RELATIONSHIPS BETWEEN OCEANOGRAPHY AND THE DEMOGRAPHY & FORAGING OF AUKLETS (CHARADRIIFORMES, ALCIDAE: AETHIA; MERREM 1788) IN THE ALEUTIAN ISLANDS, ALASKA

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#### FORAGING OF AUKLETS (CHARADRIIFORMES, ALCIDAE: AETHIA;

#### MERREM 1788) IN THE ALEUTIAN ISLANDS, ALASKA

By

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#### Abstract

Linvestigated bottom-up climate-mediated control of population of usklets (Acthic cristratelia, A. punillo, and A. pupmore) in the bering Sea over two decades of environmental variability. Broadi, this thesis comprises two parts: 1) examining the relationships between chick diet and survival in the context of large-scale climate and oceanographic patterns; and 2) using stable-isotoge analysis to infer foraging patterns throughout the annual cycle of different age classes of auklets, using information on chick diet and local oceanography to interrut the results.

I found that large-scale oceanographic patterns during the winter and spring in the North Pacific were related to auklet productivity the following breeding season. I hypothesized bottom up control of auklet productivity through food limitation, but food that chick mean composition throughout the Alexitan likuland dim cell for among vars or sites. Auklets' main prey, Necedomurs spc. copepods were most prevalent in chick diets when local sea-surface temperature (557) during the breeding season was around 4.5 ± 1.0°C, but that outside this range, the proportion of biomass represented by Necedomur copepods declined raight). There was significant overlap among Least, Crested, and Whiskered Auklets in the composition of chick meah, suggesting little topolice springetion.

Using information on moulting patterns, I found that stable-isotope ratios in the innermost primary feather (grown during incubation) indicated a shift by adult auklets to a more productive foraging location (e.g., oceanic fronts), presumably as a mechanism

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for reducing their own maintenance costs during chick maring. Then found that, among three auklet colonies in the Aleutan Islands, there were no differences among sites or years in the foraging patterns of Least Auklets during pre-breeding (breast feathers), incubation (primary 1), or post-breeding (primary 10). There appeared to be some level of ecological argregation between adult team ad Crestef Auklets at Careloi Nand.

Together, these results also indicate that local factors, such as introduced Norway rats (*Battus norwepicus*) at Kiska Island, are important factors in population' demography, and that demographic responses are not solely driven by bottom-up processes. Future studies should focus on the winter ecology and movements of authents.

#### Statement of Co-authorship

Chapters 2-6 were co-autored with Inc I. Jones, my advisor. In initiated the work on Kiaka Island in 2001, and has been a constant presence in the Alexian Islands since 1290. Jeffrey C. Williams and G. Verono Hydr of the Alaska Martime National Wildlife Reinge contributed diet and reproductive success data from Buldir and Kasatochi, and provided much-appreciated logistical support during my four years in the Alexians; Jeff and Verono are co-authors of Chapters 2 and 3. Itil Sydeman (Iranion Institute for Advanced Ecosystem Research) and Shabiro Minobe (Heikkado University) contributed greatly the oceanographic success of thir search, and are co-authors of Chapter 2 along with former graduate student Heather Major, who did Some of be initial demographic work on Kisa. Nikola Conyukhov of the Russian Academy of Sciences contributed data on Least Auklet moult from Siberia and the Pribliof Islands, and is a osubmor of Chapter 4.

#### Publications (published, or anticipated) and authorship arising from this thesis:

Chapter 2: A.L. Bond, I.L. Jones, W.J. Sydeman, H.L. Malor, S. Minobe, J.C. Williams, and G.V., Byrd. 2011. Reproductive success of planktiverous seabirds in the North Pacific is related to ocean climate on decadal scales. Marine Ecology Progress Series 42: 42:05-218. Chapter 3: A.L. Bond, I.L. Jones, J.C. Williams, and G.V. Byrd. Diet of auklet chicks in the Aleutian Islands, Alaska: similarity among islands, inter-species overlap, and relationships to ocean climate. Journal of Ornithology (in press).

- Chapter 4: A.L. Bond, N.B. Konyukhov, and I.L. Jones. Rapid but delayed primary moult in arctic-breeding Least Auklets. Condor (in revision).
- Chapter 5: A.L. Bond, and I.L. Jones. Stable isotopes reveal daily variation in foraging activity of incubating Least Auklets (*Aethia pusilla*).

Chapter 6: A.L. Bond, and I.L. Jones. Seasonal and age-class variation in the foraging of auklets in the Aleutian Islands, Alaska.

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My supervisor, Ian Jones, wrote me out of the blue, seemingly, in November 2006, and set in motion a series of events that would lead to many wonderful experiences. In the last four years, I have come to appreciate not only his ability to identify black is isotanty, but also his friendship and guidance. Lin's undrying love of the Alestians and of auklets is contagious, his encouragement in academic pursuits is unending, and his cooking is delicious. He allowed me to spread my academic wings, while guiding mecreful so I found my way.

My committee member, Ted Miller and Tom Chapman, have also been fantastic in encouraging me academically, and also in our non-academic discussions. Their guidance was instrumental in my successes, and in understanding my faults. Thanks also to my examineer, Tong Gaston, Pierre Pepin, and Ian Warkentin, for their thoughtful inght and improvements.

Some of my greatest thanks are for those who gave up summers in civilization to discover the aversome beauty, grandeur, and frustrations of the Alexitan Islands with me from 2007 2010. At Sirius Peint, Kika Island, I, Isad the pleasure of working with Grant Samion (2007). Michelle Wille (2007). Chris Barke (2008), Ein Pinnery (2009 and 2010), and Donald Prine-Hay (2010), and I shared a camp at Witchcraft Point on Kiska with Isa loses and Corey Hutchings. 2010. Our research crears on Build's Island also provided some much-needed data and company in Adak, and so I thank Stephan Iorten; Sampath Senevirum, Chris Small and Pan Woodman (2007). Lest solver and Parket. Levèque (2008), Hannah Munro (2009 and 2010), Allie Patrick (2009), and Sarah Kennedy (2010). I can only hope that you hold the sunny days, howling gales, long hikes, and delicious meals as dear as I.

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The desire for knowledge for its own sake is the one which really counts. Exploration is the physical expression of the intellectual passion. -Anslew Cherry-Garrard (1922)

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And Nahi she ketter I use to spell Nurches Who live in small caves, known on Nitches, for hutches. These Nurches have troubles, the biggest of which is The fact there are many more Nurches than Nitches. Each Nurch in a Nitch knows that some other Nurch Would like to move into his Nitch very much. So each Nurch in a Nitch has to watch that small Nitch Or Nurches who haven't a Witches.

"On Beyond Zebra", Dr. Seuss (1955)

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### List of Abbreviations and Symbols

- $\Delta^{13}$ C The discrimination factor for  $\delta^{13}$ C, or the amount of change in  $\delta^{13}$ C from a prey mixture to a consumer.
- $\delta^{13}$ C The ratio of the stable carbon isotopes  $^{13}$ C to  $^{12}$ C, compared with the international standard PeeDee Belemnite

 $\Delta^{15}N$  – The discrimination factor for  $\delta^{13}N$ , or the amount of change in  $\delta^{13}C$  from a prey mixture to a consumer.

δ<sup>15</sup>N – The ratio of stable nitrogen isotopes <sup>15</sup>N to <sup>14</sup>N, compared with atmospheric

nitrogen

ALPI - Aleutian Low Pressure Index

AIC - Akaike's Information Criteria

AIC<sub>c</sub> - Akaike's Information Criteria adjusted for small sample size

AMNWR - Alaska Maritime National Wildlife Refuge

ĉ - a measure of overdispersion of data given a statistical model

CRAU - Crested Auklet Aethia cristatella

DJF - December-January-February

JJA - June-July-August

LEAU - Least Auklet Aethia pusilla

MAM - March-April-May

NPI - North Pacific Index

NPGO - North Pacific Gyre Oscillation

P1 - the first, or innermost, primary feather

P10 - the last, or outermost, primary feather

PDO - Pacific Decadal Oscillation

PFMG - Proportion of feather mass grown

QAIC, - quasi-Akaike's Information Criteria adjusted for small sample size and extra-

model variation in error structure

SIA - Stable-isotope analysis

SLP - Sea-level pressure

SST - Sea-surface temperature

U-Z - Underhill-Zucchini moult model

WHAU - Whiskered Auklet Aethia pygmaea

w/- A model's Akaike weight.

% - Parts per thousand, per-mil notation for stable-isotope ratios

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# CHAPTER 1: INTRODUCTION AND OVERVIEW

#### Seabirds in the marine environment

Seabirds have been used as indicators of various aspects of the marine environment, including the abundance of commercially exploited fish tacks, contaminants, and climate change (Cairus 1987; Bot & & Maho 1993; Barrett 2002; Frederlssen et al. 2007; Part et al. 2009; Parsons et al. 2008; Durant et al. 2009; Indeeds, seabirds are complicatous, generally unexploited, secondary and tertiary consumers in marries systems, and can therefore provide a useful opportunity to investigate coupled climate-ecosystem variation (Durant et al. 2009).

To estabilish first if subiolity are indeed indicators of ecosystem characteristics (abiotic and biotic), there must be robust, reliable data on the process or state the seableds may be indicating, as well as the subbled for response a twing levels of the environment (Caims 1987). This inverse inference, or using a dependent variable to estimate one or more explanatory variables must be coupled with robust statistical approaches (Durant et al. 2009). The use of seableds as ecological indicators can be strengthered by simultaneous studies of multiple species at multiple sites (Chapters 2, 3).

#### Climate-mediated bottom-up control

Seabirds' responses to variation in climate and oceanography has been inferred typically to be bottom-up through control of the abundance, distribution, or availability of prey (e.g., Frederiken et ed. 2006; Fancine et el. 2001). There heve, however, been few studies seabird diet to aclimate or occanographic conditions (reviewed by Durant et el. 2009). Indeed, most studies using seabird diet as an ecological indicator focas on the relationship between seabird diet composition and prey abundrace or foraging location (e.g., Montevechi & Myers 1995; Nel et el. 2001). In many studies of seabirds' demographic responses to climate variability, a bottom- up mechanism is often invoked, but seldom tested (e.g., byrd et el. 2003; Higher 2008), although several recent studies do gregent independent data on prey availability in relation to seabirds' demographic responses to climate change (e.g., Durant et el. 2003; Waltanuki et el. 2009).

