

Influence of ocean winds on the pelagic ecosystem in upwelling regions

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Upwelling of nutrient-rich, subsurface water sustains high productivity in the ocean's eastern boundary currents. These ecosystems support a rate of fish harvest nearly 100 times the global mean and account for >20% of the world's marine fish catch. Environmental variability is thought to be the major cause of the decadal-scale biomass fluctuations characteristic of fish populations in these regions, but the mechanisms relating atmospheric physics to fish production remain unexplained. Two atmospheric conditions induce different types of upwelling in these ecosystems: coastal, alongshore wind stress, resulting in rapid upwelling (with high vertical velocity, w); and wind-stress curl, resulting in slower upwelling (low w). We show that the level of wind-stress curl has increased and that production of Pacific sardine (*Sardinops sagax*) varies with wind-stress curl over the past six decades. The extent of isopycnal shoaling, nutricline depth, and chlorophyll concentration in the upper ocean also correlate positively with wind-stress curl. The size structure of plankton assemblages is related to the rate of wind-forced upwelling, and sardine feed efficiently on small plankters generated by slow upwelling. Upwelling rate is a fundamental determinant of the biological structure and production in coastal pelagic ecosystems, and future changes in the magnitude and spatial gradient of wind stress may have important and differing effects on these ecosystems. Understanding of the biological mechanisms relating fisheries production to environmental variability is essential for wise management of marine resources under a changing climate.

California Current | sardine | wind-stress curl

Worldwide, populations of sardine (*Sardinops* spp.) and anchovy (*Engraulis* spp.) have varied greatly over time, with plentiful catches of one often alternating with the other on the scale of decades (1). These fluctuations have severe consequences for the fishing, processing, and farming (e.g., poultry, swine, and tuna) industries that depend on the fisheries' landings for income and feed. Because these small pelagic fish often dominate the intermediate trophic level in upwelling ecosystems, their populations are crucial to the transfer of energy and biomass from lower to higher trophic levels (2). Despite >50 years of effort and focused oceanographic research, a mechanistic explanation for the large variability in Pacific sardine and northern anchovy populations in the California Current Ecosystem (CCE) remains obscure. Sediment records suggest that similar fluctuations occurred over the two millennia before the development of commercial fisheries (3). The changes in sardine and anchovy abundance during the past century are therefore thought to reflect natural environmental variability, exacerbated by fishing pressure (4).

Understanding of the mechanisms relating these population fluctuations to environmental variability has not progressed past empirical observations associating sardine and anchovy biomass with temperature (1, 5). Warm periods, favorable for sardine production, occurred most recently from the 1920s to the mid-1940s and from 1977 through the present. A cool period, favorable for anchovy, occurred from the mid-1940s through 1976 (1, 5). A shift to another cool period may have occurred after the 1997–1998 El Niño (6). Identification of warm and cold

periods favorable for sardine and anchovy production has been important in describing multidecadal changes in pelagic fisheries, but this description does not provide the understanding necessary to predict how populations will vary under future conditions. Because cold periods are often associated with increased coastal upwelling and nutrient supply along the coast, the growth observed in the anchovy population during these phases is expected. However, the paradoxical growth of a massive sardine population (up to four times larger than the maximal anchovy population) during warm periods with weak coastal upwelling has puzzled fisheries oceanographers for decades (3, 7).

To investigate the relationship between climate and sardine growth in the CCE, we considered the two mechanisms by which winds supply surface waters with the nutrients required for biological production: “coastal upwelling” due to alongshore wind stress and “curl-driven upwelling” due to wind-stress curl. The importance of coastal upwelling to major fisheries production has long been recognized (8). In the traditional view of coastal upwelling ecosystems, biological productivity at all levels of the food web is attributed to persistent, alongshore, and equatorward wind stress in spring and summer. These winds force water away from the coastal boundary, a process known as Ekman transport. Nutrient-rich waters are drawn up into the euphotic zone to replace the surface waters that are forced offshore.

