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Zooplankton distribution and cross-shelf transfer of carbon in an area of complex mesoscale circulation in the northern California Current

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ABSTRACT

We conducted a research cruise in late summer (July–August) 2000 to study the effect of mesoscale circulation features on zooplankton distributions in the coastal upwelling ecosystem of the northern California Current. Our study area was in a region of complex coastline and bottom topography between Newport, Oregon (44.7°N), and Crescent City, California (41.9°N). Winds were generally strong and equatorward for >6 weeks prior to the cruise, resulting in the upwelling of cold, nutrient-rich water along the coast and an alongshore upwelling jet. In the northern part of the study area, the jet followed the bottom topography, creating a broad, retentive area nearshore over a submarine shelf bank (Heceta Bank, 44–44.4°N). In the south, a meander of the jet extended seaward off of Cape Blanco (42.8°N), resulting in the displacement of coastal water and the associated coastal taxa to >100 km off the continental shelf. Zooplankton biomass was high both over the submarine bank and offshore in the meander of the upwelling jet. We used velocities and standing stocks of plankton in the upper 100 m to estimate that $1 \times 10^6 \text{ m}^3$ of water, containing an average zooplankton biomass of $\sim 20 \text{ mg carbon m}^{-3}$, was transported seaward across the 2000-m isobath in the meandering jet each second. That flux equated to offshore transport of >900 metric tons of carbon each day, and $4\text{--}5 \times 10^4$ tons over the 6–8 week lifetime of the circulation feature. Thus, mesoscale circulation can create disparate regions in which zooplankton populations are retained over the shelf and biomass can accumulate or, alternatively, in which high biomass is advected offshore to the oligotrophic deep sea.

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1. Introduction

Coastal upwelling brings nutrient-rich water to the sea surface and creates highly productive coastal ecosystems in which zooplankton thrive. However, advective transport during upwelling is a challenge that plankton must overcome to avoid local extinction. In relatively warm

areas of the ocean, rapid growth and short generation times may offset advective losses (Escribano and Hidalgo, 2000). In the northern California Current (NCC), where the cold temperature of upwelled water leads to longer development times, diel or ontogenetic vertical migrations by zooplankton to exploit vertical current shears have been proposed as the most likely compensation mechanisms (Peterson, 1998). However, studies that have provided careful calculations of the potential impact of vertical migration on retention (e.g. Batchelder et al., 2002b; Peterson, 1998; Wroblewski, 1982) have done so in

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areas where classical Ekman upwelling cell circulation dominates. In areas of more complex three-dimensional (3-D) alongshore and cross-shelf circulation, migration patterns that are effective in controlling population losses in closed-cell circulation schemes may no longer work, and losses to the deep ocean may be large.

Both coastline and bottom topography have profound influences on circulation patterns. Examples include vortices found over submarine canyons and seamounts (e.g. Hickey, 1997; Torres et al., 2004), island wakes (Caldeira et al., 2005), and oceanic filaments generated near coastal promontories (Haidvogel et al., 1991). Such features influence biological distributions, resulting in advection and redistribution of organisms (Mackas and Coyle, 2005; Mackas and Yelland, 1999). In addition to creating advective features, topography–current interactions can also generate retentive areas where advection is low relative to surrounding areas. Examples of these are upwelling ‘shadows’ (Graham and Largier, 1997; Wing et al., 1998), re-circulation over submarine banks (Ashjian et al., 2001), and stationary eddies (Nishimoto and Washburn, 2002). In these areas, where local growth may keep pace with or exceed advective losses, biomass may be considerably elevated above surrounding areas, creating localized ‘hot spots’ important to population survival and the transfer of energy to upper trophic levels.

Several distinct regions of coastline and bottom topography occur in the NCC. Off northern and central Oregon (north of $\sim 44.5^\circ\text{N}$), the coastline and bottom topography vary gradually alongshore. Between 43.8°N and 44.5°N , a large, shallow submarine bank (Heceta Bank) extends > 100 km to sea. Farther to the south, Cape Blanco (42.8°N), Cape Mendocino (40°N), and Pt. Arena (39°N) are large coastal promontories that cause sharp deviations in the coastline. These topographic regions differ in the complexity of alongshore and cross-shelf circulation patterns that occur during summer upwelling. Coastal upwelling results in an alongshore, equatorward-flowing upwelling jet that develops through the upwelling season from a simple, alongshore current to a complex meandering current that can only maintain connection with the bathymetry in regions of relatively simple topography (Barth et al., 2005a; Gan and Allen, 2005). Where the jet is oriented along isobaths, it may act as a barrier to offshore transport (Castelao and Barth, 2005); where it deviates from the topography, it can create strong flow across isobaths. Off northern and central Oregon, the equatorward jet flows along isobaths, roughly parallel to the coastline throughout the upwelling season. The classic Ekman-driven two-dimensional (2-D) upwelling cell dominates the zonal circulation. However, when the upwelling jet encounters Heceta Bank, it deviates around the seaward margin of the bank, creating a wide area of upwelled water nearshore (Barth et al., 2005b; Gan and Allen, 2005), where flow is sluggish and re-circulation occurs (Castelao and Barth, 2005). The jet meanders strongly offshore at Cape Blanco (Barth et al., 2005b, 2000), causing strong zonal flows. At Cape Blanco and each of the more southern promontories, cold filaments of the upwelling jet frequently extend far to sea, creating deep, strong cross-isobath flow with velocities and

volume transport that far exceed those in Ekman transport (Kosro and Huyer, 1986; Strub et al., 1991).

Evidence of the importance of mesoscale circulation patterns on the distribution of plankton in the NCC is accumulating from several studies of individual advective features, all of which have demonstrated offshore transport of coastal taxa and biomass (Barth et al., 2002; Haury, 1984; Hood et al., 1990; Huntley et al., 2000, 1995; Mackas et al., 1991). Less attention has been paid to the biological impact of retentive areas (though see Largier et al., 2006; Roughan et al., 2006), even though such areas are potentially important to population growth and larval survival. Nearshore, mesoscale circulation may serve as a mechanism of either advective loss or retention of coastal populations, whereas advective features, such as upwelling filaments that cross the shelf break, may cause high transport of coastal organisms to the deep sea.

In late July–early August 2000 (hereafter August 2000), we investigated the role of mesoscale circulation on distributions of mesozooplankton in a study area between Newport, Oregon (44.7°N), and Crescent City, California (41.9°N), 2–145 km from shore. Concurrently, a fishing vessel conducted upper-ocean trawls for nekton, near-surface zooplankton (Reese and Brodeur, 2006; Reese et al., 2005), and gelatinous zooplankton (Suchman and Brodeur, 2005), while a third vessel towed instrument packages to investigate the hydrography and circulation (Barth et al., 2005b), optical properties (Cowles, unpub. data), fish and zooplankton acoustic targets (Pierce et al., 2003; Ressler et al., 2005; Sutor et al., 2005), particle distributions (Zhou and Zhu, 2002), and marine mammals and seabirds (Ainley et al., 2005; Tynan et al., 2005). The studies were part of the US Global Ocean Ecosystems Dynamics Northeast Pacific Program (GLOBEC NEP) Mesoscale Processes study (Batchelder et al., 2002a).

In this paper, we use zooplankton collected with net tows and community analysis techniques to assess the relationship between circulation features and zooplankton community structure. Our goal is to describe the relationship between mesoscale circulation and patterns of coastal zooplankton taxa in our study area, examine differences in distributions among taxa, and use zooplankton distributions and concurrent velocity fields to estimate the impact of circulation features on the advection of biomass from nearshore to the deep sea.

2. Methods

2.1. Field collections

We used conductivity–temperature–depth (CTD) profiles, vertical plankton tows, and a Multiple-Opening-Closing-Nets-and-Environmental Sensing System (MOCNESS) to study the physical and biological patterns from the R.V. *New Horizon* between 28 July and 12 August 2000. Our study area was between Newport, Oregon (44.7°N), and Crescent City, California (42°N), 2–145 km from shore. Cruise reports and data can be accessed through <http://globec.coas.oregonstate.edu/>.

