bars represent standard errors. 385 386 Figure. 2. Production of dissolved organic nitrogen (DON) and phosphorus (DOP) in the 387 16 experiments, a),b) started in January 2007 and c),d) in March 2007. The experimental 388 numbers 1 to 16 represents the same experiment numbers as shown in table 1. 389 390 Figure. 3. Significant regressions between substrate uptake (UDOC, UDIN, UDIP) and 391 produced DOM (PDOM) stoichiometery and the X–Y plots of linear relation between 392 PDOM and bioavailable DOM (BDOM) stoichiometery. With a) UDOC:UDIN with 393 PDOC:PDON, b) PDOC:PDON versus BDOC:BDON, c) UDOC:UDIP versus 394

PDOC:PDOP, d) PDOC:PDOP with BDOC:BDOP, e) UDIN:UDIP versus PDON:PDOP

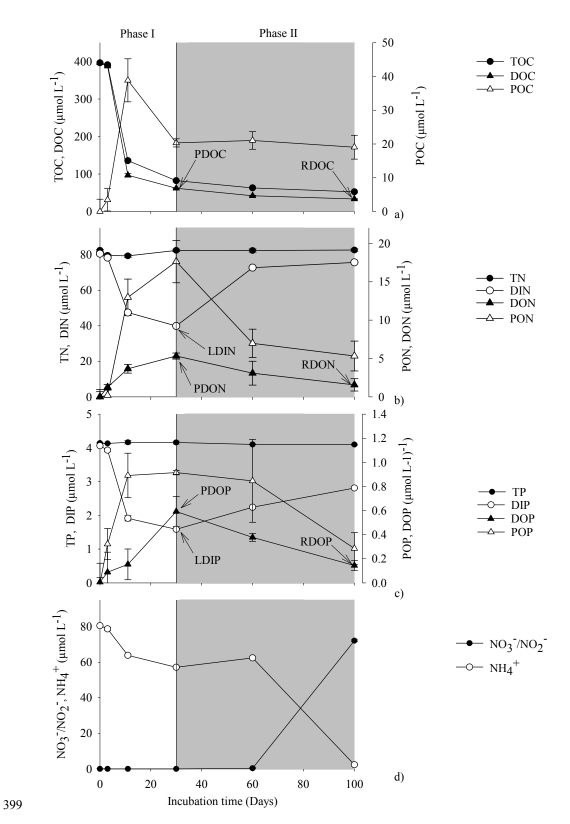
and f) PDON:PDOP with BDON:BDOP. Slope, incept, and standard error are values found

by Model II regression. R^2 = coefficient of determination, p = significant levels.

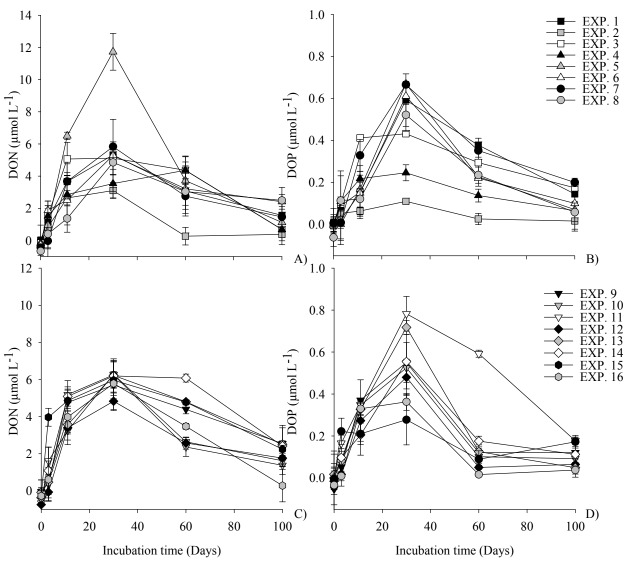
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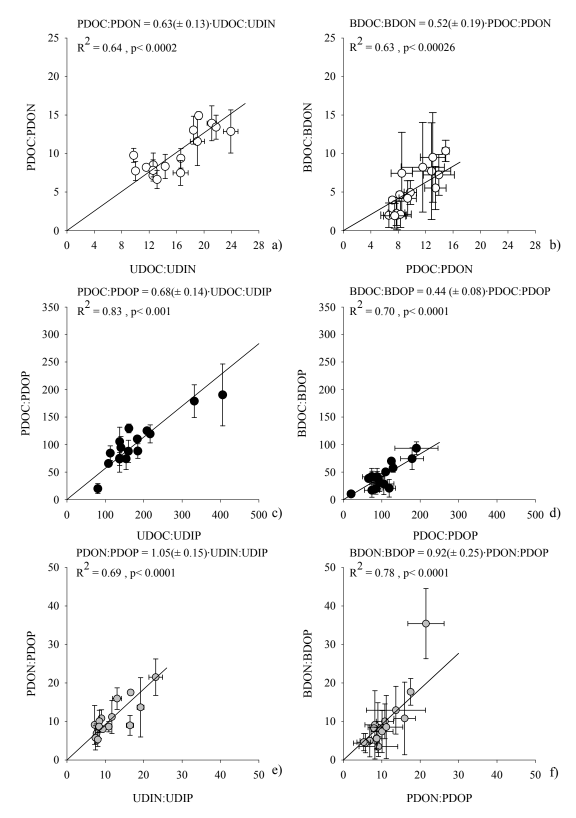
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400 Fig.1.