Curl-driven upwelling may also act as a significant source of nutrients in coastal pelagic ecosystems (9–13) and is responsible for the shoaling of isopycnals in the southern CCE during summer (14). Horizontal shear in the wind stress (wind-stress curl) over the open ocean results in a divergence of Ekman transport that is balanced by vertical transport. Although most areas of the subtropical ocean gyres are regions of anticyclonic wind-stress curl (downwelling), the eastern edge of the gyres (i.e., inshore of the wind-stress maximum) are zones of cyclonic wind-stress curl and upwelling (10, 12). Small areas of intense, positive wind-stress curl occur in the lee of major headlands leading to curl-driven upwelling with vertical velocity (w) comparable with the high w associated with coastal upwelling (13). However, the average w associated with curl-driven upwelling is slow, typically several times smaller than w associated with coastal upwelling in the CCE (15). The expansive regions of positive curl over the open ocean are the result of two characteristics of the eastern North Pacific: (i) the large-scale change in the orientation of the west coast of North America, from a coastline trending north to south poleward of 40° N latitude to a coastline oriented northwest to southeast in southern California; and (ii) the location of the maximal gradient between the

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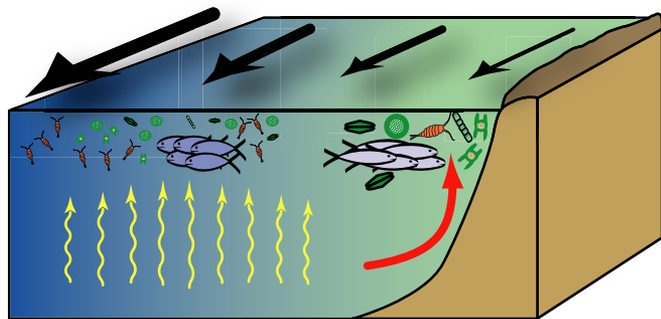


Fig. 1. Conceptual diagram displaying the hypothesized relationship between wind-forced upwelling and the pelagic ecosystem. Alongshore, equatorward wind stress results in coastal upwelling (red arrow), supporting production of large phytoplankters and zooplankters. Between the coast and the wind-stress maximum, cyclonic wind-stress curl results in curl-driven upwelling (yellow arrows) and production of smaller plankters. Anchovy (gray fish symbols) prey on large plankters, whereas sardine (blue fish symbols) specialize on small plankters. Black arrows represent winds at the ocean surface, and their widths are representative of wind magnitude.

pressure systems over the North Pacific and southwest United States (16, 17). Because these areas of positive curl are large compared with the coastal boundary region, the volume of water upwelled by slow, curl-driven upwelling is greater than that upwelled by coastal upwelling or by fast, curl-driven upwelling near the coast (15). The intensity of offshore, curl-driven upwelling is hypothesized to be related to the location of the large-scale pressure systems with respect to the coastline of California (17).

We expect the type of biological production resulting from coastal and curl-driven upwelling to differ, with high w resulting in larger phytoplankters and low w favoring smaller phytoplankters. The demand for nutrients by a phytoplankton cell is typically a function of cell volume, whereas the maximal uptake rate is a function of the cell's surface area. For this reason, smaller cells, with higher surface-area-to-volume ratios, have a competitive advantage in nutrient-limited environments (18, 19). The increased nutrient concentrations in vigorously upwelling waters (high w) reduces nutrient limitation and the competitive advantage of small cells, allowing populations of large cells with lower surface area-to-volume ratios to develop. Given that prey size correlates positively with predator size (20), larger zooplankters are favored in areas with larger phytoplankters and higher w (Fig. 1).

