Title
Foraging at a front: Hydrography, zooplankton, and avian planktivory in the northern Bering Sea

Permalink
https://escholarship.org/uc/item/5zq693n9

Journal
Marine Ecology Progress Series, 182

ISSN
0171-8630

Authors
Russell, RW
Harrison, NM
Hunt, GL

Publication Date
1999-06-11

DOI
10.3354/meps182077

License
CC BY 4.0

Peer reviewed
Foraging at a front: hydrography, zooplankton, and avian planktivory in the northern Bering Sea

Robert W. Russell*, Nancy M. Harrison**, George L. Hunt Jr

Department of Ecology and Evolutionary Biology, University of California, Irvine, California 92697-2525. USA

ABSTRACT: We studied hydrographic structure, zooplankton distributions, and foraging by planktivorous seabirds in the Anadyr Strait, northern Bering Sea, during 4 summer cruises (1984–1986, 1993). The western portion of the strait was occupied by cold, dense Anadyr water that was mixed from top to bottom. This mixed water was separated from the stratified Bering Shelf water on the eastern side of the strait by a sharp surface front (the 'Anadyr Front'). Net sampling indicated that calanoid copepods were the numerically dominant component of the zooplankton, and that densities of several species were elevated in the frontal zone, apparently due to mechanical accumulation resulting from surface convergence. Hydroacoustic surveys showed that overall zooplankton biomass was concentrated along the thermocline and at the front. Although the location of the Anadyr Front was highly variable over time scales as short as 1 d, large numbers of least auklets Aethia pusilla often flew 25 to 50 km from their breeding colonies to feed at the front. Diet samples indicated that the copepod Neocalanus plumchrus was the principal prey taken by least auklets both at the front and away from it, indicating that heavy use of the distant frontal habitat was due to the higher densities of their preferred prey (i.e. rather than absence of suitable prey species closer to shore). Whenever aggregations of least auklets were found away from the front, there was evidence that they were exploiting near-surface high-density patches of zooplankton, though the exact mechanisms responsible for the formation of such patches are unclear. In contrast to least auklets, crested auklets Aethia cristatella were usually found away from the front. In several cases, compact aggregations of crested auklets were located over acoustically observed epibenthic layers of zooplankton. Hydrographic data suggested that intense subsurface jets and/or upwelling along the eastern side of the strait might have increased the availability of the crested auklets' preferred euphausiid prey. Thus, spatial segregation of the 2 principal planktivores in Anadyr Strait likely arises because different physical mechanisms cause concentrations of preferred prey originating at different depths.

KEY WORDS: Front · Foraging · Bering Sea · Copepods · Neocalanus plumchrus · Least auklet · Aethia pusilla · Crested auklet · Aethia cristatella

INTRODUCTION

Oceanic fronts are complex fluid structures often characterized by sharp sea-surface gradients in density, temperature, and/or salinity. Frontal phenomena include, but are not limited to, boundaries between different water masses. Distributed throughout the world's oceans, fronts occur at multiple spatial scales and have highly variable kinematics and flow fields (Fedorov 1986). Fronts have been noted by marine scientists and recorded in the literature for over 2 centuries (e.g. Franklin 1786, Darwin 1845, Beebe 1926), but have not been studied in ecological detail until recently. A growing body of evidence indicates that oceanic fronts support high levels of biotic activity across a wide range of trophic levels (reviewed by Le Fèvre 1986, Olson et al. 1994). Frontal enrichment or concentration has been documented in a wide variety of marine plankton, including microbes (e.g. Floodgate et al. 1981, Fernandez et al. 1994), phytoplankton...
Fronts also appear to be important components of the marine environment of predators at higher trophic levels. Schooling fishes may aggregate in fronts to feed on the plankton concentrations (e.g. Herron et al. 1989), and also perhaps to exploit the sharp thermal gradients for thermoregulation (Brandt & Wadley 1981). Aggregations of large pelagic fishes such as tunas and swordfishes also have been documented in the vicinity of fronts (e.g. Dufour & Strelta 1973, Carey & Robison 1981, Fiedler & Bernard 1987, Podesta et al. 1993). The nature of this frontal association has been controversial (reviewed by Olson & Podesta 1987), because, although some fishes may feed at fronts (e.g. Fiedler & Bernard 1987), others may use fronts as orientation cues for migration (e.g. Laurs & Lynn 1977, Maul et al. 1984). Concentrations of pinnipeds and whales may also occur in the vicinity of fronts (e.g. Nasu 1974, Gaskin 1987, Brown & Winn 1989, Sinclair et al. 1994), but have been little studied.

Large seabird concentrations at oceanic fronts have often been documented in the literature (reviewed by Hunt & Schneider 1987, Schneider 1990, 1991), and these seem most straightforwardly interpreted as feeding aggregations. However, concentration of seabirds at fronts is often intermittent (e.g. Uda 1938, Hawke 1996). In a statistical analysis of the relationship between seabirds and fronts, Schneider et al. (1987) reported that, on average, seabird abundance was not significantly higher at fronts than in adjacent waters of the southeastern Bering Sea, though abundance was higher at strong fronts than at weaker fronts.