#### Approaches to studying bottom-up control

Ecologists interested in studying dief have adopted advances in analytical chemistry and biochemistry, including analyses of fatty add profiles (Verson et al. 2004), and ratios of stable isotopes, particularly those of carbon and nitrogen (Peterson & Fry SPZ, Selly 2000). Often, these can be used to complement traditional diet studies of put contents or equirgitates (Barrett et *el*. 2007). Recent methodological and statistical advances have made these techniques more accessible and widespread (e.g., larger et *al.* 2016; Pacologi, and the development of techniques (Martine cell Rior of results (Bond & Jone: Pacologi), and the development of techniques (Martine cell Rior of results).

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inherent assumptions and limitations (Duffy & Jackson 1986; Barrett et al. 2007). I have used two methods for studying the seabird diet – stable-isotope analysis, and prey composition of chick meals delivered by provisioning parents.

## Analysis of chick meals

Studies of food delivered or destined for next-bound thicks (chick mesil, food loads, or bill loads) are used frequently to characterize seabirds' foraging behaviour and responses to the environment (e.g., blait et al. 2000; Ballie & Junes 2003). The composition of chick meaks can often be measured with minimal disturbance to the penetra through their co observation (e.g., dendstrete & Broom 1985; Pomet et al. 1999), or through the simple capture of breeding adults en route to their nest, who regurgitate or drog the chicks' food (e.g., Montevecchi et al. 1992; Hunter et al. 2002). Chick meaks may not, however, be representative of adult diet because adults and chicks have different energetic constraints and demands (Ydenberg 1994; Barrett et al. 2007) and optimal foraging may result in adults partitioning prev items differently if the preva education of adulty on self minimerance, or provisioning their chick.

As central-place foragers, and long-lived "k-selected" species, adult seabirds should minimize the transit time to food resources (Mail 2031), balancing the trade-off between distance to foraging grounds and the nutritional quality of the resources (Elliott et al. 2003), and the evolutionary trade-off of their own survival and future reproduction (Secons 1903). Adult of some packers mouth/preceditivitizmes, balance their own and preceding the second some packers.

energetic requirements with those of their chick by a bimodal foraging strategy whereby adults make frequent, short tips to provide food for chicks, and less-frequent, long tips, to restore their own nutrient reserves (Chaurand & Weitmenskinch 1994, Weitmenskinch 1994, Weitmenskinch 1994), This strategy has been described recently in the Dovekle (Alle alle), a polar non-Procelination (Weicker et al. 2006).

## Stable-isotope analysis

There have been several advances in biochemical techniques to studying diet in the last 20 years, including stable-isotope analysis (Hobson 1987; Hobson & Clark 1992a, h) and fatty acid analysis (lyerson et al. 2004). Stable-isotope analysis offers several advantages over traditional methods, but has different drawbacks (Bond & Jones 2009b). Stable isotopes of carbon and nitrogen are used frequently in studies of seabird foraging ecology (Barrett et al. 2007: Inger & Bearhop 2008). Values are expressed as the parts-per-thousand ratio of the heavier (15N or 13C) to the lighter isotope (14N or 12C) as compared with an international standard (Pee Dee Belemnite for C. atmospheric air for N). δ<sup>15</sup>N, or the ratio of <sup>15</sup>N/<sup>14</sup>N, is a continuous measure of trophic position, and it increases between 2-5‰ with each trophic step because <sup>14</sup>N is excreted preferentially in nitrogenous waste (Steele & Daniel 1978; Minagawa & Wada 1984; Kelly 2000). There is little trophic enrichment of 813C at higher trophic levels (Rau et al. 1983; Hobson & Welch 1992), but marine (bicarbonate-based) and terrestrial carbon sources (CO2-based) differ in their δ13C values (Peterson & Fry 1987). δ13C can also provide information on

foraging area in the marine environment because of spatial "isoscapes" in the environment (Hobson 1999; Cherel & Hobson 2007; Graham et al. 2010; Jaeger et al. 2010).

Stable-isotope analysis is also useful for examining foraging behaviour when attaching data loggers is not feasible because of the difficulty in recapturing individuals to retrieve data, or because the species of interest is too small, and gear attachment would affect individuals' behaviour or even survival negatively. Isotope ratios are indicative of the diet at the time of tissue synthesis (hidson & Cakri 1992a). Feathers are sampled frequently because they offer a relatively non invusive way to sample large numbers of individuals, and if moult patterns are known, different feathers can be sampled to examine foraging throughout the annual cycle (Marra et al. 1998; Pyle 2009) because feathers are linet once grown, receiving no additional nutrient input (hobon & Cakri 1992; Chen et al. 2005a).

#### **Reproductive success**

Bottom-up influences on seakird demography will be most evident in annual reproductive success (the number of chicks reard successfully per pair). As long-lived species, seakinds will balance current and future reproduction, as well as reproduction and survival (Oobson & Jouventin 2010a, b). Consequently, reproductive success is more variable than adult's variavid literater eff. adol20. and in verser door food availability on to observation. increased predation, seabirds will abandon or skip breeding attempts (Schreiber & Schreiber 1984; Whittam & Leonard 1999).

#### Climate and auklets in the Aleutian Islands

The Aleutian Islands of Alaska span over 2000 km in the Oceanic Domain of the North Pacific. Deep oceanic waters with upwelling romes that bring nutrients to the surface, where they become available to avian consumers via prey, surround the central and wettern Aleutians (Dioder et al. 1985), territion et al. 1990, Russell et al. 1999).

Clinatically, the Akutian are influenced by basine vide clinatic patterns. (Mantua et al. 1997; Di Lorenzo et al. 2010). The Akutian Low Pressure Index (ALP) measures the strength of the Akutian Low From December-March over the area 20-70%, 120%-120% (Beamish & Boulinn 1993). Beamish et al. 1997; centred roughly over my study area. ALP is teleconnected to patterns of the El Nilo Southern Oscillation (INSO), as is the hearth Pacific Gyre Oscillation (INDO), which measures the sea-surface height and sea-surface temperature anomalies north of 20% [Di Lorenzo et al. 2010). Also related is the Pacific Decadd Docillation (PDO), a measure of sea-surface height and sea-surface area as the NPGO (Mantua et al. 1997; Mantua & Hare 2020). Finally, the North Pacific Index (NPI) is the area-weighted sea level pressure over the region bounded by 30%-65% and 150%-L40% (Tremberth & Hurrell 1994). No single index captures the range of variability of clinates in the NPC Pacific Logi course in visite or measures and Clinate and Coconceptive intercommed Head et al. 2020. Previous studies have related auklet demography to these indices (Jones et al. 2002; Jones et al. 2007).

There has been a long history of ornibhological research in the AlexLine, focused primarily on seabinds (Steller 1751; Stejineger 1885; Murie 1595; Gibson & Byrd 2007). Many seabind breeding biands are recovering following the eradication of introduced Arctic fexes (Algoe (Algoen)), Norway rats (Battra norvegicus), and other alien vertebrate species (Balley 1993; Ebbert & Byrd 2002; Buckelew *et al.* 2009). Introduced species have had a negative effect on AlexLina wilfinama (Williams *et ol.* 2003). More *al.* 2006). Ibiglinghing the need to understand local effects when interpreting relationships between climita and stability responses (Chapter 2).

#### Aethia auklets

Auklets (Achiar spp., Merrem, 1788) are the most abundant seabirds in the North Pacific numerically (Sowls et al. 1978; Gaston & Jones 1998). Auklets are accially monogamous, juy a single egg per annual breeding attempt, and their chicks have biparental care (byrd & Williams 1993), Jones 1993a, Ju, Adults freed chicks by regurgitating food stored in a blind throat pouch (Portenko 1934). Of the four species of Archina auklets, there are considered here.

Least Auklets (A. pusilla (Pallas, 1811)) are the smallest auk (85 g). Their chicks are nest-bound for about 26-29 days (Roby & Brink 1986s; Piatt *et al.* 1990b; Renner 2005), during which time they are provisioned with lipid-rich copepods (*Neocolanus* 

spp.) and smaller amounts of euphausiids (Thysanoessa spp.), gastropods, and amphipods (Bédard 1969a; Day & Byrd 1989; Harrison 1990; Gall et al. 2006). Parents provision chicks 2-3 times each per day (Roby & Brink 1986a). They are highly sensitive to disturbance at their breeding sites (Platt et al. 1990b), but on average, 50-70% of breeding pairs will rear a chick successfully (Bédard 1969b; Knudtson & Byrd 1982; Sheffield Guy et al. 2009). In North America, Least Auklets breed throughout the Bering Sea and Aleutian Islands (Sowls et al. 1978; Jones 1993b), and in Russia, they breed on the Chukotka Peninsula, on islands in the Sea of Okhotsk, and on the Kuril Islands (Dement'ev et al. 1951: Kozlova 1957: Velizhanin 1977b: Konvukhov et al. 1998; Kondratyev et al. 2000; Artukhin et al. 2001; Zelenskaya 2009). The wintering areas of Least Auklets are poorly known, but are likely in the western North Pacific, off the coast of Japan (Shuntov 1965; Velizhanin 1977a; Vyatkin 1981; Kondratyev et al. 2000; Sydeman et al. 2010). Previous work has shown that the survival of chicks was related to diet quality (Gall et al. 2006: Sheffield Guy et al. 2009), and that adult interannual survival was related to large-scale indices of winter climate in the North Pacific (Jones et al. 2002).

Created Auklets [A. cristotelle] (Pallas, 1769)) are similar to Least Auklets ecologically, but are larger (204), and their chicks are neet-bound for longer (13 days; Frazer et al. 1999). Both parents share the duties of chick nearing, together provisioning the chick 3-5 times per day (Frazer et al. 2002). They also tend to provision chicks with large proportions of explavalistic comparent to Least Auklets (Edited 1966); hum 2-10

al. 1981; Day & Byrd 1989). As with Least Auklets, between 50-70% of Created Auklet brending pairs raise a chick successfully (Frazer et al. 1999; Gall et al. 2006; Sheffield Gay et al. 2009). Chick survival in Created Auklets is also related to diet quality (Gall et al. 2006). They brend sympatrically with Least Auklets at many aites in the Bering and Okhotsk as and on the Kuril Islands (Sowls et al. 1978; Jones 1993); Konyahov et al. 1998; Kondratyev et al. 2000; Artukhin et al. 2001), and tend to winter in the eastern North Pacific Improduct Buildnamed (Sowls et al. 1978; Jones 1993); Konyahov et al.