Temporal variability in coastal and curl-driven upwelling may affect populations of planktivorous predators by influencing production of small and large plankters. Pacific sardine spawn in offshore waters, away from areas of coastal upwelling (21), and adult and juvenile sardine have a fine mesh of gill rakers with specialized denticles to retain planktonic prey as small as 10 μm in diameter (22). Even as larvae, sardine appear to specialize on small plankters and are prevented from capturing larger prey by a small mouth-gape diameter (22, 23). In comparison, anchovy spawn near the coast (21) and use coarse gill rakers to capture larger prey (22). We hypothesize that changes in sardine population growth are related to the production of small plankters and the magnitude of curl-driven upwelling in the CCE. To test this hypothesis, we posed the following questions: Is zooplankter size related to upwelling rate? Have winds favoring curl-driven and coastal upwelling changed over decades and, if so, have these changes influenced hydrography, nutrient supply, and biological production? We examined zooplankter sizes across a gradient of upwelling rates and compared estimates of historical upwelling

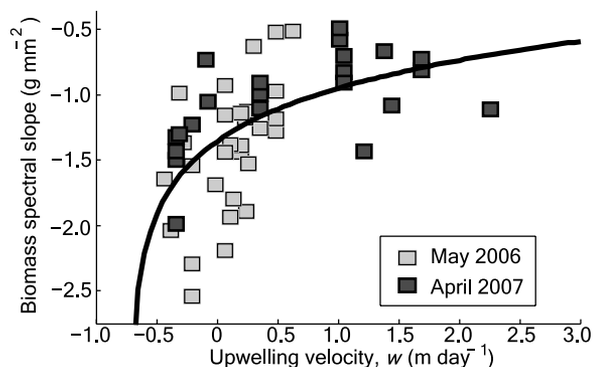


Fig. 2. Relationship between zooplankter size and upwelling rate. This relationship is modeled by using a logarithmic function ($r^2 = 0.32$, $P < 0.001$, $n = 51$, $y = 0.46 \ln(x + 0.71) - 1.20$). The mean 90% confidence interval around the slopes of the linear least-squares fits to the biomass spectra is $\pm 0.71 \text{ g mm}^{-2}$.

with concurrent measurements of water-column density, nutrient depth, chlorophyll concentration, and sardine production.

Results and Discussion

Plankter Sizes and Upwelling. A combination of data from the SeaWinds Scatterometer and shipboard and moored anemometers were used to compare w resulting from upwelling with the size of zooplankters. Zooplankton was collected during two cruises in May 2006 and April 2007 at locations spanning the CCE west of Point Conception, CA [supporting information (SI) Fig. 6]. A normalized biomass spectrum was estimated for the zooplankton collected at each station (24). The slope of the spectrum increased as w increased (Fig. 2), indicating that zooplankters are relatively larger in areas of coastal upwelling (high w) and smaller in areas of curl-driven upwelling (low w). A similar result has been found for phytoplankton in the CCE; larger phytoplankters are found nearshore where the nutricline is shallow, and smaller phytoplankters are dominant where the nutricline is deep (25).

Historic Upwelling Rates. To examine the temporal variability in coastal and curl-driven upwelling rates over the past 60 years, we used monthly averages of historic winds to calculate w of curl-driven and coastal upwelling. The geographic range of the analysis was limited to the waters off the southern and central California coasts (from Ensenada, Baja California, to Santa Cruz, CA), extending $\approx 300 \text{ km}$ offshore and encompassing the area of sardine spawning off of California (SI Fig. 6). We found that regions of intense cyclonic curl are common during summer in the lee of prominent headlands and result in small areas of high w (3–7 m day^{-1}). High rates of coastal upwelling were also present at these headlands (w of 7–12 m day^{-1}). Further offshore, large regions of positive wind-stress curl and low w (0–1 m day^{-1}) were typical (Fig. 3). These results are consistent with previous studies that have examined coastal and curl-driven upwelling over smaller temporal and spatial scales (11, 26).

