**ABSTRACT**

The Japan/East Sea (JES) contains several oceanic regions separated by dynamic boundaries. These distinct regions and the physical feature that establish and maintain the boundaries between the regions, have significant impacts on its ocean biology. Until recently, most studies of the biology of the JES have focused on nearshore regions, with few detailed studies of the interior of the JES or they dynamic features that define the different regions. In addition, the classic sampling methods used in previous work have not allowed high-resolution studies of biological-physical interactions associated with key dynamic mesoscale frontal zones, quasi-synoptic surveys of water column and biological structure in three dimensions, or broad-scale description of the seasonal cycles in the different biogeographic regions of the JES.

**SUBJECT TERMS**

JES, Japan/East Sea, dynamic boundaries, mesoscale, sea-surface temperature
BIological STrUCTURE AND SEASONALITY IN THE JAPAN/EAST SEA

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The Japan/East Sea (JES) contains several oceanic regions separated by dynamic boundaries. These distinct regions, and the physical features that establish and maintain the boundaries between the regions, have significant impacts on its ocean biology. Until recently, most studies of the biology of the JES have focused on nearshore regions, with few detailed studies of the interior of the JES or the dynamic features that define the different regions. In addition, the classic sampling methods used in previous work have not allowed high-resolution studies of biological-physical interactions associated with key dynamic mesoscale frontal zones, quasi-synoptic surveys of water column and biological structure in three dimensions, or broad-scale description of the seasonal cycles in the different biogeographic regions of the JES.

A UNIQUE BASIN

The central basin of the JES contains two primary biogeographic regions, defined by the dominant currents and hydrographic structures (Figures 1 and 2). The region is bisected by the dramatic the Subpolar Front (SPF) and its associated mesoscale circulation. The SPF persists between the cold water in the north and the warmer subtropical water in the south and flows from west to east from the Korean Peninsula to Tsugaru Strait (e.g., Isoda et al., 1991; Isoda, 1994; Preller and Hogan, 1998; Talley et al., 2004; Lee and Niiler, 2005). The Tsushima Current brings warm,
Figure 1. Geographic features and currents of the Japan/East Sea with locations of the Video Plankton Recorder (VPR) (July 4–7, 1999, green line) and SeaSoar (May 22, 1999, blue line; January 27, 2000 cyan line) data and of the net tows (black-filled circles) conducted during the July VPR cruise shown (EKWC=East Korean Warm Current, NKCC=North Korean Cold Current, SPF=Subpolar Front, TWC=Tsushima Warm Current).
tropical water into the JES through Tsushima Strait to the south and flows north along the western coast of Japan (e.g., Preller and Hogan, 1998; Hase et al., 1999; Talley et al., 2004). A branch of the Tsushima Current flows north along east coast of the Korean Peninsula as the East Korean Warm Current (EKWC) (e.g., Tomczak and Godfrey, 1994; Hase et al., 1999). The North Korean Cold Current (NKCC) flows south along the Russian coast and the northern Korean Peninsula, intersecting the EKWC at the inception of the SPF (e.g., Huh and Shim, 1987). In winter, deep low-pressure systems draw cold dry Siberian air masses over the northern JES, forming cold dense waters that result in deep convective mixing (e.g., Talley et al., 2003; this issue; Dorman et al., 2004; Khelif et al., 2005; Lee et al., this issue). To the north of the SPF, the JES follows a distinct temperate-sub-polar seasonal cycle for water temperature and associated biological characteristics (e.g., primary production, copepod species, phytoplankton type) (Figure 3). By contrast, the region to the south of the SPF is much warmer, has considerably less seasonality and is characterized by more tropical biological characteristics and species (Meshcheryakova, 1960; Nishimura, 1969; Morioka et al., 1977; Miller and Terazaki, 1989; Park et al., 1991; Hirakawa et al., 1995).

Because of its relatively small size, this semi-enclosed basin provides access over short distances to features typical of much larger ocean basins and thus is an ideal model system for intensive process studies to understand ocean dy-
Zonally averaged SST: Sep 97–Jul 99

Zonally averaged CHL: Sep 97–Jul 99

*Figure 3. Seasonal progression of satellite-derived sea surface temperature (left) and chlorophyll (right) from October 1997 to July 1999, from weekly composites, extending from the southern to northern regions and including the 1999 summer field season of the JES program. The spring bloom (A) and the fall bloom (B) are shown for 1998 and 1999.*

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April 1999 (Figure 2). The SPF separated the warmer southern basin from the cold northern waters, with an ~ 7°C temperature gradient between the two regions. Considerable mesoscale variability in the path and structure of the SPF (eddies, filaments) also is evident.

