Connecting export fluxes to plankton food-web efficiency in the Black Sea waters inflowing into the Mediterranean Sea

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Received July 23, 2009; accepted in principle December 22, 2009; accepted for publication January 15, 2010

Corresponding editor: John Dolan

The short-time scale evolution of plankton carbon partitioning and downward flux in the modified Black Sea water (BSW) mass entering the northeast Aegean Sea was studied using a Lagrangian approach (6–10 April 2008). The free-drifting sediment trap positioned at the bottom of the BSW layer and the control drifter, followed the same path within the anticyclone that circulates the BSW in the area. Zooplankton biomass increased (from 159 to 292 mg C m⁻²), as did faecal pellet production (from 5 to 8 mg C m⁻² day⁻¹), whereas a generally decreasing trend was displayed by particulate organic carbon (POC) (from 2099 to 1440 mg C m⁻²), net primary production and biomass of plankton cells (from 32 to 11 mg C m⁻² day⁻¹ and from 153 to 124 mg C m⁻², respectively). At the same time, the organic carbon flux increased (from 131 to 311 mg C m⁻² day⁻¹), due to the contribution of zooplankton detritus (from 30 to 165 mg C m⁻² day⁻¹). Normalized biomass-size spectra slopes suggest an elevated grazing pressure upon microplankton cells and a non-steady-state ecosystem. Moreover, both the overall shallow slope values and their high correlation to organic carbon flux indicate an increased efficiency of energy transfer to higher trophic levels.

KEYWORDS: plankton size spectra; downward flux; trophic transfer efficiency; Black Sea water; Mediterranean

INTRODUCTION

Plankton are a key component of the biological pump, incorporating carbon into their biomass in the surface ocean and subsequently, by their downward flux, to the ocean floor (De La Rocha and Passow, 2007). Identifying variability in plankton composition and its export out of the euphotic zone provides a basis for connecting plankton food-web functioning and biogeochemical cycles. Downward flux is mainly channelled through the sinking of faecal pellets, marine snow and phytoplankton (Turner, 2002). The strong variability of the relative contribution of these three components to the downward flux and their
remineralization rates depend on many factors including stocks, composition, size spectra, sinking velocities as well as plankton productivity and trophic interactions (Turner, 2002). The normalized biomass-size (NB-S) slopes of planktonic organisms have the potential to reveal the transfer efficiency of biomass (energy) to larger pelagic organisms and it has been suggested that these values can indicate the amount of carbon flux out of the upper ocean (San Martin et al., 2006a).

The Aegean Sea is located in the Eastern Mediterranean Sea, which is characterized as one of the most oligotrophic seas of the world (Azov, 1986; Krom et al., 2004). Oligotrophy is maintained despite the fact that the Aegean Sea receives an inflow of mesotrophic Black Sea water (BSW) mass in the northeast. This inflow consists of very light, brackish water, which is typically located in the uppermost layer (down to 20–30 m) of the Aegean Sea (Zervakis and Georgopoulos, 2002). The BSW mass is enriched in dissolved organic carbon (Sempéré et al., 2002) and nitrogen rather than inorganic nutrients (Polat and Tugrul, 1996) and induces hydrological complexity (e.g. gyres and fronts) that can be highly mobile in time scales of the order of 10 days (Zervakis and Georgopoulos, 2002). These features result in plankton production that is among the highest of the Eastern Mediterranean for both autotrophs (Ignatiades et al., 2002) and heterotrophs (Siokou-Frangou et al., 2002), and the carbon flow within the pelagic food web seems to be quite efficient, affecting higher trophic levels (Stergou et al., 1997; Siokou-Frangou et al., 2002). In addition, the front created in this area is characterized by increased plankton biomass and productivity (Isari et al., 2006; Zervoudaki et al., 2007), whereas the entire northeast Aegean Sea exhibits great spatial variability in hydrography and circulation, as well as in zooplankton communities on a scale of few kilometres (Siokou-Frangou et al., 2009).

Little is known regarding the downward flux of organic matter in the northeast Aegean except that this flux is more substantial than in the south Aegean (Lykousis et al., 2002), which was assumed to be due to a more important export of the primary production (PP) (Siokou-Frangou et al., 2002). Even less information exists on short-term variability (hours to days) of downward flux which has been proved to be as high as interseasonal flux variations, at a Western Mediterranean site (Goutx et al., 2000). Thus, an emerging hypothesis for the area under the BSW influence is that short-term variability of downward flux can be significant.

The aim of the present study was to assess the short-time scale evolution of the organic carbon downward flux out of the BSW mass entering the Mediterranean Sea (northeast Aegean) in relation to water column plankton (stocks and rates) dynamics. Therefore, in order to follow a selected BSW mass, with minimum advection and mixing terms, a short-term Lagrangian study was carried out. Plankton stocks were examined in terms of composition, NB-S slopes and C and N stable isotopic signature. Overall, these data should provide information regarding the evolution of the ecosystem efficiency (as expressed by NB-S slopes) coupled to the geochemical fluxes along the BSW course into the northeast Aegean Sea.

