Biogeosciences, 13, 989–1007, 2016 www.biogeosciences.net/13/989/2016/ doi:10.5194/bg-13-989-2016 © Author(s) 2016. CC Attribution 3.0 License.





The organic sea-surface microlayer in the upwelling region off the coast of Peru and potential implications for air–sea exchange processes

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Received: 29 May 2015 – Published in Biogeosciences Discuss.: 9 July 2015 Revised: 23 December 2015 – Accepted: 27 January 2016 – Published: 22 February 2016

Abstract. The sea-surface microlayer (SML) is at the uppermost surface of the ocean, linking the hydrosphere with the atmosphere. The presence and enrichment of organic compounds in the SML have been suggested to influence airsea gas exchange processes as well as the emission of primary organic aerosols. Here, we report on organic matter components collected from an approximately 50 µm thick SML and from the underlying water (ULW), $\sim 20 \,\mathrm{cm}$ below the SML, in December 2012 during the SOPRAN ME-TEOR 91 cruise to the highly productive, coastal upwelling regime off the coast of Peru. Samples were collected at 37 stations including coastal upwelling sites and off-shore stations with less organic matter and were analyzed for total and dissolved high molecular weight (>1 kDa) combined carbohydrates (TCCHO, DCCHO), free amino acids (FAA), total and dissolved hydrolyzable amino acids (THAA, DHAA), transparent exopolymer particles (TEP), Coomassie stainable particles (CSPs), total and dissolved organic carbon (TOC, DOC), total and dissolved nitrogen (TN, TDN), as well as bacterial and phytoplankton abundance. Our results showed a close coupling between organic matter concentrations in the water column and in the SML for almost all components except for FAA and DHAA that showed highest enrichment in the SML on average. Accumulation of gel particles (i.e., TEP and CSP) in the SML differed spatially. While CSP abundance in the SML was not related to wind speed, TEP abundance decreased with wind speed, leading to a depletion of TEP in the SML at about 5 m s^{-1} . Our study provides insight to the physical and biological control of organic matter enrichment in the SML, and discusses the potential role of organic matter in the SML for air-sea exchange processes.

1 Introduction

The sea-surface microlayer (SML) is the uppermost layer of the water column and the interface between the ocean and the atmosphere. The accumulation of organic matter, distinct physical and chemical properties and a specific organismal community (neuston) distinguish the SML as a unique biogeochemical and ecological system. It has been suggested that the SML has a gel-like nature (Cunliffe and Murrell, 2009; Sieburth, 1983) of varying thickness (20-150 µm, Cunliffe et al., 2013) with dissolved polymeric carbohydrates and amino acids present as well as gel particles, such as transparent exopolymer particles (TEP) of polysaccharidic composition, and Coomassie stainable particles (CSPs) of proteinaceous composition. These gelatinous compounds originate from high molecular weight polymers that are released form phytoplankton and bacterial cells by exudation and cell break up (Chin et al., 1998; Engel et al., 2004; Verdugo et al., 2004). Polysaccharide-rich gels, like TEP, were attributed mainly to phytoplankton exudation (Passow, 2002), while the production of protein-containing gels, such as CSPs, has been related to cell lysis and decomposition, as well as to the absorption of proteins onto non-proteinaceous particles (Long and Azam, 1996). Gels are transported to the SML by rising bubbles (Azetsu-Scott and Passow, 2004; Zhou et al., 1998) or are produced from dissolved precursors directly at the air-sea interface during surface wave action (Wurl et al., 2011). Gel particles can promote microbial biofilm formation (Bar-Zeev et al., 2012) and mediate vertical organic matter transport, either to the atmosphere (Leck and Bigg, 2005; Orellana et al., 2011) or to the deep ocean (Passow, 2002).

Accumulation of organic matter in the SML may be tightly coupled to phytoplankton abundance in the water column (Bigg et al., 2004; Galgani et al., 2014; Gao et al., 2012; Matrai et al., 2008). Thus, organic matter accumulation and composition in the SML may also reflect the sensitivity of marine microorganisms in the surface ocean to environmental changes, which was shown during previous mesocosms studies (Engel et al., 2013; Riebesell et al., 2009; Schulz et al., 2013). Distinct from the SML and on top of it lies the nanolayer, a monomolecular film, which, like the SML, shows seasonality features with carbohydrate-rich polymeric material being most abundant during the summer months and possibly related to a combination of primary production (phytoplankton abundance) and photochemical and/or microbial reworking of organic matter (Laß et al., 2013).

In our study we focused on the upper micrometers of the water-air interface that we operationally define as SML, whose compositional changes and accumulation of organic matter may influence two air-sea interface processes necessary to understand oceanic feedbacks on the atmosphere: sea-spray aerosol (SSA) emission and air-sea gas exchange (Cunliffe et al., 2013). During biologically productive periods, a high amount of SSA with a predominant organic composition is emitted from the ocean's surface (O'Dowd et al., 2004). These compounds primarily reveal a polysaccharidic, gel-like composition, suggesting that the abundance and size of dissolved polysaccharides and marine gels in the sea surface may influence the organic fraction of SSA (Orellana et al., 2011; Russell et al., 2010). It has also been shown that the presence of biogenic surface active substances (surfactants) in the SML leads to capillary wave damping, alters the molecular diffusion of gases (Frew et al., 1990; Liss and Duce, 2005) and thereby affects gas exchange rates particularly at lower wind speed (Jähne and Haußecker, 1998). In this respect, the understanding of sources, composition and fate of biological components in the SML becomes of particular relevance for environments, where biological productivity is high like in coastal upwelling regimes.

Off the coast of Peru, the coastal upwelling region extends between approximately 4 and 40° S. In this area, upwelling processes are sustained by winds throughout the year but feature high inter-annual variability induced by the El Niño– Southern Oscillation (ENSO) cycle (Tarazona and Arntz, 2001). Eastern boundary upwelling systems (EBUSs) like the system off the coast of Peru are characterized by high biological productivity supported by deep upwelling of nutrients and often associated with subsurface oxygen minimum zones (OMZs). The supply of oxygen to the OMZ is largely controlled by physical (i.e., diffusive and advective) mechanisms, whereas biological processes (i.e., respiration of organic matter) provide sinks (Lachkar and Gruber, 2011).

OMZs are significant source regions for major climaterelevant gases such as carbon dioxide, methane, hydrogen sulfide and nitrous oxide (Paulmier et al., 2008, 2011). Processes affecting gas exchange in these regions need to be understood in order to accurately estimate trace gas fluxes from the ocean to the atmosphere and consequences on climate. In 2008, the VAMOS Ocean-Cloud-Atmosphere-Land Study Regional Experiment (VOCALS-REx) and the Chilean Upwelling Experiment (VOCALS-CUpEx) conducted between Southern Peru and Northern Chile focused on the link between aerosols, clouds and precipitation as well as on physical and chemical couplings between the upper ocean and the lower atmosphere (Garreaud et al., 2011; Wood et al., 2011). During the SOPRAN cruise METEOR91 (M91), we studied organic matter components at the very sea surface since properties of the SML may represent a major uncertainty for gas, heat and aerosol fluxes in this specific region and in other oceanic environments. During our cruise, organic matter concentration and composition of the SML and the underlying seawater were studied on 37 different stations, providing the first SML data set for the upwelling system off the coast of Peru, including the first data set on gel particles in EBUSs so far.

2 Material and methods

2.1 Field information and sampling

The R/V *METEOR* cruise M91 studied the upwelling region off the coast of Peru (Bange, 2013). Samples were collected between 4.59° S and 82.0° W, and 15.4° S and 77.5° W from 3 to 23 December in 2012. The overall goal of M91 was to conduct an integrated biogeochemical study on the upwelling region off the coast of Peru in order to assess the importance of OMZs for the sea-air exchange of various climate-relevant trace gases and for tropospheric chemistry. Salinity and temperature were measured with a CTD at each station. Global and UV radiation and wind speed data were retrieved from the DShip database for the time of sampling based on the sensors installed on board.