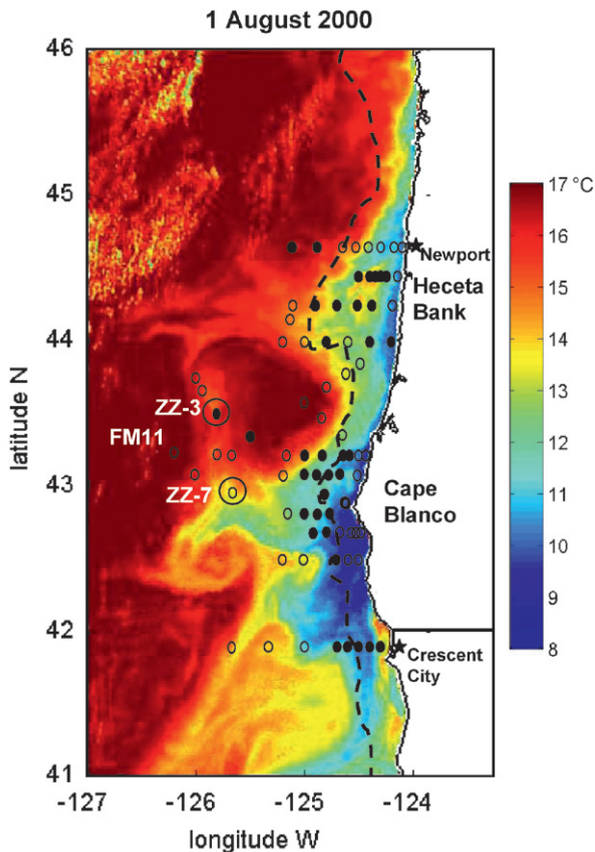


Fig. 1. Sea surface temperature image from the AVHRR on 1 August 2000 with locations of vertical net tows and CTD casts overlaid. Day (○) versus night (●) sampling is indicated by the symbols. The two stations analyzed for depth-distributions from MOCNESS sampling are labeled as ZZ-3 and ZZ-7 and circled and the location of the statistical outlier station, FM11, is shown. The seaward location of the 13.5 °C (yellow) isotherm roughly indicates the position of the core of the equatorward jet as defined in Barth et al. (2005b). The black dashed line is the 200-m isobath.

At each station (Fig. 1), zooplankton were collected with a 50-cm diameter, 202- μm mesh net towed vertically from near bottom (or 100 m depth in deep water) to the surface at a rate of 30 m min^{-1} . A TSK flowmeter was used to monitor the volume of water filtered. Samples were preserved in a 5% buffered formalin/seawater solution. In the laboratory, zooplankton samples were diluted to 5–10 times the settled volume and subsampled with a 1.1-ml stempel pipette. Two to four subsamples were counted for a total of typically 500–700 individuals per sample and 125–250 individuals of the dominant taxa. Copepods and euphausiids were identified by species and developmental stage. Other taxa were identified to genus when possible or larger taxonomic group otherwise. Euphausiid furcilia, chaetognaths, and pteropods (mainly *Limacina helicina*) were counted and their lengths were measured. The abundance of each taxonomic group (number of individuals m^{-3}) was calculated. Species diversity was calculated both as the total number of taxa at a location (species richness) and using H' , the Shannon–Weiner

diversity index (MacArthur and MacArthur, 1961; McCune and Grace, 2002):

$$H' = - \sum_i p_i \ln p_i$$

where p_i is the proportion of individuals from a sample unit belonging to species or taxon i . Biomass values (in mg carbon m^{-3}) were calculated by multiplying abundances with length-weight or stage-specific values found in the literature or generated in our laboratory (e.g. Chisholm and Roff, 1990; Peterson et al., 2002; Uye, 1982). With the exception of doliolids, gelatinous forms were included in abundance data, but not biomass calculations. Each taxon was examined for differences between daytime and nighttime densities using all samples collected. Those which are known to perform diel migration below the depth of our 100 m net casts (euphausiids older than furcilia stage III, shrimp larvae, crab larvae, chaetognaths, and fish larvae) were excluded. No others showed significant differences between day and night, even those that are able to migrate over >100 m ranges (e.g. the copepods *Calanus* spp., *Metridia pacifica*), so were retained in analyses. Other studies in our area have also shown little influence of diel migration on upper-100 m abundances (Peterson, 1972; Peterson and Keister, 2002; Shaw and Robinson, 1998).

In addition to vertical tows, a 1 m^2 mouth area MOCNESS equipped with 335 μm mesh nets was used to collect depth-stratified samples at several stations. Of those stations, two in the offshore areas of the upwelling filament (Fig. 1) were sorted for several dominant coastal and offshore copepod species to determine whether horizontal patterns seen in upper-100 m distributions were representative of those at depth.

At each location where zooplankton were collected, we also performed a CTD cast to 5 m off the bottom (maximum depth of 200 m in deep water) using a Sea-Bird SBE 911 plus CTD. At most stations, water samples were taken from multiple depths for analyses of chlorophyll and nutrient (NO_2 , NO_3 , PO_4) concentrations. Nutrients were analyzed by Oregon State University's chemical oceanography laboratory following the protocols of Gordon et al. (1994). Water for chlorophyll analysis was filtered onto Whatman GF/F filters. Chlorophyll- a was extracted with 90% acetone and analyzed using a Turner model 10 AU fluorometer. All maps of water and biological properties (Figs. 4, 5, 9, and 10) were generated by kriging the discrete data using spherical models with Surfer 8.03 software (Golden Software, Inc., 2002).

2.2. Satellite images

Sea surface temperature data from the Advanced Very High Resolution Radiometer (AVHRR) were captured with 1 km resolution and processed by Ocean Imaging of Solana Beach, California (www.oceani.com) for the US GLOBEC program. All are single-pass images, selected because they had the lowest cloud cover within the time period.

2.3. Velocity and transport estimates

Shipboard acoustic Doppler current profiler (ADCP) velocity data were collected from the R.V. *Wecoma*. We used an RD Instruments 153.6-kHz hull-mounted narrow-band profiler, with 8 m vertical bins and 2.5 min ensemble averaging. The shallowest data were from 17 m. Velocities were detided using the Erofeeva et al. (2003) tidal model. After removal of the barotropic tidal current, a baroclinic tidal signal may still be present. The baroclinic tide is challenging to estimate and highly variable in time and space. Semi-diurnal baroclinic tidal currents off Oregon can be as large as 0.10 m s^{-1} (Kurapov et al., 2003). For other ADCP processing details, see Barth et al. (2005b). Volume transports were estimated from profiles by integrating the observed velocities down to 100 m. Velocities shallower than 17-m were estimated by extending the 17-m value to the surface, i.e. a “slab” model. Biomass transports were estimated by multiplying interpolated vertical net biomass data by the volume transport, point by point approximately every 3 km along the chosen boundary (e.g. the 2000 m isobath) and normal to the local boundary orientation. See Fig. 3 caption for more detail.

2.4. Statistical analyses

We used non-parametric multivariate analyses to analyze zooplankton community structure (McCune and Grace, 2002). The original main “species” matrix had 82 stations \times 64 taxa, with values in the matrix equal to the density ($\# \text{ m}^{-3}$) of each taxon at each station. Prior to analyses, data were $\text{Log}_{10}(Y+0.01)+2$ transformed to normalize the variance while preserving distances among low values (McCune and Mefford, 2005). Rare taxa (those present in <3 samples) were removed to prevent them from strongly influencing results. The final matrix consisted of 82 stations \times 52 taxa. All multivariate analyses were conducted in PC-ORD for Windows 5.05 (McCune and Mefford, 2005).

We sought groupings based on zooplankton community structure using agglomerative hierarchical cluster analysis with a Euclidean distance measure and Ward's Linkage Method (McCune and Grace, 2002). For presentation, the cluster dendrogram is scaled both by Wishart's (1969) objective function and by percentage of information remaining. Wishart's objective function is a measure of information loss as clustering proceeds, and is calculated as the sum of the error sum of squares from the centroid of each group to the items in that group. No rigorous criteria exist for deciding how many cluster groupings to retain for discussion, but we used a multi-response permutation procedure (MRPP) (Mielke, 1984) and Indicator Species Analysis (ISA) (Dufrene and Legendre, 1997) to help decide the optimal number of groupings. The A-statistic from MRPP (Field et al., 1982) assesses within-group homogeneity; thus, when calculated for groups identified by varying numbers of clusters, it can indicate the number of clusters at which within-group homogeneity peaks or plateaus—i.e. the number of

clusters above which separation of the groups is not meaningful. From ISA (described below), a maximum total number of significant indicator species and a minimum overall averaged *p*-value (i.e. more significant results) help determine the optimal number of clusters.

A separate cluster analysis was performed on stations in ‘environmental space.’ The clustering method was identical to the zooplankton community clustering described above except that groupings were based only on the hydrographic variables (temperature, salinity, density) measured at 3, 20, and 50 m depths. Results of this cluster analysis identified groups of stations with similar hydrographic characteristics. A comparison of how stations grouped “hydrographically” versus in “taxonomic/community” structure helps elucidate the mechanisms of variability in community structure.

Following cluster analysis, ISA was used to investigate which taxa were driving the differences among the identified clusters. The indicator value (IV) for a species or taxon in a cluster group is calculated as the product of the relative abundance of the species in the group (the mean abundance in the group divided by the sum of the mean abundances in all groups) and the frequency of occurrence of the species in samples in the group. The statistical significance of each species IV was determined by a Monte Carlo method, in which sample units were randomly reassigned 1000 times to test if the IV was higher than expected by chance (Dufrene and Legendre, 1997).