**METHOD**

**Experimental design**

The strategy of the cruise (6–10 April 2008) was to use a multidisciplinary approach following the track of the BSW with Lagrangian drifters. The position of the BSW core was determined by satellite images of sea surface temperature coupled with thermosalinograph records and CTD casts along the track of Lagrangian drifters (24.95–25.50°E, 40.10–40.55°N) (Fig. 1a and b). The latter consisted of a drifting sediment trap and a “control” drifter. The depth of the surface mixed layer (SML) occupied by the BSW was confined in the 0–16 m layer, as revealed by the CTD casts (Fig. 2), and therefore, the sediment trap was deployed at 16 m depth. Drifters were designed following the TELEFOS drifter (Zervakis et al., 2005) based on the standard Davis–CODE design (Davis, 1982) (George Messaritis Co., Greece). The second drifter (control, i.e. without trap) was deployed next to the drifting sediment trap in order to have an undisturbed control of the BSW track. Furthermore, comparison of their two tracks verified that the two instruments showed no windage, faithfully remaining in the same water body.

Several biogeochemical variables were studied in the sediment trap material and in the water column at four stations (3, 5, 6, 8); each station was sampled at the start of four successive trap deployment periods (A, B, C, D) that lasted from 0.3 to 1.3 days (Fig. 1c). Samples were additionally collected at station 2 prior to the first trap deployment at a nearby site. Water column samples were collected at 3, 10 and 20 m depth by means of a CTD-rosette. However, in order to compute the vertical fluxes out of the SML and given that the sediment trap was deployed at 16 m depth, the respective values of exported material at the bottom of the SML were obtained as the average of 10 and 20 m depth values. Mesozooplankton (200–2000 µm) samples were collected by vertical tows of a WP2 200 µm mesh net (0–20 m layer). Special care was taken during sampling of sensitive organisms such as ciliates.
C and N elemental and isotopic analysis

For the determination of suspended particulate organic carbon (POC) and particulate organic nitrogen (PON) concentrations, 1.0–5.0 L of seawater was filtered through pre-combusted (450°C), pre-weighed GF/F filters, then stored at −20°C in the dark and analysed in the laboratory with a Thermo Scientific FLASH 2000 CHNS elemental analyser (Verardo et al., 1990; Cutter and Radford-Knoery, 1991). Inorganic carbon was removed using 2 mol L⁻¹ HCl.

For C and N stable isotopic analysis of sinking particulate organic matter (POM), two to three subsamples from sediment trap material were filtered through pre-combusted (450°C), pre-weighed GF/F filters and then stored at −20°C until analysis. In the laboratory, GF/F filters were lyophilized until dry. For organic C and N elemental and isotopic analyses, filters were then sealed in a desiccator and exposed to an HCl vapour solution for >18 h. Samples were packaged into tin cups, and C and N content and stable isotopic composition were analysed using a PDZ Europa elemental analyser interfaced with a stable isotope ratio mass spectrometer (Stable Isotope Facility, University of California Davis, USA).

Picoplankton to mesozooplankton abundance in water column samples

For counts of heterotrophic bacteria and autotrophic cyanobacteria (Synechococcus spp. and Prochlorococcus spp., cell size 0.2–2 μm), 2 mL duplicate water samples were...
fixed with paraformaldehyde (final concentration 1%), deep-frozen in liquid nitrogen and kept at 
−80°C. Analysis was performed using a FACSCalibur (Becton Dickinson) flow cytometer equipped with a standard laser and filter set and using deionized water as sheath fluid. SYBRGreen I stain (Molecular Probe) was used to stain heterotrophic bacterial DNA.

For counts of auto- and heterotrophic nanoflagellates (AN and HN, respectively; cell size: 1.5–20 μm), 30 mL water samples were fixed with borax-buffered formalin (final concentration 2%), stained with DAPI (Porter and Feig, 1980), and filtered onto black polycarbonate filters of 0.6 μm pore size. AN and HN were distinguished using UV and blue excitation, then further classified in size categories and the biovolume was calculated.

Counting of larger nano- and microplankton (>5 μm cell size) was performed on samples preserved in alkaline (for diatoms, coccolithophorids and dinoflagellates) or acid (for ciliates) Lugol’s solution (final concentration 2%) and refrigerated. Subsamples of 100 mL were analysed after sedimentation for 24 h (Utermöhl, 1958). Cells were counted, distinguished into size-classes and major taxonomic groups and identified down to genus or species level. Cell sizes were measured and converted into cell volumes using appropriate geometric formulae and image analysis (Image-pro plus 6.0).

Mesozooplankton samples were preserved in 4% buffered formaldehyde. Taxonomic identification was performed microscopically on subsamples (of known volume), further scanned for abundance and size determination by image analysis (Image-pro plus 6.0). Macrozooplankton (i.e. zooplankton >2000 μm) was also captured by the net; however, since many empty size classes were present above 2000 μm, the latter was chosen as upper limit for the size spectra analysis.