On 37 different stations between 5 and 16° S off the Peruvian coast (Fig. 1), a total of 39 SML samples was collected from a rubber boat using a glass plate sampler according to the original approach described by Harvey and Burzell (1972). Different methods have been developed to sample and investigate the SML. These methods do not only differ in terms of application but also with respect to the thickness of the SML sampled as well as to selective removal of certain components. Several studies evaluated these methods against each other. A recent summary can be found in the "Guide to best practices to study the ocean's surface" (Cunliffe and Wurl, 2014). During this study, we applied the glass plate technique because it allows for sampling of a relatively large volume needed to analyze different organic components while keeping the simultaneous sampling of ULW minimal. Two stations were sampled twice in a time frame of 24 h (stations 12_1 and 12_3, 16_2 and 16_3). Our glass plate with the dimensions of 500 mm (length) $\times 250 \text{ mm}$

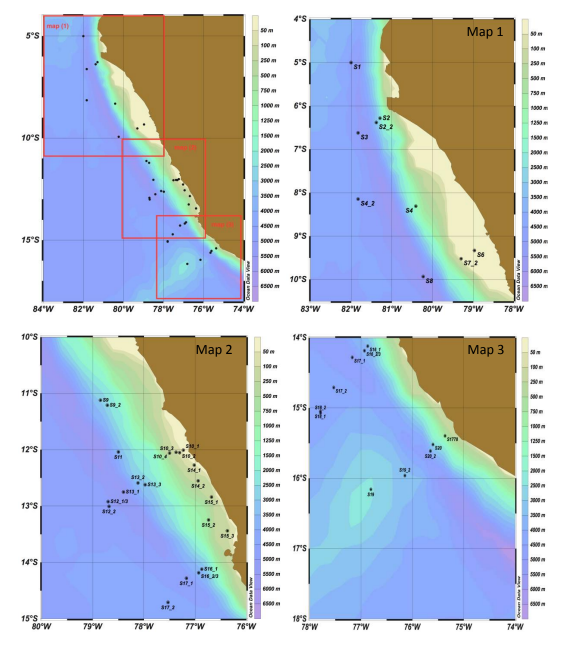


Figure 1. Maps of stations where sampling for the sea-surface microlayer (SML) and underlying seawater (ULW) was conducted during the SOPRAN Meteor 91 cruise along the coastal upwelling area off the coast of Peru in 2012.

(width) × 5 mm (thickness) was made of borosilicate glass and had an effective sampling surface area of 2000 cm² (considering both sides). For each sample, the glass plate was inserted into the water perpendicular to the surface and withdrawn slowly at a rate of approximately 20 cm s^{-1} . The sample, retained on the glass because of surface tension, was removed with the help of a Teflon wiper. Samples were collected as far upwind of the ship as possible and away from the path taken by the ship to avoid contamination. For each sample the glass plate was dipped and wiped about 20 times. The exact number of dips and the total volume collected were recorded. Samples were collected into acid-cleaned (HCl, 10%) and Milli-Q-washed glass bottles, and the first milliliters were used to rinse the bottles and then discarded. Prior to each sampling, both the glass plate and wiper were washed with HCl (10%) and intensively rinsed with Milli-Q water. At the sampling site, both instruments were copiously rinsed with seawater in order to minimize any possible contamination with alien material while handling or transporting the devices.

The apparent thickness (d) of the layer sampled with the glass plate was determined as follows:

$$d = V/(A \times n), \tag{1}$$

where V is the SML volume collected (i.e., 60-140 mL) A is the sampling area of the glass plate ($A = 2000 \text{ cm}^2$) and n is the number of dips (Cunliffe and Wurl, 2014). We will use d (µm) as an operational estimate for the thickness of the SML.

At the same stations, after sampling the SML, about 500 mL samples were collected from the underlying seawater (ULW) at ~ 20 cm depth by holding an acid-cleaned (HCl 10%) and Milli-Q-rinsed borosilicate glass bottle. The bottle was open and closed underwater to avoid simultaneous sampling of SML water. For safety reasons, sampling for the SML from a rubber boat could be made only during daylight hours.

2.2 Chemical and biological analyses

2.2.1 Total organic carbon (TOC) and dissolved organic carbon (DOC)

Samples for TOC and DOC (20 mL) were collected in combusted glass ampoules, DOC after filtration through combusted GF/F filters (8 h, 500 °C). Samples were acidified with 80 µL of 85 % phosphoric acid, heat sealed immediately, and stored at 4 °C in the dark until analysis. DOC and TOC samples were analyzed by applying the high-temperature catalytic oxidation method (TOC-VCSH, Shimadzu) modified from Sugimura and Suzuki (1988). The instrument was calibrated every 8-10 days by measuring standard solutions of 0, 500, 1000, 1500, 2500 and 5000 μ g C L⁻¹, prepared from a potassium hydrogen phthalate standard (Merck 109017). Every measurement day, ultrapure (MilliQ) water was used to determine the instrument blank, which was accepted for values $< 1 \,\mu$ mol C L⁻¹. TOC analysis was validated on every measurement day with deep seawater reference (DSR) material provided by the Consensus Reference Materials Project of RSMAS (University of Miami) yielding values within the certified range of $42-45 \,\mu\text{mol}\,\text{C}\,\text{L}^{-1}$. Additionally, two internal standards with DOC within the range of those in samples were prepared each measurement day using a potassium hydrogen phthalate (Merck 109017). DOC and TOC concentration was determined in each sample from 5 to 8 injections. The precision was < 4% estimated as the standard deviation of replicate measurements divided by the mean. Particulate organic carbon (POC) was determined as the difference between TOC and DOC.

2.2.2 Total nitrogen (TN) and total dissolved nitrogen (TDN)

TN and TDN were determined simultaneously with TOC and DOC, respectively, using the TNM-1 detector on the Shimadzu analyzer. Nitrogen in the samples is combusted and converted to NOx, which chemiluminesces when mixed with ozone and can be detected using a photomultiplier (Dickson et al., 2007). Calibration of the instrument was done every 8–10 days by measuring standard solutions of 0, 100, 250, 500 and 800 μ g N L⁻¹, prepared with potassium nitrate Suprapur[®] (Merck 105065). Particulate nitrogen (PN) was determined as the difference between TN and TDN. Deep seawater reference (DSR) material provided by the Consensus Reference Materials Project of RSMAS (University of Miami) was used on every measurement day and yielded values within the certified range of 31–33 µmol N L⁻¹. The precision was <2% estimated as the standard deviation of 5–8 measurements divided by the mean.

2.2.3 Total, dissolved and free amino acids

For total hydrolysable amino acids (THAA), 5 mL of sample were filled into pre-combusted glass vials (8 h, 500 °C) and stored at -20 °C until analysis. Samples for dissolved hydrolysable (DHAA) and free amino acids (FAA) were additionally filtered through 0.45 µm Millipore Acrodisc[®] syringe filters and then stored in the same way as samples for THAA. Analysis was performed according to Lindroth and Mopper (1979) and Dittmar et al. (2009) with some modifications. Duplicate samples were hydrolyzed for 20 h at 100 °C with hydrochloric acid (suprapur, Merck) and neutralized by acid evaporation under vacuum in a microwave at 60 °C. Samples were washed with water to remove remaining acid. Analysis was performed on a 1260 HPLC system (Agilent). Thirteen different amino acids were separated with a C18 column (Phenomenex Kinetex, $2.6\,\mu\text{m}, 150 \times 4.6\,\text{mm}$) after in-line derivatization with ophthaldialdehyde and mercaptoethanol. The following standard amino acids were used: aspartic acid (AsX), glutamic acid (GlX), serine (Ser), arginine (Arg), glycine (Gly), threonine (Thr), alanine (Ala), tyrosine (Tyr), valine (Val), phenylalanine (Phe), isoleucine (Ileu), leucine (Leu), γ - amino butyric acid (GABA). α - amino butyric acid was used as an internal standard to account for losses during handling. Solvent A was 5 % acetonitrile (LiChrosolv, Merck, HPLC gradient grade) in sodium-di-hydrogen-phosphate (Merck, suprapur) buffer (PH 7.0). Solvent B was acetonitrile. A gradient was run from 100 % solvent A to 78 % solvent A in 50 min. FAA were determined from DHAA samples without prior hydrolysis in separate analyses. Particulate hydrolysable amino acids (PHAA) were determined by subtracting DHAA from THAA. The detection limit for individual amino acids was 2 nmol monomer L^{-1} . The precision was < 5 %, estimated as the standard deviation of replicate measurements divided by the mean.