Finally, Whikered Auklets (A, pugmeet (Gmelin), 1289) differ from comparen in that they are active at breeding colonies nocturnally. Because adults feed their chicks only once or twice per night (Konyukhov et al. 2000), they tend to provision chicks with larger meaks, and for a longer period (Hemer et al. 2002), Hutter et al. 2002, Chapter 3). Whikered Auklet chick diet tends to be similar to that of Least Auklets (Day & Byrd 1989); Humer et al. 2002). Unlike Least ar Cleated Auklets, however, after fieldinge, young and adult Whikered Auklets return to the colony at night for several weeks after the breeding season (Konyukhov & Zubakin 1994); Zubakin & Konyukhov 1994), which suggess that they winter in the vicinity of breeding colonis (Enginger 1185; Byrd & Williams 1993); Gibson & Byrd 2007). Their breeding distribution and population numbers are less known than that of their congeness because of their nocturnal habits, but colonies are found throughout the Alexian and northern Kuril blands in areas free of easier year or rout (Aramahin 1931); Swite et al. 3070; Charlister et al. 2007; Distribution and population of easier year or rout of throughout the Alexian and northern Kuril blands in areas free of easier year or rout for damahin 1931; Swite ed et . 3078; Konsthery et al. 2000;

Artukhin et al. 2001). Adult survival has been negatively related to ALPI (Jones et al. 2007).

#### Study sites

I studied auklets at four breeding colonies in the central and western Aleutian Islands - Buldir, Kiska, Gareloi, and Kasatochi Islands. I also analysed data on auklet moult from St. George and St. Paul islands in the Pribilofs, and from Cape Ulyakhpen on Russia's Chukotka Peninsula. Buldir Island (52°23'N 175°31'E), the westernmost study site, is approximately 2000 ha, and the most isolated of the Aleutian islands, being the sole landmass in a 200 km pass (Byrd & Day 1986). Approximately 140.000 Least. 280 000 Crested, and 30 000 Whiskered Auklets nest on the island (Byrd et al. 2005). although these numbers should be treated with caution as there are currently no reliable methods to census auklets (Sheffield et al. 2006; Renner et al. 2010). Alien predators such as foxes or rats were never introduced to Buldir, and so it is one of the most diverse seabird breeding colonies in the world (Byrd & Day 1986). While Buldir is volcanic, both volcanoes on the island have been inactive in the Holocene (Coats 1953; Wood & Kienle 1990), and vegetative succession may be restricting access to breeding sites, causing the auklet populations to decline (IL Jones pers, obs., sensu Roby & Brink 1986b).

Approximately 120 km east of Buldir lays Kiska Island (52'08'N, 177'36'E), the largest island in the Rat Islands group of the Aleutians (28,711 ha). The large auklet

colony is situated at Siriua Point at the extreme northem ling of the Island and at the base of Kiaka Volcano. The colony covers 1.8 km<sup>2</sup> (Joens *et al.* 2001), with 1.16 million Least and 332,000 Crested Auklets breeding, although there is demographic evidence that the Least Auklet population is declining (Byrd *et al.* 2005; HL Mijor *et al.* upublished data). The colony is situated on the north face of active Rika Volcano, and on a parasitic lives cone that emerged in 1962 (Coats *et al.* 1961; Miller *et al.* 1998) creating abundant auklet breeding habitat (Jones & Hart 2006). Arctic fores were introduced to Kiaka for fur farming in 1885 (Johbrook & Walker 1925), and removed in 1966-1987 (Deines & McClellan J97a, b). Horwy ratis were introduced accletally during military occupation of the Island from 1941-1946, are still present (Murie 1959; Bond *et al.* 

Garebic liabid (51'4/647, 137'45'5'0, 370 km east of buildi) is the northermost, and largest (67,200 hi) sland in the Delarof hiands group of the Alexitans, and also home to an active volcano. Gareliol Volcano is one of the most active volcanoes in the to alexitans, singuing several lines in the late century (Coomb et al. 2000). It is home to large auklet colony, estimated at 1866,000 Crested and 402,000 Least Auklets (Byrd et al. 2005). Article Toeses were introduced in 1925, and eradicated in 1996 (Panagi 1996). Gareloi is now likely the largest auklet colony in the Alexitan Islands, with > 1 million being present (Done's Net 2006).

Kasatochi Island (52°11'N, 175°31'W, 585 km east of Buldir) in the Andreanof Islands group of the central Aleutians was a 287 ha Island home to 20,000 Crested and

15,000 Least, Auklets until a catastrophic emption of Kastanchi Volcano in August 2008 buried the island in pyroclastic lava floes and ash (Williams et al. 2010). Although the eruption occurred after most auklets had finished breeding, most breeding crevices remain inaccessible as of the 2010 breeding season. Arctic foxes were present on Kastanchi from 1527.1991 [Charl et al. 1996].

St. George Island (56°35'N, 169°35'W) is a 90-km<sup>2</sup> island in the Pribilof Islands in the Bering Sea with a high diversity of breeding seabirds (Preble & McAtee 1923), and an estimated 250,000 Least Auklets breeding in 1977 (Hickey 1977). At the Ulakaia colony, a photographic count revealed about 88.000 birds (44.000 pairs; Renner & Renner 2010), a decline from the 129,000 birds reported in the 1970's (Hickey 1977). Vegetative succession, which cuts off access to breeding crevices, and predation by native Arctic foxes have caused the number of auklets to decline in the last century (Roby & Brink 1986b), but because of the challenges of censusing auklets at breeding colonies, it is not possible to compare the two counts directly (Renner & Renner 2010). St. Paul Island (57\*11'N. 170\*16'W) is a 104-km<sup>2</sup> island also in the Pribilof Islands. The island is home to more than 23.000 Least Auklets (Hickey 1977). The Pribilofs are near the continental shelf break (Hunt Jr. et al. 2008), and while sea ice reached the Pribilofs during most winters before 1977, sea ice extent has been considerably variable more recently, and now frequently reaches its southern extent north of the islands (Stabeno et al. 1999; Overland & Stabeno 2004).

Finally, Cape Unylahgen (Bussian: Mace Yatema) on the Chukotik Peninitula (6423%), (23754%) is on the eastern end of Bezympannaya Bay, and is one of only four mainland aukket colonies on Chukotka. 1.2 million Least and Crested Aukkets netted here in the late 1950s (Konyukhov et al. 1998). This area of the northern Bering Sea is covered by sea ice annually (Fettere et al. 2010).

## Thesis outline and rationale

In this thesis, I investigate climate-mediated control of reproductive success (Chapter 3) and chick diet (Chapter 3), and attempt to relate foraging patterns of auklets (Chapter 5 & 6), and other life history stages (moult, Chapter 4) to oceanographic conditions in the Bering Sea.

# CHAPTER 2: REPRODUCTIVE SUCCESS OF PLANKTIVOROUS SEABIRDS IN THE NORTH PACIFIC IS RELATED TO OCEAN CLIMATE ON DECADAL SCALES

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#### Abstract

Growing evidence indicates relationships between seabird demography and both large- and small-scale variation in climate and oceanography, yet few studies have examined multiple species and locations simultaneously. As secondary consumers, least, whiskered and crested auklets (Aethia pusilla, A. pygmaea and A. cristatella), congeneric planktivorous seabirds endemic to the Bering and Okhotsk seas, are expected to respond to changes in ocean climate due to their low trophic positioning. From 1990 through 2008, we measured reproductive success (productivity) and breeding phenology (mean hatching date) of auklets at Buldir, Kiska and Kasatochi, three islands spanning 585 km across the Aleutian Islands, Alaska. A model including island, species and the winter Aleutian Low Pressure Index best explained productivity, with reproductive success decreasing among all species with increasing ALPI (B = -0.273 ± 0.0263 S.E.). likely through control of water temperature and prev (zooplankton) availability. Auklet productivity also increased with increasing winter sea surface temperature (SST) in the western North Pacific and western Bering Sea (and correspondingly decreased with increasing SST in the Gulf of Alaska) and was correlated negatively with spring sea level air pressure in the North Pacific. These responses are reflective of positive values of the Aleutian Low. Despite the short time span of our data, we found similar correlations between climate and auklet productivity among all species and islands. Together, our results suggested that ocean climatic conditions and reproductive success of planktivorous auklets are related significantly.

#### Introduction

Climate and oceanographic conditions in the North Pacific Ocean covary on decadal time scales or longer (Overland et al. 1999; Biondi et al. 2003), often decadal time scales or longer (Overland et al. 1999; Biondi et al. 2003), Biondi et al. 2001; d'Orgeville & Peitier 2009). A well inown regime shift occurred in 1976-77, with warmer sea surface temperature (IST), lower sea level pressure (SLP), and a more intense Alculan low pressure system (Hare & Manita 2000; Roldiover et al. 2005). In the North Pacific and Bering Sau, this shift resulted in greater climatic variability (Rond et al. 2005), Hare & Elin 2005, Hondow et al. 2005, greater statification of the water column, and increased primary productivity (Trenberth & Hurrell 1994, Iida & Saitoh 2007). Other purported ecosystem shifts have occurred in 1989-1990 and 1989-1999, but thes patiative shifts in system state have not been well documented (Overland et al. 1999): Hare & Manta 2007).

Seahn's, as compacious, generally unequioted, scondary and tertary consumers in marine systems, provide a unique opportunity to investigate coupled climate-ecosystem variation [Durant et al. 2009]. Indeed, the relationship between ocean climate, ranging from direct measures of SIT to multivariate climate indices (e.g., Pacific Decadul Oscillation, Mantua & Hare 2002), and seabird breeding performance has been investigated frequently in the North Pacific (e.g., Gjerdrum et al. 2005; Aronaham & Sydeman 2006; Buyle et al. 2006; Lawres et al. 2007; Some fort agenitization of the state of the source state al. 2007. relationhips between large-scale climate indices and seabird demographic rates (Kitaysky & Golubova 2000, Byrd et al. 2008, Sandvik et al. 2008), whilst others do not (Wanless et al. 2009). In the North Pacific, variation in seabird breeding performance is thought to result from "bottom-up" climate forcing of seabird prey abundance (Lehodey 2004).

While linkage between climatic and oceanographic variability has been found in many marine organisms (Beamish et al. 1997; Mantua et al. 1997; Chavez et al. 1999; Hunt Jr. et al. 2002; Durant et al. 2006; Hynt et al. 2006; Hunt Jr. et al. 2008), most studied so not address spatial components in climatic and oscenargeniki variability. Integrating data over large areas can mask smaller-scale variation over space and time. More recently, there has been a focus on examining relationships between seabirds' demography and oconographic variability or spatial terms, providing a better understanding of how geographic patterns of climate affect marine animals (Byrd et al. 2008; Vatamal et a 2009).