We used non-metric multi-dimensional scaling (NMDS) ordination using Sorensen's (Bray–Curtis) distance measure to examine similarities in zooplankton community composition among stations and to compare cluster groups to environmental (e.g. temperature, density, salinity, chlorophyll) gradients. NMDS was chosen for this analysis because it is considered one of the most robust ordination methods when dealing with zero–zero species density pairs (Field et al., 1982; Gray et al., 1988), which are common in our data. The final ‘stress’ (a measure of the goodness-of-fit between the data and the final ordination) was examined in relation to the ordination dimensionality to help choose the fewest dimensions necessary to adequately describe the data. Dimensionality was increased if the addition of an axis reduced the stress by >5 , to a maximum of three dimensions, above which interpretation is prohibitively difficult. Stress values approaching or exceeding 20 are considered high; a value of zero indicates perfect monotonicity (McCune and Grace, 2002). The coefficient of determination (r^2) of each axis determines the proportion of the variation in the original distances that is represented by the axis. The Pearson correlation between each ordination axis and individual species and environmental parameters was used to measure the strength and direction (in relation to the axes) of relationships.

We correlated environmental variables with the NMDS ordination axes to examine which variables were important in the alignment of the zooplankton communities. The environmental variables we tested were hydrographic measurements at 3, 20, and 50 m depth (temperature, salinity, density) and chlorophyll and NO_3 measured from

water collected between 1 and 2 m depth (hereafter, 'surface').

3. Results

3.1. The physical setting

In the NCC, upwelling occurs seasonally in spring and summer. In 2000, the spring transition from poleward winds and downwelling, to equatorward winds and upwelling, occurred in mid-March. Prior to our cruise in August, upwelling winds had been exceptionally strong for >6 weeks (Barth et al., 2005b) resulting in cold, upwelled water all along the coast (Fig. 1). During our cruise, the meandering path of the jet caused expanded areas of cold, dense water over both the wide, shallow (<200 m) Heceta Bank, where the jet deviated westward to follow the bottom topography, and in a filament of water that extended seaward off of Cape Blanco (Fig. 1). A time series of satellite sea surface temperature (SST) images from AVHRR shows the development of upwelling over Heceta Bank area and the offshore filament (Fig. 2). Between 11 July and 23 July, the area of upwelled water over Heceta Bank expanded to the 200-m isobath, where it remained until early September. Off Cape Blanco, an offshore deviation of the jet had developed by 11 July, and by 23 July had turned anticyclonically to the northwest. Cold water was carried over the slope >100 km to sea, partially enclosing a warm pool to the north.

Measured ADCP velocities revealed a complex field of mesoscale circulation which included areas of retention and offshore advection (Fig. 3). Velocity vectors show strong flow, particularly along the offshore edge of the warm, anticyclonic water ($\sim 43.5^\circ\text{N}$ 125.5°W) where it is nearly enclosed by the upwelling filament: depth-averaged (0–100 m) velocities reached $>50\text{ cm s}^{-1}$ and 25 m velocities reached $>70\text{ cm}^{-1}$ there (not shown). Strong westward flow occurred offshore of the northern and central edge of Heceta Bank (44.5°N 125.2°W), in the northwest-turning filament off Cape Blanco (42.9°N

125.3°N), and in the small filament present at the southwest corner of our study area (41.8°N 125.6°W). Velocities were very low over Heceta Bank, at the inshore edge of our two southernmost transects, and nearshore along 42.9°N . Velocities inshore of the 200-m isobath were nearly uniformly equatorward or shoreward north of Cape Blanco, but less uniform south of the Cape. The 250 m velocity field revealed a poleward undercurrent that hugged the seaward edge of the 200 m isobath (Fig. 3B).

Near-surface temperatures mapped using the CTD data (Fig. 4A) agree well with satellite SST although the features were not as well defined using discrete observations. Both the expanded area of upwelled water over Heceta Bank and off Cape Blanco were evident to >50 m depth (Fig. 4B). Near the surface of the filament (Fig. 4A and C), the gradients in temperature and density between the oceanic and upwelled water were stronger and occurred farther offshore than at depth (Fig. 4B and D), indicating deep mixing of the upwelled and oceanic water and weaker flow at depth. Station groups identified by hydrographic cluster analysis (using temperature, salinity, and density at three depths, (Fig. 4C) indicate that stations along the outer edge of the Heceta Bank complex, in the center of Heceta Bank, and off Cape Blanco from the nearshore out toward the edge of the filament (Group B) were hydrographically similar and differed from the offshore oceanic water (Group A). The region that hydrographic Group B occupied in the center of Heceta Bank had elevated temperatures and lower density values than surrounding stations indicating the influence of offshore water. That influence was not apparent at 50 m depth (Fig. 4B and D).

3.2. Biomass distribution

Zooplankton biomass was high in the cold, coastal water shoreward of the core of the upwelling jet (indicated in Fig. 1) all along the coast and offshore of Cape Blanco (Fig. 5). Biomass was low in the warm

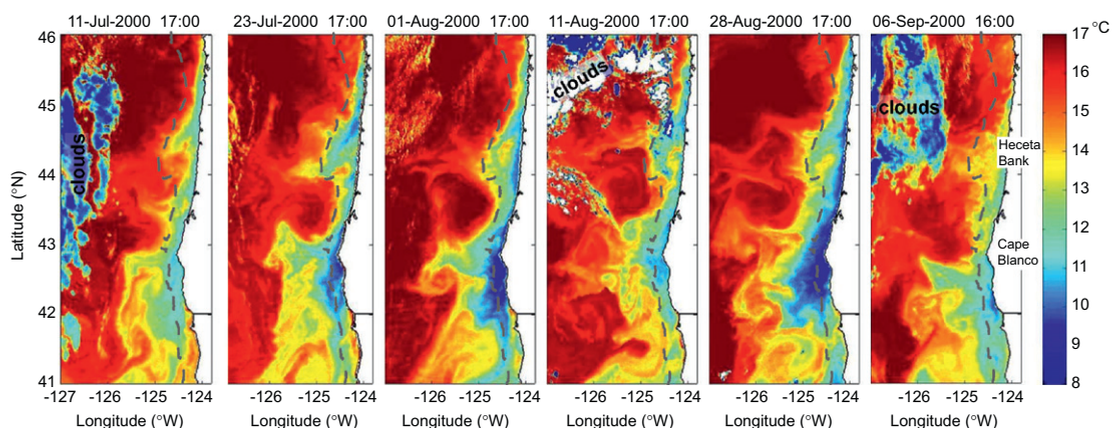


Fig. 2. Sea surface temperature images from the AVHRR showing the development of the upwelling system and the upwelling filament off of Cape Blanco labeled by local time and date. Images are the clearest single passes within 1–2 week intervals. Note that cloud masks are not applied. The 200-m isobath is shown as a dashed line.