2.2.4 Total and dissolved combined carbohydrates

For total and dissolved combined carbohydrates > 1 kDa (TCCHO and DCCHO), 20 mL were filled into pre-

combusted glass vials (8h, 500°C) and kept frozen at -20 °C until analysis. Samples for DCCHO were additionally filtered through 0.45 µm Pall Acrodisc[®] syringe filters. The analysis was conducted according to Engel and Händel (2011) applying high performance anion exchange chromatography coupled with pulsed amperometric detection (HPAEC-PAD) on a Dionex ICS 3000. Samples were desalinated by membrane dialysis (1 kDa MWCO, Spectra Por) for 5 h at 1 °C, hydrolyzed for 20 h at 100 °C with 0.4 M HCl final concentration, and neutralized through acid evaporation under vacuum and nitrogen atmosphere (1 h, 60 °C). Two replicate samples were analyzed. The retention of carbohydrates on exchange columns, and thus the reproducibility of results are highly sensitive to changes in temperature (Panagiotopoulos et al., 2001; Yu and Mou, 2006). For our system, best resolution of sugars was obtained at 25 °C and therefore applied constantly during all analyses. In order to minimize degradation of samples before analysis, the temperature in the auto-sampler was kept at 4 °C. The system was calibrated with a mixed sugar standard solution including (a) the neutral sugars: fucose ($4.6 \,\mu$ M, Fuc), rhamnose ($3.1 \,\mu$ M, Rha), arabinose (2.0 µM, Ara), galactose (2.4 µM, Gal), xylose/ mannose (3.1 µM, Xyl/ Man), glucose (2.4 µM, Glc), (b) the amino sugars: galactosamine (2.0 µM, GalN), glucosamine (2.8 µM, GlcN), and (c) the acidic sugars: galacturonic acid (2.8 µM, Gal-URA), gluconic acid (5.1 µM, Glu-Ac), glucuronic acid (3.0 µM, Glc-URA) and muramic acid (1.9 µM, Mur-Ac). Regular calibration was performed by injecting 12.5, 15.0, 17.5 and 20 µL of mixed standard solution. Linearity of the calibration curves of individual sugar standards was verified in the concentration range 10 nM- $10 \mu \text{M}$. Therefore, the standard mixture was diluted 10, 20, 50, and 100 fold with Milli-Q water. Injection volume for samples and for the blank was 17.5 µL. To check the performance of carbohydrate analysis and stability of the HPLC-PAD system, a 17.5 µL standard solution was analyzed after every second sample. The detection limit was 10 nM for each sugar with a standard deviation between replicate runs of < 2%. Milli-Q water was used as blank to account for potential contamination during sample handling. Blanks were treated and analyzed in the same way as the samples. Blank concentration was subtracted from sample concentration if above the detection limit. Particulate combined carbohydrates (PC-CHO) were determined as the difference between TCCHO

2.2.5 Gel particles

and DCCHO.

Total area, particle numbers and equivalent spherical diameter (d_p) of gel particles were determined by microscopy after Engel (2009). Therefore, 20 to 30 mL were filtered onto 0.4 µm Nuclepore membranes (Whatmann) and stained with 1 mL Alcian Blue solution for polysaccharidic gels (i.e., transparent exopolymer particles (TEP)) and 1 mL Coomassie Brilliant Blue G (CBBG) working solution for proteinaceous gels (i.e., Coomassie stainable particles (CSP)). Filters were mounted onto Cytoclear©slides and stored at -20 °C until microscopy analysis. The size– frequency distribution of gel particles was described by the following:

$$\frac{\mathrm{d}N}{\mathrm{d}\left(d_{\mathrm{p}}\right)} = k d_{\mathrm{p}}^{\delta},\tag{2}$$

where d*N* is the number of particles per unit water volume in the size range d_p to $(d_p + d(d_p)$; Mari and Kiørboe, 1996). The factor *k* is a constant that depends on the total number of particles per volume, and δ ($\delta < 0$) describes the spectral slope of the size distribution. The value δ is related to the slope of the cumulative size distribution $N = ad_p^\beta$ by $\delta = \beta+1$, where *N* is the total number of particles per unit water volume. The less negative is δ , the greater is the fraction of larger gels. Both δ and *k* were derived from regressions of $\log(dN/d(d_p))$ vs. $\log(d_p)$ over the size range 1.05–14.14 µm ESD.

Formation of exopolymeric gel particles (e.g., TEP) can be described in terms of coagulation kinetics (Engel et al., 2004; Mari and Burd, 1998). Aggregates can be described using a fractal scaling relationship (e.g., $M \sim L^D$), where M is the mass of the aggregate, L the size and D is the fractal dimension, which is controlled by the size of particles that form the aggregate as well as by the processes of particle collision, e.g Brownian motion, shear, or differential settlement (Meakin, 1991). Assuming that TEP are formed by shear-induced coagulation D can be estimated from δ (Mari and Burd, 1998):

$$D = \frac{(64 - \partial)}{26.2}.$$
 (3)

2.2.6 Heterotrophic bacteria

For bacterial cell numbers, 4 mL samples were fixed with $200 \,\mu\text{L}$ glutaraldehyde (25 % final concentration) and stored at $-20 \,^{\circ}\text{C}$ until enumeration. Samples were stained with SYBR Green I (molecular probes). Heterotrophic bacteria were enumerated using a flow cytometer (Becton & Dickinson FACSCalibur) equipped with a laser emitting at 488 nm and detected by their signature in a plot of side scatter (SSC) vs. green fluorescence (FL1). Heterotrophic bacteria were distinguished from photosynthetic prokaryotes (e.g., *Prochlorococcus*) by their signature in a plot of red fluorescence (FL2) vs. green fluorescence (FL 1). Yellow-green latex beads (Polysciences, $0.5 \,\mu\text{m}$) were used as internal standard. Sampling bacterioneuston with a glass plate does not bias cell abundance measurements (Stolle et al., 2009).

2.2.7 Phytoplankton

For photoautotrophic cell numbers $< 20 \,\mu\text{m}$, 4 mL samples were fixed with $20 \,\mu\text{L}$ glutaraldehyde (25 % final concentration), and stored at $-80 \,^{\circ}\text{C}$ until enumeration. Phytoplankton counts were performed with a FACSCalibur flow-cytometer (Becton Dickinson) equipped with an air-cooled

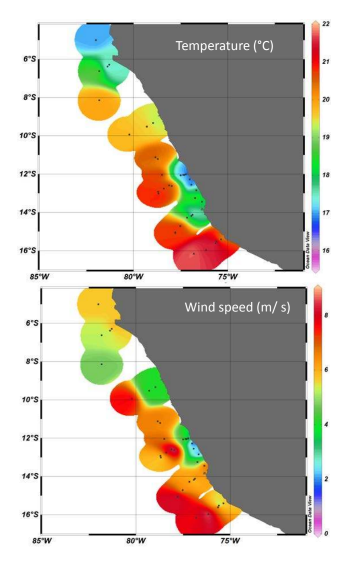


Figure 2. (a) Surface water (1 m depth) temperature ($^{\circ}$ C) and wind speed (m s⁻¹) (b) during M91.

laser providing 15 mW at 488 nm and with a standard filter set-up. The cells were analyzed at high flow rate (\sim 39–41 µL min⁻¹) with the addition of 1 µm-fluorescent beads (Trucount, BD). Autotrophic groups were discriminated on the basis of their forward or right angle light scatter (FALS, RALS) as well as from chlorophyll and phycoery-thrin (characteristic for cyanobacterial, mainly *Synechococcus* populations) fluorescence. Cell counts were analyzed using BD CellQuest Pro-Software. Two groups were distinguished: non-cyanobacterial-type phytoplankton (NCPL) and cyanobacterial-type phytoplankton (CPL).

2.3 Data analysis

The relative concentration of a substance A in the SML was compared to the underlying water (ULW) by the enrichment factor (EF), defined by the following:

$$EF = (A)_{SML} / (A)_{ULW}, \tag{4}$$

where (*A*) is the concentration of a given parameter in the SML or ULW, respectively (GESAMP, 1995). Because the concentration of a component is normalized to its values in the underlying water, EFs for different components can be readily compared. Enrichment of a component is indicated by EF > 1, depletion by EF < 1.