Auklets (Charadmönres, Alcidae: *enthio*) are a group of planktivorous seabilds occurring in the Bering and Ohkots Seas that consume and provision offspring with zooplanktsn, primarily euphausild crustaceans and large coopeods (Day & Byrd 1989; Filint & Golowin 2002; Gail et al. 2006). Annual reproductive output is reflected by success in the hatching and rearing offspring from a single egg (Byrd & Williams 1993; Jones 1993a, b). Here, we test the hypothesis that the productivity (and timing of breeding) of auklets covaries with occan climate, and that the corolation is constituent. among species and Breeding colonies in the western Akustan halans. We considered the correlation between productivity, large-scale ocean climate indices (Akustian Low Pressure Index, PCO), and direct measures of sea surface temperature and sea level at pressure based on satellite data for three auklet species: least, whishered and cretted auklets (Aethia pusility, A pygmozo, and A cristaterilo) breeding at Budir, Kiska, and Kastatochi in the central and western Akustan fusiland, Asakaa (Figure 2-1). Our objectives were to: 1) quantify relationships between auklet reproductive success and indices and direct measures of cost and indice; 2) electric hoceanographic regions where productivity-climate correlations are strongest; 3) examine the covariance in productivity among different alund threeding colonies and among different auklet species; and 1 to better understand underlying mechanisms determining demographic variation in planktives sushel's poecies in the North Pacific.

## Methods

## Auklet productivity

Each year, we checked a sample of auklet breeding nest sites (rock crevices) from late May until early August, encompassing the entire breeding season for all three species using established protocols (for details seeFraser et al. 1999; Hunter et al. 2002; Major et al. 2006). To account for potential variability among islands, years, and observers, one trained observer was stationed on each island in each year to ensure consistency. Within each year, the same individual would check the same nest sites, and all observers used the same methods and criteria (below) for determining nest success (Fraser et al. 1999: Major et al. 2006). We visited breeding sites every 4-7 days, and determined their status (empty, egg, chick, adult, unknown). New sites were located and included in the sample each year, to account for previously monitored sites that were no longer in use, but the majority of crevice nests in most years were sites that had been monitored in previous years. We assumed that our annual sample of breeding sites monitored at each colony was representative and reflective of productivity at each colony site. We considered sites active when we found an adult on two consecutive visits, or we saw an egg. We scored sites as successful in fledging a chick if the chick disappeared after 25 days (Least Auklets, Roby & Brink 1986a), 26 days (Crested Auklets, Fraser et al. 1999) or 32 days (Whiskered Auklets, Hunter et al. 2002).

## Timing of breeding

We estimated date of harthing for a sub-sample of crevice breeding sites monitored. We included only crevices that we scored as an egg on one visit, and as a chick on the very next visit (4-7 days later), and we assumed hatching date to have occurred at the midpoint between the two visits. We then used the mean hatching the reach specification combination in each year as a measure of timing of breeding.

## Climate and oceanographic data

We used multiple large-scale atmospheric and oceanographic climate indices, and evaluated their relationship to auklet reproductive success. While metrics of climate are interrelated, as might be expected, we chose a variety of indicators since no one index reflexts the variability of the North Pacific adequately (Bond et al. 2003). The Adduction Low Pressure index (ALPL, Beaminh & Boullon 1995; Beaminh et al. 1997) is the anomaly from the 1590-1997 mean of the area with pressure 5 100.5 kPa over the area 20-70%, 120°C-120°W; positive ALPI values indicate a relatively strong Alexian lowpressure system. ALPI is centered on our study area (approximately 51.53%), 175°C-175°W). The related North Pacific Index (NPI) is the area-weighted sea-level pressure over the region bounded by 30°A 65% and 150°C-140°W (Trenberth & Hurrell 1994). The Pacific Decadd Unitation PDOI is the areaponaly (Boo 100 Aprox) (Boo 160 oscillation based on the leading principle component of sea surface temperature (S3T) north of 20°H (Mantue et al. 1997). Finally, the North Pacific Gyre Oxillation (NPGO) is derived output from a model; it represents the second principal component of sea surface height anomalies measured over the same areas as the PDO (D) Lorenzo et al. 2008). Several of these indices (PDO, ALP), APG have been related to auklet demography (Jones et al. 2002). For sea-surface temperature (SST), we used the 4 km AVMRP Rhihlmed version 5 dataset (available at

http://www.node.noaa.gov/SatelliteData/pathfinder4km/available.html). We measured SST in s 50m<sup>2</sup> radius around each colony reflecting the likely foraging range of auklets (Obst et al. 1995; Thayer et al. 2008). Wolf et al. 2009b). We assessed SST during June and July when auklets are present at the breeding colonies. For our geospatial approach (see below), lurge-scale climate data, (SST and SLP) over the North Pacific were taken from the KCH-NACH analysis (Slawy et al. 1996).

## Statistical procedures

All statistical tests were performed in SPSS 16.0.2 (SPSS Inc., Chicago, USA), and were weighted by the sample size of next sites studied each year for each species at each sland. To investigate whether there were linear trends in productivity or timing of breeding over time, we used a linear regression for each species-sland combination. We used nonparametric correlations (Spearmar's p) to look for covariance among and between species and blands (Bmd et al. 2008. To control for multiels consultions. we used the False Discovery Rate (Benjamini & Hochberg 1995), which is less restrictive and more powerful than a Bonferroni-type fumly-wise correction (Sarcia 2004; Grozbols et al. 2008). We use p < 0.10 to determine significance. Setting a higher of lovel is desired when the sample size (in this case, number of years) is low, to increase the power to detect a relationship when one is present (Lebreton et al. 1992; Field et al. 2004; Grozbols et al. 2008). This approach has been used regularly in other studies of stabild's in the North Partic (e.g., Arsham & Sydeman 2004; Byder (al. 2008).

We used a three-step process to investigate reliationships among occan clinate and aukier productivity. First, we used Spearman rank correlations to search for potential relationships to include in a statistical model (see below). Scood, as this analysia does not account for spatial variability in variables (e.g., SST and SJP), we examined spatial correlations to provide inference regarding spatial variability (Szép & Maller 2005; Schneeder *et al.* 2009; Watanuki *et al.* 2009); we limited this analysis to the Budiar and Eastatoch time series which were > 1Dy in duration. Spaarman correlations coefficients between reproductive success and SJP or SST were mapped over the North Pacific and Berling sca and raves of significant correlations (or 0.3D) were delimeted. We used generalized linear models with a binomial logit-link function (a logistic regression approach), to select the most important climate variables. Variable selection was based upon quasi-Akake's thormation Criteria adjutated for smill sample isse (QAC) and extra-binomial variations by including an estimate of model deviance (2

approach rather than a null-hypothesis testing because the variation in the dependent variable (productivity) was generally small (9-16% among all species and all islands: Grosbois et al. 2008). We considered the model with the lowest QAIC- value to be the best fitting model to the data (Burnham & Anderson 2002). We constructed 12 a prioiri models composed of biologically plausible combinations of five variables of interest, including a Null Model (intercept only) and a Global Model that included all additive terms and their interactions for species, and islands. Once the best-fitting model was identified, the climate and oceanographic covariates of interest, and biologically important interaction terms were added to subsequent models to examine the effect of climate, a similar procedure to the modelling of recapture rate and then survival rate in mark-recapture studies (Lebreton et al. 1992; Grosbois et al. 2008). We only included potential climate covariates that were correlated significantly with auklet reproductive performance in our preliminary assessment (see above). This decreases the number of candidate models, and reduces the risk of multi-collinearity in covariates (Grosbois et al. 2008). Models with  $\Delta QAIC_c > 2$  were considered to have substantially less support, and overall model support was assessed using Akaike weights (wa Burnham & Anderson 2002). This was done on both the full dataset, and again using data from Buldir and Kasatochi only, as in the geospatial analysis. This is because the dataset from Kiska is shorter (7 years), and because Kiska is anomalous in that is also has introduced predators that affect auklet reproductive performance. Finally, we used the estimated marginal means (EMMs) from the top-ranked generalized linear model to examine

differences among species and islands, and differences were considered significant if 95% confidence intervals did not overlap.

#### Results

We studied between 23-215 breeding pairs of each species on each sland in each year (total = 7733 breeding sites over all years and all species, Appendix 1, Figure 2-2). Aukle productivity ranged from 0.10 to 0.45 chicks/nest for most species for most years, with 2001 and 2002 being unusually low at Kiska and 2003 being an unusually low at bluich (Appendix 2).

Linear regression identified only one significant trend in reproductive success (or phenology Appendix 2), that being an increase in reproductive success over time for least auklets at Kiska, our shortest dataset (p = 0.012, all other p < 0.11), we used raw data, not determedavious. For further comparison.

Reproductive success was correlated more frequently than phenology, and all significant correlations were positive (in phase; Table 2-2). There was a general pattern of correlation among species within Islands, and using our shortest time series (Kika), nine of ten correlations are positive despite the small sample size (n = 7 years), providing reasonably strong evidence of covariation (Figure 2-2).

Timing of breeding was rarely correlated among species or islands; crested auklet timing of breeding at Buldir and Kiska was correlated positively (Table 2-1).

#### Relationships to climate indices

Productivity was related frequently and significantly to cean climate indices (Table 2-3). Of the climate covariates we considered, the Aleutian Low Pressure Index (ALPI) and the winter and spring North Pacific Gyre Oscillation (NPGO) were most often related to audite productivity.

## Geospatial mapping of climate and reproductive success correlations

In frour cases, genepatial correlation analysis found that reproductive success was correlated with winter (December-January February) SST (Figure 2-3), but in different areas of the North Pacific. Productivity of least and created auklets at likeliar was negatively correlated with SST in the Guil of Jaaka region, while that of whiskered auklets on bulkir was positively related to SST in the western Bering Sea. The spatial correlation pattern for least auklets on Bulkir is characterized by a negative correlation along the American west coast and a positive correlation in the central North Pacific, reminicent of the SST pattern of a negative PDC (Mattua ceri Casy), albrough the PDC pattern does not have another action center with the opposing sign in the western North Pacific. Reproductive success of created auklets on Kasatochi were positively relationship between reproductive success of created auklets on Kasatochi were positively relationship between reproductive success and spring (March, April, May) S5T was for whishered auklets between the gattern was denoted by the YDC pattern does not between the pattern was identiced. The relation to the for witter the author by another pacific Ceasa between 40.570 (Figure 2-1). The only relationship between reproductive success and spring (March, April, May) S5T was for whishered auklets the degreed and the pattern was identiced to that for witter SST. This pattern is similar to the second mode of SST variations (Bond et al. 2003), which is closely related to the NPGO (DI Lorenzo et al. 2009). This is consistent with the relatively high correlation between productivity and clinitate indices (Table 2-3). No significant relationships were found between reproductive success and winter SPLP, and only two for spring SLP. Whiskered auklet reproductive success on Buldir was negatively related to SLP in the eastern Pacific, while that of least auklets on Kasitochi was equilively related to the spring SLP in the western Pacific (Figure 2-4).