With local phytoplankton growth rates as high as 16 g C m⁻² d⁻¹ and estimated annual carbon production of ~500 g m⁻², the northwestern Bering Sea (Fig. 1) is one of the most productive marine ecosystems in the world (Sambrotto et al. 1984, McRoy et al. 1987, Springer & McRoy 1993). Together with this high primary productivity, the region is characterized by large standing stocks of zooplankton (Springer et al. 1989), an abundant benthic fauna (Highsmith & Coyle 1990), large numbers of transient marine mammals (Fay et al. 1984, Johnson & Nelson 1984, Nerini 1984, Nelson & Johnson 1987, Obst & Hunt 1990), and immense breeding colonies of seabirds on the area's islands (Fay & Cade 1959, Sowls et al. 1978). This biological richness has been attributed in part to the northward transport of nutrient-rich oceanic water from the continental slope into the northern Bering Sea (Hansell et al. 1989, 1993, Springer & McRoy 1993), which is caused by sea level differences between the Arctic Ocean and the Bering Sea (Stigebrandt 1984, Coachman 1993). The northward flow, known as the 'Anadyr Current', passes along the Siberian coast on the western side of the Bering Sea and ultimately through the Bering Strait (Fig. 1; Coachman et al. 1975, Kinder et al. 1986, Overland et al. 1996). As it flows to the north, it carries with it an entire zooplankton community, including species that are more characteristic of deep oceanic waters (e.g. Eucalanus bungii and Neocalanus spp.; Johnson 1963, Springer et al. 1989). Indeed, the northern Bering Sea ecosystem has been likened to a chemostat (Sambrotto et al. 1984) or a conveyor belt (Springer & Roseneau 1985).

The principal avian planktivores in the northern Bering Sea are auklets, of which the least auklet Aethia pusilla and crested auklet Aethia cristatella are the most abundant. Immense breeding colonies are located on St. Lawrence Island, King Island, and the Diomede Islands (Sowls et al. 1978). During the chick-rearing period, these birds fly to sea to forage and return to colonies carrying loads of zooplankton in their sublingual pouch. Previous studies that have examined auklet diets and foraging habitat selection have suggested that these birds are dependent on prey found principally in Anadyr water and adjacent Bering Shelf water influenced by the Anadyr Current (Bédard et al. 1979, Franks 1992a, Laubscher et al. 1993, Yoder et al. 1994), and zooplankton (e.g. Smith et al. 1986, Epifanio 1992).
precluded sampling west of the International Date Line, and all of our transects were terminated prior to a complete crossing of the strait.

Hydrographic data were collected by lowering a NeilBrown conductivity-temperature-depth (CTD) probe to within 5 m of the bottom while the ship was stopped at sampling stations along transects. Vertical and horizontal cross-sections of physical properties were contoured using the minimum curvature gridding algorithm in the Surfer Version 5.02 Surface Mapping System. For simplicity, we present only temperature contours here, as the distributions of temperature, salinity, and density were always very similar.

Subsurface distributions of zooplankton along the cruise transects were studied using a BioSonics echosounding system while the ship was underway between stations. The system was composed of a 200 kHz model 101 echosounder and a model 120 echointegrator. The receiver gain was set at 6 dB, the band width at 5 kHz, the absorption coefficient for 200 kHz, the range at 150 m, the transmitter on 0 dB, and the pulse width at 0.5 m. Before each transect was initiated, the system was calibrated with an internal signal. The transducer was towed beside the ship and outside the wake in a V-fin depressor. Sonic surveys were conducted at a speed of ~6 knots. Signals were blanked to 5 m below surface to reduce noise, and echoes were integrated vertically in 2 m bins from 5 m below surface to the bottom. Each integration consisted of 100 pings and examined approximately 200 m of horizontal distance. Incoming signals were corrected for time-varied gain, digitized, and transferred to a microcomputer for further processing and analysis. We did not attempt to convert echo return voltages into estimates of absolute biomass because of uncertainties about the target strengths of the zooplankton constituents. Contour plots of zooplankton distribution were created from volume scattering measurements using Surfer with minimum curvature gridding.

Zooplankton community composition and species abundance were sampled by conducting vertical tows from just above the bottom to the surface with a 1 m diameter, 505 μm mesh net at selected hydrographic stations. Zooplankton samples were preserved in 5% formaldehyde solution for later laboratory analysis. Identification of the taxa in these samples was completed before Neocalanus flemingeri was identified as a species distinct from Neocalanus plumchrus (Miller 1988), so densities of N. plumchrus reported in this paper may include an unknown proportion of the very similar N. flemingeri.

The distribution of auklets was studied by making continuous counts of birds on the water within 300 m of one side of the ship while the ship was underway. Many of our interpretations of observed auklet distrib-
utions rely on the assumption that the majority of birds have commuted from St. Lawrence Island. Unfortunately, little information is available on the distribution or abundance of auklets on the Russian side of Anadyr Strait. We cannot discount the possibility that some of the birds observed during our transects had commuted from Russian colonies. However, we routinely observed vast flights of auklets flying northwestward away from St. Lawrence Island toward the Anadyr Front and southeastward from the front toward the island, indicating at least that many of the birds were Alaskan in origin.

Diets of least auklets were examined by collecting birds from a small boat at selected locations using a 12-gauge shotgun. Contents of the stomach and gular pouch were removed immediately following collection and preserved in 80% ethanol. Prey samples were later sorted and identified in the laboratory.

The energetic value of selected potential prey species was determined from zooplankton samples collected in the southern Bering Sea, just north of the central Aleutian Islands, during a cruise in June 1995. Following determinations of chemical composition, the energy value of fresh prey biomass was calculated assuming that the metabolizable energy yields of protein and fat are 18.04 kJ g⁻¹ and 39.54 kJ g⁻¹, respectively, for uricotelic organisms (Bell 1990).

**RESULTS**

**Distributions of auklets and copepods along transects**

We conducted 2 replicate transects along Line A northwestward from Gambell on July 9-10, 1984. On the first transect (afternoon of July 9), a concentration of least auklets was evident at the island-ward edge of the Anadyr Front, ~25 km from Gambell, although a number of birds were also present closer to the island on the adjacent, stratified, Bering Shelf water (Fig. 2A). Net tows indicated that high densities of *Neocalanus plumchrus* were present in the vicinity of the concentration of auklets at the front (Fig. 2A). On our second transect along this line the following morning, the front had moved closer to St. Lawrence Island and occupied a broad zone in the strait. A concentration of least auklets was again present in the frontal zone; however, while the front had moved closer to the birds’ breeding areas, the birds were feeding farther away than during the previous transect, in an area where hydrographic data revealed the presence of fine-scale structure in the water column (Fig. 2B).