Differences in data as revealed by statistical tests (*t* test) were accepted as significant for p < 0.05. Average values for total concentrations are given by their arithmetic mean, averages for ratios by their geometric mean. Average values are reported with ± 1 standard deviation (SD). Calculations, statistical tests and illustration of the data were performed with the software packages Microsoft Office Excel 2010, Sigma Plot 12.0 (Systat) and Ocean Data View (Schlitzer, 2013). Weighted-average gridding was used in ODV to display data in the SML according to data coverage with automatic scale lengths (53 permille *x* scale length, 40 permille *y* scale length).

3 Results

3.1 The physical environment

Coastal upwelling off the coast of Peru can occur throughout the year (Carr and Kearns, 2003). During the M91 cruise upwelling and upwelling velocities were determined from ³He/⁴He disequilibrium (Steinfeldt et al., 2015). High upwelling velocities of > $3 \times 10^{-5} \text{ m s}^{-1}$ were observed south of Lima (stations 10, 14, 15; Fig. 1). The coastal upwelling of deep water resulted in strong gradients of surface seawater temperature and salinity along the Peruvian shelf as well as with increasing distance to the shelf during M91. Salinity measured at about 1 m depth corresponding to the ship's keel varied between 32 and 35 with the lowest values occurring close to the coast at stations 10_1 to 10_4, 14_1 and 14_2 and 15_1 to 15_3. Here, temperatures were below the average of all surface stations $(19.25 \pm 1.7 \,^{\circ}\text{C})$, indicating the colder, upwelling deep water (Table 1, Fig. 2). Wind speed encountered during the cruise ranged between 0.6 and $9.0 \,\mathrm{m\,s^{-1}}$ with the lower wind speeds also observed closer to the coast, i.e., between 12 and 14° S and at the northern stations (Fig. 2). Thus, higher wind speed was observed at the more off-shore stations having higher surface water temperatures, leading to significant co-variation between surface water temperature and wind speed (Fig. 3). Global radiation and UV radiation varied between 10 and 1103 W m^{-2} , and between 0.8 and 71 W m⁻², respectively, with no significant impact of SML organic matter accumulation.

Table 1. Hydrographic conditions encountered during SML sampling off the coast of Peru in 2012 (M91). Data on air temperature, wind speed, global and UV radiation were obtained from the ship's DShip database for the time of sampling.

	Temperature (°C)	Salinity	Air temperature (°C)	Wind speed $(m s^{-1})$	Global radiation $(W m^{-2})$	UV radiation (W m ⁻²)
Average	19.25	34.87	19.67	5.66	570	37 935
SD	1.70	0.50	0.89	2.14	366	23 384
Min	15.91	32.02	17.30	0.60	10	0.812
Max	21.90	35.32	21.50	9.00	1103	71.10

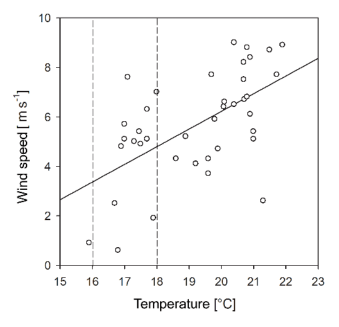


Figure 3. Direct relationship between surface water temperature and wind speed during M91 SML sampling, p < 0.001, r = 0.58, n = 37. Data between 16 and 18° were selected for analysis of wind speed effects at similar temperatures, see Fig. 7.

3.2 SML properties and organic matter accumulation

Estimates for SML thickness are depending on the method applied to sample the SML (Carlson, 1982; Zhang et al., 1998). For the glass plate technique, Zhang et al. (1998) showed that SML thickness decreases with increasing withdrawal rates; i.e., from 50-60 µm for a withdrawal rate of 20 cm s^{-1} , to $10-20 \mu \text{m}$ at rate of $5-6 \text{ cm s}^{-1}$. Their results confirmed earlier studies that generally revealed thinner SML layers at slower withdrawal rates (Carlson, 1982; Harvey and Burzell, 1972; Hatcher and Parker, 1974). During this study, the SML was sampled with the glass plate at $\sim 20 \,\mathrm{cm} \,\mathrm{s}^{-1}$, yielding a thickness between 45 and 60 $\mu\mathrm{m}$, with an overall mean value of $49 \pm 8.89 \,\mu\text{m}$ (n = 39). This value is in good accordance with the proposed apparent sampling thickness of $50 \pm 10 \,\mu\text{m}$ (Zhang et al., 1998) and fits well to previous observations for the SML sampled with a glass plate at the same withdrawal rate (Cunliffe et al., 2011;

Table 2. Concentration of various organic components in the SML during M91, given as average (avg.) and standard deviation (SD), as well as minimum (min) and maximum (max); n = number of observations. For abbreviations see text.

	Unit	Avg.	SD	min	max	п
DOC	μ mol L ⁻¹	94	13	71	122	39
TOC	μ mol L $^{-1}$	127	33	82	199	39
POC	μ mol L ⁻¹	33	25	2.3	96	39
TEP number	$\times 10^{6} \mathrm{L}^{-1}$	19	15	1.8	63	39
TEP area	$\mathrm{mm}^{2} \mathrm{L}^{-1}$	100	106	6.9	408	39
DCCHO	$nmol L^{-1}$	1111	550	507	2668	39
PCCHO	$nmol L^{-1}$	1084	1300	41	5156	34
TN	μ mol L $^{-1}$	16	4.9	8.7	28	39
TDN	μ mol L ⁻¹	12.5	4.0	7.7	25	39
PN	μ mol L ⁻¹	3.3	3.7	bd	16	39
CSP number	$\times 10^{6} L^{-1}$	118	72	19	311	39
CSP area	${\rm mm}^2 {\rm L}^{-1}$	1024	728	137	3051	39
FAA	$nmol L^{-1}$	151	104	49	531	37
DHAA	$nmol L^{-1}$	770	359	423	2017	30
PHAA	$nmol L^{-1}$	1176	774	208	3956	29
NCPL	$\times 10^3 \mathrm{mL}^{-1}$	45	53	5.4	300	35
CPL	$\times 10^3 \mathrm{mL^{-1}}$	27	35	3.7	193	35
Het. bacteria	$\times 10^4 \mathrm{mL^{-1}}$	195	206	3	854	36

Galgani and Engel, 2013; Galgani et al., 2014; Zhang et al., 1998; Zhang, 2003). Using direct pH microelectrode measurements, Zhang (2003) later confirmed an in situ thickness of $\sim 60 \,\mu\text{m}$ for the SML, which they defined as the layer of sudden change of physico-chemical properties.

We therefore assume that samples obtained from the SML during this study well represented the SML, as defined by Zhang (2003). Thickness of the SML as determined during this study increased significantly with amount of organic substances in the SML, determined as TOC concentration (p < 0.005; n = 39). This corroborates earlier findings from experimental studies showing that organic matter produced by phytoplankton increases the thickness of SML sampled with a glass plate (Galgani and Engel, 2013). No correlation instead was observed between SML thickness and wind speed (r = -0.11, n = 39) or between SML thickness and temperature (r = -0.06; n = 39).

Unless stated otherwise, all observations described in this paragraph relate to the SML. In general, concentration of or-

Table 3. Correlation coefficients (*r*) between concentrations of various organic components in the SML and their concentration in the underlying seawater (ULW), temperature (*T*, °C), and wind speed (*U*, m s⁻¹) at the time of sampling. Correlations yielding a significance level of p < 0.01 are marked bold. For abbreviations see text.

SML	<i>r</i> ULW	r_T	r_U	п
DOC	0.75	-0.04	0.06	39
TOC	0.79	-0.53	-0.35	39
POC	0.68	-0.67	-0.48	39
TEP number	0.51	-0.58	-0.69	39
TEP area	0.58	-0.65	-0.69	39
DCCHO	0.94	-0.44	-0.29	39
PCCHO	0.77	-0.59	-0.38	34
TDN	0.24	-0.18	-0.05	39
PN	0.59	-0.55	-0.43	39
CSP number	0.53	-0.04	0.15	39
CSP area	0.68	-0.36	-0.31	39
FAA	0.34	-0.34	0.19	37
DHAA	0.30	-0.47	-0.37	30
PHAA	0.56	-0.64	-0.53	29
NCPL	0.70*	-0.24	-0.21	35
CPL	0.90*	-0.21	-0.31	35
Het. bacteria	0.92	-0.33	-0.37	36

* Only 30 samples were analyzed for NCPL and CPL from the ULW.

ganic components in the SML showed spatial distribution patterns resembling those of temperature and wind speed (Figs. 3, 4, 5). Highest concentration values for nearly all organic components were observed at the upwelling stations 10_1 to 10_4, 14_1 and 14_2 and 15_1 to 15_3 (Fig. 1) in accordance with high estimated primary production rates (Steinfeldt et al., 2015) and high Chl *a* concentrations (Hu et al., 2015) determined in surface waters at these sites during M91.