#### Factors affecting auklet productivity

Based on the results from our examination of correlation coefficients and spatial correlation analysis, we included only ALPL, NRGO (winter and spring) along with local breeding season SST as covariates in the generalized linear models. The best-fitting model for productivity of all auklets from 1990-2008 included differences among species and islands, ALPI and the island × ALPI interaction term. No other model had AQAU< < 2, and the top model received 72% support, 3 times more than the second-best model atter adjusting for < 25.5 (Table > 4.2). Parameter existints are represented in Table > 25.5 (Table > 4.2). Parameter existints are represented in Table > 25.5 (Table > 4.2). Parameter existints are represented in Table > 25.5 (Table > 4.2). Parameter existints are represented in Table > 25.5 (Table > 4.2). Parameter existints are represented in Table > 25.5 (Table > 4.2). Parameter existints are represented in Table > 25.5 (Table > 4.2). Parameter existints are represented in Table > 25.5 (Table > 4.2). Parameter existints are represented in Table > 25.5 (Table > 4.2). Parameter existints are represented in Table > 25.5 (Table > 4.2). Parameter existints are represented in Table > 25.5 (Table > 4.2). Parameter existints are represented in Table > 25.5 (Table > 4.2). Parameter existints are represented in Table > 25.5 (Table > 4.2). Parameter existints are represented in Table > 25.5 (Table > 4.2). Parameter existints are represented in Table > 25.5 (Table > 4.2). Parameter existints are represented in Table > 25.5 (Table > 4.2). Parameter existints are represented in Table > 25.5 (Table > 4.2). Parameter Parameter > 4.5 (Table > 4.2). Parameter = table > 5.5 (Table > 4.2). Parameter = table > 5.5

Using data from only Buldir and Kasatochi, the top two models included effects of species and AIPI, with one including their interaction. These models were almost equally supported, but were more than 7 times more supported than the third-ranked model after correcting for  $\ell = 4.2$  Table 2-6.

Based on estimated marginal means (LMMs) from the full dataset, Budir and Ksaatochi had similar reproductive success rates during the study and both were higher than Kiska. Among islands, created and whikered auklets had higher reproductive success than losat auklets [Lable 2-7]. An increased ALP was associated with decreased productivity among all species on all listing (§ n-0.273 ± 0.023) [S.L.F. Jeure 2-8].

## Discussion

Our results indicate that auklet reproductive success varied with large-scale climate indicate, and that even with a short time series for multi-indic comparisons (7 years), the majority (9/20) of correlations were in phase. The combination of only 7 years' data from Kiska, and the potentiary comming ended phase interformation of control of complicate the interpretation, but performing analysis using data from only Buddin and Kasatochi, combined with independent spatial correlation analysis yielded similar patterns. We note, however, that the oscillation between 'warm' and 'cold' phases of the Pacific Cosan may be masked or accentuated by increasing global temperatures (d'Orgovelle & Petier 2003), and so our correlations should be treated as minimum estimates.

Interestingly, we found no evidence of linear trends in our reproductive success or phenology data, contrary to that found in piscivorous seabirds in the Pribilof Islands (Byrd et al. 2008) and North Atlantic (Gaston et al. 2005; Moe et al. 2009). It is likely

that our relatively short time series overall (19 years) was insufficient to document potential climate-linked trends in productivity.

Reproductive success tended to be more correlated among species on the same island than for any single species among islands. Least and created auklets tend to provision their chicks with diets consisting of euphausids and calamid corpopads (Bédard 1969a, Day & Byrd 1989; Huart J. & Farrison 1990; on the observed pattern suggests that general prey availability differences among breeding sites was more relationships between climate indices or local oceanographic conditions and timing of breeding in the Acidase, and found tends for earlier breeding (Gaston et al. 2005; More et al. 2009; Just Preveding (Byrd et al. 2006; Warless et al. 2009), or to relear over the Acidase and Sund tends for earlier breeding (Gaston et al. 2005; More et al. 2009; Just Preveding (Byrd et al. 2006; Warless et al. 2009), or to relear over the Acidase and Sund tends to the Acidate statistication (Acidates & Syleman 2004; Durant et al. 2004; Mais study). While snow cover, and consequently air temperature, may delay adulet neiting at northern colonies (Seal) 1975; More et al. 2009; Justeng et al. 2006; Warless et al. 2006; Maries et al. 2005; More et al. 2009; Justeng et al. 2006; Justeng et al. 2006; Warless et al. 2006; Maries 1975; More et al. 2009; Justeng et al. 2006; Maries 1975; More et al. 2009; Justeng et al. 2006; Warless 1975; More et al. 2009; Justeng et al. 2007; Justen

#### The influence of climate on reproductive performance

Overall, a higher Albutian Low Pressure Index (ALPI), which corresponded with a stronger, more intense Aleutian low-pressure system, was associated with decreased productivity in least, whiskered and crested auklets across their range in the Aleutian liabd, Assia, albuton the magnitude of this relationship was variable among species

and islands. A stronger Aleutian low is supposed to correspond with a larger spring algal bloom in the Bering Sea (lida & Saitoh 2007), but also possibly with a decrease in zooplankton biomass due to cold temperature limitation (Walsh & McRoy 1986; Huntley & Lopez 1992; Pinchuk et al. 2008). In the eastern Bering Sea, warmer SST (and positive values of PDO) were predicted to result in a decrease in zooplankton biomass in surface waters during the auklets' breeding season (Coyle et al. 2008; Jin et al. 2009), likely caused by the copepods' early descent in the water column to undergo diapause (Miller et al. 1984; Miller & Nielson 1988). In the northern Bering Sea, auklets are less selective in the prev species they choose when zooplankton abundance is low (Springer & Roseneau 1985: Hunt Jr. & Harrison 1990; Russell et al. 1999). Under these conditions, prev species that are not typically exploited by auklets because of their lower energy content (e.g., Calanus marshallae) are more prevalent in adult and chick diet. The same could be true in the Aleutian Islands during a stronger Aleutian low, leading to a decrease in productivity. Survival of auklet chicks has been shown to be related to the quality of previtems delivered by their parents (Gall et al. 2006). In the northern California Current System, Cassin's auklets have low reproductive success in warm years when biomass of their main prev species, Neocolonus cristatus, was early, and consequently not abundant in chicks' diet (Sydeman et al. 2006). We do not believe that such a mismatch occurred with Aethia auklets in the Aleutian Islands, as chick diet composition has been relatively constant over our study period (authors' unpubl. data).

Aside from indirect effects of ocean climate via effects on productivity, direct effects are also possible and non-mutually esclusive to indirect effects Jones et ol. 2007). Heavy rainfall during early chick rearing can flood nesting crevices, resulting in chick death (author') personal observation). The relative importance of such local, atypical effects as compared with pery availability is unknown.

Another direct affect of stormy weather could operate via increased wave action, that is thought to decrease availability of zooplankton to prediators (Wroblewski & Richma 1979). Unfortunately, land based field observational studies (such as ours) alone do not provide direct information on plankton availability, and are therefore inadequate to confirm the relative roles of direct and indirect effects of ocean climate on auklet breeding success.

#### ALPI and NPGO

Conveniently, our three study islands lie nearly at the centre of the geographic area used for ALPI calculations. ALPI is also measured over the period December-March, and other studies have found significant relationships between seabler deropolative performance and winter climate (Jabanam & Sydeman 2004; Jacobi et al. 2006; Byrd et al. 2008; Sandvik et al. 2008), presumably through climate-mediated controls on prey development, abundance and distribution (Ditaryky & Goldonou 2000; Frederiksen et al. 2006; Byrd et al. 2008). We previously found that adult survival in whiskered asklets counsel with ALPI are well, through the face (negative effects of a Stormy North Pacific) or indirect effects (i.e., bottom-up control of food supplies, Jones et al. 2007). This contrasts partly with recent findings that reproductive success in Cassin's auklet (Ptychoramphus aleuticus) is driven by local factors rather than large-scale climatic variables (Wolf et al. 2009b). Cassin's auklet chicks are fed only at night by their nocturnally-active parents, grow slower and take much longer to fledge than Aethia auklets (Ydenberg 1989), meaning that there is a larger timeframe over which nocturnal species' reproductive success may be affected by climatic variation. Cassin's Auklets breed from southern California to the Aleutian Islands, and so large-scale climate variation is likely to be inherently more variable over their range than that of Aethia auklets in our study. Pacific-wide climate patterns affect the coasts of California, British Columbia, and Alaska differently and span several oceanic domains (California Current, North Pacific Current, Alaska Coastal Current). Our study of auklets in the Aleutian Islands spans one oceanic domain where large-scale climate effects would likely be more consistent than those experienced by Cassin's Auklets in the study by Wolf et al. (2009b).

The NPGO is an index of sea surface height and temperature anomalies over 110-180°W and 25-62°N (Di Lorenzo et al. 2008), and tends to explain more of the SST anomaly than the PPOO [Bond et al. 2003]. It is controlled largely by basin-wide variations in upswelling and advection in the ocean, and therefore influences nutrient abundance and salinity directly (Di Lorenzo et al. 2009), which in turn effects the abundance of advance (Di Lorenzo et al. 2009). In all cases where the relationship

between Aleutian auklet productivity and either winter or spring NPGO was significant. the relationship was negative such that a higher NPGO index corresponded with lower reproductive success. A high NPGO is associated with a stronger Aleutian Gyre in the Gulf of Alaska, and a stronger North Pacific Current, the main warm-water current that runs west-to-east along the south of the Aleutian chain at about 40-50°N. Furthermore, the NPGO mode has been strengthening since about 1993 (Bond et al. 2003; Di Lorenzo et al. 2008), and may be the result of anthropogenic climate change (Di Lorenzo et al. 2008). Furthermore, the NPGO and the Aleutian low-pressure system are linked to El Niño Southern Oscillation (ENSO) dynamics. There is an atmospheric teleconnection between the Aleutian low and ENSO, and the North Pacific Oscillation (which is related to the NPGO) leads ENSO by approximately 8-12 months (Di Lorenzo et al. 2010). Biologically, this means that there should be concern about the Aleutian Islands population of auklets, as both increased ALPI and NPGO are related to decreased reproductive success, likely through a bottom-up mechanism of limiting prev availability during either the pre-breeding period (Schroeder et al. 2009), or during chick rearing (Gall et al. 2006). Combined with vegetative succession (Roby & Brink 1986b), and the detrimental effects of rats at Kiska (Major et al. 2006), the effects of oceanography or climate may be masked or dampened by colony effects.