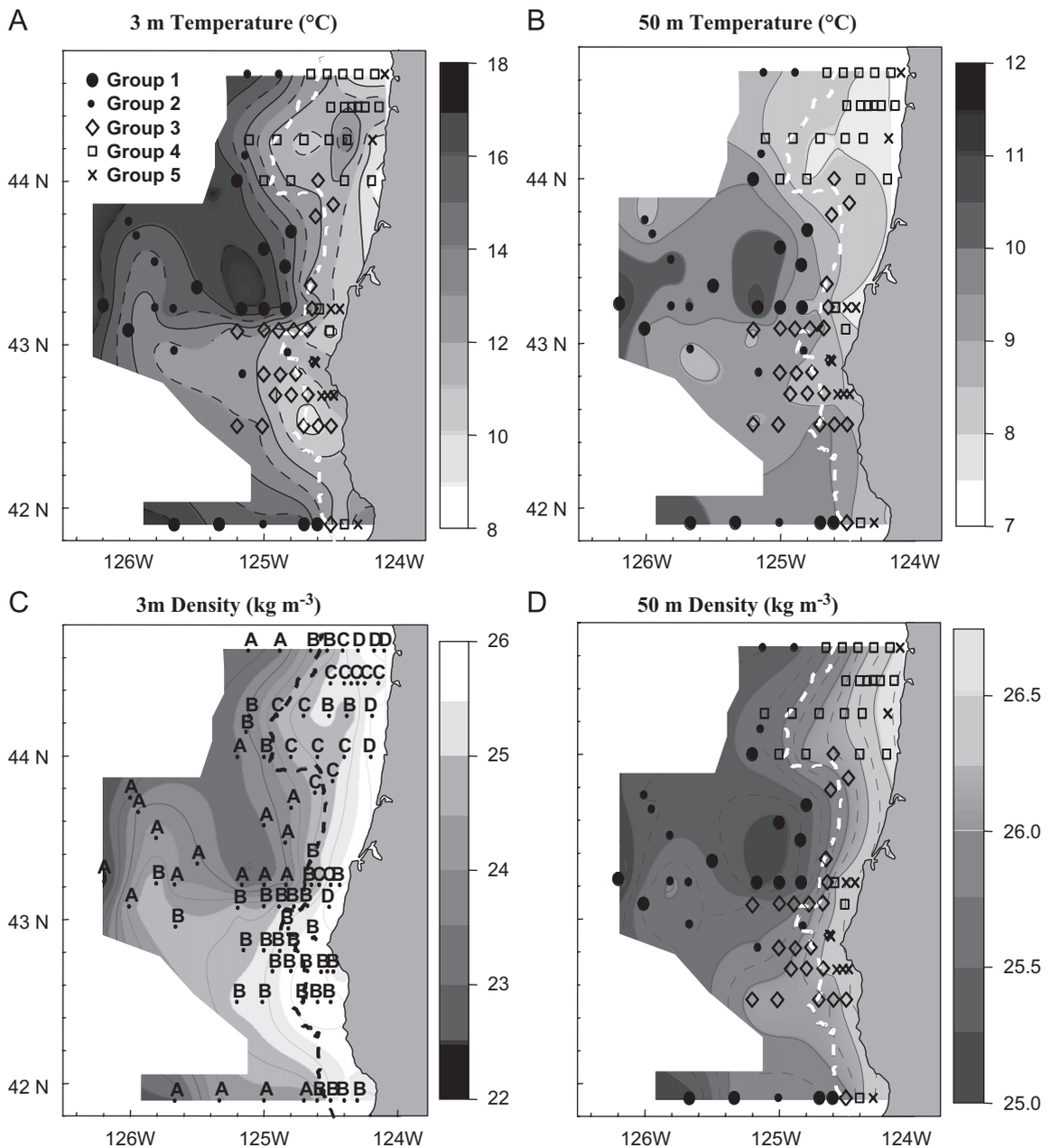


Fig. 4. Temperature and density at 3 and 50 m depths from CTD data. The 200-m isobath is shown as a dotted line. Symbols in A, B, and D represent zooplankton community cluster groupings identified by cluster analysis. Letters in C represent cluster groups defined by the hydrography rather than by the zooplankton taxonomic composition, where (A) 3 m Temperature ($^{\circ}\text{C}$); (B) 50 m Temperature ($^{\circ}\text{C}$); (C) 3 m Density (kg m^{-3}); (D) 50 m Density (kg m^{-3}).

Sorenson distance of 4.1 standard deviations from the grand mean. Station FM11 resided on an edge of the NMDS ordination plot, separated from other stations. However, its outlying position was not driven by unusual total abundances or distributions of any one, or few, taxa. This was the most offshore station we sampled, and highest surface ocean temperature and lowest salinity occurred there. Therefore, we determined that the station's position as an outlier most likely reflected its position at the extreme of the environmental gradients we measured,

and hence brings valuable information to the analyses. Experimental removal of the station from analyses did not noticeably change the ordination of the remaining stations; therefore, station FM11 was retained in all multivariate analyses reported here.

Density, salinity, and temperature at all depths were strongly correlated with Axis 1 of the ordination (Table 1, Fig. 8). Of the environmental variables we measured, the water density at 50 m was most strongly related to the axis ($r=0.87$) though near-surface and 20-m

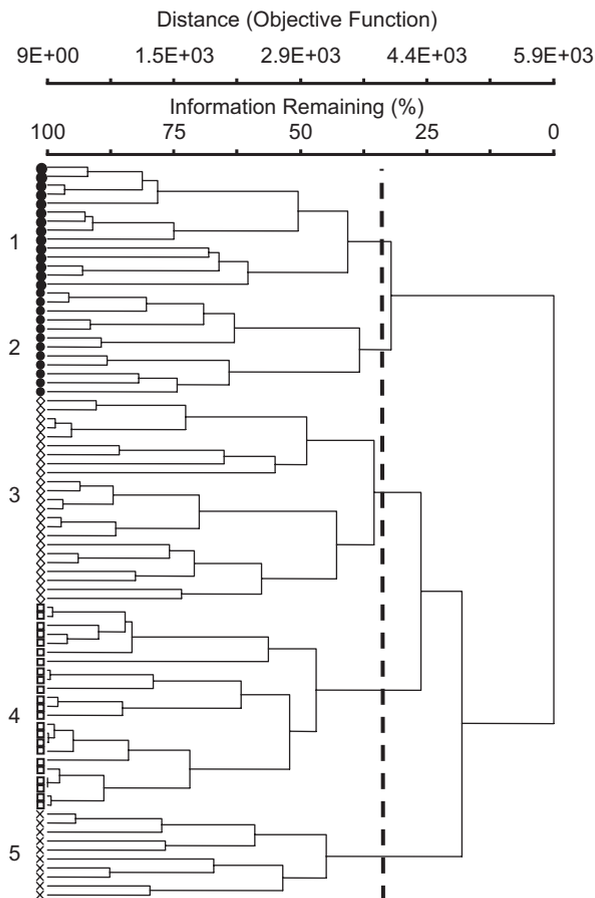


Fig. 7. Zooplankton community dendrogram of stations clustered in species space. Symbols are the same as in Fig. 4. Cluster number is identified along the left.

warmer at 3 m depth than where other groups occurred. The hydrography where Groups 3 and 4 occurred (around Cape Blanco vs. over Heceta Bank) differed only slightly and is reflected in the lack of separation of the two groups along Axis 1. The groups separated along Axis 3 of the ordination (Fig. 8), indicating a stronger correlation with latitude than hydrography. Two warm-water, oceanic species, *Acartia danae* and *Calocalanus styliremis*, were found almost exclusively in Group 1 and were moderately negatively correlated with Axis 1.

Of the 52 taxa examined, 29 showed strong patterns in abundance along hydrographic variables and among cluster groups (Table 2, Figs. 9 and 10). Several of the most abundant species (e.g. *Pseudocalanus mimus*, *Acartia longiremis*, *Calanus marshallae*, and *Centropages abdominalis*) showed gradations in abundance from Group 5 through Group 1: abundances were highest in the nearshore cluster (Group 5), high over Heceta Bank (Group 4), and decreased steadily from the nearshore portion of the upwelling filament to the offshore tip, to the warm, deep ocean (Group 3 to Group 1). All of these taxa were strongly positively correlated with Axis 1 of the ordination (Table 3). Because Axis 1 represents low-to-high temperature and high-to-low salinity and

density gradients, positive species correlations along that axis indicate increasing abundances from the warm, fresh offshore water to the cold, dense nearshore water. Several taxa (e.g. *Calanus pacificus*, *Clausocalanus arcuicornis*, and *Oncaea* spp.) showed the reverse pattern—they were most abundant in Group 1, least abundant or absent in Group 5, and negatively correlated with Axis 1.

We separate the study area into four dynamic biogeographic regions based on the zooplankton communities identified by the taxonomic cluster analysis: the very nearshore (Cluster Group 5), the most offshore (Groups 1 and 2), and the mid-shelf communities north (over Heceta Bank, Group 4) and south (off Cape Blanco, Group 3). The offshore, warm-water groups (Groups 1 and 2) had the highest number of taxa, highest Shannon's diversity indices, and highest number of significant indicator species (Table 2). The very nearshore (Group 5) and most offshore (Groups 1 and 2) communities were the most dissimilar. Several taxa were good indicators of those contrasting communities (Table 2), meaning that those taxa were found most consistently in highest abundances at the locations that made up the clusters. Strongest indicators of the offshore groups (Groups 1 and 2) were *Calocalanus styliremis*, several species of *Clausocalanus*, *Mesocalanus tenuicornis*, *Oncaea* spp., *Paracalanus parvus*, and siphonophores. Strongest indicators of the very nearshore group (Group 5) were *Acartia hudsonica*, *Acartia longiremis*, *Calanus marshallae*, *Centropages abdominalis*, the cladocerans *Evadne* sp., and *Podon* sp., and gastropod and bivalve larvae.

Mid-shelf communities north (over Heceta Bank, Group 4) and south (off Cape Blanco, Group 3) separated from each other, but examination of the densities of animals in those clusters (Table 2) reveals few differences. Euphausiid eggs and nauplii were in uniquely high abundance over Heceta Bank, whereas euphausiid calyp toes and the small cyclopid copepod, *Oithona similis*, were in highest abundance to the south. Those were the only taxa whose abundances were well correlated ($r > 0.6$) with Axis 3 (Table 2, Fig. 10). Their correlations with Axis 3, along which Groups 3 and 4 separated, and their contrasting distributions in relation to Heceta Bank suggest that those taxa drove the separation of the two cluster groups. Other abundances between Groups 3 and 4 were similar.

Lastly, we examined the depth-distributions of several of the dominant copepods (*Calanus marshallae*, *Calanus pacificus*, *Acartia longiremis*, and *Pseudocalanus mimus*) at two offshore locations in the upwelling filament using MOCNESS collections to 350 m depth. Because the MOCNESS mesh size (335 μ m) was too large to collect *Oithona similis*, we do not have vertical distribution data for that species. Depth-distributions varied by species (Fig. 11). The smaller, neritic copepods *Acartia longiremis* and *Pseudocalanus mimus* were restricted primarily to the upper 100 m, as has been noted at other locations in the NCC (Lamb and Peterson, 2005; Mackas and Galbraith, 2002), whereas the larger copepods, *Calanus marshallae* and *Calanus pacificus*, were distributed deeper. There were not apparent differences in vertical distributions due to the day/night sampling difference between the stations.