Conversions of abundance data to carbon units

Heterotrophic bacterial abundance data were converted using 20 fg C cell⁻¹ (Lee and Fuhrman, 1987), 250 fg C cell⁻¹ for Synechococcus (Kana and Glibert, 1987) and 50 fg C cell⁻¹ for Prochlorococcus (Campbell et al., 1994). The biovolume-carbon conversion factor was 183 fg C μm⁻³ for flagellates (Caron et al., 1995) and 190 fg C μm⁻³ for ciliates (Putt and Stoecker, 1989). For larger phytoplankton, cell volumes per species were converted to carbon content applying appropriate conversion factors (Verity et al., 1992; Montagnes et al., 1994). The sizes of zooplankters were converted to carbon based on literature size-carbon relationships (Uye, 1982; Alcaraz et al., 2003).

Primary production

Photosynthetic carbon fixation rates were estimated by means of the ¹⁴C technique (Steemann Nielsen, 1952), as modified for the oligotrophic Eastern Mediterranean (Psarra et al., 2000). Samples were placed in 250 mL polycarbonate bottles (three light and one dark per depth), inoculated with 5 μCi of NaH¹⁴CO₃ tracer and incubated in situ for ca. 2 h around midday, yielding maximum PP rates. Then samples were filtered onto 0.2 μm pore-size polycarbonate filters at <100 mmHg vacuum pressure. Size fractionation onto 0.2, 2.0 and 5.0 μm polycarbonate filters (parallel filtration) allowed estimation of pico- (0.2–2.0), small nano- (2.0–5.0) and larger nano- and microphytoplankton (>5 μm) contributions. To remove excess ¹⁴C-bicarbonate, filters were soaked in 1 mL of 0.1 N HCl in uncapped polyethylene vials overnight and then counted using a Packard
Liquid Scintillation Analyzer, after the addition of 4 mL of BSF scintillation cocktail. Daily (24 h) net PP rates integrated in the 0–16 m layer were estimated (Moutin et al., 1999).

Mesozooplankton faecal pellet production

For the measurement of mesozooplankton faecal pellet production, additional WP2 200 μm net vertical tows were performed in the upper 20 m. Animals were incubated in prefiltred (GF/F) seawater (four 250 mL bottles with a 100 μm mesh near the bottom) for 1 h at the mean BSW temperature. Then, animals were separated from pellets and preserved with 4% formalin. Pellets (number and size) were measured by inverted microscopy and mesozooplankton as described above (Image-pro plus 6.0). The largest mesozooplankton faecal pellet value (610 μm length and 1.43 × 10^7 μm^3) was assumed to be the minimum size of mesozooplankton pellets and was used to distinguish meso- from macrozooplankton pellets in the trap. Faecal pellet volume was converted to carbon using Mediterranean literature values of faecal pellet density (Komar et al., 1981), dry weight to wet weight ratio (Elder and Fowler, 1977) and carbon content per dry weight (Marty et al., 1994).

Sediment trap parameters

The sediment trap design was based on Knauer et al. (Knauer et al., 1979) and consisted of a set of eight PVC cylinders (8 × 65 cm) mounted on a stainless steel cross-frame and connected to one of the above-described drifters. All cylinders were previously filled with filtered seawater (0.2 μm) of increased salinity (plus ca. 3 psu), whereas a solution of ~4% buffered formalin was added as a preservative in three of them. The other five cylinders were left unpreserved to avoid the “swimmer effect”. Here we present analysis only from the unpreserved cylinders. The trap was deployed during four successive periods (varying from 0.3 to 1.3 days) (Fig. 1c). At the end of each deployment period, the content of the unpreserved cylinders was gently mixed. Water subsamples were taken in Lugol’s alkaline solution and formalin for phytoplankton and ciliate microscopic analyses, respectively, and also filtered onto GF/F filters for elemental and stable C and N isotopic composition analyses (as described above). The remaining sample was examined for swimmers, which were removed (always <6% of the total zooplankters) and then concentrated (reverse filtration over a 20 μm mesh) and preserved with formalin for larger particle analysis.

Low-dilution subsamples were scanned, whereas high-dilution subsamples were examined using an inverted microscope. Carcasses and faecal pellets sizes were converted to carbon as described above and corrected for carbon losses due to degradation (Lee and Fisher, 1994) considering the half-time of the trap deployment period.

Biomass-size spectra data processing

The above plankton biomass data from pico- to mesozooplankton (0.2–2000 μm) were integrated over the depth of the BSW water column and organized into NB-S, expressed in carbon units and plotted on a double logarithmic plot of normalized biomass versus particle size (Platt and Denman, 1978). Data were processed as in San Martin et al. (San Martin et al., 2006a, b); the size classes with zero biomass and the inflection points at the interfaces between different methodologies were not included, as they can be subject to error resulting in curvature of the spectrum. Although the Platt and Denman model is sensitive to missing size ranges in the size spectrum, all depth-integrated complete community spectra had an r^2 > 0.98 and a significant regression slope (ANOVA, P < 0.001) when anomaly data (see results) were disregarded. A one-way ANOVA was applied to investigate differences among NB-S slopes.