402 Fig.2.



404 Fig.3.

Table 1. Concentrations of DOC (UDOC), DIN (UDIN) and DIP (UDIP) utilized during the experiments, DOC (PDOC), DON (PDON), and DOP (PDOP) produced during the experiments, and residual DOC (RDOC), DON (RDON) and DOP (RDOP) at the end of the experiments. Experiments 1 to 8 were commenced in January and experiments 9 to 16 in March 2007. Values are means of three replicates \pm standard error given in μ mol L⁻¹.

Exp.	DOC	DIN	DIP	UDOC	UDIN	UDIP	PDOC	PDON	PDOP	RDOC	RDON	RDOP
1	396 ± 1	80.6 ± 0.6	4.06 ± 0.02	396 ± 1	40.7 ± 0.9	2.48 ± 0.10	52 ± 1	5.3 ± 1.4	0.59 ± 0.13	33 ± 1	1.6 ± 0.8	0.14 ± 0.04
2	440 ± 6	80.2 ± 0.4	1.42 ± 0.02	440 ± 6	20.8 ± 0.4	1.08 ± 0.08	51 ± 1	5.7 ± 0.8	0.11 ± 0.04	31 ± 1	0.4 ± 0.1	0.02 ± 0.05
3	425 ± 1	25.1 ± 0.1	4.18 ± 0.03	425 ± 1	19.5 ± 0.3	2.32 ± 0.01	47 ± 2	5.1 ± 1.0	0.43 ± 0.04	35 ± 1	1.2 ± 0.2	0.17 ± 0.02
4	163 ± 1	77.6 ± 0.3	4.45 ± 0.02	163 ± 1	8.6 ± 0.4	1.19 ± 0.02	26 ± 2	2.2 ± 0.9	0.25 ± 0.04	21 ± 1	1.6 ± 0.4	0.07 ± 0.05
5	789 ± 3	93.4 ± 0.2	4.73 ± 0.03	789 ± 3	62.8 ± 0.3	3.78 ± 0.08	84 ± 1	11.7 ± 0.8	0.67 ± 0.05	42 ± 1	1.1 ± 0.5	0.07 ± 0.02
6	411 ± 6	157.5 ± 0.4	4.47 ± 0.04	411 ± 6	32.7 ± 0.6	3.00 ± 0.04	45 ± 4	5.3 ± 0.9	0.61 ± 0.06	24 ± 1	2.4 ± 0.5	0.10 ± 0.01
7	396 ± 2	79.2 ± 0.1	8.11 ± 0.06	396 ± 2	34.2 ± 0.3	3.66 ± 0.08	48 ± 1	4.3 ± 0.8	0.67 ± 0.08	28 ± 1	1.5 ± 0.6	0.20 ± 0.02
8	433 ± 1	88.1 ± 0.4	4.59 ± 0.04	433 ± 1	18.1 ± 0.8	2.35 ± 0.10	42 ± 1	4.8 ± 0.8	0.52 ± 0.09	28 ± 1	1.3 ± 0.8	0.06 ± 0.04
9	543 ± 2	84.0 ± 0.6	7.06 ± 0.07	543 ± 2	29.4 ± 0.7	3.95 ± 0.15	74 ± 1	5.7 ± 0.8	0.55 ± 0.10	38 ± 1	1.9 ± 0.9	0.12 ± 0.07
10	440 ± 7	82.1 ± 0.3	4.09 ± 0.02	440 ± 7	35.0 ± 0.8	3.90 ± 0.06	49 ± 3	6.3 ± 0.9	0.52 ± 0.07	39 ± 1	1.4 ± 0.5	0.09 ± 0.03
11	411 ± 9	51.0 ± 0.4	7.69 ± 0.06	411 ± 9	41.1 ± 0.8	5.10 ± 0.07	48 ± 2	6.3 ± 0.8	0.78 ± 0.08	38 ± 1	1.6 ± 0.3	0.18 ± 0.04
12	409 ± 3	88.5 ± 0.2	7.57 ± 0.07	409 ± 3	24.6 ± 0.6	2.91 ± 0.15	45 ± 1	4.8 ± 1.0	0.48 ± 0.09	32 ± 1	1.8 ± 0.4	0.06 ± 0.02
13	867 ± 2	84.1 ± 0.3	7.43 ± 0.10	867 ± 2	45.1 ± 0.6	5.37 ± 0.10	93 ± 2	6.2 ± 0.8	0.72 ± 0.06	54 ± 3	2.5 ± 0.4	0.05 ± 0.03
14	449 ± 2	164.7 ± 0.4	7.28 ± 0.05	449 ± 2	34.1 ± 1.0	2.91 ± 0.09	41 ± 2	6.2 ± 0.9	0.55 ± 0.13	34 ± 1	2.4 ± 1.1	0.11 ± 0.04
15	442 ± 1	74.8 ± 0.4	13.64 ± 0.05	442 ± 1	30.9 ± 0.5	1.33 ± 0.09	50 ± 3	6.0 ± 1.0	0.28 ± 0.12	42 ± 1	2.2 ± 0.5	0.17 ± 0.03
16	448 ± 3	96.5 ± 0.9	7.57 ± 0.03	448 ± 3	27.0 ± 1.6	2.06 ± 0.08	43 ± 1	5.8 ± 1.1	0.36 ± 0.08	27 ± 2	2.3 ± 0.9	0.04 ± 0.02