We repeated a single transect along Line A on August 9, 1985. During this survey, the surface manifestation of the front occurred west of the International Date Line and auklet densities were relatively low in the survey area (Fig. 3). The hydroacoustic survey
showed a dense layer of zooplankton along the thermocline, and least auklets were concentrated in an area where the hydroacoustic data indicated that zooplankton were being entrained from the thermocline to the surface (Fig. 3).

We conducted 3 replicate transects along Line A northwestward from Gambell on August 11-12, 1986. On the first run during the afternoon of August 11, the front apparently was located close to Russia and our transect did not reach it. Nevertheless, we found a large concentration of least auklets over the deepest portion of the strait, ~33 km from Gambell (Fig. 4A). A concentration of crested auklets was evident closer to the island at a point where the thermocline domed upward (Fig. 4A). The concurrent hydroacoustic survey indicated that 2 layers of zooplankton were present: a midwater layer along the thermocline away from the island, and an epibenthic layer below 30 m near the island. There were indications of surface entrainment of zooplankton as the front was approached, and the concentration of least auklets was over an apparent surface concentration of prey (Fig. 4A). The crested auklets were near but not directly over another apparent surface concentration of zooplankton, and instead seemed to be more closely associated with the epibenthic layer (Fig. 4A).

The second transect was run on the same day but later in the evening. The hydroacoustic survey showed a continuous scattering layer along the thermocline (Fig. 4B), with no evidence of the epibenthic layer detected during the previous survey (cf. Fig. 4A). As during the previous survey, a dense surface patch of zooplankton attracted large numbers of least auklets, but a smaller concentration of least auklets was also evident at the Russian end of the line, where CTD data indicated that the thermocline was beginning to rise to the surface and the front was rapidly moving eastward into the study area (Fig. 4B). Also at this time, the upward deformation of the thermocline near the island had apparently relaxed; although a few crested auklets were still present in the vicinity as before, the concentration had clearly dispersed (Fig. 4B).

The third and final transect in this series was conducted on the afternoon of the following day. At that time, the front had moved into our study area and was apparent at the northwest end of the line, ~45 km from Gambell. A massive concentration of least auklets was found feeding at the front (Fig. 4C). Few crested auklets were observed during this transect. Five net tows along the line indicated that the highest densities of *Neocalanus plumchrus* (~78 ind. m$^{-3}$) were present in the vicinity of the front (Fig. 4C). Least auklets were apparently overflying *N. plumchrus* densities of ~37 ind. m$^{-3}$ and even higher densities of *Calanus marshallae* (~59 ind. m$^{-3}$) to reach the frontal concentration of *N. plumchrus* (Fig. 4C).

Late in the day on August 12, 1986, we conducted a fine-scale study of horizontal frontal structure and auklet distribution near the end of Line C (Fig. 5). Extremely large numbers of least auklets were foraging directly within the frontal zone, and appeared to be concentrated in a frontal meander (Fig. 5).

We returned to the Anadyr Strait in 1993 and performed 1 transect along Line B northwestward from Gambell on June 15. At that time, much of the frontal zone was apparently located slightly beyond the International Date Line and we were forced to stop just short of crossing it. Nevertheless, it was clear that large numbers of least auklets were
Concentrated at the front (Fig. 6). Crested auklets were also present in large numbers on this transect, and there was a remarkably sharp spatial separation between the 2 auklet species. All crested auklets were feeding to the St. Lawrence Island side of the concentration of the least auklets, near the deepest part of the strait (Fig. 6).

We conducted our first transect along Line C running northwestward from Savoonga on the north side of St. Lawrence Island during the afternoon of August 4, 1985. A sharp front was present ~70 km from Savoonga, but densities of least auklets were not obviously elevated in the frontal zone (Fig. 7). Peak densities of least auklets occurred only ~30 km from the island, over the point closest to the island where the thermocline still was strong (i.e. the CTD data suggested mixing closer to island). A net tow indicated high densities of *Neocalanus plumchrus* (171 ind. m$^{-3}$) in the vicinity of the least auklet concentration (Fig. 7). Two small aggregations of crested auklets were also present beyond the peak concentration of least auklets, one over the shallow thermocline and one at the nearshore edge of the front (Fig. 7).
We ran a series of 3 transects along Line C on August 11, 12, and 13, 1985. On August 11, the front was encountered ~90 km from Savoonga, and therefore had moved ~20 km west since the previous survey 1 wk before (Fig. 8A; cf. Fig. 7). Least auklets were not abundant along this transect, but a sharp concentration nevertheless was evident in the frontal area (Fig. 8A).

In contrast to most of the other transects, peak numbers of least auklets apparently were feeding on the side of the front abutting Anadyr water, rather than near the shelf-water edge of the front (Fig. 8A). Small numbers of crested auklets were present along the offshore half of the transect, with a suggestion of a concentration in the frontal zone but separated from and slightly islandward of the least auklets (Fig. 8A).

The second transect in this series was conducted on August 12 at a finer spatial resolution. The geographic distribution of birds was very similar to the previous transect but the front had moved >5 km closer to the island. As a result, the concentration of least auklets was ~5 km northwest of the front, over fully mixed Anadyr water, whereas the crested auklets were closer to the front (Fig. 8B). The concurrent hydroacoustic survey showed a strong concentration of zoo-plankton in the front (Fig. 8B).