Although average w resulting from coastal upwelling is approximately an order of magnitude larger than w resulting from open-ocean, curl-driven upwelling, curl is more important to total upwelling transport because it covers a spatial area 18–22 times larger than the area of coastal upwelling. In our analysis, we found that wind-stress curl has been responsible for at least 60% (and up to 80%) of the annual, wind-forced upwelling transport in the southern CCE. Monthly time series of the two upwelling processes are significantly correlated ($P < 0.001$) and indicate that both coastal and curl-driven upwelling have increased since 1948. However, there are important distinctions

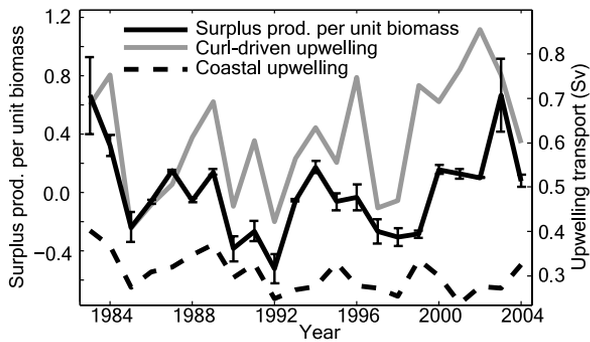


Fig. 5. Upwelling and surplus production per unit biomass of Pacific sardine. Curl-driven upwelling from May through July showed the highest correlation with surplus production per unit biomass ($r = 0.62$, $P < 0.005$, $n = 22$). Coastal upwelling during the same period was not significantly correlated with production ($r = 0.40$, $P = 0.067$, $n = 22$). Error bars are \pm one standard deviation of the sardine production estimates.

0.05). Curl-driven and coastal upwelling records were not correlated over this time period (Fig. 5).

In addition, we compared the influence of coastal upwelling, curl-driven upwelling, and sea-surface temperature (SST) on sardine production using environmentally dependent surplus production (EDSP) models during two periods for which environmental and fisheries data are available: 1948–1962 and 1983–2004. SST has been shown to be reasonably effective in explaining the dynamics of sardine biomass in EDSP models (31), and we were interested in testing whether SST or a measure of upwelling produced the best estimate of sardine production. We found that use of curl-driven upwelling in the model produced the best fit to observed production during both periods (SI Fig. 8). The model using SST as the environmental variable was more successful than that using coastal upwelling, and all three environmental variables performed better than the null model (which did not include environmental variability). The sum of squared deviations for the model using curl-driven upwelling was 0.30 MT^2 . Values for the models using SST and coastal upwelling were 0.44 MT^2 and 0.47 MT^2 , respectively. The null model resulted in a sum of squared deviations equal to 0.51 MT^2 .

Although use of curl-driven upwelling in the EDSP models produced the best estimates of sardine production during both time periods, the months during which curl produced the best estimate shifted between 1948–1962 and 1983–2004. In the more recent period, use of curl-driven upwelling during May–July was optimal for estimating production and suggests that conditions during these months were most influential in determining sardine production. Since 1983, extensive spawning has occurred off of central and southern California in April (32). May–July corresponds to the early life-history stages during which the environment has the strongest influence on survival (27). Use of curl-driven upwelling during October, November, and December produced the best model performance during the 1948–1962 period. This result is consistent with the observation that spawning in autumn offshore of Baja California is more important to sardine production during periods of low population size (32). The months of SST and coastal upwelling that produced the best estimates of production also shifted from late summer and autumn during the 1958–1962 period to spring and early summer during the 1983–2004 period.

Consideration of Other Upwelling Systems. The CCE is the only upwelling system with the environmental time series required to investigate wind-forced upwelling and the response of water-column properties and fisheries production at decadal and

multidecadal scales. Observations of historic, oceanic winds depend on ship traffic, and the number of observations in the CCE is high compared with other eastern boundary currents (16). Atmospheric models offering high-resolution estimates of wind stress over the past 60 years have not yet been developed in other regions of the globe. In addition, the long-term hydrographic datasets provided by the CalCOFI program are unique.