Table 2. Significant linear regressions between used DOC (UDOC) and produced DOC (PDOC);
bioavailable (BDOC) and PDOC; used dissolved inorganic nitrogen (UDIN) and produced DON (PDON);
bioavailable DON (BDON) and PDON); used phosphate (UDIP) and produced DOP (PDOP);, and
bioavailable DOP (BDOP) and PDOP. Slope, intercept, and standard error (SE) are values found by Model II
regression. R² = coefficient of determination, p = level of significance, n.s. – not significant.

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X	Y	Slope (±SE)	Intercept (±SE)	R^2	р
UDOC	PDOC	0.11 ± 0.01	n.s	0.90	< 0.00001
BDOC	PDOC	1.4 ± 0.2	27 ± 3	0.80	< 0.00001
UDIN	PDON	0.18 ± 0.02	n.s	0.85	< 0.0001
BDON	PDON	1.0 ± 0.2	1.7 ± 0.3	0.92	< 0.001
UDIP	PDOP	0.17 ± 0.02	n.s	0.79	< 0.00001
BDOP	PDON	1.1 ± 0.1	0.08 ± 0.03	0.82	< 0.00001

Table 3. a) Absorption coefficients (a375) at day 0 (a375₀), 30 (a375₃₀) and 100 (a375₁₀₀). b) Initial (FDOMt₀), produced (PFDOMt₀) and refractory (RFDOMt₀) protein–like fluorescence; c) Initial (FDOMm₀), produced (PFDOMm) and refractory (RFDOMm) marine humic–like fluorescence; and Experiments 1 to 8 were commenced in January and experiments 9 to 16 in March 2007, corresponding to the experiments described in table 1. Values are means of two replicates \pm standard error.

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Experiment	$a375_0 (m^{-1})$	$a375_{30} (m^{-1})$	$a375_{100} (m^{-1})$	_ a
1	0.05 ± 0.05	0.44 ± 0.08	0.67 ± 0.02	
2	0.01 ± 0.02	0.27 ± 0.03	0.61 ± 0.09	
3	0.06 ± 0.06	0.25 ± 0.10	0.92 ± 0.12	
4	0.07 ± 0.08	0.32 ± 0.08	0.60 ± 0.09	
5	0.01 ± 0.01	0.36 ± 0.03	0.52 ± 0.10	
6	0.07 ± 0.08	0.50 ± 0.05	0.84 ± 0.17	
7	0.01 ± 0.04	0.38 ± 0.10	0.45 ± 0.18	
8	0.02 ± 0.02	0.13 ± 0.07	0.64 ± 0.09	
9	0.07 ± 0.08	0.31 ± 0.08	0.41 ± 0.04	
10	0.08 ± 0.09	0.42 ± 0.10	0.69 ± 0.07	
11	0.08 ± 0.09	0.24 ± 0.05	0.34 ± 0.07	
12	0.03 ± 0.04	0.24 ± 0.04	0.38 ± 0.05	
13	0.02 ± 0.02	0.32 ± 0.08	0.41 ± 0.16	
14	0.04 ± 0.04	0.39 ± 0.04	0.53 ± 0.03	
15	0.01 ± 0.01	0.34 ± 0.05	0.34 ± 0.09	
16	0.01 ± 0.01	0.52 ± 0.03	1.33 ± 0.09	