By the final transect in this series on the following morning, the front had moved even closer to St. Lawrence Island and was ~75 km from Savoonga. Least auklets were scattered along the line but most birds were loosely concentrated near the front (Fig. 8C). The hydroacoustic survey revealed a strong
scattering layer along the shallow thermocline, and peaks in least auklet density were clearly associated with 2 strong surface concentrations of zooplankton in the vicinity of the front. A small concentration of crested auklets was evident just island-ward of the front and away from the majority of least auklets (Fig. 8C). These crested auklets appeared to be exploiting an epibenthic concentration of zooplankton (Fig. 8C).

We returned to Line C in 1986 and conducted transects during the mornings of August 12 and 13. On August 12, the front was present -90-95 km from St. Lawrence Island and a small concentration of least auklets was evident there (Fig. 9A). A larger concentration of crested auklets was located closer to the island, though no distinctive hydrographic features were evident (Fig. 9A). The front had not changed position by the following day, and a large concentration of least auklets was located near the edge of the front abutting Anadyr Water (Fig. 9B). Very few crested auklets were observed during this transect.

Zooplankton abundance

Net tows indicated that the copepod *Eucalanus bungii* was the most abundant species in the front as
Fig. 9. Transects run along Line C on August 12–13, 1986. Upper panels show auklet abundance, and bottom panels show the thermal structure of the water column (°C) from CTD casts. Note the horizontal scale difference between the 2 transects. Locations of hydrographic stations are marked by triangles along the bottom of the panels showing CTD contours. (A) Morning of August 12 (04:51–06:53 h local time). (B) Morning of August 13 (02:38–07:25 h local time).

well as in both of the adjacent water masses (Table 1). Among the principal components of the zooplankton, the copepods *Neocalanus plumchrus* and *E. bungii* as well as the chaetognath *Sagitta elegans*—a copepod predator—occurred at the front in significantly elevated densities. *Metridia pacifica* appeared to be about equally abundant in Anadyr water and at the front, but was significantly less abundant in shelf water. *Pseudocalanus* spp. appeared to be much more abundant at the front than elsewhere, but these smaller copepods were not sampled adequately by our nets so we did not conduct statistical tests. In contrast, copepodids were most abundant in shelf water. Densities of other species did not vary significantly among water masses.

**Auklet diets**

Diet samples collected from least auklets feeding in Bering Shelf water and at the Anadyr Front are summarized in Table 2. A diverse array of zooplankters occurred in least auklet diets, but *Neocalanus plumchrus* was the dominant prey item in auklet diets both at the front and elsewhere. Crested auklets were not sampled for diet.

Mean metabolizable energy values of the principal copepod species available to least auklets were 5.36 kJ g⁻¹ for *Neocalanus cristatus* (±1.31 SE, n = 4 determinations), 4.95 kJ g⁻¹ for *N. plumchrus* (±0.52 SE, n = 4), and 2.02 kJ g⁻¹ for *Eucalanus bungii* (±0.49 SE, n = 4). Energy value differed significantly among these 3 species (1-way ANOVA: $F_{2,9} = 4.45, p = 0.045$).

**DISCUSSION**

Our repeated crossings of the Anadyr Strait revealed common patterns in hydrography, zooplankton distribution, and auklet distribution. The western portion of the strait is occupied by cold dense Anadyr water from top to bottom. Where this mixed water interfaces with the stratified Bering Shelf water, the thermocline rises and forms a sharp surface front. Consistent with previous studies

<table>
<thead>
<tr>
<th>Species</th>
<th>Anadyr water</th>
<th>Anadyr Front</th>
<th>Bering Shelf water</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Neocalanus plumchrus</em></td>
<td>43.4 (10.4)</td>
<td>66.4 (16.1)</td>
<td>31.0 (16.1)</td>
<td>0.051</td>
</tr>
<tr>
<td><em>Neocalanus cristatus</em></td>
<td>2.3 (1.2)</td>
<td>2.3 (1.5)</td>
<td>1.4 (0.6)</td>
<td>0.83</td>
</tr>
<tr>
<td><em>Eucalanus bungii</em></td>
<td>58.0 (34.9)</td>
<td>343.0 (133.7)</td>
<td>52.1 (30.0)</td>
<td>0.010</td>
</tr>
<tr>
<td><em>Calanus marshallae</em></td>
<td>8.4 (3.8)</td>
<td>22.4 (11.1)</td>
<td>15.5 (6.0)</td>
<td>0.64</td>
</tr>
<tr>
<td><em>Metridia pacifica</em></td>
<td>13.1 (6.1)</td>
<td>10.1 (3.7)</td>
<td>2.0 (1.4)</td>
<td>0.052</td>
</tr>
<tr>
<td>Calanoid copepods</td>
<td>18.6 (6.4)</td>
<td>33.3 (7.8)</td>
<td>132.1 (40.5)</td>
<td>0.059</td>
</tr>
<tr>
<td>Euphausiid furcilia</td>
<td>1.7 (0.7)</td>
<td>34.5 (17.8)</td>
<td>25.6 (11.1)</td>
<td>0.19</td>
</tr>
<tr>
<td><em>Aglantha digitalis</em></td>
<td>0.1 (0.1)</td>
<td>7.4 (6.6)</td>
<td>14.2 (11.1)</td>
<td>0.33</td>
</tr>
<tr>
<td><em>Sagitta elegans</em></td>
<td>0.8 (0.6)</td>
<td>13.4 (4.1)</td>
<td>5.6 (3.8)</td>
<td>0.006</td>
</tr>
<tr>
<td>Sample size (no. of tows)</td>
<td>6</td>
<td>7</td>
<td>10</td>
<td></td>
</tr>
</tbody>
</table>
Table 2. Diet samples obtained from least auklets collected from feeding flocks on Bering Shelf water and at the Anadyr Front. All birds were collected from 00:02-01:22 h local time on August 13, 1986, immediately following the completion of the fine-scale 2-dimensional study (Fig 5). Values shown are the numbers of identifiable prey items in each sample. X: present but not quantifiable