Wind-forced upwelling results in high primary and fisheries production in eastern boundary currents around the world (8), and the concepts presented here are applicable to each of these regions. However, the effects of coastal and curl-driven upwelling may vary with conditions specific to each area. For instance, latitudinal differences may have a significant influence on w and production of plankton and fish. The rate of curl-driven upwelling in the Humboldt Current Ecosystem (HCE) off the coast of Peru will be more than three times that in the CCE for a given wind-stress curl because of the difference in the Coriolis parameter with latitude. A wind-stress curl of $0.5 \cdot 10^{-6} \text{ N m}^{-3}$ at 35° N latitude in the CCE will result in w of 0.5 m day^{-1} . The same level of cyclonic wind-stress curl at 10° S latitude in the HCE will create w of 1.7 m day^{-1} . Much larger plankton sizes may result from curl-driven upwelling in the HCE. Anchovy feed most effectively on large plankton (22), and we would expect the HCE to be dominated by anchovy and support a larger anchovy population than the CCE under similar curl conditions. Consistent with this hypothesis, anchovy is the dominant fish species in Peru, whereas sardine is dominant in the CCE. Annual anchovy landings in Peru peaked at $>13 \text{ MT}$ in the mid 1970s, whereas maximal landings in California peaked at $\approx 0.4 \text{ MT}$ in the early 1980s (1).

Our results demonstrate a mechanism, from physics to fish, relating variability in production of Pacific sardine to environmental changes over interannual and decadal scales. We show that the level of production in a large, marine ecosystem depends on wind-stress curl. Coupling predictions of atmospheric winds with a simple hydrographic model will allow forecasting of sardine production in the CCE. Such forecasts have increasing ecological and economic value as globalization of commerce and industrialization of fisheries continue in response to growing demand and utilization of marine resources (33). Simultaneously, predictions of future climate conditions are becoming more precise (34) and offer an opportunity to more effectively manage fisheries if the biological responses to physical variability are understood. Credible mechanistic hypotheses relating atmospheric physics to variability in the ocean's biota are essential to prudently manage marine resources under a changing climate.

Materials and Methods

May 2006 and April 2007 Cruise Data. Zooplankton samples were collected during research cruises in May 2006 and June 2007 as part of the CCE Long-Term Ecological Research program. Cruises were structured to sample across the CCE, ranging from areas of coastal upwelling to offshore, oligotrophic areas. Zooplankton was sampled by using a BONGO net of $202\text{-}\mu\text{m}$ -Nitex mesh, towed obliquely to 210 m (depth permitting) between 2100 and 0400 hours following the strict protocol of the CalCOFI program (35). Three-eighths of the sample from one BONGO codend was wet-sieved through nested screens of 5,000, 2,000, 1,000, 505, and $202 \mu\text{m}$, and dry masses in each size class were determined (36). A linear least-squares line was fit to approximate the biomass-size spectrum for each sample according to the following formula:

$$\log \frac{B_x}{\Delta x} = m[\log(x)] + b,$$

where B_x is the sample biomass retained on a filter of mesh size x , Δx is the size interval for each fraction (taken here as 5,000, 3,000, 1,000, 495, and $303 \mu\text{m}$), and m and b are the slope and y -intercept of the linear, best-fit line.

Wind-stress data collected by the SeaWinds Scatterometer were used to estimate w . For two sampling stations near the coast where scatterometer

performance were determined by minimizing the sum of squared deviations between the modeled and observed ASP.

In the stepwise regression procedure, the environmental time series explaining most of the variability in surplus production per unit biomass was included in as the first explanatory time series in the model. Additional environmental time series were included only if inclusion significantly improved model fit ($P < 0.05$).