RESULTS

Hydrography, particle and plankton dynamics in the BSW layer

Both the control drifter and the drifting sediment trap followed an anticyclonic track (Fig. 1b) characterized by high speeds (40 ± 22 cm s⁻¹). Analysis of the salinity–temperature (S/T) characteristics of the water column along the drifter tracks revealed the presence of three distinct water types (Fig. 2). The surface layer was occupied by BSW having recently entered the Aegean Sea. Below the BSW mass, the water column was occupied by waters of Levantine Sea origin (S > 38.5 psu). Between these two layers, and only at the northern part of the drifter track, was also a layer characterized by low salinity and temperature (Fig. 1b and 2). This layer is a remnant of BSW exiting the Dardanelles in winter (winter BSW), which remains in the North Aegean and gradually subsides under the lighter waters exiting the Dardanelles in spring (Zervakis and Georgopoulos, 2002).
The fully Lagrangian behaviour of both the sediment trap and the control drifter are confirmed not only by their identical tracks, but also by the fact that these tracks followed the core of the sea-surface salinity minimum, a clear index of “fresh” BSW in the region, as revealed by the ship’s thermosalinograph (Fig. 1b). The evolution of the mean T/S characteristics of the surface water mass in the above-mentioned core is presented in Fig. 2, overlaid on a T/S diagram from all the CTD casts. As expected in a Lagrangian experiment, the T/S properties remain largely constant; the small changes that are observed are attributed to mixing with the underlying water masses of Levantine waters (mostly for stations 2, 3 and 8), and winter BSW (for stations 5 and 6). Hence, although the Lagrangian drifters followed the water parcels of the surface layer, the water characteristics in the subsurface layers changed due to the shear evident between the upper and lower layers.

Suspended POC decreased progressively along the drifter track from 2099 mg C m⁻² to 1440 mg C m⁻² (Fig. 3a). Similarly, the C:N ratio in suspended POM, with the exception of low values recorded at the first station, displayed a progressive decreasing trend from 9.3 to 7.7, though variability among stations was rather low (coefficient of variation 7.7%).

Phytoplankton carbon biomass (>5 μm) in the BSW was always dominated by dinoflagellates >10 μm, followed by small (5–10 μm) nanoflagellates (Fig. 3b), whereas coccolithophores and diatoms represented only <4 and <1% of total autotrophic biomass, respectively. A progressive decrease in all four phytoplankton groups’ biomass was observed until station 5. After this station, diatom biomass decreased further reaching minimum values (>4-fold decrease), whereas coccolithophores, nanoflagellates and small dinoflagellates increased again, leading to an overall increase in total phytoplankton biomass to slightly lower than the initial levels.

Meso- and macrozooplankton carbon biomass was always dominated by copepods, followed by chaetognaths and salps (Fig. 3c). A general tendency of biomass increase was observed for: (a) copepods >2 mm (*Calanus helgolandicus* adults and juveniles) and for mesozooplankton copepods (with the exception of station 6) and (b) for chaetognaths and total zooplankton biomass, after the first station. At station 2, only juvenile *C. helgolandicus* were collected (<2 mm). Big salps (maximum 5 cm) were seen at the sea surface all along the drifter track except at the last station; however, WP2 net sampling collected them efficiently only at station 6, where their abundance was estimated to be greater than 1 specimen m⁻².

Primary productivity by larger nano- and microphytoplankton (>5 μm), representing cells mostly prone to sinking, was rather elevated at the first stations (32.53 mg C m⁻² day⁻¹, at station 2), then displayed an almost 3-fold decrease from station 2 to station 6 (11.57 mg C m⁻² day⁻¹), and, a 2-fold increase thereafter (21.12 mg C m⁻² day⁻¹ at the last station) (Fig. 3d). These >5 μm phytoplankton cells were responsible for 24–34% of total PP (Fig. 3d). Mesozooplankton faecal pellet production increased along the drifter track from 5.3 to 7.9 mg C m⁻² day⁻¹ (Fig. 3d).

### Biogeochemistry and plankton dynamics of the exported material out of the BSW layer

The POC export out of the BSW layer ranged from 131 mg C m⁻² day⁻¹ at the first deployment period to 311 mg C m⁻² day⁻¹ at the last (Fig. 3e). The material identified was plankton cells, faecal pellets, zooplankton carcasses and marine snow. Plankton cells consisted of diatoms, coccolithophores, dinoflagellates and ciliates. Temporal dynamics of total phytoplankton C flux matched those of total POC flux, with plankton cells constituting always between 16 and 24% of the POC flux (Table I), and both displaying an overall 3-fold increase from period A to period D (Fig. 3f). Overall, phytoplankton community composition in the trap in terms of biomass was not significantly different from the overlying water column, with dinoflagellates again dominating, followed by coccolithophores and diatoms (Fig. 3f). Between the first and the last deployment periods, there was a 4-fold decrease in the percentage diatom contribution accompanied by a slight increase in coccolithophores and dinoflagellates, similar to the water column trends.