<table>
<thead>
<tr>
<th>Prey species</th>
<th>Bering Shelf water</th>
<th>Anadyr Front</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1 2 3 4 5 6 7 8</td>
<td>9 10 11 12 13 14 15</td>
</tr>
<tr>
<td>Neocalanus plumchrus</td>
<td>360 418 350 490</td>
<td>30 85 x</td>
</tr>
<tr>
<td>Neocalanus cristatus</td>
<td>2 2 3 3 3 0 0 0</td>
<td>3 4 x 0</td>
</tr>
<tr>
<td>Parathemisto libellula</td>
<td>0 0 0 0 0 1 0</td>
<td>2 0 0 0</td>
</tr>
<tr>
<td>Gammarid amphipod</td>
<td>0 0 0 0 0 1 0 0</td>
<td>0 0 0 0</td>
</tr>
<tr>
<td>Unidentified amphipod</td>
<td>0 0 1 0 0 0 0 0</td>
<td>0 0 0 0</td>
</tr>
<tr>
<td>Euphausiid larvae</td>
<td>0 0 0 0 0 0 0 0</td>
<td>0 0 0 0</td>
</tr>
<tr>
<td>Unidentified Pleustidae</td>
<td>0 2 0 1 0 0 0 0</td>
<td>0 0 0 0</td>
</tr>
<tr>
<td>Lithode crab larvae</td>
<td>0 0 0 0 0 0 0 0</td>
<td>0 0 0 0</td>
</tr>
<tr>
<td>Boreogadus sp.</td>
<td>0 0 0 0 0 4 0 0</td>
<td>0 0 0 0</td>
</tr>
<tr>
<td>Theragra sp.</td>
<td>3 0 0 0 0 0 0 0</td>
<td>0 0 0 0</td>
</tr>
<tr>
<td>Unidentified fish larvae</td>
<td>0 0 1 0 0 0 0 0</td>
<td>0 0 1 0</td>
</tr>
<tr>
<td>Unidentified fish bones</td>
<td>0 0 0 0 0 0 0 0</td>
<td>0 0 0 0</td>
</tr>
<tr>
<td>Unidentified gadid oolith</td>
<td>0 1 0 0 0 0 0 0</td>
<td>0 0 0 0</td>
</tr>
</tbody>
</table>

*N Sample included large quantity of unidentifiable Neocalanus mush

(Nihoul et al. 1993, Gawarkiewicz et al. 1994), we found the location of the Anadyr Front to be variable; it was not present in the same position among cruises, nor even among days within the same cruise. The distribution of least auklets responded to the movements of the front throughout our study. Large concentrations of least auklets often occurred at the front, and their use of the front appeared to be related to high levels of food availability.

**Concentration of zooplankton along the thermocline and in the frontal zone**

Acoustic surveys in both 1985 and 1986 revealed a high biomass of plankton in the vicinity of the front, and also indicated that there was an abrupt change in the vertical distribution of plankton at the front. Whereas most of the acoustically observed plankton biomass was concentrated along the thermocline on the stratified side of the front, available evidence suggests that plankton was thoroughly mixed in the turbulent waters west of the front. Concentration of copepods along the thermocline is consistent with previous observational and experimental studies, which have demonstrated that aggregations of zooplankton often occur at property interfaces (Harder 1968, Turner & Dagg 1983).

Direct net sampling indicated that calanoid copepods were the numerically dominant component of the zooplankton, and that copepod densities were significantly enhanced in the frontal zone. Previous studies have found copepod concentrations in association with a wide variety of frontal features. For example, Mackas & Louttit (1988) documented concentrations of Neocalanus plumchrus along the margin of the Fraser River outflow plume. Herman et al. (1981) found that copepod abundance was 3 to 4 times greater at the shelfbreak front south of Nova Scotia than in adjacent waters. Cooney (1981) showed that copepod biomass was enhanced along the shelfbreak front in the southeastern Bering Sea, and Wishner et al. (1995) found peak densities of Calanus finmarchicus associated with a salinity front off Cape Cod, Massachusetts. Copepod concentrations have also been documented in the vicinity of upwelling fronts off California (Smith et al. 1986).

The causes of such frontal concentrations have been widely debated, and may vary geographically and/or temporally. Sustained chemical fluxes and nutrient regeneration in frontal zones can maintain persistent phytoplankton blooms (e.g. Traganza et al. 1987, Dewey & Moum 1993), which may potentially drive an upward cascade of trophic interactions through food chains (i.e. via aggregation of organisms at higher trophic levels to feed on growing biomass concentrations at lower levels; Atkinson & Targett 1983). However, zooplankton concentrations can also be generated in the absence of enhanced primary production purely through mechanical accumulation resulting from the hydrodynamic convergence associated with fronts (Olson & Backus 1985, Franks 1992b, Govoni & Grimes 1992).

Concentration of copepods in the Anadyr Front likely was not due to enhanced production, because the ontogenetic development of most of the copepod species present occurs well to the south of the study area in early spring (Cooney & Coyle 1982, Vidal &
Pods were presumably dispersed uniformly throughout the water column by the intense turbulence, and were probably difficult to exploit profitably. Haney (1991) asserted that least auklets preferentially occurred in mixed waters in the Anadyr Strait, but his analyses did not discuss the role of the front, rendering his conclusions difficult to assess critically.