Complex patterns in biology in April 1999 also were associated with SST or physical structures. High chlorophyll (~ 2 mg m⁻³) on the western boundary was associated with coastal upwelling in the EKWC; this elevated chlorophyll propagated northward in the EKWC and then eastward where the flow separates from the coast at ~ 40°N in a series of undulating rings and eddies as part of the SPF. A quasi-permanent eddy located south of where the EKWC and the East Korea Warm Current (EKCC) merge near 40°N (Kim et al., 1991, 1996; Shin et al., 1995; Suh et al., 1999; Ashjian et al., 2005; Mitchell et al., 2005) also may be associated with elevated chlorophyll.

By contrast, much lower chlorophyll (< 0.3 mg m⁻³) was observed in the eastern portion of the southern basin off the Japan coast, where the oligotrophic Tsushima Current moves northward and warming began in the south in March but later (May-June) in northern regions. Maximum temperatures occurred in all regions in September.

Phytoplankton blooms developed in both spring and fall of both years (Figure 3). The spring phytoplankton bloom began in the southern basin in March and propagated northward, reaching peak chlorophyll (~ 5 mg m⁻³) at ~ 47°N in late May. The spring bloom was followed by a rapid decay during the summer. A particularly strong (> 1 mg chl m⁻³) phytoplankton bloom occurred at the SPF (~ 42°N), perhaps associated with the complex mesoscale structure of the front (anticyclonic eddies, filaments) (Figure 2). The autumn phytoplankton bloom began in November and persisted until January. This autumn bloom was observed each year, but the strength and timing varied spatially and temporally. The bloom was quite strong (>1 mg chl m⁻³) in the northern basin, but considerably lower in the southern basin.

VARIATIONS IN PRIMARY PRODUCTION
Spatial and temporal variations in primary production were further explored by considering satellite-derived chlorophyll in different regions over a ten-year period (1996–2006). The analysis was based on a 10-year time series of satellite-derived chlorophyll from the Ocean Color and Temperature Sensor (OCTS) and Sea-viewing Wide Field-of-view Sensor (SeaWiFS) satellites for which monthly means were calculated. Monthly anomalies from the mean then were constructed by calculating the ratio of the monthly mean chlorophyll for each month in the time series relative to the long-term mean for that month. The major regions in the basin then were identified using principal components and empirical orthogonal functions based on the 10-year series of chlorophyll anomalies. The first principal component objectively separated the north-
ern and southern regions and showed the frontal area in between (Figure 4). Based on this objective partitioning, four characteristic spatial domains were defined for which large-scale interannual variations in various biological parameters were explored.

Net primary production (NPP) by phytoplankton for each of the four spatial domains over the 10 years was estimated with an empirical model (Behrenfeld and Falkowski, 1997) using corresponding monthly satellite-derived chlorophyll, photosynthetically available radiation (PAR), and SST. The methods of chlorophyll and NPP time-series analysis within discrete geographic domains were described in Kahru and Mitchell (2000, 2002).

Time series of NPP (Figure 5) show distinct maxima during the spring bloom in April–May in all four domains. The fall bloom is smaller in magnitude but is also a recurrent feature in both the transition (2) and the southern (3) domains. In the far north domain (4), the spring bloom lasts longer and may merge with the fall bloom. Note also that in 2003 the fall bloom in this region was of greater magnitude (i.e., NPP) than the spring bloom. Primary production in northern regions (1, 4) usually exceeded that observed in the southern region (3). Seasonality in NPP in the southern region also was less than in other regions, with sustained levels of moderate NPP throughout the year. Interannual variability in the magnitude of the blooms was observed, however, a marked feature for all four areas was the especially low primary production during the El Niño year of 1997.