Phytoneuston abundances (< 20 μ m) varied between 3.7 × 10³ and 1.9,× 10⁵ mL⁻¹ for cyanobacterial-type phytoplankton (CPL; mainly *Synechococcus* spp.) and between 5.4 × 10³ and 3.0 × 10⁵ mL⁻¹ for other non-cyanobacterialtype phytoplankton (NCPL). Generally, highest abundance was determined on and close to the upwelling stations (Fig. 4). On all other stations, cell abundance of CPL and NCPL differed spatially, with higher abundance of NCPL at the southern stations and higher numbers of CPL at the northern stations (Fig. 4). NCPL and CPL were closely related to cell abundance in the ULW (Table 3).

Heterotrophic bacteria were determined in abundances between 3.0×10^4 and $8.5 \times 10^6 \text{ mL}^{-1}$ with the highest numbers observed at the upwelling stations and southeast of the upwelling (Fig. 4). Heterotrophic bacteria in the SML were highly positively correlated to abundances in the ULW (r = 0.94; n = 36; p < 0.001) and negatively influenced by wind speed, although less clearly (r = -0.37; n = 36; p = 0.01). No significant influence on heterotrophic bacterria abundance was detected with respect to global radiation or UV radiation.

TOC concentration ranged between 82 and 199 μ mol L⁻¹, and was clearly higher than DOC concentration on all stations. Particulate Organic Carbon (POC) concentration was calculated as the difference between TOC and DOC and ranged from 2.3 to $96 \,\mu mol \, L^{-1}$. Highest POC concentration was observed at the upwelling stations (Fig. 5). In general, POC concentration was highly correlated to temperature (r = -0.67, n = 39 p < 0.001) and to wind speed (r = -0.48, n = 39 p < 0.001; Table 3). DOC concentration ranged between 71 and 122 μ mol L⁻¹ (Table 2) and, in contrast to POC, was not significantly related to temperature or wind speed (Table 3). Relatively high DOC concentrations of about $100 \,\mu\text{mol}\,\text{L}^{-1}$ were observed at stations 9 and 9 2 (Fig. 5), but excluding these stations from analysis did not reveal a correlation to temperature or wind speed either. DOC is a bulk measure and is quantitatively dominated by refractory compounds that are independent from recent biological productivity. More closely linked to productivity and likely stimulated by the upwelling of nutrients along the Peruvian coast are labile and semi-labile compounds such as dissolved combined carbohydrates and amino acids. Indeed, both DCCHO and DHAA reached highest concentrations at the upwelling stations (Fig. 5). Thereby, maximum concentration of DCCHO of $2670 \text{ nmol } \text{L}^{-1}$ (mean: $1110 \pm 550 \,\mathrm{nmol}\,\mathrm{L}^{-1}$) was observed at station 15_2, slightly south of the station 14 1 exhibiting highest DHAA concentrations of 2020 nmol L⁻¹ (mean: 770 \pm 360 nmol L⁻¹; Table 2). In general high DCCHO concentration was more focused to the upwelling, and exhibited strong horizontal gradients to the northern and southern stations.

DHAA concentration was on average lower than DC-CHO concentration (Table 2) and horizontal differences were less pronounced than for DCCHO. Both components of semi-labile DOC were inversely correlated to temperature (DCCHO r = -0.44, n = 39, p < 0.001; DHAA: r = -0.47, n = 30, p < 0.001), linking their accumulation in the SML to productivity in the cold upwelling waters.

Concentrations of carbohydrates and amino acid in particles, and in gels (i.e., TEP, CSP) in particular, were highest at the coastal upwelling stations also. Particulate carbohydrates and amino acids (PCCHO, PHAA) were highly correlated to POC concentrations (PCCHO: r = 0.70, n = 39, p < 0.001; PHAA: r = 0.81, n = 30, p < 0.001).

In general, numerical abundance as well as total area were about 10-fold higher for CSP than for TEP (Table 2). Spatial variability of gel particles abundance was high, and yielded lowest values of total TEP area of $6.9 \text{ mm}^2 \text{ L}^{-1}$ at station 13_1 and highest values of $408 \text{ mm}^2 \text{ L}^{-1}$ at station 15_1 , about 100 nautical miles apart. The highest abundance of both TEP and CSP was observed close to the coastal upwelling, but apart from these stations, the distribution of TEP in the SML clearly differed from that of CSP (Fig. 5). While higher TEP abundance was observed at the northern stations,

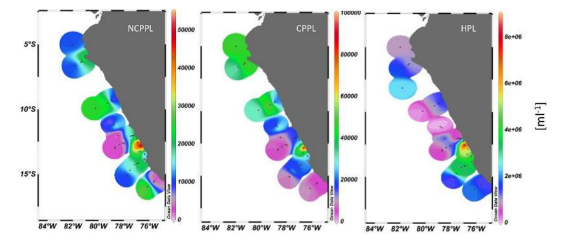


Figure 4. Phyto- and bacterioneuston ($< 20 \,\mu$ m) abundance (number mL⁻¹) in the SML off the coast of Peru during M93: NCPL: "Noncyanobacterial-type" phytoplankton; CPL: "cyanobacterial-type" phytoplankton; HPL: heterotrophic bacterioplankton.

CSP abundance was more pronounced at the southern stations. Moreover, stations of highest and lowest concentration of CSP were different from those of TEP. Lowest value of CSP total area of $137 \text{ mm}^2 \text{ L}^{-1}$ was observed at station 11_1 and highest values of $3051 \text{ mm}^2 \text{ L}^{-1}$ at station 14_1 .

3.3 Accumulation patterns in the SML

For almost all components investigated during this study, concentration in the SML was significantly related to the respective concentration in the ULW (Table 3). Thereby, correlations between SML and ULW were strongest for combined carbohydrates, particularly for DCCHO. Close correlations were also observed for bulk organic carbon measurements, i.e., TOC and DOC, and POC is a combination thereof. For dissolved nitrogenous compounds (i.e., TDN, FAA and DHAA), no relationship between SML and ULW concentrations was observed, suggesting that loss or gain of these compounds in the SML were faster than exchange processes with the ULW. Temperature had an effect on most organic compounds in the SML, with generally higher concentrations at lower temperature (Table 3). This can largely be attributed to the higher production of organic matter at the colder upwelling sites. Concentrations of particulate components POC, TEP, PHCCHO, PHAA and particulate nitrogen (PN) were also inversely related to wind speed, whereas DC-CHO and DHAA were inversely related to temperature but not to wind speed. Clear differences were observed for the two different gel particle types determined in this study. In contrast to TEP, neither abundance nor total area of CSP were related to wind speed, nor to seawater temperature. Instead abundance of CSP in the SML was mostly related to their abundance in ULW. However, with the exception of CSP, particulate components in the SML were affected by changes in wind speed more than concentration of dissolved compounds (Table 3).

Enrichment factors indicated a general accumulation of organic matter in the SML with respect to the ULW (Fig. 6), which happened at most stations. Thereby, clear differences were observed between EF values of different components. The highest enrichment was observed for FAA that were enriched more than 10-fold at some stations. Moreover, FAA were consistently enriched in the SML, except for one station where the lowest FAA concentration was determined (49 nmol L^{-1}). The largest variability of EF was observed for abundance and total area of gel particles. For TEP total area, values of EF ranged between 0.2-12, with highest EF observed at the coastal upwelling station 14_1, where the wind speed recorded was $0.6 \,\mathrm{m \, s^{-1}}$. In proximity of this station, the lowest EF of TEP was determined (station 15_3) indicating a clear depletion at wind speed of 7 m s^{-1} . The EFs of CSP total area ranged between 0.4 and 4.8. Thus highest EF of CSP was clearly lower than for TEP, and in contrast to TEP it was observed at the more offshore station 18 2 at a higher wind speed rate of $9.2 \,\mathrm{m \, s^{-1}}$. Total and dissolved hydrolysable amino acids (THAA, DHAA) were enriched in the SML at almost all stations (Fig. 6), with EFs in the range 0.8-4.6 (DHAA) and 0.4-3.4 (THAA). Median EFs were 1.7 and 1.4 for DHAA and THAA, respectively.