Our net sampling indicated that Neocalanus plumchrus occurred abundantly east of the front in stratified water. These oceanic copepods are presumably introduced into the eastern side of the strait via the intrusion of Anadyr water, and may subsequently be injected into the upper layer of Bering Shelf water via mixing across the thermocline, as suggested by some of our hydroacoustic surveys (e.g. Figs. 3 & 4B). Despite the occurrence of N. plumchrus in stratified water, many least auklets overflew this water to feed at the front. Enhanced availability of N. plumchrus therefore must have been the principal reason why these birds usually travelled so far from their colonies to feed at the front.

Diet and habitat selection by least auklets

Our finding that least auklets were feeding preferentially on Neocalanus plumchrus is in agreement with most previous trophic studies of this bird (Bedard 1969, Searing 1977, Bradstreet 1985, Springer et al. 1986, 1987, Harrison 1987, 1990, Day & Byrd 1988, Hunt & Harrison 1990, Hunt et al. 1990, 1993, Piatt et al. 1990b, Obst et al. 1995). There was no evidence that least auklets took Eucalanus bungii at the front, despite the fact that it outnumbered N. plumchrus by a factor of 5 (Table 1) and these copepods are about the same size (~6 mm length). However, the energy value of N. plumchrus is 2.5 times greater than that of E. bungii. Selection of copepod prey by least auklets therefore appears to be dictated primarily by the energy content of individual prey items rather than the relative abundance of the different prey species, which is consistent with predictions about diet selection from optimal foraging models (Stephens & Krebs 1986). E. bungii is also more transparent than N. plumchrus, and might be more difficult to see underwater. Some workers have suggested that seabirds are generally opportunistic, taking whatever prey are available at a given place and time (e.g. Ainley et al. 1992). Our data, together with previous studies (Hunt & Harrison 1990, Hunt et al. 1990, Obst et al. 1995), clearly indicate that least auklets are not random samplers of their marine environment, but instead are highly selective in their diet.

Use of the front by least auklets

The Anadyr Current is the proximate source of the copepods upon which least auklets preyed. However, west of the front in unstratified Anadyr water, copepods were presumably dispersed uniformly throughout the water column by the intense turbulence, and were probably difficult to exploit profitably. Haney (1991) asserted that least auklets preferentially occurred in mixed waters in the Anadyr Strait, but his analyses did not discuss the role of the front, rendering his conclusions difficult to assess critically.

Our net sampling indicated that Neocalanus plumchrus occurred abundantly east of the front in stratified water. These oceanic copepods are presumably introduced into the eastern side of the strait via the intrusion of Anadyr water, and may subsequently be injected into the upper layer of Bering Shelf water via mixing across the thermocline, as suggested by some of our hydroacoustic surveys (e.g. Figs. 3 & 4B). Despite the occurrence of N. plumchrus in stratified water, many least auklets overflew this water to feed at the front. Enhanced availability of N. plumchrus therefore must have been the principal reason why these birds usually travelled so far from their colonies to feed at the front.

Least auklets, like other alcids, have a specialized flight morphology (specifically, reduced wing area) that results in a high energetic cost of flight; these birds therefore are expected to forage as close to their colonies as is economically feasible (Obst et al. 1995). In all transects where we observed large concentrations of least auklets away from the front, net samples and/or hydroacoustic surveys demonstrated high levels of food availability where the birds were feeding. In the one instance where we conducted a net tow in the vicinity of a least auklet concentration away from the front, the net sample indicated that the birds were exploiting an extremely dense patch of Neocalanus plumchrus in stratified Bering Shelf water (Fig. 7). When we exclude this data point, the mean density of N. plumchrus in waters overflown by least auklets averaged 15.5 ind. m⁻². This suggests that there may be a threshold density of N. plumchrus below which it is economically infeasible for least auklets to forage, and that this threshold is ≥16 ind. m⁻². The precise threshold value is difficult to estimate, because our nets sampled deeper parts of the water column that were inaccessible to least auklets (maximum; diving depth estimated to be about 15 m; Obst et al. 1995), and the integrated water-column densities therefore may not accurately reflect the availability of copepods to foraging auklets.

While concentrations of least auklets were frequently associated with the Anadyr Front, their abundance at the front varied dramatically. Some of this variation was undoubtedly attributable to diel patterns of foraging and colony attendance (Piatt et al. 1990a), which we did not study. We suspect, however, that some of the variation may also reflect frontal dynamics.
Fronts have been popularly conceptualized as unitary structures with predictable dynamics, but this view is probably oversimplified. Recent workers have recognized that frontal zones are characterized by complex internal structure and may incorporate features across a wide range of spatial scales (Rodionov 1994). For example, recent studies of the Mississippi River plume front have indicated that the frontal system consists of a large-scale (~2–20 km width) frontal zone within which are embedded small-scale (~10–50 m width) convergence zones (Govoni & Grimes 1992). Furthermore, although there was evidence of overall concentration of larval fishes in surface waters in the frontal zone, exceptionally high densities were not observed regularly at the small-scale features (Govoni & Grimes 1992). The small-scale convergence zones were sinuous in shape and were ephemeral, forming and dissipating over time scales of 2 to 6 h. They were not always observed during transects across the large-scale zone. Govoni & Grimes (1992) suggested that 'differences of larval fishes within the [large-scale] frontal zone are probably the result of their accumulation along ephemeral convergence zones and subsequent dispersal and mixing during relaxation of convergence', and that the spatial distribution of larvae in the vicinity of the front is consequently 'the aggregate result of the repeated formation and degeneration of [small-scale] convergence zones.' Olson et al. (1994) also distinguished between large-scale frontal zones and fronts, arguing that 'the primary biological response ... is tied to the dynamics of the smaller scale features, i.e., the individual fronts, that are characteristic of these frontal zones'.