## Geospatial mapping of climate and reproductive success

The strongest geospatial correlations provided evidence that reproductive success of least auklets on Buildr might be related to the PDD based on winter SST correlations (Figure 2-3). Similarly, the correlation map for whiskered auklet reproductive success on Buildr and winter/spring SST (Figure 2-3) resembled by Victoria mode of SST anomalies (Bond et al. 2003). The Victoria mode is the atmospheric expression of the NPGD in the North Pacific Oscillation, an atmospheric SJP pattern, athough we again found weak correlations possible because of unit ont time scale.

The negative relationship between aukler productivity and winter SST in the Gulf of Alaska further suggests a bottom-up mechanism. Warmer SST is related to earlier timing of life history concerns by cooperado (Mukcias *et al.* 2007), Indiangi dipanase, which may render the copepods unavailable to auklets (Goldblatt *et al.* 1999; Richardson 2008). Data from continuous planistor recorders in the Neuth Facilito: showed a marked decrease in copepod abundance in writer (December-March), abhough smaller numbers may remain accessible to auklet to scalable predactor (For *et al.* 2010).

Critical to an understanding of geospatial patterns of demography/oceanography relationships in seabrick is an understanding of the birds' Spatial distribution withing the time of interest, which in our case is winter and spring. Knowledge of the non-breeding distributions of Aethia auklets, especially from the ice-free Aleutians, is poor. Whiskered auklets are believed to be non-migratory, spending winters in the vicinity of the breeding colony (Steppinger 1855; Konyukhov & Zubakin 1947; Jubakin & Konyukhov

1954; Gibson & Byrd 2007). Created auklets winter in the eastern Alexians, particularly in productive upwelling areas used a Unimak Pass (Renner et al. 2006; Sydenm et al. 2010), and least auklets are more abundant in the western Pacific during the winter and sporing (Vyatkin 1961; Sydemm et al. 2010), and some winter in the ses of Japan off the coast of Primore, Rusia (Shuntor 1965; Veichhanin 1977a; Rondrafayer et al. 2000), although the extent of mixing between Russian and North American populations is unknown. Nevertheless, large-scale clinate patterns would be expected to influence the species differently during the winter, but as the relationships between ALPI and productivity among all species on all blands was similar, the differing effects may manifest in a similar way.

#### Other factors affecting reproductive success

All species' reproductive success was negatively related to the ALP across all ialands generality, but local factors must alion have been important. These factors included variable abundance and predation by aikan predation such as Glaucous-winged Guid (aurus glaucecase), Peregrine Factors (facto peregrinius), and/or fabil Eagles (Helioretus leucocephala), and rainstorms during peak hatching periods when chicks were vulnerable to chilling, and earthquakes collapsing crevices. An extreme example of local effects was the August 7, 2008 eruptions at Kastotoh buriet the entire auket colony ale funder thousands of toos of hot voicanic abi, entombing late auket chicks and any accompanyture bath (Villiams er 2010). Acotter example is introduced

Norway rats that have been present at Kiska since the 1940's (Murie 1959), which were thought to be implicated in severe reproductive failure in least auklets there in 2001 and 2002 (Major et al. 2006). Auklets form a considerable portion of the rats' diet, and caches of dead birds were found annually, being especially large in the years of breeding failure (Major et al. 2007).

## Conclusions

Our results indicated that ocean climate and associated effects on ocean productivity and storminess influences variation in Aleutan Aethia auklet reproductive performance. Colony-specific effects, such as variation in the abundance and behaviour of naturally occurring and introduced predators, vegetative succession, and even seismic and valanic activity must also be considered.

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# Figures





sites at Buldir, Kiska, and Kasatochi islands, Aleutian Islands, Alaska.



Figure 2-2. Productivity anomalies (differences from the overall mean) of least, crested and whiskered auklets at Buldir, Kiska and Kasatochi, Aleutian Islands, Alaska during 1990-2008.


Figure 2-3. Biogeography of Spearman correlation coefficients between least, crested and whistered auklet productivity at Budir and Katastochi and whiter sea-surface temperature (S51). Contours indicate significant (p < 0.10) Spearman correlations, with the assumption that each vers's productivity measure was independent.



Figure 2-4. Biogeography of Spearman correlation coefficients between least, created and whiskered aukter reproductive success at Buildr and Kasatochi, and spring sea-level pressure. Contours indicate significant (p < 0.10) Spearman correlations, with the assumption that each wars' productivity measure was independent.



# Aleutian Low Pressure Index

Figure 2-5. Productivity (number of fledglings/number of eggs laid) of least, crested and whiskered auklets at Buldir, Kiska, and Kasatochi during 1990-2008 showing the generally negative relationship with the Aleutian Low Pressure Index (ALPI), least auklets. Buldir (slopes: -0.0096). Kiska (-0.0657). Kasatochi (-0.0045); crested auklets, Buldir (-0.0281). Kiska (-0.0320). Kasatochi (-0.0169): whiskered auklets. Buldir (-0.0372).

# Tables

Table 2-1. Spearman's p correlation matrix for breeding phenology (mean hatch date) of least (LLAU), created (CRAU), and whiskered auklets (WHAU) at Buildi, Kuka, and Kaatacchi in the Alextian Islands from 1990-2008. Significant correlation after correction for false dioxeeved date (see tota) are indicated with \*.

	Buldir	Kiska	Kasatochi	Buldir	Kiska	Kasatochi
	LEAU	LEAU	LEAU	CRAU	CRAU	CRAU
Kiska LEAU	-0.239					
Kasatochi LEAU	0.458	-0.373				
Buldir CRAU	0.581	-0.108	0.378			
Kiska CRAU	0.667	0.200	0.616	0.900*		
Kasatochi CRAU	0.388	-0.248	0.519	0.208	0.051	
Buldir WHAU	0.452	0.673	-0.196	0.445	0.821	0.092

Table 2-2. Spearman's p-correlation matrix for productivity of least (LEAU), crested (CRAU), and whiskered audiest (WAU) at Budley, Kiska, and Kasatechi in the Alexitan Islands from 1990-2008. Significant correlations after correcting for false discovery date (see text) are indicated with \*.

	Buldir	Kiska	Kasatochi	Buldir	Kiska	Kasatochi
	LEAU	LEAU	LEAU	CRAU	CRAU	CRAU
Kiska LEAU	0.206					
Kasatochi LEAU	-0.260	0.216				
Buldir CRAU	0.569	0.090	-0.249			
Kiska CRAU	0.581	0.643	-0.601	0.219		
Kasatochi CRAU	0.162	0.865*	0.373	0.127	0.543	
Buldir WHAU	0.375	0.793	0.018	0.469	0.402	0.565

Table 2-3. Spearman's p correlations for productivity and climate indices for least (LRU), created (CRU), and whistered auklets (WAU) at Budir, Kiska, and Kasatochi in the Aleutan Islands from 1990-2008. Significant correlations after correcting for false discovery date for text are indicated with \*.

		NPGO	NPGO	NPI	NPI	PDO	PDO	1.2
		DEC-	MAR-	DEC-	AUG-	DEC-	AUG-	
	ALPI	FEB	MAY	MAR	APR	MAR	APR	SST
Buldir								
LEAU	-0.161	-0.127	-0.148	0.258	0.35	-0.532	-0.411	-0.436
Kiska								
LEAU	-0.707	-0.729	-0.528	0.021	0.029	0.281	0.054	0.406
Kasatochi								
LEAU	-0.103	0.283	0.271	-0.133	-0.278	-0.128	-0.142	0.368
Buldir								
CRAU	-0.313	-0.387	-0.297	0.227	0.264	-0.192	-0.295	-0.027
Kiska								
CRAU	-0.469	-0.945*	-0.835*	0.287	0.436	0.029	-0.086	0.091
Kasatochi								
CRAU	-0.373	-0.169	-0.038	0.191	0.128	-0.122	-0.324	0.198
Buldir								
WHAU	-0.529	-0.474	-0.415	0.307	0.231	0.114	0.033	0.053

Table 2-4. Summary of the model set for predicting productivity of least, crested, and whiskered auklets at Buldir, Kiska, and Kasatochi, Aleutian Islands, Alaska from 1990-2008.

Model*	# Parameters	QAIC	$\Delta \text{QAIC}_{c}$	Wi
I+S+ALPI+I*ALPI	8	1818.624	0.000	0.72
I+S+ALPI+I*ALPI+S*ALPI	11	1820.790	2.166	0.24
I+S+ALPI	6	1825.154	6.529	0.03
I+S+ALPI+S*ALPI	8	1826.500	7.876	0.01
I+S+NPGO/DJF	7	1832.111	13.486	0.00
I+S+NPGO/MAM	7	1839.563	20.939	0.00
1+S	5	1844.406	25.782	0.00
I+S+SST	6	1845.380	26.756	0.00
I+S+I*S	8	1847.243	28.619	0.00
s	3	1854.478	35.854	0.00
1	3	1856.643	38.018	0.00
Intercept Only	1	2496.983	678.359	0.00

\*Note: Models are sorted by increasing QAIC, (quasi Akaike's information criteria corrected for small sample size) value with the most parimonisus model at the top. Akaike weight (w) is the likelihood that a given model of the model set is the best somorimation of the data. Model parameters: is listed. SectoreA. JP: Neutral Corr

Pressure Index, NPGO: North Pacific Gyre Oscillation (DJF: December-March; MAM:

March-May), SST: Sea Surface Temperature. Model corrected for c = 5.62.

Table 2-5. Parameter estimates from the top-ranked generalized linear model (Table 2-4) for productivity of least, crested, and whiskered auklets at Buldir, Kiska, and Kasatochi, Aleutian Islands from 1990-2008.

			Lower	Upper			
			95%	95%	Wald		
Parameter	β	S.E.	C.I.	C.I.	χ²	df	p-value
Intercept	0.391	0.1009	0.193	0.588	14.975	1	< 0.001
Crested auklet	0.203	0.0794	0.047	0.358	6.508	1	0.011
Least auklet	-0.274	0.0784	-0.428	-0.121	12.246	1	< 0.001
Whiskered auklet	0*						
Buldir	0.200	0.0824	0.039	0.362	5.903	1	0.015
Kasatochi	0.148	0.0849	-0.018	0.315	3.049	1	0.081
Kiska	0 <sup>a</sup>						
ALPI	-0.273	0.0263	-0.325	-0.222	108.252	1	< 0.001
Buldir * ALPI	0.168	0.0317	0.106	0.23	27.926	1	< 0.001
Kasatochi * ALPI	0.220	0.033	0.155	0.285	44.49	1	< 0.001
Kiska * ALPI	0*						

<sup>a</sup>This parameter is set to 0 because it is redundant.