Concentration of TCCHO and DCCHO in the SML were often similar to the ULW, with EF values ranging between 0.6 and 1.4 (DCCHO) and between 0.3 and 1.7 (TCCHO), respectively.

In general, variability of EFs was smaller for dissolved than for particulate organic compounds, suggesting differences in the accumulation dynamics.

In contrast to all organic chemical compounds, bacteria were found to be depleted in the SML at almost all stations (Fig. 6), having a median EF of 0.8

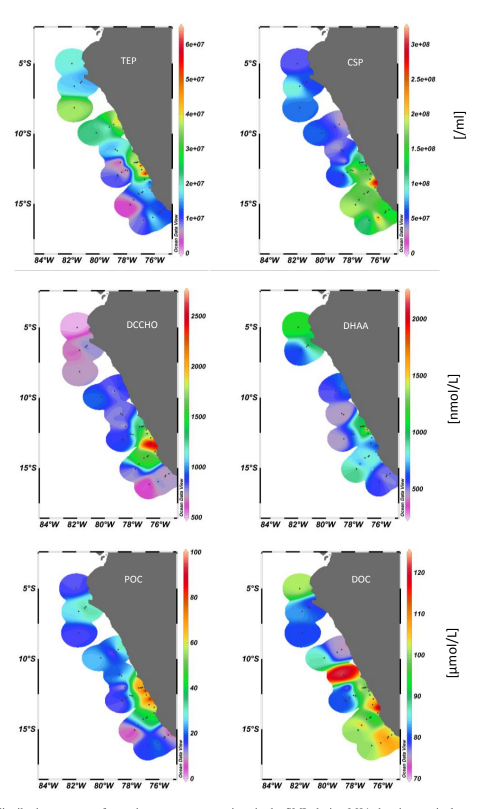


Figure 5. Surface distribution patterns of organic matter concentrations in the SML during M91 showing particulate organic carbon (POC, μ mol L⁻¹), dissolved organic carbon (DOC, μ mol L⁻¹) dissolved combined carbohydrates (DCCHO, nmol L⁻¹), dissolved hydrolysable amino acids (DHAA, nmol L⁻¹) and abundance of TEP (L⁻¹) and CSP (L⁻¹).

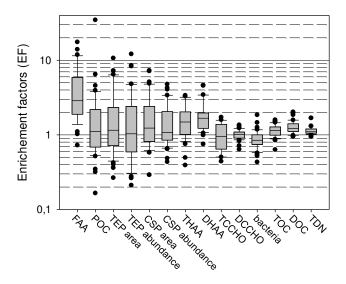


Figure 6. Box and whisker plot of enrichment factors (EFs) calculated for various particulate and dissolved components during M91. Each box encloses 50% of the data with the median value of the variable displayed as a line. The bottom of the box marks the 25%, and the top the 75% limit, of data. The lines extending from the top and bottom of each box marks the 10 and 90% percentiles within the data set and the filled circles indicate the data outside of this range. For abbreviations, see text.

3.4 Size distribution of gel particles within the SML

Abundance of gel particles in the SML and ULW decreased with increasing particle size according to the power law function given in Eq. 2 (Fig. 8). The parameter δ describes the slope of the particles size spectrum. Lower values of δ indicate relatively higher abundance of smaller particles. Data fits to the function were very well described for each sample with $r^2 > 0.90$, yielding a standard error for δ of < 20 %. For TEP, δ varied between -2.63 and -1.38 (mean value: -1.86, SD: 0.27) for particles in the SML and between -2.25 and -1.25 (mean value: -1.70, SD: 0.30) for particles in the ULW. To compare the size distribution of TEP in the SML and the ULW, we calculated the slope ratio $(\delta^* = \delta_{\text{SML}} / \delta_{\text{ULW}};$ Fig. 9). Size distributions of TEP in the SML and ULW were generally quite similar yielding δ^*_{TEP} in the range of 0.78-1.42, with a median value of 1.1. Neverthe less, spatial differences were observed, with $\delta^*_{\text{TEP}} \le 0.95$ at the more coastal northern stations and $\delta^*_{\text{TEP}} > 1.1$ more offshore at the southern stations (Fig. 9). At the upwelling stations with high TEP abundance slopes of SML and ULW were very similar, yielding δ^*_{TEP} in the range 0.95–1.1. This showed a relatively higher abundance of smaller TEP in the SML at the offshore stations, whereas relatively more, larger-sized TEP were present close to the coast in the northern part of the study region. This comparison also showed that sampling of TEP from the SML with a glass plate does not bias TEP size distribution, e.g., by inducing parti-

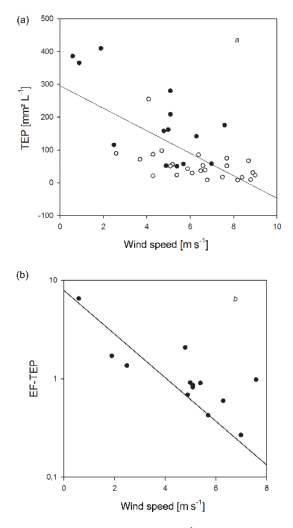


Figure 7. Influence of wind speed $(m s^{-1})$ on the total area concentration of TEP $(mm^2 L^{-1})$ in the SML at all stations (**a**) and relationship between TEP enrichment factors (EFs) and wind speed $(m s^{-1})$ for only those stations that showed similar sea surface temperature as indicated in Fig. 3. Filled dots indicated data from stations of similar sea surface temperature. Data in plot (**b**) were fitted by a power law function.

cle aggregation during sampling. Such a bias would be expected, especially at stations where TEP were highly abundant, like at the upwelling stations. However, particularly at those stations, no differences in size distributions of TEP in the SML and ULW were observed. Fractal scaling exponents of TEP were estimated from Eq. 3 and yielded D = 2.51 for both SML and ULW samples ($D_{SML} = 2.51 \pm 0.015$; $D_{ULW} = 2.51 \pm 0.011$). The very similar fractal dimension for TEP in the SML and ULW suggests that TEP in the SML and in the bulk water are formed by similar aggregation processes. The value of D = 2.51 estimated in this study is close to 2.55 proposed by Mari and Burd (1998) for seawater TEP.

In the SML, the number of TEP in the smallest size class (1.25–1.77 $\mu m)$ ranged from 96 to $1.38\times 10^4\,mL^{-1},$

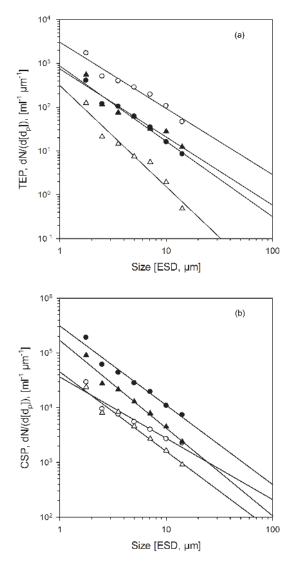


Figure 8. Size–frequency distribution of TEP (**a**) and CSP (**b**) observed during the M91 cruise for samples collected from the SML (open symbols) and in the ULW (filled symbols) at the stations with lowest wind speed of 0.6 m s^{-1} (circles) and highest wind speed of 9.0 m s^{-1} (triangles). Linear regression of $\log(dN/d(d_p))$ vs. $\log(d_p)$ was fitted to the particles in the size range of 1.05–14.14 µm ESD.

and included on average $61 \pm 5.2\%$ of all TEP. For CSP, variability of abundance in the $1.25-1.77\,\mu\text{m}$ size class was much smaller and ranged between 1.46×10^4 and $2.33 \times 10^5 \,\text{mL}^{-1}$. Although CSP thus represented the largest fraction of small gel particles, the relative abundance of CSP in the smallest size fraction was lower, yielding an average contribution of $52 \pm 6.0\%$ of all CSP. Similar to TEP, size distribution of CSP followed the power law relationship of Eq. (2), yielding δ values between -1.12 and -2.01 (mean value: -1.44, SD: 0.20) for particles in the SML and between -1.11 and -1.88 (mean value: -1.39, SD: 0.17) for particles in the ULW. With $D = 2.50 \pm 0.008$, the fractal dimension of CSP was almost identical to that of TEP, suggesting that similar processes, i.e., shear-induced aggregation, are responsible for CSP formation. The slope ratio, δ *, for CSP varied between 0.77 and 1.32, with a median value of 1.0. No spatial pattern was observed for the distribution of δ *_{CSP}. Slopes of the size distribution of CSP in the SML and ULW were not significantly different (p = 0.176, n = 39, paired t test), indicating that CSP size distribution, similarly to TEP, is not biased by the sampling approach of the glass plate.