Faecal pellets were primarily cylindrical in shape (>99%), followed by few of rectangular shape. Zooplankton carcasses derived mostly from copepods, followed by chaetognaths and salps. Phytoplankton carbon flux exceeded both that of faecal pellets and zooplankton carcasses only in the first deployment period (Fig. 3f, g and h and Table I). Afterwards, faecal pellet and carcasses flux increased, with carcasses becoming the dominant carbon flux over the last two deployment periods (~10-fold increase). Carcass flux was dominated by mesozooplankton, except during the last deployment period, where large copepod carcasses (*C. helgolandicus*) took over, with an overall >10-fold increase. Faecal pellets constituted the highest carbon flux during the second deployment period (Fig. 3h and Table I), and then decreased progressively, due to a decrease in macrozooplankton pellet flux. In contrast,
Fig. 3. Comparison of water column production and composition to vertical flux (organisms and particles > 5 μm) in the BSW layer (0–16 m integrated values). (a) POC concentration and C:N ratio of suspended POM; (b) unicellular plankton groups biomass; (c) zooplankton biomass; (d) net PP > 5 μm (percentages are ratios of net PP > 5 μm to total net PP), mesozooplankton fecal pellet production; (e) POC flux and C:N ratio of sinking POM; (f) unicellular plankton C flux; (g) zooplankton carcasses C flux; (h) fecal C flux. See the correspondence between stations and trap deployment periods in Fig 1c. f.p., fecal pellet; AN, autotrophic nanoflagellates.
the carbon flux of mesozooplankton faecal pellets increased progressively throughout the entire study (Fig. 3h).

A summary of export fluxes of organic carbon and nitrogen and the associated stable isotope compositions are reported in Table II. We observed a large range of carbon and nitrogen fluxes between sediment trap deployments, and also high variability between replicate samples of individual deployments. The coefficient of variation ranged from 15 to 44% for POC fluxes and from 5 to 19% for PON fluxes. The POC flux during deployment A was significantly lower than that of deployment B \((P < 0.01)\) while the final sediment trap deployment captured a significantly higher POC flux than all previous deployment periods \((P < 0.01)\). Deployment A also exhibited significantly lower PON export fluxes than deployments B and D \((P < 0.04)\). We observed no significant differences in the carbon to nitrogen ratios \((C:N)\) between any of the deployments; however, an overall decreasing trend was observed (Fig. 3e).

The average stable carbon isotopic composition of exported POC \(\delta^{13}C\) was similar for each sediment trap deployment \(\approx -24.0 \pm 0.6\)‰, and the variability between deployments was less than that observed between replicate traps of individual deployments \(\approx < 1.0 \pm 0.6\)‰. The average of stable nitrogen isotopic composition \(\delta^{15}N\) of exported PON ranged from 4.8 to 0.4‰ and exhibited a decreasing trend over the course of the experiment. However, there was again considerable variability between replicate samples of individual deployments, whereas \(\delta^{15}N\) compositions were not significantly different between deployments. This variability could have resulted from the inclusion, for example, of a single, additional large copepod in a replicate sample.

Community normalized carbon biomass-size spectra
The NB-S slopes of the plankton community (pico- to mesozooplankton) in the BSW varied from \(-1.09\) to \(-1.00\), the shallowest slope being found at the last station (Fig. 4a and b). These slopes were not significantly different between stations \((P > 0.1)\). Interestingly, a trough anomaly in the slopes of these size spectra was present (Fig. 4a) at the size classes corresponding to large plankton cells \(>50 \mu m ESD\) which were

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<th>Table I: Percentages of plankton components and detritus carbon (estimated) to total carbon (measured POC) in water column stations and sediment trap deployment periods</th>
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<td>Plankton component or detritus</td>
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<td>Dinoflagellates</td>
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<td>Other flagellates</td>
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<td>Diatoms</td>
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<td>Coccolithophores</td>
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<td>Ciliates</td>
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<td>Salps &gt; 2 mm</td>
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<td>Chaetognaths &gt; 2 mm</td>
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<td>Copepods &gt; 2 mm</td>
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<td>Total zoo</td>
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<td>Total faecal pellets</td>
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Sediment trap mesozooplankton (values with asterisks) include copepods <200 µm. UP, unicellular plankton; NM, not measured.

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<th>Table II: Elemental composition of sediment trap material collected during the four deployment periods</th>
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<td>Deployment period</td>
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The values shown are averages (± s.d) of pooled sediment trap samples \(n = 4\) for deployments A and B, and \(n = 3\) for deployments C and D. Stable isotope values are in the per mil (‰) notation.
dominated by ciliates. This anomaly was more or less present at all stations except station 3 (Fig. 4a), where the lowest meso- and macrozooplankton biomass was observed (Fig. 3c). After disregarding these anomalies, \( r^2 \) in all stations increased above 0.98 and slope values were slightly changed (<1%).

The temporal evolution of the BSW NB-S slopes (Fig. 4b) was not significantly correlated to that of the water column parameters (Fig. 3a–d) except with the biomass of total zooplankton (excluding salps) and that of copepods (both \( r^2 > 0.94, P < 0.01 \)). The NB-S slopes obtained at the station corresponding to the beginning of each trap deployment period were plotted against estimates of total exported POC (i.e. the sum of cells, carcasses and faecal pellets carbon) captured during that period and found to be highly correlated \( (r^2 = 0.93, P < 0.01) \) (Fig. 4c).