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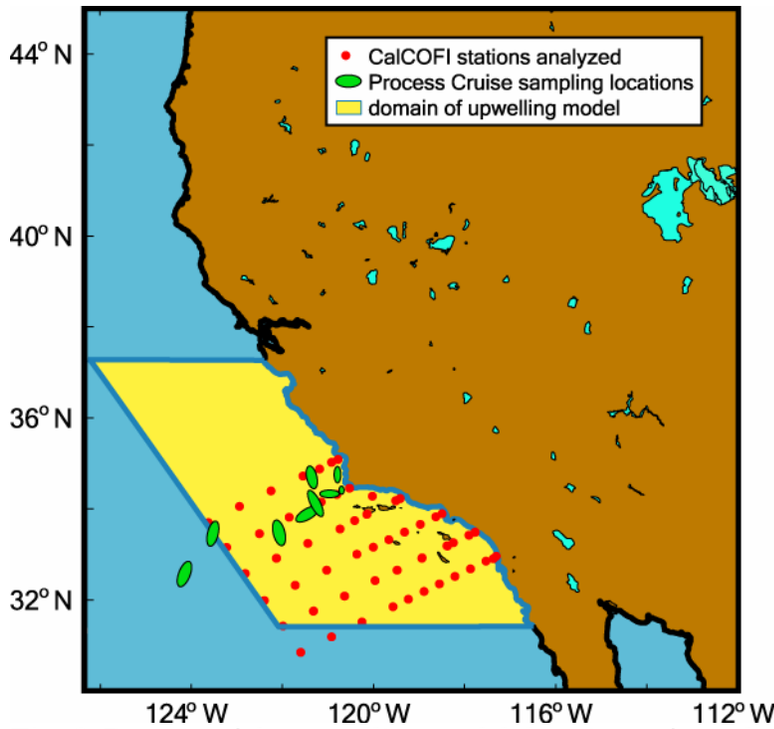


Fig. 6. Domain of the upwelling model, locations of zooplankton collections, and CalCOFI stations. The domain of the upwelling model is from Ensenada, Baja California to Santa Cruz, CA and extends ≈ 300 km offshore. Each process cruise station was occupied for 3-5 days, and zooplankton was collected between dusk and dawn.