Although our CTD sections indicated that the front itself was very sharp, we have no information on the flow dynamics and circulation patterns. However, our visual observations while crossing the Strait indicated that the presence of a surface manifestation of the front was variable. During one crossing of the front in 1986, we observed surface-feeding birds foraging in a narrow line along a slick which apparently represented the surface expression of the front. During most other crossings, there were few visual cues marking the presence of the front. Variability in the surface manifestation of the front may adversely affect the auklets' ability to locate it and/or track it accurately. Supporting evidence comes from our observation of a least auklet concentration that persisted at the same site for 2 consecutive days, despite the fact that the front had moved out of the area by the second day (Fig. 8A, B). We also suspect that circulation patterns associated with the Anadyr Front are more complex than is otherwise suggested by the strikingly uniform CTD sections. Variability in kinematic structure may affect the extent to which copepod concentrations develop and persist in the frontal zone, and consequently affect the birds' use of the front. A cross-front hydroacoustic survey supported these ideas, demonstrating a complex multimodal pattern of zooplankton concentration near the surface (Fig. 8C).

### Distribution of crested auklets

In contrast to least auklets, crested auklets, when found, were usually away from the front in areas without any striking hydrographic features. On transects where they were encountered, crested auklets consistently occurred in compact aggregations that were clearly separated from least auklets and on the nearshore side of the front. This spatial segregation of the 2 species could have resulted from (1) differences in flight capabilities, (2) interspecific competition, or (3) dietary differences.

Within any group of bird species that differ in size but are otherwise morphologically and ecologically similar, flight mechanics theory (Pennycuick 1989) dictates that larger species can fly faster and therefore have larger potential foraging radii. Thus, there will be some habitats potentially exploitable by larger species that cannot be used by smaller species. This foraging radius hypothesis can clearly be rejected here, because crested auklets are larger (~280 g vs ~80 g) and therefore have a larger potential foraging radius than least auklets, yet crested auklets were usually found closer to their breeding sites.

Previous workers have proposed that differences in foraging zones at sea have evolved as an ecological isolating mechanism in response to historic interspecific competition in alcid communities (Cody 1973; but see Bédard 1976). In other words, differences in foraging distributions may reflect evolutionary niche differentiation, or the 'ghost of competition past' (sensu Connell 1980). Fixed foraging zones or preferred foraging distances also seem unlikely in our study, because the pattern of separation between the 2 species sometimes reversed (i.e., crested auklets were found farther from St. Lawrence Island than least auklets in several instances). Spatial segregation could, however, result from competition in the present. This possibility is difficult to evaluate because we have no direct evidence either for or against competition, and furthermore, there is little consensus in the literature about whether interspecific competition is an important factor underlying patterns of space use in seabirds. The problem is especially difficult for diving birds because their submarine behavior during foraging bouts is invisible to shipboard observers. However, observations of alcids by Duffy et al. (1987) in an artificial environment indi-
cated that aggressive interactions are common under-
water, and Duffy et al. suggested that underwater
interspecific interactions may be involved in the par-
titioning of space among different species. Whether
competition can cause spatial segregation in planktiv-
orous seabirds awaits further study.

The most likely explanation for the observed pat-
terns of spatial segregation is that different physical
factors influenced the availability of the 2 species’ pre-
fected prey. In 2 instances, concentrations of crested
auklets were located over acoustically observed epi-
benthic scattering layers (Figs. 4A & 8C), which prob-
ably consisted primarily of euphausiids. Although we
were not able to collect diet samples from crested auk-
lets in this study and do not know with certainty what
the birds were eating, available information on the diet
of crested auklets elsewhere in the northern Bering
Sea generally points to euphausiids as their principal
prey during the chick-rearing period. On St. Lawrence
Island, Bédard (1969) found that crested auklets fed
mostly on Thysanoessa spp. Similarly, collections of
foraging crested auklets in the Chirikov Basin indi-
cated that T. raschii was the most important prey type,
though copepods and amphipods were also taken
(Harrison 1987). Platt et al. (1988) and Haney & Solow
(1992) also reported that Thysanoessa spp. were the
dominant prey taken by crested auklets off St. Law-
rence Island. Further south, Hunt et al. (1981) exam-
ined auklet diets at the Pribilof Islands and found that
crested auklets took mostly euphausiids (69% by vol-
ume, primarily T. inermis), together with substantial
numbers of amphipods (30% by volume, primarily
Parathemisto libellula).

Euphausiids may be concentrated by topographi-
cally forced upwellings when they attempt to remain at
depth by swimming downward (Simard et al. 1986),
and in these circumstances may then be exploited
easily by seabirds (e.g. Coyle et al. 1992). On 1 of our
transects, a strong upward excursion of the thermo-
cline was evident in CTD data in the area where we
observed crested auklets foraging over a distinct epi-
benthic scattering layer (Fig. 4A). If this ‘dome’ was
caused by subsurface upwelling, it may have been a
source of enhanced euphausiid density via the mecha-
nism previously described. Alternatively, domes of
deep Anadyr water intruding into the overlying Bering
Shelf water may represent discrete areas where
Anadyr water enters the Chirikov Basin as a strong
subsurface jet. This interpretation is supported by our
observations of the plankton net’s wire angle during
some vertical tows, which suggested considerable cur-
rent shear. We hypothesize that crested auklets may
exploit euphausiids that are advected by rapidly mov-
ing subsurface jets of Anadyr water along the eastern
side of the strait. By positioning themselves over such
jets, crested auklets may realize an increase in the
effective availability of their euphausiid prey—even in
the absence of any real variation in prey density—
because the rate of contact with individual prey items
should increase. Interestingly, when the deformation
in the thermocline relaxed following the completion of
the transect just described, the epibenthic scattering
layer disappeared completely and the concentration of
crested auklets dispersed (Fig. 4B; c.f. Fig. 4A).