**DISCUSSION**

The BSW entering the Aegean Sea is known to create a thin surface layer of brackish water, overlying the highly saline waters of Levantine origin (Zervakis and Georgopoulos, 2002). In the present study, this water mass was observed in the upper 16 m of the water...
column. In addition, the hydrography in the region is characterized by the presence of a fast permanent anticyclone around the island of Samothraki which circulates the surface BSW layer within the North Aegean (Zervakis and Georgopoulos, 2002). The anticyclonic circulation increases the residence time of the BSW mass by “trapping” it in the interior of the anticyclone. Both drifters deployed followed the same path along the periphery of this permanent anticyclone and the CTD profiles obtained along this track indicated a progressive salinity modification of the BSW; however, this modification through vertical mixing was delayed by the presence, at the northern edges of the anticyclone, of a layer characterized by low salinity and temperature, which is a remnant of a winter BSW layer (Zervakis and Georgopoulos, 2002) (Fig. 2).

**Biogeochemistry, food-web and export dynamics of the BSW layer**

Most of the biochemical parameters displayed strong temporal variability with rather progressive trends. Similar overall increasing trends were only observed for zooplankton biomass and mesozooplankton faecal pellet production with their respective downward fluxes. Owing to the shallow, short-time sediment trap deployments (i.e. reduced degradation) and the high sinking velocities of zooplankton carcasses and faecal pellets (>25 m day$^{-1}$) (Frangouls et al., 2005), most of the latter were quickly exported to the bottom of the BSW. Moreover, the C:N ratios both in suspended and exported POM displayed an overall decreasing trend. Minimum C:N ratios recorded at the end of the experiment denote a relatively “fresher” material (Gordon and Cranford, 1985), indicative of enhanced in situ production and increased plankton metabolic and growth rates. This is confirmed by the simultaneous increase of zooplankton biomass, faecal pellet production and zooplankton detritus flux. On the other hand, opposite POC trends were observed, with suspended POC values decreasing gradually along the drifter track (except for a slight increase at the last station), whereas the respective POC flux in the trap significantly increased.

Suspended POC concentration (79–131 mg C m$^{-3}$) was higher than that reported (8–56 mg C m$^{-3}$) for different regions in the E. Mediterranean (Scritti et al., 2003; Ediger et al., 2005), but was within the range (20–196 mg C m$^{-3}$) of that measured in the N. Aegean during Spring (Krasakopoulou et al., 2002). Previous studies suggest that these high POC values may originate from river runoff or local biogenic production rather than from the BSW inflow (Karageorgis et al., 2003). POC flux values in this study were higher than those recorded down to 50 m for the rest of the Aegean Sea and other open Mediterranean waters (e.g. Moutin and Rainbault, 2002). This high export may be attributed to the elevated PP in the northeast Aegean, especially during Spring, through an efficient omnivorous planktonic food web in this area (Siokou-Frangou et al., 2002; Zervoudaki et al., 2007). It has been well documented that sedimentation in the form of fresh phytoplankton material is highest during periods of new production in shelf seas (Smetacek, 1984). However, although the total net PP values measured here for the 0–16 m layer were elevated (74.6 ± 32.6 mg C m$^{-2}$ day$^{-1}$, deriving from Fig. 3d), typical for spring values in the area (41.5–157.1 mg C m$^{-2}$ h$^{-1}$, for the 0–100 m layer) (Ignatiades et al., 2002), they are insufficient to explain the high values of POC flux, which far exceed those of PP by cells >5 µm. Neither can these PP values be attributed to mineral nutrient concentrations that were very low and displayed a decreasing trend in the BSW layer during this study (NO$_3$ + NO$_2$: 0.11 ± 0.07 µmol L$^{-1}$, PO$_4$: 4.4 ± 0.5 nmol L$^{-1}$; unpublished results), unless these low nutrient concentrations are due to uptake processes. Moreover, the percentage contribution of daily PP (>5 µm) (Fig. 3d) to the respective phytoplankton C flux (Fig. 3f) displayed a sharp decrease from 93.5% at the first deployment to 28.4%, at the last deployment. This might be attributed to the progressive decrease in PP which in turn could be related to the progressive exhaustion of dissolved inorganic nitrogen. A most probable explanation, however, could be that phytoplankton cells becoming less active after mineral nutrient exhaustion tend to sediment at higher rates, thus contributing to the increase of POC flux. It is worth noting, however, that a direct comparison of phytoplankton biomass and/or daily PP to the respective phytoplankton C flux would be misleading due to low sinking velocities of phytoplankton natural assemblages compared with larger organisms and particles, namely zooplankton cells and carcasses, as described above. Whole phytoplankton community sinking velocities in experimental mesocosms during a spring diatom bloom ranged from 0.29 to 1.53 m day$^{-1}$ (Riebesell, 1989). In this study, with mixed natural nano- and microphytoplankton assemblages dominated by smaller cells (namely dinoflagellates), lower sinking velocities would be anticipated. Therefore, even if maximum sinking rates of 1.53 m day$^{-1}$ were be applied in our data, and depending on the vertical distribution of phytoplankton cells, a minimum of 5–10 days would be required for them to sink to the bottom of the BSW. Given the time scale of the present study, this point leads to a decoupling of PP and suspended
phytoplankton biomass from the respective phytoplankton C flux. Apparently, temporal dynamics of the latter reflect the state of the BSW phytoplankton dynamics, at a time period some days prior to the beginning of our experiment.

Overall, phytoplankton C flux constituted a rather consistent component of the total POC flux (16–24%), whereas zooplankton detritus (faecal pellets and carcasses) constituted a higher but more variable contribution, always exceeding 60% of the POC flux, after the first deployment period.