No overall relationship was established between the slope of the size distribution of TEP and wind velocity (δ_{TEP} vs. wind speed: r = -0.19, n = 37, p = 0.20). However, TEP size distribution was much steeper at the station with highest wind speed compared to the one with lowest wind velocity (δ_{TEP} at 0.6 m s⁻¹ = -1.51, $r^2 = 0.95$, n = 7; δ_{TEP} at 9.0 m s⁻¹ = -2.31, $r^2 = 0.95$, n = 7; Fig. 8a). In particular, at the high wind speed a loss of larger TEP, i.e., > 7 µm was observed in the SML compared to the ULW and relative to the low wind speed station.

For CSP a significant inverse relationship was observed between the slope δ and wind speed (δ_{CSP} vs. wind speed: r = -0.61, n = 37, p < 0.001). A loss of larger CSP was also observed by direct comparison between low and high wind speed stations (δ_{CSP} at 0.6 m s⁻¹ = -1.12, $r^2 = 0.92$, n = 7; δ_{TEP} at 9.0 m s⁻¹ = -1.45, $r^2 = 0.97$, n = 7; Fig. 8b).

4 Discussion

It has been suggested that the presence of organic matter in the SML influences a series of processes relevant to airsea exchange of gases, dissolved and particulate components. EBUSs are characterized by high biological productivity and strong across shelf gradients of organic matter concentration (Capone and Hutchins, 2013). Therefore EBUSs are ideal model systems to study the linkages of biological productivity and SML properties, with respect to characteristics of organic matter composition and factors controlling organic matter enrichment in the SML.

4.1 Organic matter characteristics of the SML in the upwelling region off the coast of Peru

Strong horizontal gradients in organic matter concentration of the SML were observed for the coastal and shelf-break region off the coast of Peru with generally higher organic matter concentrations in the SML towards the area of upwelling of colder, nutrient-rich deep water. Hence, increasing ecosystem productivity is one likely factor responsible for higher concentrations of organic components in the SML. Significant correlations between organic matter concentration in the SML and in the ULW were determined and showed that the SML basically reflects the underlying seawater system. The close connectivity between SML organic properties and biological development was also shown during a recent meso-

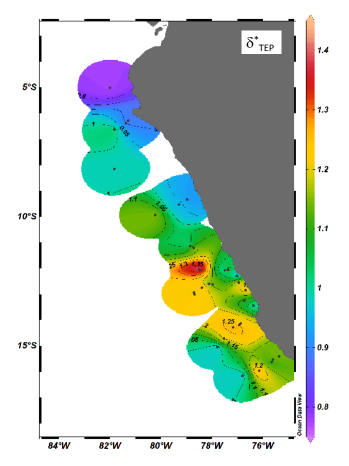


Figure 9. Spatial distribution of the slope ratio, $\delta *$, for TEP in the upwelling region off the coast of Peru during M91.

cosm study, indicating that ecosystem changes impact SML organic matter composition and concentration (Galgani et al., 2014). Despite this finding that relates to a more general characteristic of the SML, clear differences in the accumulation behavior of different organic matter components were determined during this study and are in good accordance with previous observations. A generally higher SML accumulation was observed for amino acids compared to carbohydrates. Significant enrichment of amino acids in the SML has been determined previously for coastal as well as open ocean sites, and higher accumulation of FAA compared to DHAA and THAA, as also observed during this study, appears to be a consistent SML feature (Carlucci et al., 1992; Henrichs and Williams, 1985; Kuznetsova and Lee, 2002, 2001; Kuznetsova et al., 2004; Reinthaler et al., 2008). As for this study, wind velocity and temperature have not been identified as physical factors responsible for amino acid enrichment in the past (Kuznetsova et al., 2004). FAA and DHAA are labile to semi-labile substrates and taken-up by heterotrophic microorganisms (Keil and Kirchman, 1992). Turnover times of these components in the water column are usually in the range of minutes to days (Benner, 2002; Fuhrman and Ferguson, 1986). The observed accumulation of FAA and DHAA in the SML may therefore be related to a reduced activity of bacteria. For different coastal Baltic Sea sites, Stolle et al. (2009) determined a lowered bacterial biomass production in the SML, despite bacterial cell numbers being similar to those in the ULW. During M91 bacteria were mostly depleted in the SML compared to the ULW supporting the idea of the SML being an "extreme environment" for bacteria. Earlier studies showed that some bacteria may be adapted to UV radiation in the SML as well as in the ULW (Agogué et al., 2005; Carlucci et al., 1985). Amino acid consumption by bacterioneuston under UV-B stress may be reduced (Santos et al., 2012), which may give an explanation for the higher concentrations of FAA and DHAA in the SML during M91. However, no significant correlation between bacterial abundance and UV radiation or between UV radiation and amino acid concentrations in the different pools was observed during this study, suggesting that at most stations history rather than instantaneous UV radiation, if at all, responsible for controlling bacteria and organic matter components in the SML.

SML thickness during this study was significantly related to TOC concentration, but not to wind speed. A thickening of the SML with increasing wind speed up to 8 m s^{-1} has been observed by Falkowska (1999) from samples collected in the Baltic Sea and explained by increased advective transport of organic matter to the SML (e.g., through bubble adsorption) at higher turbulence. During M91, accumulation of organic matter in the SML was higher at the upwelling stations where wind speed often was quite low. Hence, a higher source of organic matter in the ULW may have counterbalanced the wind speed effect.

Wind speed, however, was determined as a factor controlling accumulation of particulate material, in particular TEP, in the SML in addition to the dynamics occurring in the ULW. TEP are marine gel particles hypothesized to be neutrally or positively buoyant thanks to their high water content (Azetsu-Scott and Passow, 2004; Engel and Schartau, 1999). TEP were moreover suggested to form within the SML, either by wind-shear-induced aggregation of precursors or due to coalescence of pre-cursor molecules, primarily polysaccharides, when entrained air bubbles burst at the sear surface (Wurl et al., 2011). Adsorption of DOM onto bubble surfaces and TEP formation by bubble bursting have been determined during experimental flotation and bubbling studies using surface seawater from different locations (Wallace and Duce, 1978; Zhou et al., 1998). Bubble scavenging of DOM in the upper water column may thus be responsible for high concentrations of TEP at the SML, because more TEP precursors are lifted up the water column (Gao et al., 2012; Wurl et al., 2011). In addition, compression and dilatation of the SML due to capillary waves may increase the rate of polymer collision, subsequently facilitating gel aggregation (Carlson, 1993). During M91, TEP enrichment in the SML was inversely related to wind speed, supporting earlier observations

of Wurl and colleagues (Wurl et al., 2009, 2011). However, in contrast to earlier observations showing EF values > 1 for TEP in the SML also at higher wind speed, we found the SML to be depleted of TEP at wind speed of $\sim 5 \text{ m s}^{-1}$ and above. It has been suggested that TEP aggregation rates in the SML are higher than in the ULW, due to enhance collision rates by shear or bubble bursting. TEP have been shown to control coagulation efficiencies of solid particles, such as diatoms and coccolithophores (Chow et al., 2015; Engel, 2000; Logan et al., 1995). At higher wind speed, increased aggregation rates of TEP with solid particles, eventually containing mineral ballast, may thus favor the formation of aggregates that become negatively buoyant and sink out of the SML. This, may explain the observed loss of larger TEP (> $7 \mu m$) from the SML relative to the ULW and to the SML at low wind speed. Enhanced aggregation rates could then also explain the inverse relationship between POC and wind speed, observed during this study.