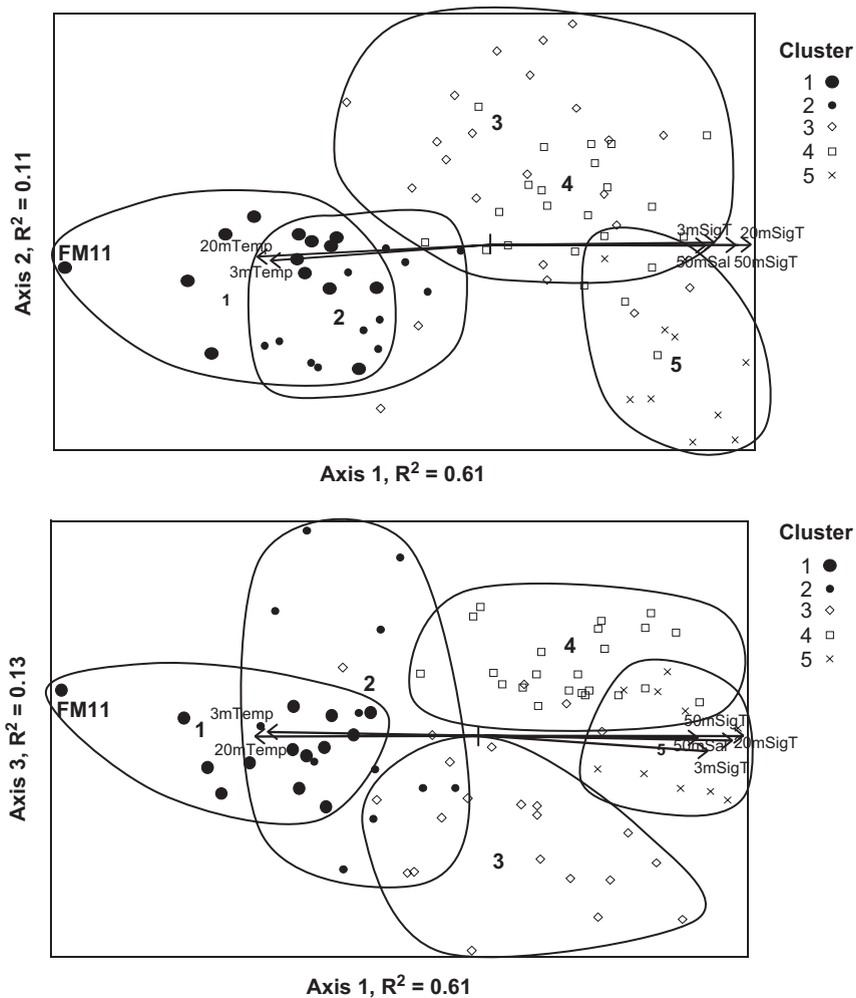


Fig. 8. Non-metric multidimensional scaling (NMS) ordination of stations based on species composition. Three-dimensional results shown as: Top, Axis 1 vs. Axis 2; Bottom, Axis 1 vs. Axis 3. Stations are coded by their cluster grouping as identified in cluster analyses (Fig. 7). The environmental gradients that correlated with the axes $R^2 > 0.6$ are shown; the direction of the correlation is indicated by arrows; the magnitudes of the correlations are given in Table 1.

4. Discussion

In late summer 2000, the meandering path of the equatorward upwelling jet created local areas of onshore retention and offshore advection. Where the jet deviated from the coastline to flow around Heceta Bank, it left an inshore area where current velocities were low and thus retention times were high. South of Heceta Bank, AVHRR images (Fig. 2) indicate that a meander of the jet developed off of Cape Blanco, extended > 100 km offshore and broke down over an approximately 8-week period. Seaward ADCP velocities were high in the filament, indicating large offshore transport of water to the deep sea.

The effects of the hydrography on zooplankton distributions can easily be seen: zooplankton biomass was high where cold surface water was present nearshore, over the wide Heceta Bank area, and in the cool water advected offshore in the upwelling filament, but was low where surface waters were warm. The warm-water, oceanic taxa, which are commonly found throughout the

area during winter (Keister and Peterson, 2003), were held offshore by the upwelling system in late summer. These taxa include the copepods *Clausocalanus* spp., *Paracalanus parvus*, and *Calanus pacificus*—they were in high numbers only well offshore and in the warm eddy off Cape Blanco (Fig. 9). In contrast, the cold-water taxa, which are the typical nearshore dominants during summer upwelling (Keister and Peterson, 2003), were abundant on the shelf, in the retentive area over Heceta Bank, and offshore in the advective filament off Cape Blanco. The patterns in zooplankton distributions seen result both from species affinities for different water masses (i.e. cold, upwelled water versus warm, oceanic water masses) and from the circulation patterns, which are indicated by horizontal gradients in density, salinity, and temperature.

4.1. Heceta Bank retentive area

Heceta Bank is known as a ‘hot spot’ where high standing stocks of chlorophyll, nekton, and upper trophic

Table 1

Pearson correlations between environmental variables and NMS ordination axes

	Axis 1		Axis 2		Axis 3	
	<i>r</i>	<i>r</i> ²	<i>r</i>	<i>r</i> ²	<i>r</i>	<i>r</i> ²
Latitude	0.22	0.05	-0.09	0.01	-0.58	0.33
Longitude	0.74	0.55	-0.21	0.04	-0.10	0.01
Depth	-0.67	0.45	0.29	0.08	0.09	0.01
Distance	-0.76	0.58	0.17	0.03	-0.15	0.02
On/off code	0.75	0.56	-0.14	0.02	-0.22	0.05
D/N code	-0.11	0.01	0.33	0.11	-0.01	0.00
3 m temperature	-0.80	0.64	0.21	0.05	-0.11	0.01
3 m salinity	0.70	0.50	0.07	0.01	0.30	0.09
3 m σ_θ	0.81	0.65	-0.08	0.01	0.22	0.05
20 m temperature	-0.82	0.67	0.18	0.03	0.05	0.00
20 m salinity	0.75	0.56	0.13	0.02	0.24	0.06
20 m σ_θ	0.85	0.72	-0.02	0.00	0.12	0.02
50 m temperature	-0.75	0.56	0.06	0.00	0.40	0.16
50 m salinity	0.79	0.62	0.04	0.00	0.09	0.01
50 m σ_θ	0.87	0.76	0.01	0.00	0.00	0.00
1 m Chla	0.58	0.33	0.13	0.02	-0.09	0.01

D/N code is time of day, coded as day or night. On/off code is station depth coded as >200 or <200 m. Distance is distance from shore.

levels occur (e.g. Ainley et al., 2005; Reese and Brodeur, 2006; Ressler et al., 2005; Tynan et al., 2005). Low velocities over the bank have been noted in several previous studies (Barth et al., 2005a; Castelao and Barth, 2005; Kosro, 2005). Modeling and observations during a 2001 study of Heceta Bank indicated that low velocities over the bank result from the offshore movement of the upwelling jet where the shelf widens, leaving a lee inshore of the jet (Barth et al., 2005a; Gan and Allen, 2005; Kosro, 2005). In 2000, the water in this lee was retained long enough to warm (Barth et al., 2005a,b) and for primary production to draw down the ambient nutrients. Similar processes were noted in a 1995 study by van Geen et al. (2000), who reported biological draw-down of $p\text{CO}_2$ over Heceta Bank that did not occur south of Cape Blanco. The different advective regimes—active circulation in the south versus sluggish circulation over the bank—were implicated in the biological response.

The high abundance of euphausiid eggs on Heceta Bank reflects the location of spawning adults. Ressler et al. (2005) used acoustics and MOCNESS tows to study adult euphausiid distributions during this cruise. They report high abundances over Heceta Bank and directly offshore of Cape Blanco in cold chlorophyll-rich areas. Abundances elsewhere were very low. In euphausiid fecundity experiments (Gómez-Gutiérrez et al., 2007), only females collected from the Heceta Bank region produced large broods—females collected elsewhere produced small broods or did not spawn. From the distributions of spawning adults, eggs, and early life stages of euphausiids, it seems plausible that Heceta Bank was a primary spawning ground in August, from which larval euphausiids were advected south and offshore. Based on velocities measured where egg abundances were highest ($\leq 1 \text{ km d}^{-1}$ to the south, Fig. 3), eggs would not have been advected off the Bank before hatching into nauplii $\sim 40 \text{ h}$ after spawning. We estimate the path length between peak naupliar abundance and peak calyptopis abundance as $\sim 200 \text{ km}$ (Fig. 10), along which velocities of $17 \pm 7 \text{ cm}$

s^{-1} occurred; thus the journey would take $\sim 8\text{--}20$ days to complete. Because euphausiid nauplii develop to calyptopes in $\sim 6\text{--}18$ days (Feinberg et al., 2006), the distributions of the different euphausiid life stages support the hypothesis that the calyptopes found offshore were spawned over Heceta Bank. The distributions also indicate near-continuous spawning over the bank: only a small region between the stations where nauplii and calyptopes were abundant was devoid of larval euphausiids. If spawning was synchronized and infrequent, we likely would not have sampled all of the early life stages during the few (~ 6) days required to sample the study area. Furthermore, we returned to the transect at 44.3°N 1 week later (not shown) where we again sampled equally high abundances of euphausiid eggs indicating frequent high spawning over Heceta Bank that was not noted elsewhere.