Previous studies (Yamada et al., 2005; Yamada and Ishizaka, 2006) have shown that the timing of the spring bloom depends on the strength of wind mixing and the vertical stratification. With relatively weak winds and a shallow upper mixed layer, the spring bloom starts and peaks early but the total primary production is relatively low. By contrast, in years with strong winter mixing and a deep thermocline, the spring bloom starts later but total primary production tends to be higher. Such mechanisms should drive the annual variations observed here.

**FLUORESCENCE SEASONALITY ACROSS THE SPF**

The role of the SPF in structuring the biogeography of the JES basin is clear in the broad-scale analyses of temperature,
fluorescence, and primary production. These strong gradients are maintained on finer spatial scales at the SPF where the northern and southern water masses juxtapose. High-resolution sampling of the upper ocean was used to describe the anatomy of biological-physical associations at this dynamic boundary.

The cross-frontal structure of the SPF and its impact on the distribution and magnitude of chlorophyll fluorescence was described in early summer (May 1999) and winter (January 2000) using a SeaSoar towed vehicle. The SeaSoar was equipped with a conductivity-temperature-depth sensor (CTD), fluorometer, photosynthetically active radiation (PAR) sensor, multi-wavelength optical backscatter sensor, and multi-wavelength absorption/attenuation meter. Because the instrument sampled in the upper 250 m of the water column, subsurface detail was revealed that could not be resolved by satellite-sensed ocean color.

The front was clearly evident in temperature, salinity, and density during both the spring and strongly wind-forced winter conditions (Figure 6). Phytoplankton biomass was enhanced at the front during both seasons. During the spring (May 1999) survey, the SPF was centered near 39.5°–40°N, as it had been in April 1999 (Figure 2), and was most clearly delineated by salinity. Chlorophyll concentration and suspended particulate material, indicated by optical backscatter (bb_ssp), were both elevated north of the front, at about 25-m depth, relative to south of the front where concentrations were reduced and chlorophyll and particles were deeper (40 m). We observed dramatic subduction along the

Figure 5. Time series of net primary production in the four domains defined in Figure 4. Panel 2 shows the Northern Oscillation Index, an El Niño index adapted for the North Pacific (Schwing et al., 2002). El Niño periods are shown as black-masked areas. The year label is centered at January 1.
Figure 6. Cross-frontal SeaSoar sections for May 22, 1999 and January 27, 2000 of temperature, salinity, density (black contour lines over temperature), chlorophyll fluorescence, and backscatter at 488 nm (white contour lines over chlorophyll, May 1999) or beam attenuation coefficient at 660 nm (black contour lines over chlorophyll, January 2000). May 1999 section from along 134°E longitude and January 2000 section from along 134.23°E longitude. Subduction of water from the northern side of the front is clearly seen where chlorophyll extends downward from the surface.
Figure 7. Distribution of large copepods, small copepods, and fluorescence along a transect crossing the SPF on July 4–7, 1999 (Figure 1). Temperature isotherms are overlain as cyan lines. Periods of day and night are indicated by black and white bars across the top of each panel. Large copepods were quantified using the low-magnification camera of the Video Plankton Recorder, while small copepods were quantified using the high-magnification camera.

27 kg m$^{-3}$ isopycnal of water, chlorophyll biomass, and particles from north of the front to a depth of nearly 200 m, below a layer of warmer, saltier water. The low salinity and elevated oxygen concentration (not shown) along the 27 kg m$^{-3}$ isopycnal indicated that the source of the subducted water was cooler, fresher surface water from the north of the front. The deepening of this water may have occurred simply by overlaying it with warmer, saltier, less-dense Tsushima Current water advecting from the south or by active southward, downward transport. Regardless of the mechanism, the process resulted in transportation of primary production from the surface layer to nearly 200 m, well below the euphotic zone.

During the winter (January 2000), increased chlorophyll again was observed at the SPF (Figure 6). This increased biomass, 0.75–1 mg m$^{-3}$, was associated with enhanced stratification in the frontal region. As for May 1999, downward transport of chlorophyll along the frontal boundary occurred, although not as dramatically. Unlike May 1999 where the mechanism of downward transport was ambiguous, vertical circulation during the winter was driven by cold, downfront winds that resulted in subduction on the dense side of the front and upwelling on the frontal interface (Thomas and Lee, 2005). High-chlorophyll water was transported from the upper 50 m to a maximum depth of only ~100 m, in contrast to the subduction to ~250 m observed during May. Although chlorophyll at the front was elevated (0.75–1 mg m$^{-3}$), primary productivity was limited overall by lower light and increased vertical mixing.
ZOOPLANKTON DISTRIBUTIONS AND THE SPF