In contrast to TEP, no impact of wind speed was determined for CSP accumulation, or for CSP enrichment in the SML. Moreover, clear spatial differences were observed for the distribution of TEP and CSP in the SML. Although both TEP and CSP are gel particles that form from dissolved organic precursors released by microorganisms, their spatial and temporal occurrence in marine systems can be quite different (e.g., TEP accumulate towards the end of phytoplankton blooms, while CSP rather co-occur with maximum phytoplankton abundance (Cisternas-Novoa et al., 2015; Engel et al., 2015). Moreover, the depth distribution of TEP and CSP was shown to be different for open ocean sites (Cisternas-Novoa et al., 2015). These spatial and temporal differences in the occurrence of TEP and CSP in the water column may explain the spatial separation of both types of marine gels in the SML observed during this study. However, the observed differences in relation to wind speed suggest that additional factors control the enrichment of TEP and CSP in the SML. It has been shown that CSP are less prone to aggregation than TEP (Engel et al., 2015; Prieto et al., 2002). Similarly, CSP may be less involved in aggregation formation and sinking out of the SML at higher wind speed. Yet, similarly to TEP, larger CSP were observed in the SML at low wind speed suggesting that both kind of gels may be involved in slick formation that becomes disrupted when wind speed increases.

4.2 Implications of organic matter accumulation in EBUSs

4.2.1 Air–sea gas exchange

Although the SML and surface active substances (surfactants) within are widely believed affecting the exchange of gases and heat at the air-sea interface (Davies, 1966; Frew, 1997; Salter et al., 2011), particularly at lower wind speed (Liss, 1983), we still have little quantitative knowledge on how natural organic components at the immediate sea-surface alter the gas transfer velocity in water (kw). Our data showed a depletion of the SML with respect to TEP and POC at wind speeds $> 5 \text{ m s}^{-1}$, suggesting that an effect of these "insoluble" components on gas exchange is, if any, operating only at low wind speed. Due to their fractal scaling, gel particles have a relatively large surface to volume ratio and may act as a cover, reducing molecular diffusion rates at the interface between air and sea.

Accumulation of dissolved organic components in the SML during M91 was not related to wind speed. DCCHO and DHAA concentration representing fresh DOM were highest at the upwelling sites and therefore negatively related to seawater temperature. DOM, such as DCCHO and chromophoric dissolved organic matter (CDOM), have demonstrated surfactant properties and reduced gas transfer velocity in water (kw) at low wind speed in laboratory and field experiments (Frew et al., 2004, 1990). The reduction of kw is thereby believed to be related to a dampening of small, capillary waves. Salter et al. (2011) recently showed that artificial surfactants can suppress gas transfer velocity by up to 55 % at sea. Suppression of k666 (i.e., kw normalized to a Schmidt number of 666) during their field study was dependent on wind speed, but was detected up to 11 m s^{-1} , encompassing the full range of wind speed determined during M91. Thus, accumulation of natural DOM particularly in upwelling regimes with high biological production and coastal wind shelter as observed during this study may have an influence on gas exchanges rates as well.

Across the SML, the diffusivity of climate-relevant gases such as methane (CH₄), has been proposed being mediated by SML bacteria, as possible sink (Upstill-Goddard et al., 2003) or source of this compound (Cunliffe et al., 2013). About $\sim 30\%$ of the atmospheric concentration of nitrous oxide (N₂O), one of the strongest greenhouse gases and responsible for ozone depletion, is supported by oceanic sources (Solomon et al., 2007). Of total oceanic N2O production, oxygen minimum zones (OMZs) contribute about 25-75 % (Bange et al., 2001). In EBUSs, high primary production and induced high aerobic remineralization associated with large-scale circulation maintain the presence of OMZs (Gutknecht et al., 2013; Paulmier and Ruiz-Pino, 2009), which, in the last decades, have been expanding and intensifying due to enhanced stratification and reduced ventilation (Keeling et al., 2010; Stramma et al., 2008). During M91, N₂O concentration in surface waters was highly supersaturated at the upwelling sites and in particular at station 14_1 (Arevalo-Martinez et al., 2015). Although a direct influence of organic matter in the SML on gas-exchange was not investigated during M91, it can be assumed that the high enrichment of organic components in the SML observed the upwelling sites was one factor contributing to N2O supersaturation.

Our study was intended to understand how organic matter accumulates in the SML, which might mediate the transfer rate of trace- and greenhouse gases such as N_2O in oceanic regions like OMZs affected by a changing climate. A recent laboratory study reported π non-covalent interactions of N₂O with phenols, suggesting a possible important role of N₂O in biological processes by specifically binding to phenolic groups as those of the amino acids tyrosine and phenylalanine (Cao et al., 2014). Tyrosine and phenylalanine in the SML of our study represented a small molar percentage of total amino acids pool (data not shown), but were present. As we found evidence of overall accumulation of amino acids in the SML during our cruise, for those amino acids in particular the median EF both in the total (THAA) and in the dissolved (DHAA) fraction was > 1, suggesting a possible interaction of specific SML organics with N2O in the coastal upwelling region off the coast of Peru. Although the experiment conducted by Cao and colleagues cannot be directly translated to our setting, it provides interesting ideas for the interaction of N2O with biological macromolecules worth further investigation.

Overall, our results showed that accumulation of organic substances occurs in EBUSs and is related to the increased biological production. Hence, the organic SML may play a particularly important role for exchange of climate-relevant gases that are associated to high organic matter production and resulting anoxia in upwelling systems like the one off the coast of Peru.

4.2.2 Organic aerosol production

The structure of sea-spray aerosols (SSA), originating by bubble bursting at the sea surface, is a function of biological, chemical and physical properties of the SML, which may comprise a vast array of organic surface-active compounds, microorganisms and exopolymer gels (Leck and Bigg, 2005; Quinn and Bates, 2011; Wilson et al., 2015). Despite recent evidences showing that high levels of chlorophyll a are not directly related to the organic carbon content of SSA (Quinn et al., 2014), still organic SSA largely derive from the oceanic surface layer and therefore are also subject to the effects of climate change on marine systems (Andreae and Crutzen, 1997). Polysaccharides and polysaccharidic nanogels (Orellana et al., 2011; Russell et al., 2010) as well as particulate amino acids and proteinaceous compounds (Kuznetsova et al., 2005) are present in organic SSA particles. During M91, we found a different accumulation behavior of TEP and CSP in the SML. TEP showed a close inverse relationship to wind speed, being depleted in the SML above 5 m s^{-1} , while particulate proteinaceous compounds (CSP) accumulated independently of wind speed. Submicron gels embedded in sea spray may represent an important source for primary organic aerosols in the more offshore wind exposed regions. TEP as well as dissolved polysaccharides include sugars with carboxylic groups such as uronic acids and may contribute to the relatively high fraction of carboxylic acid that was observed in the organic matter component of marine aerosols (Hawkins et al., 2010). In the upwelling region off the coast of Peru the wind-driven export of polysaccharidic components to the atmosphere thus might represent a loss-pathway of these organic compounds from the SML that would then contribute to a larger extent to the organic SSA mass. Proteinaceous compounds, including CSP, are probably more stable at the sea surface and may contribute to organic mass in aerosols even at higher wind speed.

However, future studies that investigate gel particles within the SML and in SSA are needed to clarify if the observed loss of TEP from the SML at higher wind speeds is indeed related to a transport of TEP to the atmosphere, or if CSP contribute to organic aerosol mass.

The accumulation of organic matter in the SML, and the distinct behavior of certain compounds at the water–air interface is certainly an important issue for all exchange processes between the ocean and the atmosphere that needs to be further exploited.

Acknowledgements. We thank the captain and crew of R/V *ME*-*TEOR* during cruise leg M91 for logistic support during sampling, especially help related to the rubber boat operation, as well as H. Bange as chief scientist and all the scientific crew. A great acknowledgement goes to J. Roa for helping with SML sampling on board and for TOC/TN and carbohydrates analysis, respectively. Further technical help was provided by R. Flerus, S. Manandhar and N. Bijma for amino acids and microscopy analysis, as well as T. Klüver for flow-cytometry counts. This work was supported by BMBF project SOPRAN II and III (Surface Ocean Processes in the Anthropocene, 03F0611C-TP01 and 03F0662A-TP2.2) and is a contribution to the international SOLAS program.

Edited by: B. Ward

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