The low velocities and high chlorophyll biomass over the Bank also provided suitable habitat for early life stages of other zooplankton. *Calanus marshallae* nauplii (the only copepod nauplii we quantitatively sampled) were abundant over the Bank area (Fig. 10). They were less abundant south of Cape Blanco and off the shelf, even south of the tip of Cape Blanco where chlorophyll was high (Figs. 5B and 10). The Heceta Bank region seems to be an example of an 'ocean triad' (Bakun, 1996) where zooplankton are concentrated (through high spawning), retained (by the slow circulation), and enriched (via upwelling and high chlorophyll biomass), thus creating the 'hot spot' of upper trophic activity (Reese and Brodeur, 2006).

4.2. Advective upwelling filament

In strong contrast to the retentive area over Heceta Bank, the seaward meander of the jet off Cape Blanco created a region of offshore, cross-isobath advection. Fastest seaward flow in the filament occurred along the northern margin, in the core of the cold water observed in the satellite image

Table 2
Characteristics of five station groupings identified by cluster analysis

Cluster group	1	2	3	4	5
Number of samples	14	12	23	23	10
Average number of taxa	24	18	15	14	16
Shannon's diversity index	3.13	2.87	2.62	2.57	2.73
Hydrographic characteristics					
Depth	1557 (1064)	1920 (1144)	599 (780)	182 (292)	62 (24)
Temperature at 3 m (°C)	16.1 (1.2)	14.6 (1.7)	11.7 (1.3)	11.6 (1.5)	11.2 (1.3)
Salinity at 3 m	32.4 (0.5)	32.5 (0.5)	33.1 (0.5)	32.9 (0.5)	33.6 (0.1)
σ_θ at 3 m	23.7 (0.6)	24.1 (0.7)	25.2 (0.5)	25 (0.6)	25.6 (0.3)
Temperature at 20 m (°C)	14 (1.7)	12.1 (1.6)	10.1 (1.1)	9.2 (0.7)	9.2 (0.6)
Salinity at 20 m	32.6 (0.3)	32.7 (0.4)	33.2 (0.4)	33.1 (0.4)	33.8 (0.1)
σ_θ at 20 m	24.3 (0.6)	24.8 (0.5)	25.6 (0.4)	25.6 (0.4)	26.1 (0.1)
Temperature at 50 m (°C)	9.8 (0.7)	9.3 (0.6)	9 (0.4)	8 (0.3)	8.2 (0.5)
Salinity at 50 m	33 (0.3)	33.1 (0.3)	33.6 (0.3)	33.6 (0.3)	33.9 (0.1)
σ_θ at 50 m	25.5 (0.3)	25.6 (0.2)	26 (0.2)	26.2 (0.3)	26.4 (0.1)
Chlorophyll- <i>a</i> at 1 m ($\mu\text{g ml}^{-1}$)	0.5 (0.4)	0.7 (0.5)	1.8 (1.2)	4.7 (3.1)	8.8 (5.9)
Zooplankton Biomass (gC m^{-3})	6.7 (3.1)	11.8 (6.5)	17.9 (10.2)	25.9 (17.7)	63.2 (31.4)
Species abundances (gC m^{-3})					
Copepods					
<i>Acartia danae</i>	3 (5)*	0 (1)	0 (0)	0 (0)	0 (0)
<i>Acartia hudsonica</i>	0 (0)	0 (0)	19 (37)	6 (12)	803 (1795)*
<i>Acartia longiremis</i>	88 (79)	133 (74)	412 (280)	480 (297)	1084 (706)*
<i>Acartia tonsa</i>	0 (1)	0 (0)	0 (1)	0 (1)	1 (4)
<i>Calanus marshallae</i>	5 (7)	8 (11)	79 (78)	91 (80)	232 (235)*
<i>Calanus pacificus</i>	4 (6)*	3 (4)	1 (3)	0 (1)	0 (0)
<i>Centropages abdominalis</i>	1 (2)	1 (2)	9 (15)	20 (27)	290 (495)*
<i>Calocalanus styliremis</i>	9 (10)*	0 (1)	0 (1)	1 (3)	0 (0)
<i>Clausocalanus arcuicornis</i>	17 (13)*	7 (16)	0 (1)	0 (0)	0 (0)
<i>Clausocalanus parapergens</i>	3 (5)*	0 (0)	1 (5)	0 (0)	0 (0)
<i>Clausocalanus pergens</i>	12 (17)*	1 (2)	4 (11)	1 (2)	0 (0)
<i>Corycaeus</i> spp.	0 (1)	4 (10)*	0 (0)	0 (0)	0 (0)
<i>Ctenocalanus vanus</i>	20 (15)*	7 (8)	16 (38)	1 (4)	3 (5)
<i>Eucalanus</i> spp.	1 (1)	1 (1)	0 (0)	0 (0)	0 (0)
<i>Mesocalanus tenuicornis</i>	13 (8)*	2 (2)	1 (3)	1 (4)	3 (6)
<i>Metridia pacifica</i>	13 (7)	16 (12)*	20 (15)	4 (5)	2 (8)
<i>Oithona similis</i>	679 (393)	742 (454)	835 (378)	404 (198)	1237 (690)*
<i>Oncaea</i> spp.	11 (8)*	2 (3)	3 (5)	0 (1)	0 (0)
<i>Paracalanus parvus</i>	159 (158)*	61 (50)	4 (8)	17 (20)	8 (10)
<i>Pseudocalanus mimus</i>	760 (664)	1157 (1063)	2319 (1083)	2412 (2091)	6827 (4073)*
<i>Scolecithricella minor</i>	2 (3)	0 (1)	4 (5)	3 (4)	2 (4)
Other taxa					
<i>Euphausiid</i> eggs	3 (3)	5 (10)	6 (12)	239 (285)*	125 (343)
<i>Euphausiid nauplii</i>	1 (2)	1 (2)	4 (12)	107 (166)*	129 (211)
<i>Euphausiid calyptopes</i>	109 (140)	147 (190)	11 (22)	35 (78)	174 (397)
<i>Euphausiid furcilia</i>	7 (9)	7 (6)*	4 (9)	2 (4)	48 (50)
Amphipods	0 (0)	0 (0)	1 (3)	0 (0)	6 (9)
<i>Evadne</i> sp.	0 (0)	0 (0)	1 (2)	3 (11)	242 (455)*
<i>Podon</i> sp.	0 (0)	0 (0)	1 (2)	0 (2)	242 (267)*
Gastropods	2 (5)	1 (2)	1 (5)	1 (2)	14 (22)*
Bivalves	0 (0)	0 (1.2)	2 (5)	2 (10)	288 (526)*
Polychaetes	1 (2)	1 (1)	0 (1)	3 (12)	66 (200)
Radiolarians	12 (13)	14 (13)*	3 (5)	0 (2)	0 (0)
Medusae	0 (0)	0 (0)	0 (0)	1 (3)	13 (34)
Larvaceans	130 (85)*	52 (69)	4 (13)	49 (67)	32 (27)
Doliolids	3 (7)	24 (50)*	0 (1)	0 (0)	0 (0)
Siphonophores	20 (18)*	16 (21)	0 (1)	0 (0)	0 (0)

Mean values with standard errors in brackets.

Only taxa that showed pattern across hydrographic variables or were significant indicators of a cluster group are shown. A * designates the cluster group for which a taxon's indicator value (IV) was significant.

west of the 2000 m isobath at $\sim 42.7^\circ\text{N}$ 125.3°W , and in the northwestward moving water in the northern extension of the filament. Weaker shoreward flow occurred along the

southern margin of the filament. In the 'root' of the filament between the 200 and 2000 m isobaths, flow was not strongly directional and was generally weak.