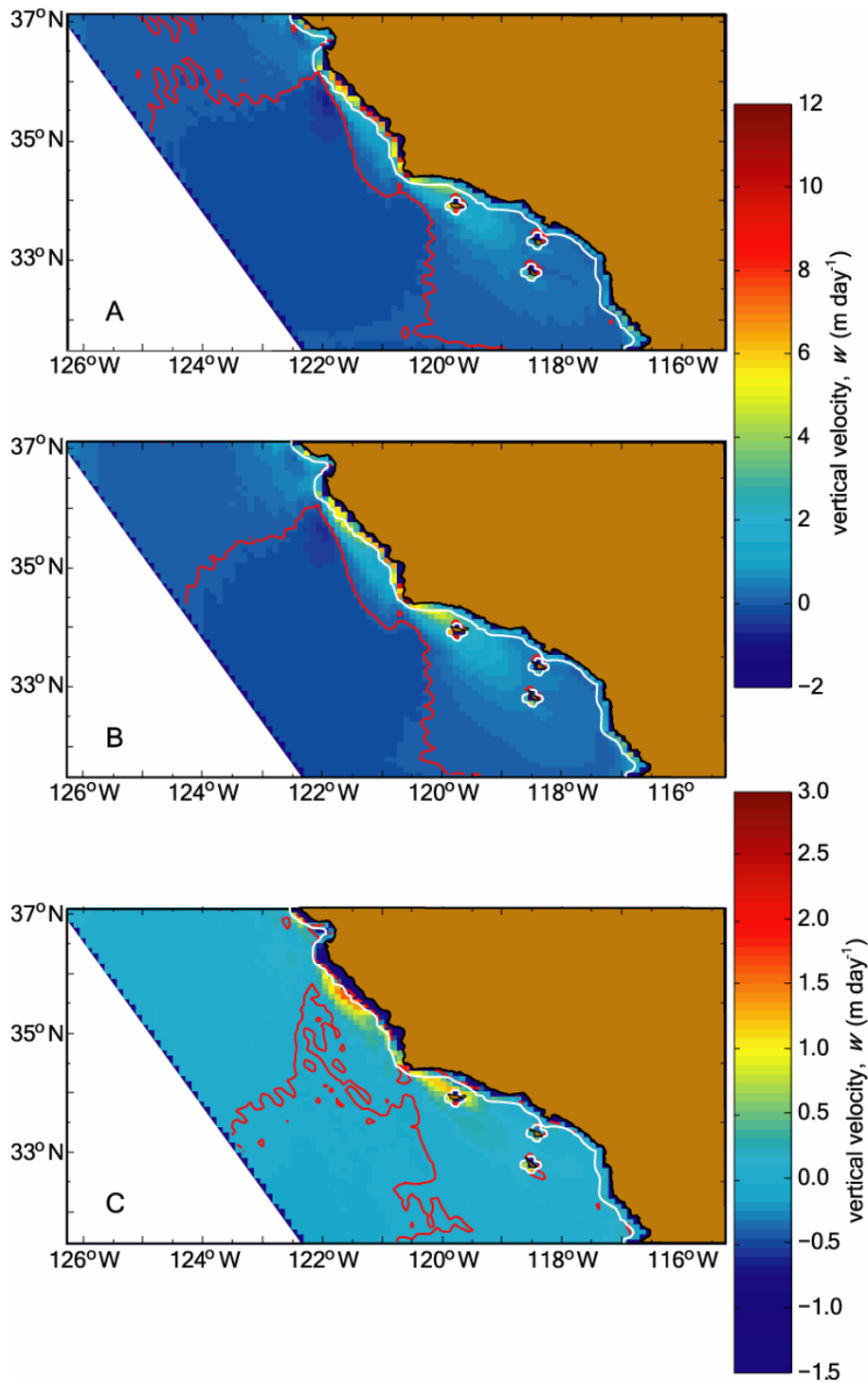


Fig. 7. Upwelling in the CCE. Wind-stress curl and coastal, alongshore winds were used to estimate mean w resulting from coastal upwelling and curl-driven upwelling in May, June, and July during 1950-1959 (A), 1990-1999 (B), and the difference in upwelling between the two periods (1990-1999 minus 1950-1959, C). The zero contour is noted by the red line. The white line along the coast separates the domains of coastal upwelling and curl-driven upwelling. Note the increase in curl-driven upwelling and the decrease in coastal upwelling in the 1990s relative to the 1950s.

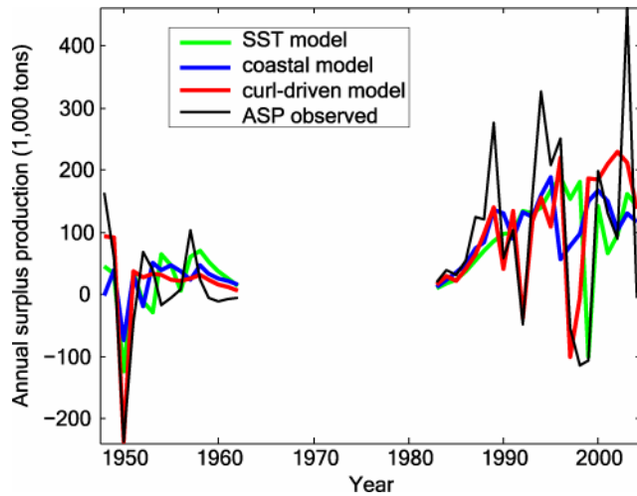


Fig. 8. Observed and modeled surplus production in the Pacific sardine population for two periods (1948-1962 and 1983-2004). The surplus production model incorporating curl-driven upwelling produced the best fit to observed production during both periods of stock assessment. Coastal upwelling and mean SST were less successful at estimating surplus production.