Table 2-6. Summary of the model set explaining productivity of least, crested, and whiskered auklets in the Aleutian Islands from 1990-2008 using only data from Buldir and Kasatochi.

Model	# Parameters	QAIC	$\Delta QAIC_c$	w
S+ALPI+S*ALPI	6	1942.986	0.000	0.49
S+ALPI	4	1943.267	0.281	0.43
S+NPGO/DJF	4	1947.378	4.391	0.05
S+NPGO/MAM	4	1949.853	6.866	0.02
s	3	1951.288	8.302	0.01
S+SST	4	1952.190	9.203	0.00
I+S+I*S	5	1953.267	10.281	0.00
Intercept Only	1	1964.425	21.439	0.00
1	2	1965.426	22.440	0.00

\*Note: Models are sorted by increasing QAIC, [quail Akalier's information criteria corrected for small sample site) value with the most parsimonious model at the top. Akalie weight (w) is the likelihood that a given model of the model set is the best approximation of the data. Model parameters: I: taland, S: Species, ALPI. Alexidian Low Pressure Index, NPGO. North Pacific Gyre Oscillation (DIF: December-March; MAM: MAM: Model Surface Temoersture, Model corrected for c' = 4.33 Table 2-7. Estimated marginal mean (EMM) productivity for least, crested and whiskered auklets at Budily, Kask, and Kastochi in the Aleutian Islands, 1990-2008 was derived from the top-ranked generalized linear model. Differences were considered significant when confidence intervals do not overlap.

Parameter	EMM	95% Confidence Interval
Least auklet	0.45	0.43-0.47
Crested auklet	0.60	0.56-0.64
Whiskered auklet	0.62	0.59-0.65
Buldir	0.62	0.60-0.63
Kiska	0.35	0.29-0.41
Kasatochi	0.61	0.59-0.63

# CHAPTER 3: DIET OF AUKLET CHICKS IN THE ALEUTIAN ISLANDS, ALASKA: SIMILARITY AMONG ISLANDS, INTER-SPECIES OVERLAP, AND RELATIONSHIPS

# TO OCEAN CLIMATE

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## Abstract

Seabirds are effective samplers of the marine environment, and can be used to measure resource partitioning among species and sites via food loads destined for chicks. We examined the composition, overlap, and relationships to changing climate and oceanography of 3216 food loads from Least, Crested, and Whiskered Auklets (Aethia pusilla, A. cristatella, A. pygmaea) breeding in Alaska, during 1993-2006. Meals comprised calanoid copepods (Neocalanus spp.) and euphausiids (Thysanoessa spp.) that reflect secondary marine productivity, with no difference among Buldir, Kiska, and Kasatochi islands across 585 km of the Aleutian Islands. Meals were very similar among species (mean Least-Crested Auklet overlap C = 0.68: Least-Whiskered Auklet overlap C = 0.96) and among sites, indicating limited partitioning of prev resources for auklets feeding chicks. The biomass of copepods and euphausiids in Least and Crested Auklet food loads was related negatively to the summer (June-July-August) North Pacific Gyre Oscillation, while in Whiskered Auklets food loads this was negatively related to the winter (December-January-February) Pacific Decadal Oscillation, both of which track basin-wide sea-surface temperature (SST) anomalies. We found a significant quadratic relationship between the biomass of calanoid copepods in Least Auklet food loads at all three study sites and summer (June-July) SST, with maximal copepod biomass between 3-6°C (r<sup>2</sup> = 0.71). Outside this temperature range, zooplankton becomes less available to auklets through delayed development. Overall, our results suggest that auklets are able

to buffer climate-mediated bottom-up forcing of demographic parameters like

productivity, as the composition of chick meals has remained constant over our study.

# Introduction

Seability are useful models for investigating distancy resource use and overlap in a community setting. In particular, seabind nesting diets reflect availability, desinability, and perhaps competition for prey in a changing marine environment (caims 1987; Platt ed. 2007). During the breeding season, partners are teid to their breeding sites, as chicks cannot feed themselves for several weeks in most species (Hamer et al. 2002). Forced into becoming central-place foragers, adults must balance their own emergetic investment with the need to obtain and deliver food to the chick in environment where food resources are potentially limiting, and where adults may experience bottom up control (furners & Binkhand 1984).

Ecologists have long been interested in how multiple similar species coexist in stable communities (Darwin 1895), Wallace 1876), leading to the development of the concepts of "niche" and "niche overlap" (Grinnell 1917, Geisel 1955, Hutchinson 1957; Planka 1974). The idea that two or more species canont occupy the same niche in space and time ("competitive exclusion principle") Gause 1934) has led to innumerable studies of how species accommodate their ecological similarity (Hubbell 2001; Chase & Leibidd 2003). This overlap can be reduced by exploiting different resources, or by exploiting the same resources but in different areas or at different times, especially during peak resource demand (Echwernner et al. 2008). Decreasing competition would be expected to benefit individual survival and reproductive success (Hutchinson 1957), and is herefore cernal to taikes of dimorganya. The considerce of the small plankihorous auklet (Alcidae, genera Aethia and Ptychoramphus) in the Aleutian Islands (Jones 1999; all five breeding at Buldir Island, Alaska) thus presents a potential test case for evaluating hypotheses about food resource competition.

Seabilistic sould act as sentimetis of climate change over multiple spatial and temporal scales (Parsons et al. 2008; Durant et al. 2009), a possibility that has led to some studies relating demographic parameters (e.g., breending success, adult survival) to large-scale multivariate climate indices such as the North Atlantic Oscillation, Pacific aceadal Oscillation, or el NiHo Southern obcillation (e.g., lones et al. 2002; Sandwä et al. 2005). Some studies invoked a bottom-up control mechanism whereby oceanographic conditions limit prey availability (e.g., Durant et al. 2003; Irons et al. 2000; Mills et al. 2008) or cause a temporal mismatch between consumer requirements and prey availability (e.g., Highrer 2000; Gaston et al. 2007). Audite foraging may provide an ideal subject for testing the "sentime" hypothesis as foraging by these planktivores would be expected or criate clevel to secondary ocean productivity.

In the North Pacific Ocean, changes in climate and oceanography have been well documented (Overland *et al.* 1999; Hare & Mantua 2000; Bionifi *et al.* 2001). Following a marked change in climate and oceanography in 1976-77, climatic variability increased (Biond *et al.* 2003), sea surface temperature (537) increased, and sea-level pressure decreased (Hare & Mantua 2000; Rodionov *et al.* 2005). These changes had demographic consequences for several seabird species (e.g., Gjerdrum *et al.* 2003); Hashamah Sydema 2004; Chapter 27, Jones *et al.* 2007; Brit *et al.* 2008).

Least (Aethia pusilla, mean adult mass 85 g), Crested (A. cristatella, 260 g) and Whiskered (A. pygmaea, 108 g) auklets are planktivorous alcids endemic to the Bering. and Okhotsk seas of the North Pacific. They breed sympatrically in dense colonies on remote islands, are socially monogamous, and share incubation and chick rearing (Byrd & Williams 1993; Jones 1993a, b). Auklet chick diet consists mainly of calanoid copepods (Neorgianus son.) and euphausiids (Thysanoessa son.), with some amphipods, decapods, and other invertebrates (Day & Byrd 1989: Harrison 1990: Gall et al. 2006). The quality of chick diet (e.g., the amount of lipid-rich zooplankton; Roby et al. 1986) is related to chick survival in auklets (Gall et al. 2006: Sheffield Guy et al. 2009), and other seabirds (e.g., Durant et al. 2003). Reproductive success of auklets in the Aleutian Islands may be linked to foraging success through climate- and oceanographic-mediated processes (Chapter 2). To explore this possibility further, we quantified auklet chick diet over the same period as Bond et al. (Chapter 2), and investigated the relationship of auklet chick diet to indices of North Pacific climate and oceanography. Sometimes it is assumed that auklet adult and chick diets are the same, but Crested Auklet adults may feed heavily on cephalopods, which were never fed to chicks (Hunt Jr. et al. 1998); our study therefore focuses on food loads captured by adults and destined for their nest-bound chick. Previous work suggested that breeding Aethia auklets reduce inter-specific competition through dietary differences (e.g., Hunt Jr. et al. 1998) or spatial separation (Russell et al. 1999), ideas that have not been tested at multiple colony sites over time.

Our objectives were its: 1) quantify diet of Least, Crested, and Whikered Auklet chicks ampled across a SSS im span of the Alexitain blands from 1993-2006; 2) determine diet variation among species, islands, and stage of the breeding seasor; 3) examine relationships diet to key indices of annual oceanographic and climatic variability; and 4) investigate correlations between chick det and productivity as measured by flow of et al. (Oupster 2).

## Methods

## Study sites & field collection

We performed this study at three colonis: Main Takus, Buller Island (\$2:23%, 1375557; 1993-2006, Crested, Least, and Whiskerel Auklets); Srink Point, Kish Island (\$2:00%, \$1773167; 2001-2006, Crested and Least Auklets); and Tundreing Taku, Kasatochi Island (\$2:11%, 137511%); 1996-2006, Crested and Least Auklets) perdominating, while at Kaka about 10<sup>6</sup> briefs were present, mostly Least Auklets (authors unpublished data). These colonies span 585 km of the Aheutian chain (Figure 3-1), and because auklets forage within 50 km of their breeding colony (Hunz Jr. & Harrison 1990; Obst et al. 1995; Find & Goloxkin 2002); we assume that there was Ittle spatial overlap of foraging birds from different sites. There was no significant difference in 537 around auklet toolonies listen at the two; see Clinitical and Sterengenghic Indices below), we assumed that auklets from all colonies experienced similar oceanographic conditions in a given year.