Table 3
Pearson correlations between taxa and NMS ordination axes

	Axis 1		Axis 2		Axis 3	
	<i>r</i>	<i>r</i> ²	<i>r</i>	<i>r</i> ²	<i>r</i>	<i>r</i> ²
<i>Acartia danae</i>	–0.55	0.31	0.11	0.01	–0.02	0.00
<i>Acartia hudsonica</i>	0.68	0.46	0.39	0.15	0.20	0.04
<i>Acartia longiremis</i>	0.73	0.53	–0.02	0.00	0.16	0.03
<i>Calanus marshallae</i>	0.83	0.68	–0.09	0.01	0.21	0.04
<i>Calanus pacificus</i>	–0.48	0.23	0.27	0.07	0.17	0.03
<i>Calocalanus styliremis</i>	–0.56	0.32	0.12	0.01	0.02	0.00
<i>Centropages abdominalis</i>	0.58	0.34	0.20	0.04	0.01	0.00
<i>Clausocalanus arcuicornis</i>	–0.73	0.53	0.20	0.04	0.08	0.01
<i>Clausocalanus parapergens</i>	–0.41	0.17	0.03	0.00	0.09	0.01
<i>Clausocalanus Pergens</i>	–0.47	0.22	–0.05	0.00	0.06	0.00
<i>Corycaeus</i> spp.	–0.29	0.08	0.02	0.03	–0.05	0.00
<i>Ctenocalanus vanus</i>	–0.70	0.49	0.19	0.04	0.33	0.11
<i>Mesocalanus tenuicornis</i>	–0.62	0.39	0.21	0.05	0.18	0.03
<i>Metridia pacifica</i>	–0.55	0.30	–0.33	0.11	0.26	0.07
<i>Oithona similis</i>	0.15	0.02	0.29	0.08	0.61	0.38
<i>Oncaea</i> spp.	–0.55	0.31	0.08	0.01	0.33	0.11
<i>Paracalanus parvus</i>	–0.55	0.30	0.22	0.05	–0.44	0.19
<i>Pseudocalanus mimus</i>	0.70	0.49	0.06	0.00	0.35	0.12
<i>Scolecithricella minor</i>	–0.10	0.01	–0.46	0.21	0.19	0.04
Euphausiid eggs	0.19	0.04	–0.27	0.07	–0.60	0.36
Euphausiid nauplii	0.44	0.19	0.03	0.00	–0.61	0.37
Euphausiid furcilia I–III	–0.01	0.00	0.59	0.35	–0.01	0.00
<i>Evadne</i> sp.	0.48	0.23	0.33	0.11	–0.07	0.01
<i>Podon</i> sp.	0.50	0.25	0.48	0.23	0.12	0.02
Bivalve larvae	0.42	0.18	0.40	0.16	0.03	0.00
Radiolarians	–0.70	0.48	0.28	0.08	0.19	0.04
Gastropod larvae	0.05	0.00	0.45	0.20	–0.01	0.00
Larvaceans	–0.34	0.11	0.59	0.35	–0.44	0.20
Doliolids	–0.51	0.26	0.26	0.07	–0.33	0.11
Siphonophores	–0.69	0.48	0.19	0.04	0.09	0.01

The highest correlation with any axis is in bold. Only taxa that were significant indicators of a cluster group are shown.

as on the shelf, whereas, relative to nearshore abundances, *Calanus marshallae* were found offshore in very low abundances. Possibly, the larger taxa that were not found offshore in the upper layers of the filament exhibited behaviors that led them to migrate below the depth of our nets (>100 m) when they sensed increasing water temperatures or decreased chlorophyll offshore. Alternatively, because larger taxa may have higher food requirements (Vidal, 1980) or be subject to higher predation by visual predators (Luo et al., 1996), those taxa may have experienced higher mortality off the shelf than smaller taxa. This species-specific response to offshore advection has been noted in previous studies in this region (Peterson and Keister, 2002) and in eddies off British Columbia (Mackas and Galbraith, 2002). Because larger taxa are likely important prey items for visual predators, the mechanisms generating the patterns are of ecological interest.

The deeper vertical distribution of *Calanus marshallae* observed with the MOCNESS tows compared to the smaller coastal taxa (Fig. 11) suggests that the relative ‘loss’ of large taxa from the offshore filament may partly have been due to their migration below the depth of our vertical nets (max. depth of 100 m). Interestingly, *Calanus marshallae* were found almost exclusively in the deepest layers sampled whereas *C. pacificus* (75 µg C female^{–1}) were collected primarily from the upper 100 m. This

contrast likely resulted from the different habitat preferences of the two species: *C. pacificus* is a warm-water species that usually inhabits the oligotrophic offshore waters whereas *C. marshallae* is a boreal, coastal species that thrives in cold, productive areas (Keister and Peterson, 2003). *C. marshallae*’s occurrence at depth may indicate that they descended to avoid the warm upper layers when carried off the shelf. However, the cues for such migration are not clear, and the low numbers even at depth (<20 m^{–3}) compared to nearshore prevent us from ruling out other explanations (i.e. predation or starvation) for their low numbers in the surface layer.

The relatively high abundances of *Oithona similis* in the Cape Blanco area compared to the Heceta Bank region is interesting (Fig. 10) and may indicate an affinity of that species to advective versus retentive regions. In the Benguela Current, *O. similis* is abundant in the highly advective Lüderitz upwelling region where most copepods cannot sustain high population densities (Huggett, 2008). Elevated production and biomass of *Oithona* in frontal regions has been reported (Castellani et al., 2007) theoretically because their ciliate prey are abundant there, but their relative absence from the Heceta Bank region remains unclear. Because *Oithona similis* is a numerically dominant species worldwide, the precise mechanisms allowing them to prosper in such advective regions warrants further study.

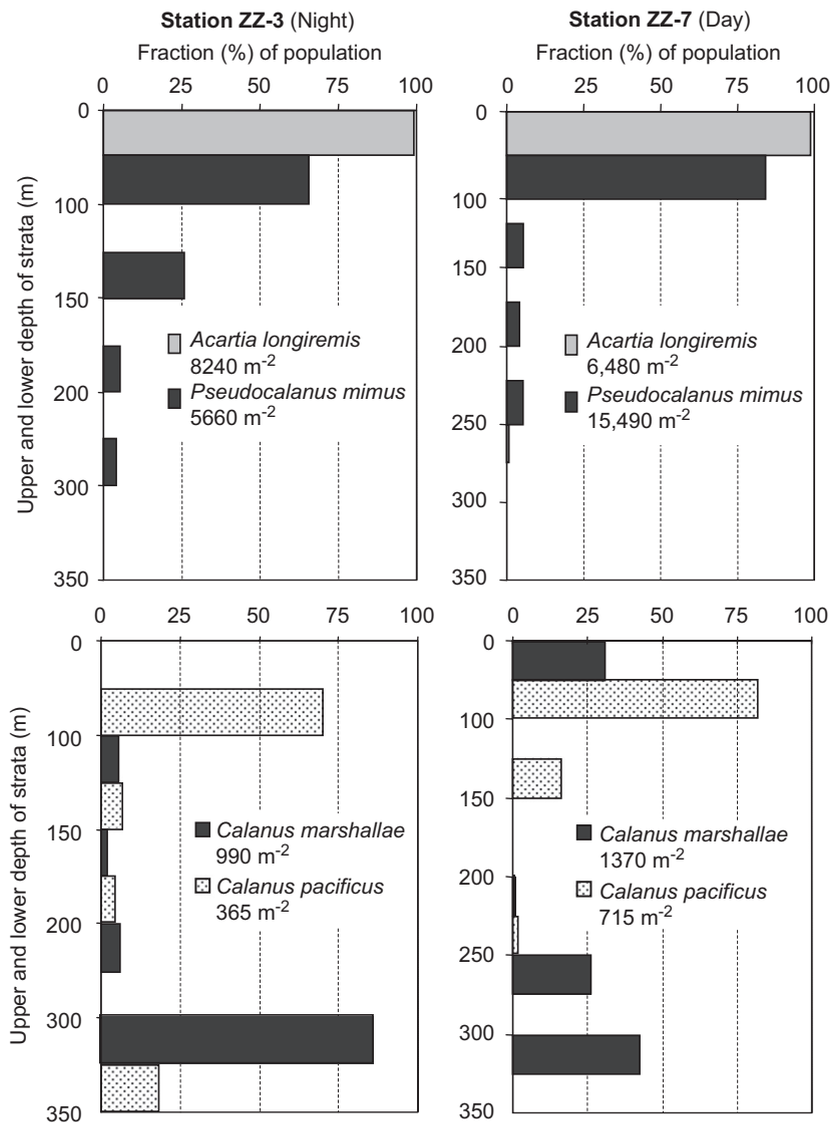


Fig. 11. Depth-distributions of copepods at stations ZZ-3 (left) and ZZ-7 (right) from MOCNESS tows expressed as the fraction of the species total upper 350 m abundance found in each depth interval. Station ZZ3 was sampled at night, ZZ7 during day. Total abundance m⁻² in the upper 350 m is given for each species/station combination. The upper and lower depths of each sampling stratum are shown on the y-axes.

4.3. Transport estimates

Upwelling filaments can transport considerable volumes of water offshore, and often persist for several weeks through fluctuating wind conditions. We assessed the offshore transport of zooplankton biomass in the filament surveyed during this cruise using ADCP velocities and zooplankton biomass interpolated to points along the 2000-m isobath (black vectors and color circles in Fig. 3). For this purpose, we defined the alongshore range of the filament as 42.5–43.2°N based on satellite SST. That range includes the northern margin of the filament, where seaward velocities were high and zooplankton biomass was moderate, and the southern margin where velocities were weakly seaward or shoreward and biomass was very high.