In most cases where we encountered crested auklets,
no striking hydrographic features were evident (Figs. 6,
7, 8A,C & 9). We speculate that on some transects, our
CTD casts may have been too coarsely spaced to re-
solve the fine structure associated with hydrographic
features important to crested auklets.

Role of the Anadyr Front in the northern
Bering Sea ecosystem

Some previous workers have suggested that fronts act as boundaries between marine communities (Brandt
& Wadley 1981; but see Sournia 1994). This does not
appear to be the case in the Anadyr Strait, because dif-
fferences in zooplankton species composition between
water masses are manifested primarily in relative dif-
fences in densities of the constituent species, rather
than in absolute taxonomic composition (Table 1; see
also Springer et al. 1989, Piatt et al. 1992). While the
Anadyr Front cannot be characterized as a bioge-
ographic boundary, it is a feature that supports elevated
zooplankton populations. In turn, enhanced avail-
ability of preferred copepod prey associated with this
front attracts large numbers of foraging least auklets.
Crested auklets generally did not occur at the front,
and instead may have been exploiting enhanced fluxes
of euphausiids being advected by subsurface jets on
the eastern side of the strait. Thus, spatial segregation
of the 2 principal planktivores in the Anadyr Strait
likely arises because of their use of different prey
originating at different depths. All prey taken by auk-
lets probably originates within same system (i.e. the
Anadyr Current), but apparently different physical
mechanisms are responsible for the localized enhance-
ment of availability of the 2 prey types. Hunt et al.
(1998) reached qualitatively similar conclusions con-
cerning a physical oceanographic basis to niche differ-
etiation among auklets around the Aleutian Islands.

Large auklet colonies are distributed mostly along
the western tip and northwestern side of St. Lawrence
Island (Fig. 1). We suspect that this distribution reflects
the predictability and economy of prey availability,
and that the Anadyr Front may be important for sup-
porting the large local populations. Despite the vari-
ability of the front over short time scales, the avail-
ability of the front (and associated concentrations of plankton) appears to be highly predictable over the long term. The presence of this front may therefore be one of the most important factors controlling the reproductive performance of least auklets in the region. Auklets need the Anadyr Current for successful feeding, because this current carries their preferred prey. However, they also need a structured water column, because strong turbulence and intense mixing undoubtedly preclude the formation of plankton patches in pure Anadyr water.

Unfortunately, little is currently known about the ecological consequences of the intense interannual variability in physical and biological processes that affects the region. Physical variability is manifested in year-to-year changes in the eastward extent of excursions of the Anadyr Current into the Chirikov Basin and around St. Lawrence Island (Coachman & Aagaard 1988, Coachman 1993, Gawarkiewicz et al. 1994). There is undoubtedly also intense interannual variability in zooplankton source populations (Sambrotto & Goering 1983, Smith & Vidal 1986), but little is currently known about how (or whether) this affects the northern Bering Sea ecosystem. Regardless of how interannual variability is manifested (i.e. eastward excursions of Anadyr water vs changes in the composition and/or abundance of zooplankton populations), we hypothesize that the proximity of the Anadyr Front may buffer least auklets breeding along the western side of St. Lawrence Island against depressions in prey availability compared to auklets breeding farther east on the island. Indeed, we discovered from conversations with the native people of Savoonga that auklets suffered a near-total reproductive collapse on central St. Lawrence Island in 1984: few birds laid eggs, and most of the occupied nests were subsequently deserted. Additionally, many birds washed ashore dead late in the summer of 1984. In the following year (1985), large numbers of birds bred and their nesting appeared to be successful. Bédard (1969) also observed large numbers of starving crested auklets on the north side of St. Lawrence Island in one year. This anecdotal evidence of occasional large-scale reproductive failure and adult mortality suggests that perturbations in the Anadyr Current may have profound effects on auklet populations, though the exact mechanisms are not clear. Long-term multidisciplinary studies will be required to understand fully how the interplay between physical and biological processes in the region affects the foraging success and reproductive performance of these interesting planktivorous birds.

Acknowledgements. We thank K. Coyle, Z. Eppley, B. Obst, R. Veit, M. Whitehouse, C. Williams, and B. Young for field assistance during the 1984–1986 cruises, and J. M. Grebmeier for providing the opportunity for us to join the 1993 cruise. We also thank the captain and crew of the RV ‘Alpha Helix’ for outstanding logistical support, and R. Veit and 4 anonymous reviewers for comments on an earlier draft of the paper. Financial support was provided by NSF grants DPP-8308232 and DPP-9321636 (to G.L.H.) and DPP-9306694 (to J. M. Grebmeier and L. W. Cooper).

LITERATURE CITED


Darwin C (1845) Journal of researches into the natural history and geology of the countries visited during the voyage of H. M. S. Beagle round the world, under the command of Capt. Fitz Roy, R. N. John Murray, London


Fedorov KN (1986) The physical nature and structure of oceanic fronts. Springer-Verlag, Berlin


Franklin B (1786) A letter from Dr. Benjamin Franklin, to Mr. Alphonsus le Roy, member of several academies, at Paris. Containing sundry maritime observations. Trans Am Philos Soc 2:294–329


Hawke DJ (1996) Relatively infrequent seabird aggregation at nearshore fronts and tidal plumes at locations around Banks Peninsula, New Zealand. Notornis 43:66–70


Piatt JF, Roberts BD, Hatch SA (1990a) Colony attendance and population monitoring of Least and Crested Auklets on St. Lawrence Island, Alaska. Condor 92,97–106


Editorial responsibility: Otto Kinne (Editor), Oldendorf/Luhe, Germany

Submitted: December 30, 1997; Accepted: September 28, 1998
Proofs received from author(s): May 6, 1999