Experiment FDOMt₀ (ppb Trp) PFDOMt (ppb Trp) RFDOMt (ppb Trp) b) 1 3.5 ± 0.6 17.7 ± 0.7 21.1 ± 0.1 2 10.0 ± 0.9 3.2 ± 0.6 13.2 ± 0.3 3 2.6 ± 0.4 13.0 ± 1.0 15.6 ± 0.7 4 2.3 ± 0.2 6.1 ± 0.7 8.4 ± 0.5 5 2.5 ± 0.2 19.9 ± 0.7 22.4 ± 0.5 6 3.9 ± 0.1 17.7 ± 0.6 21.5 ± 0.5 7 3.0 ± 0.3 21.2 ± 0.4 24.2 ± 0.1 12.2 ± 0.2 8 1.3 ± 0.5 10.9 ± 0.7 9 2.3 ± 0.6 16.3 ± 0.8 18.6 ± 0.2 10 3.2 ± 0.3 7.9 ± 0.9 11.1 ± 0.5 11 2.6 ± 0.3 10.6 ± 0.4 13.2 ± 0.2 12 2.3 ± 0.6 7.9 ± 1.1 10.2 ± 0.5 13 4.5 ± 0.9 15.2 ± 1.0 19.7 ± 0.1

14	2.9 ± 0.1	11.4 ± 0.3	14.3 ± 0.2
15	1.0 ± 0.2	14.5 ± 0.6	15.5 ± 0.4
16	2.3 ± 0.2	9.4 ± 0.6	11.7 ± 0.4

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_	Experiment	FDOMm ₀ (ppb Trp)	PFDOMm (ppb Trp)	RFDOMm (ppb Trp)	c)
	1	2.58 ± 0.3	9.0 ± 0.5	11.7 ± 0.1	
	2	2.5 ± 0.5	10.6 ± 0.8	13.1 ± 0.3	
	3	2.0 ± 0.6	8.9 ± 0.7	10.8 ± 0.1	
	4	2.5 ± 0.3	4.7 ± 0.7	7.2 ± 0.5	
	5	2.9 ± 0.1	14.0 ± 0.2	16.8 ± 0.2	
	6	2.2 ± 0.3	5.0 ± 0.9	7.2 ± 0.6	
	7	1.9 ± 0.4	5.4 ± 0.5	7.3 ± 0.2	
	8	1.8 ± 0.2	4.8 ± 0.2	6.6 ± 0.2	
	9	2.8 ± 0.3	7.8 ± 1.0	10.5 ± 0.7	
	10	2.3 ± 0.5	3.7 ± 0.6	6.0 ± 0.1	
	11	1.8 ± 0.6	4.0 ± 0.7	5.8 ± 0.2	
	12	2.4 ± 0.3	3.2 ± 1.1	5.6 ± 0.8	
	13	2.6 ± 0.1	13.1 ± 0.2	15.7 ± 0.1	

 2.8 ± 0.5

 5.2 ± 0.5

 5.1 ± 0.3

 5.0 ± 0.3

 7.2 ± 0.1

 6.9 ± 0.1

 2.2 ± 0.3

 2.0 ± 0.4

 1.8 ± 0.2

- Table 4. Matrix of the correlation coefficient (R^2) of the significant (p < 0.05) linear regressions
- between DOM and the produced protein– (PFDOMt) and marine humic–like (PFDOMm)
- 428 flourophores. n.s. not significant.

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X–Y	$FDOM_T$	$FDOM_M$
UDIN	0.60	0.33
UDOC	0.53	0.64
RDOC	n.s.	0.45
BDOC	0.77	0.53
BDON	0.30	0.38
BDOP	0.44	n.s.