The faecal pellet C flux and its contribution to the total POC flux in this study (10–53%) was higher than that reported in the oligotrophic South Aegean (<6% at 200 m depth) (Wassmann et al., 2000) and in the open Western Mediterranean (6–30% at 50 m; Fowler et al., 1991). This faecal flux was initially dominated by macrozooplankton faecal pellets and at the end by mesozooplankton faecal pellets. Considering the dominance of cylindrical pellets in the trap and that of copepods in the water column, the origin of mesozooplankton faecal pellet flux appears to be copepods. However, the origin of the cylindrical macrozooplankton faecal pellets is less evident. Salp faecal pellets are excluded, as they are known to produce rectangular pellets (Frangoulis et al., 2005). Although large copepods (C. helgolandicus) could explain a fraction of the large pellets, the size of some cylindrical pellets exceeded the size of C. helgolandicus body. The larger cylindrical pellets could originate from chaetognaths (Dilling and Alldredge, 1993), which were found in the water column, and/or other larger crustacean (e.g. euphausiids). The latter were not captured with the WP2 net but may have been present in the water column and/or migrated into the BSW only during the night.

Sinking zooplankton carcasses also constituted an important C flux, exhibiting a progressive increase over the course of the experiment, becoming the dominant organic carbon flux (>40%) in the last two trap deployment periods. This increase could be related to the progressive increase in zooplankton biomass observed in the water column and/or to a parallel progressive increase in non-predatory mortality rate. The latter may be due to the progressive exhaustion of larger phytoplankton (namely diatoms), resulting in the higher zooplankton detritus export recorded at the last deployment period. Comparison of carcass flux with other studies is difficult as there are few studies that accurately measure this flux. To our knowledge, no such measurements exist in the open Mediterranean, and the only flux estimates (at 200 m) derived from a model study applied in the NW Mediterranean, which estimated a maximum carcass C flux of 6.3 mg C m$^{-2}$ day$^{-1}$ (Andersen and Nival, 1988). This flux was lower than our minimum values (14.3 mg C m$^{-2}$ day$^{-1}$).

The above elevated C flux originating from zooplankton faecal pellets and carcasses, together with the observed trends in the water column zooplankton, suggests that the latter were favoured by the environment. These trends involve an overall increase in the mesozooplankton, chaetognaths and macrozooplankton copepod biomass, as well as that of mesozooplankton faecal pellet production. As the latter probably results from increased food consumption, this might partially account for the observed decrease in PP and the biomass of larger phytoplankton cells during the first half of the study, thus resulting in a respective increase in heterotrophic to autotrophic biomass ratio (for organisms >5 μm ESD: from 4.2 at the first station to 9.5 at the last one). The increase in this ratio coupled to the increase in zooplankton biomass and faecal pellet production could suggest that zooplankton inhabiting the BSW layer enhanced their production (biomass) and grazing along the anticyclonic track of the incoming water mass.

As stated above, we observed increasing trends of POC and PON fluxes, and a decreasing, though not significant, trend in C:N composition of sinking particles over the course of the experiment. The above trends coincided with an increasing trend in phytoplankton sinking flux (3-fold), but mostly with that in zooplankton carcasses (~10-fold) and its relative contribution to the total POC export (Fig. 3). Significant shifts in the stable isotopic composition of sinking POC or PON were not observed during the sediment trap deployment period. The 13C signatures of all sediment trap deployments were similar throughout the experiment (−24.0 to −24.5‰). These values are slightly lighter than typical marine values (ca. −21‰) (e.g. Michener and Shell, 1994) and may be indicative of terrigenous inputs to the system. The 15N signatures showed a general decreasing trend during the sampling period (from 4.6 ± 2.8 to 0.4 ± 3.4‰; Table II) but were highly variable both between deployments and between replicate samples of the sediment traps. Given the substantial variability between replicates, it is difficult to interpret the 15N data. The changes in the relative contribution of phytoplankton, zooplankton and faecal pellets to the sinking flux (Table I) captured by individual traps of each deployment appear to have dramatically altered the integrated 15N signal. We note, however, that the significant increases in sinking zooplankton carcasses during the sampling period did not coincide with similar increases in the 15N signature of the sinking PON flux, as predicted by the ~3‰ isotopic enrichment of 15N that is expected as N is
transferred to higher trophic levels (Checkley and Miller, 1989). Suspended PON and mesozooplankton collected from the water column at three stations during this study also exhibited decreasing trends in δ15N signatures, ranging from 4.0 ± 1.8 to 1.3 ± 0.1‰ and from 5.4 ± 0.3 to 4.1 ± 1.2‰, respectively (unpublished results). The spatiotemporal trends in both suspended and sinking PON of the evolving water mass suggest a shift in the δ15N composition of N that was initially assimilated into the food webs of these regions. This shift, from an isotopically enriched to depleted N source, coincided with increased fluxes of POC, PON and zooplankton carcasses. This could have resulted from the removal (i.e. sinking) of 15N-enriched PON and zooplankton carcasses. Given the low migration capacity of these organisms, the most probable explanation for this anomaly is high mortality, namely due to grazing pressure. This phenomenon appears to be evident in the northeast Aegean, where a strong grazing impact by copepods on ciliates was recorded together with an important consumption of PP (Zervoudaki et al., 2007), supporting a previous hypothesis of strong top-down control of the ciliate population in this area (Pitta and Giannakourou, 2000). The absence of this anomaly at station 3 (Fig. 4a), where the lowest meso- and macrozooplankton biomass was encountered (Fig. 3c), further enhances this hypothesis.