Integrating upper 100 m velocities along the entire 2000-m isobath within the study area (41.9–44.6°N) resulted in a net offshore volume transport of 1.0 Sv (1 Sv = 10⁶ m³ s⁻¹) and a flux of 1200 metric tons carbon per day in zooplankton biomass moving seaward. Transport across the isobath within the filament itself (42.5–43.2°N), though occupying only ~25% of the full latitudinal range, was about 50% of the total offshore transport (0.5 Sv seaward) and 75% of the biomass flux (900 tons C d⁻¹). During the 6–8 week lifetime of the filament, 4–5 × 10⁴ tons of C may have been delivered in the form of zooplankton biomass to the deep ocean. Even recognizing the difficulty in estimating fluxes from biomass estimates in a dynamic system and hence the errors they include (e.g. errors introduced by temporal changes in biomass and flow fields and by integrating

across discrete observations), the transport is significant and resulted in localized, elevated biomass offshore. Even > 100 km off the shelf, biomass was 3–4 times higher within the filament than in surrounding oceanic water (Fig. 5).

After accounting for the flux of ~ 700 tons Cd^{-1} that moved into the study area along our northernmost and southernmost transects (Fig. 3), and the ‘instantaneous’ standing stock of zooplankton inside the 2000 m isobath during the cruise ($\sim 3.2 \times 10^4$ tons C), a net seaward transport of $\sim 2\%$ of the biomass occurred per day in the upper 100 m. This is a conservative estimate because we have not accounted for velocity shear across the upper 17 m, or transport that occurred below 100 m depth, both potentially large fractions of the total. Considering that crustacean zooplankton (the dominant taxa in our samples) grow at $\sim 0.1 d^{-1}$ (Peterson et al., 2002), advective losses were 15–20% of growth. This estimated flux is a larger portion of the inshore production than measured by Mackas and Yelland (1999) (<10%) in filaments off Vancouver Island, B.C., although their estimates included phytoplankton so are not directly comparable. In both regions, the offshore flux removed a small enough portion of the total that populations could have increased (depending on other losses such as predation) even while high advective losses occurred.

Studies of individual upwelling filaments in the California Current System indicate that circulation dynamics within features differ spatially and temporally. Filaments can attain longer east–west spatial scales and higher volume transports than those measured here (Barth et al., 2000; Kosro and Huyer, 1986). So, because mesoscale circulation varies spatially and temporally (Keister and Strub, 2008) and wind-forcing can be stronger than it was in 2000 (Pierce et al., 2006), the offshore transport in filaments could easily be stronger in other years and at other locations.

Upwelling filaments transport coastal taxa to the deep sea in all four of the Eastern Boundary Current upwelling systems, but few quantitative measures of biomass fluxes have been reported to compare with the current study. In the Humboldt Current, Marin et al. (2003) used SeaWiFS data to estimate that $\sim 1.6 \times 10^3$ metric tons of carbon per day was advected offshore in filaments, resulting in export of 5–12% of the coastal carbon production. That flux is on the order of that estimated here. Filaments in the Canary Current region are shallower (<100 m compared to >300 m) and lower velocity than those in the California and Humboldt Current systems, resulting in volume transports of 0.9–1.5 Sv compared to 1 to ≥ 3 Sv in the California Current (Ramp et al., 1991; Barth et al., 2000; Leth and Shaffer 2001; Barton and Arístegui, 2004; Hernández-León et al., 2007; Sanchez, Relvas et al., 2008). Seaward carbon flux is likely to be comparatively low. Indeed, nutrients (Joint et al., 2001), chlorophyll, and copepod biomass (Halvorsen et al., 2001) reported within one Canary Current filament were all low compared to those measured in this study. Similar measurements in the Benguela Current have not yet been reported.

4.4. Transport pathways

Retention or return to the nearshore is ecologically necessary for maintenance of nearshore populations; thus, many coastal zooplankton have life history strategies that slow advective losses from the shelf to allow life cycle closure in upwelling ecosystems. These strategies can include ontogenetic and diel vertical migrations (Peterson, 1998) or high population growth rates (Escribano and Hidalgo, 2000). On Heceta Bank, low velocities indicated retention and small advective losses which may have allowed for high population growth and the high biomass we sampled there.

Whether plankton that are advected offshore in upwelling filaments can return to the shelf to complete their life cycle is unknown. Because velocities in the filament did not tend to strongly change direction with depth in the upper 300 m (not shown), migrations within the depths of most zooplankton’s diel range (<100 m) would not substantially diminish their offshore transport or allow return to the shelf. However, one potential path of return from the offshore filament to nearshore is notable from hydrographic and zooplankton data collected during our study period: Barth et al. (2005b) noted that near-surface drifters did not closely track the deeper circulation. A drifter released over Heceta Bank traveled south alongshore, seaward to the northwest tip of the offshore filament, then completed an anticyclonic path when it crossed dynamic height contours and was entrained into the eastward moving jet on the southwestern edge of Heceta Bank at $\sim 44^\circ N$ $125.25^\circ W$ (Barth et al., 2005b). Although the drifter did not return onto the shelf, it traveled >100 km shoreward from the deep ocean in <1 week. The disconnect between the surface and deep circulation indicates a potential way for zooplankton to return towards the shelf that was not available at depth.

Distributions of the small copepod *Oithona similis* (Fig. 10) seem to illustrate this pathway: abundances at the southern offshore edge of Heceta Bank were more similar to abundances in the northern offshore tip of the filament than to other locations within the bank region. It seems likely that the relatively high abundances at the edge of the Bank reflect a delivery of high numbers of individuals in near-surface circulation from offshore. Many of the dominant copepods in our study area reside just below the 10–20 m thick surface Ekman layer (Lamb and Peterson, 2005) and therefore would not have been able to exploit this return path. However, *O. similis* in our area typically occupy the upper 20 m Ekman layer (Lamb and Peterson, unpublished data) and so may have moved shoreward during a short-term wind reversal. Mackas and Galbraith (2002) noted that strong winds periodically move surface waters over eddies independently of the underlying geostrophic flow. Hence, taxa residing near-surface can be lost from geostrophic circulation features more easily than those beneath the Ekman layer.

It is not known whether the animals that do not return to the shelf contribute to secondary production in the offshore regions to which they have been moved. They may be subject to high predation over the continental slope and in the deep ocean where zooplanktivorous

euphausiids and fish reside. However, copepod egg production measurements made in upwelling filaments off California (Smith and Lane, 1991) and in the Canary Current (Hernández-León et al., 2002; Yebra et al., 2004) indicate that zooplankton may actively feed and reproduce in offshore expressions of features. In coastally generated eddies off Vancouver, Canada, coastal taxa are found offshore >15 months after separating from the shelf (Mackas and Galbraith, 2002). Hence, advected coastal populations may continue to contribute to offshore production for substantial lengths of time.

4.5. Trophic interactions

Biological relationships among the chlorophyll, zooplankton, and hydrography noted in this study corresponded to patterns exhibited by organisms in higher trophic levels. Reese and Brodeur (2006) identified Heceta Bank and the nearshore area around the tip of Cape Blanco as 'hot spots' of nekton biomass and species diversity. Abundances of juvenile coho and chinook salmon (Brodeur et al., 2004) and seabirds (Ainley et al., 2005) were also elevated in those regions. Many of the seabirds, particularly Shearwaters and Cassin's Auklets, were strongly related to frontal areas and locations of high chlorophyll biomass during this cruise. Humpback whales and harbor porpoises were strongly associated with areas of high chlorophyll biomass, whereas Dall's porpoises were found offshore around the edges of the filament (Tynan et al., 2005).

5. Conclusions

Mesoscale circulation features can advect huge volumes of coastal water offshore (Hormazabal et al., 2004; Kosro and Huyer, 1986; Mackas and Yelland, 1999). A single filament may advect as much water offshore as all of the Ekman transport along 1000 km of coastline (Kosro and Huyer, 1986). Several filaments simultaneously occur along the Oregon and California coast each summer. Their net offshore transport in coastal upwelling systems is clearly indicated by the dominance of neritic taxa within them (e.g. Haury, 1984; Mackas et al., 1991; Rodríguez et al., 1999). Here, we measured offshore transport in a meander of the upwelling jet as ~ 0.5 Sv, and a biomass flux of >900 tons of carbon per day in zooplankton alone. Coastal taxa, which are typically most abundant over the continental shelf (e.g. *Calanus marshallae*, *Acartia longiremis*, *Pseudocalanus mimus*), were abundant >100 km offshore in the filament, whereas warm-water, oceanic taxa were less abundant there.

Both the timing and intensity of mesoscale circulation (Keister and Strub, 2008) and peak nearshore zooplankton biomass (Peterson and Keister, 2002; Peterson and Miller, 1975) vary spatially and interannually in the NCC, so the biomass delivered to the deep sea, and hence the productivity of the ecosystem as a whole, are also likely to vary. Eastern boundary currents are highly productive in terms of primary and secondary production, so variability in the transfer of water and the associated

planktonic communities between the shelf and deep ocean may have dramatic consequences for ecosystem dynamics and global carbon cycling, survival of oceanic zooplanktivores (e.g. Logerwell et al., 2001; Yen et al., 2006), and upper trophic level organisms (Palacios et al., 2006).

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