Different zooplankton communities also juxtapose at the dynamic SPF. Zooplankton species compositions of the northern and southern regions of the JES are distinct and associated with particular water-mass types (e.g., Meshcheryakova, 1960). The northern JES contains larger temperate and sub-polar colder water species, such as the large copepod species Neocalanus plumchrus (see references in Ashjian et al., 2005), while the southern JES contains smaller, more tropical species and genera such as the small copepod Oithona spp.

High-resolution sampling of zooplankton taxa distributions and abundance in association with hydrography in the southern JES, the SPF, and the northern JES just to the north of the SPF were conducted using a Video Plankton Recorder (VPR) during July 1999 (Ashjian et al., 2005, 2006) to describe how the SPF structures and separates these distinct communities (Figure 7). The VPR is essentially an underwater video microscope that images plankton and measures coincident environmental variables (e.g., temperature, salinity, fluorescence) at high temporal, and hence, spatial resolution (Davis et al., 1992a, b, 1996, 2004; Tang et al., 1998; Hu and Davis, 2005, 2006). For this study, the VPR was mounted on a V-fin that was towed behind the ship at 8-10 knots, surveying from the near-surface to ~80 m. In-focus plankton images were detected, measured, identified, and counted automatically; these counts then were divided by the calibrated image volume and merged with environmental variables by time to yield concentrations of plankton (# L⁻¹) at specific depths. The JES study was one of the first VPR surveys to be at least in part analyzed in near-real time on board the ship. The VPR was equipped with both high-resolution and low-magnification cameras that imaged 3 ml and 29.6 ml, respectively. Because of the different image volumes, each camera effectively imaged and quantified different sizes of plankton and yielded separate distributions of large and small copepods.

Clear segregation of habitat by large and small copepods was seen across a summertime transect that extended from just to the north of the SPF to near the coast of Japan in the southern JES (Figure 1). Large copepods were most abundant just to the north of the SPF located at 150-km along-track distance (defined according to T-S characteristics [Ashjian et al., 2005, 2006] and seen here as the subducting thermocline), while small copepods were most abundant just to the south of the front. In the vertical, large copepods were found below the upper water column thermocline in water of <10°C, while the smaller copepods were confined primarily to warmer water (>12°C) above the thermocline. Both large and small copepods demonstrated some evidence of diel vertical migration; the range of migration of the smaller copepods was defined by the depth of the thermocline so that a shallower day distribution (20 m) was observed to the north of the front (~100-120-km distance) where the thermocline was shallower, and deeper day distribution (50 m) was observed to the south of the front (300-km distance). For large copepods, the diel signal was very subtle, manifested only as an increase in abundance above the thermocline during the night at ~200-km distance. Large copepods may have migrated to the near-surface at night to the north of the SPF (0-50-km distance); however, differentiation of that behavior from the association of large copepods with the thermocline is difficult. Elevated relative fluorescence was observed in the thermocline, with a peak located at the SPF consistent with what was observed using the SeaSoar (Figure 6). Both large and small copepods were associated with the fluorescence peak during the day while the small copepods migrated to the near-surface during the night, presumably to... statistical analyses... yielded insight into seasonal cycles, regional variability, biogeography, and the biological response to physical... forcing not achieved previously.

Utilize an alternative food supply (e.g., microzooplankton, small particles).

Examination of net tows taken in the different water types across the transect showed that to the north of the...
...this suite of observations and this approach represent a substantial contribution towards understanding biological processes in highly dynamic ocean regions.

CONCLUSION
The high-resolution, nearly synoptic description of biological distributions and processes, and their association with physical features, presented here is representative of the exciting present and future direction of oceanography. Although significant understanding of the broad-scale distributions of hydrography, chlorophyll, primary production, and plankton were achieved in earlier studies of the JES, these conclusions were based on climatology, on widely spaced and temporally disparate measurements, and on low-resolution or integrated vertical sampling. By contrast, the field

sampling methods used in the ONR JES Program were of sufficiently high horizontal and vertical spatial resolution and synopticity to permit not only the identification of biological-physical as-

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