Another peculiarity of the NB-S slopes determined in this study was the anomaly encountered in the size spectra (Fig. 4a). Such an anomaly (i.e. the shift towards lower biomass values) may correspond to excess mortality or migration losses (Zhou, 2006) at that particular section of the spectrum. This anomaly appeared in the carbon size classes that correspond to larger phytoplankton and ciliates biomass. Given the low migration capacity of these organisms, the most probable explanation for this anomaly is high mortality, namely due to grazing pressure. This phenomenon appears to be enhanced in the northeast Aegean, where a strong grazing impact by copepods on ciliates was recorded together with an important consumption of PP (Zervoudaki et al., 2007), supporting a previous hypothesis of strong top-down control of the ciliate population in this area (Pitta and Giannakourou, 2000). The absence of this anomaly at station 3 (Fig. 4a), where the lowest meso- and macrozooplankton biomass was encountered (Fig. 3c), further enhances this hypothesis.

The theoretical value predicted by a model characterizing a near-steady-state open ocean pelagic system, in terms of carbon, is −1.22 (Platt and Denman, 1978). The NB-S slopes of the present study (−1.09 to −1.00) being shallower than the above theoretical value imply a non-steady-state ecosystem. However, it is important to emphasize that it is more the regularity and the linearity of the NB-S spectrum, and not the numerical value of the slope alone that may characterize a close-to-steady-state system (Quinones et al., 2003). Therefore, the above-mentioned anomaly recorded in the NB-S further supports a non-steady-state condition of the ecosystem.

Characteristics of community normalized carbon biomass-size spectra

The overall shallow NB-S slopes (−1.09 to −1.00) indicate that there is more biomass transferred to higher trophic levels than might have been expected at equilibrium. This confirms a previous suggestion that the northeast Aegean ecosystem has a higher efficiency of carbon (energy) transfer channelled to mesozooplankton compared with the rest of the Aegean Sea (Siokou-Frangou et al., 2002).

Interestingly, NB-S slopes in this study confirm their potential to reveal the efficiency of carbon flux not only to higher trophic levels but also as sinking out of the upper mixed layer (Richardson et al., 2004). This is shown by the strong relationship between NB-S slopes and the total estimated organic carbon flux (Fig. 4c): higher trophic transfer efficiency indicated by shallower slopes corresponded to higher downward carbon export (end phase), relative to a lower trophic transfer efficiency indicated by steeper slopes, coinciding with a lower downward carbon export out of the BSW (initial phase). In addition, the higher C export values (shallower slopes) corresponded to a higher percentage contribution from higher trophic levels (i.e. zooplankton faecal pellets and carcasses) than under lower C flux values (steeper slopes). The observed relationship between NB-S slopes and carbon fluxes could be due to the fact that both constitute integration over recent time. In contrast, the lack of relationship between the NB-S slopes and the stock/production data (e.g. biomass, PP, as found in the present study (except for zooplankton biomass) and in San Martin et al. (San Martin et al., 2006b), could be attributed to the fact that the latter represent instantaneous measures.

CONCLUSIONS

During this experiment, the substantial variability observed for most biogeochemical variables confirms our original hypothesis that the area under the BSW influence exhibits significant variability in downward flux on short-time scales. The strongest signal was the increase in carbon export out of the BSW, essentially due to a sharp increase in the contribution of zooplankton detritus. A parallel increase in zooplankton biomass and faecal pellet production in the overlying BSW layer, in combination with the overall shallow NB-S slopes, confirmed the hypothesis of a more efficient energy transfer to higher trophic levels in the northeast Aegean Sea, as suggested by previous studies. The recorded anomaly in the NB-S slopes hints at an elevated grazing pressure upon ciliates and larger phytoplankton and a non-steady-state condition of the ecosystem. Overall,
this study provided evidence for connecting BSW plankton food-web efficiency (as expressed by NB-S slopes) to carbon export fluxes in a Mediterranean area affected by the BSW inflow.

ACKNOWLEDGEMENTS

We are grateful to A.C. Banks for providing satellite images and to K. Tsiaras for model predictions of current fields. We thank T. Zoulas, T. Moutso Poupolous for support in the field and the captain and the crew of R/V “Aega" for shipboard assistance. Thanks also to S. Stavrakakis, K. Christodoulou and M. Pettas for their help in the design and construction of the sediment traps as well as to I. Magiopoulos for flow cytometry help in the design and construction of the sediment traps as well as to I. Magiopoulos for flow cytometry analysis and K. Charalambous for CHN analysis. We also thank three anonymous reviewers and the Associate Editor for helpful comments on the manuscript.

FUNDING

Research for this paper was supported by the SESAME project (contract no. 036949), supported by the European Commission’s Sixth Framework Programme on Sustainable Development, Global Change and Ecosystem.

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