Adults carry food for the chick in a throat pouch (Portenko 1934), and regurgitate this food when captured in a noose carpet or mist net (Jones et al. 2002: Jones et al. 2004; Jones et al. 2007). We sampled food regurgitated by adults throughout the chickrearing period (late June-early August) in each year. Each year, we collected food samples in the same 24-hour period over 2-calendar days during early, mid, and late chick rearing on each island, weather permitting. Early was defined as roughly when 50% of chicks had hatched (late June or early July): mid was defined as the period when all chicks had hatched (mid July, about 10 days following the first collection period); and late was defined as 10 days after the second collection (just before the start of fledging). For a subset of samples where reguralitated food was clearly delineated (e.g., not overlapping with another regurgitated food load, or where portions were regurgitated down rock crevices), we estimated the percentage of food recovered from regurgitated contents to the nearest 5%, and stored samples in 75% ethanol. We identified prev to the lowest taxonomic level possible and counts of individuals were converted into wet biomass (boreafter "biomass") using our own measurements and published data (Appendix 4), and use biomass (or % biomass) throughout. The mass of some food loads was estimated based on the proportion (±5%, as noted above) of each food load collected. To compare food load mass to adult body mass, we used published data on

adult body mass from auklets in the Aleutian Islands (Byrd & Williams 1993; Jones 1993a, b)

#### Prey availability

As a measure of prey availability, we used data from a continuous plankton recorder (OP) during the period of chick rearing (late hane to sarly August) 20002 between 173°W - 173°E and 52-64°N. Data on prey availability were unavailable south of this mars, so while some addets may forage outside the C<sup>\*</sup>R<sup>\*</sup>R<sup>\*</sup>area<sup>\*</sup>, we believe the data are sufficient to capture inter-annual variability in mesoranglankton abundance. The Pacific CPR program of the Sir Alster Hardy Foundation for Ocean Science (SAHFOS) provided the data. Nets were towed for 18 km at a depth - 15m; further details are in Richardson et al. (2006), Maximum estimated dilegaths for all three species are < 20m, and most dires are likely valabilower (Haner (2011).

## Climatic and oceanographic indices

We used multiple large-scale atmospheric and oceanographic climate indices, as no single index reflects the entire variability of climate or oceanography in the North Placific Ocean (Bond et al. 2003). The Aleutian Low Pressure Index (ALPI, Beaminh & Bouillon 1993; Beamish et al. 1997) is the anomaly from the 1950-1997 mean of the area with pressure 1030-54 woret the care 30-700, 12072; L2072, Worklew ALPI values and the pressure 1030-54 woret the care 30-700, 12072; L2072, USPC 1000; MDI et ALPI values and and the pressure 1030-54 woret the care 30-700, 12072; L2072, USPC 1000; ALPI values ALPI values and the pressure 1030-54 woret the care 30-700, 12072; L2072, USPC 1000; MDI et ALPI values 1030-54 MI values and the pressure 1030-550 means and the pressure 1030-560 means and the care and the pressure 1030-560 means and the pressure 1030-500 means and the pressure 1030-560 means and the pressure 1030-56

indicate a relatively strong Aleutian low-pressure system. ALPI is centred on our study area (approximately 51-53\*N, 175\*E-175\*W). The related North Pacific Index (NPI) is the area-weighted sea-level pressure over the region bounded by 30"N-65"N and 160"E-140°W (Trenberth & Hurrell 1994). The Pacific Decadal Oscillation (PDO) is a long (20-30 years) oscillation based on the leading principal component of sea-surface temperature (SST) north of 20"N (Mantua et al. 1997). Finally, the North Pacific Gyre Oscillation (NPGO) is the second principal component of sea surface height anomalies measured over the same areas as the PDO, but also matches the second empirical orthogonal function (EOF: a spatial statistic similar to a principal component score) of SST anomalies closely (Di Lorenzo et al. 2008). Several of these indices have been related to auklet demography in previous studies (Jones et al. 2002; Jones et al. 2007; Chapter 2). These metrics of climate are interrelated, but each is based on slightly different primary components (Trenberth & Hurrell 1994; Bond et al. 2003; Di Lorenzo et al. 2008). We used seasonal averages of three monthly means (e.g., winter: December-January-February, or DIE) for all indices except ALPL which is an annual value.

For sea-surface temperature (SST), we used the 4 km AVHRR Pathfinder Version 5 dataset (available at

http://www.nodc.noaa.gov/SatelliteData/pathfinder4km/available.html). We extracted SST in a 50 km radius around each colony site, which is the likely foraging range of auklets during June and July (Hunt Jr. & Harrison 1990; Obst et al. 1995; Thayer et al. 2009; Wolf et al. 2009b;.

# Statistical analyses

All statistical tests were conducted in SPS3 16.0.2 (SPS3 Inc., Chicago, IL, USA). For food load mass, we fitted a generalized linear mixed model for each species, using island and food-sampling period (early, middle, or late chick rearing) as predictors of load size for each species, and year as a random factor. We used the estimated marginal means and 95% confidence intervals from the parameter estimates to determine significant pairwise differences; pairs with non-overlapping confidence limits were considered to differ statistically.

To measure pair-wise distary overlap between species within islands and within species among islands in a given year, we used Horri's (1966) modification of Morista's Index (1959), as recommended by Diamond (1983). This provides an annual index 0 s C 1. U were identical dists have C + 1 and completed visionilar directs have C = 0.

We used nonparametric correlations (Spearmar's o) to look for covariance between overlap indices, load composition, prey availability and oceanographic or climatic variables. The same approach was used to examine changes in load composition over its Co control Gr multiple comparison, we used the False Discovery Rate and p < 0.10 (Benjamini & Hochberg 1995; García 2004; Grosbois et al. 2008). Setting a higher o: level is desired when the sample size (in this case, number of years) is low, as this increases the power for detecting a genuine relationship between climate and ecological variables (Leberton et al. 1992; Field et al. 2004; Grosbois et al. 2008).

After significant diet-climate relationships were identified, we included those climate variables in a series of generalized linear models with a gamma function error structure and identity link, and used quasi-Akaike's Information Criteria adjusted for small sample sizes and extra-binomial variation (QAIC<sub>c</sub>) for model selection. Using only a subset of covariates decreases the number of candidate models, and reduces the risk of multicollinearity (Grosbois et al. 2008). We considered the model with the lowest QAIC. value to be the best-fitting model to the data (Burnham & Anderson 2002). We constructed a Null Model (intercept only) and models that included additive terms and their interactions for sampling periods (early, middle, or late chick rearing), and islands (the Global Model). Once the best-fitting model was identified, the climate and oceanographic covariates of interest, and biologically significant interaction terms were added to subsequent models to examine the effect of climate (a procedure similar modelling recapture rate and then survival rate in mark-recapture studies; Lebreton et al 1992: Groshois et al 2008). Models with ADAIC, > 2 were considered to have substantially less support, and overall model support was assessed using Akalke weights (denoted as wi, Burnham & Anderson 2002). This was done on both the full dataset and on data from Buldir and Kasatochi islands because the dataset from Kiska Island is briefer (5 years), and because Kiska is anomalous because it has introduced predators that might affect provisioning behaviour (Major et al. 2006).

## Results

We identified 40 prey taxa categories in taxat Adulet food back (n = 810, 29 in CresteR Judiet food loads (n = 1110), and 32 in Whikkered Judiet food loads (n = 480; Appendices 5-7). Food loads were composed mainly of Necosithur coepools and Thynanoesson euphausilist (Appendices 5-7). Fine taxonomic resolution was not possible in all years, so larems were grouped for subsequent analysis into higher-level taxa as "Nocealonus" or "rephausilid", other taxa (e.g., amplipodi, decapodi, Culonus matchallar) were not included in analysis because they often comprised < 5% biomass in any given year. Notable exceptions were for taxat Adulet on Kasatochi in 1997, 1999, and 2006 (18,9%, 38.5%, and 13.1%) decapodir, respectively, Crested Auklets on Buldir in 2003 (12.9% Themiota emplopodi), and on Kasatochi in 2004 (20.5% Themiota amphipodir, Equera 3-2, 3-3, 4-1).

After correcting for the False Discovery Rate, we found no significant annual trends in the proportion of *Neocalanus* or euphausilds in loads of Least, Crested, or Whiskered auklets (all P > 0.12).

#### Food load mass

Food load mass was estimated for 683 Least, 1071 Crested, and 316 Whiskered Auklet food loads (64% of the total). Meal size differed significantly among sampling periods (all P < 0.01), but not among islands (all P > 0.32). The random effect of year explained less than 5% of the variance in all three species. Based on 95% confidence intervals of marginal means, load sizes increases significantly throughout the chickrearing period for Crested, and Least Auklets. In Whiskered Auklets, load size was smaller in the early than in the mid period, and load size in the late period din onc differ from that in other periods (Table 3-1). As a percentage of adult body mass, Least Auklet loads averaged 3-56.11% [Jadult mass: 26 g), Crested Auklets chick meals 3-25.7% (adult mass: 260 a), and Whiskered Auklet chick meals 7-41-03% (adult mass: 188 a).

## Dietary overlap and similarity among species and islands

Within-sland dietary overlap between paciet was considerable. At Bulki, overlap was highest between Least and Whikered auklets, followed by Crested and Whikered auklets, and in many vara, orweing between species are 300 (Tolle 3-2). At Kasatochi, overlap between Least and Crested Auklets was higher than that on Bulkir, and was > 0.50 in 5 of 11 years (Table 3-2). At Kinka, overlap between Least and Crested Auklets was low, but based on only three years' data, and small sample size (Table 3-2). Appendices 5, 6). Based on 95% confidence intervals given above, we conclude that at Bulkir three was more distary overlap between Least and Crested Auklets makes more distary overlap between Least and Crested Auklets that between other species, and that overlap between Least and Crested Auklets on Bulkir microverlap between Least and Crested Auklets on Bulkir are positively correlated with overlap between Least and Crested Auklets on Bulkir are positively correlated with overlap between Least and Microwerkie Auklets on Bulkir are correcting for the Table Rocewerk Ret (P o 2001). We also used the overlap index: C as a massure of diretary similarity within species among islands. Lead composition was consistent among islands for all species. Least Auklet diet was identical on all ialmads (range of mean C for pairwise island comparisons = 0.97.087; Table 3-3). Similarity in Crested Auklet loads was lower than that for Least Auklets, but still high (range of mean C for pairwise island comparisons = 0.72.050). We conclude that load composition of Least and Crested auklets did not differ among islands (Table 3-3).

## Prey availability: CPR data

After correcting for multiple comparisons using the False Discovery Rate, we found no significant relationships between prey availability (Neocolanus stage V biomass, and explausial biomass) and oceanographic of climatic indices. We also found no relationship between auklet load composition (proportion of Neocolanus or explausial biomass in chick meah) and total zooplankton biomass. Neocolanus (stage V copepodite) biomass, or explausial biomass, in plankton tows during chick rearing in 2000-2006.

## Relation to climate/oceanographic indices

After correcting for the False Discovery Rate, we identified only two significant correlations between load composition and climate indices. The proportion of exphasiids in Least Auklet loads was positively correlated with winter (December-January-February) NPSC on Bulder ( $\mu = 0.7$ , P = 0.006), and with spring (March-April-May) NPI on Kiska ( $\rho = 0.58$ , P = 0.005). Given the paucity of significant monotonic correlations, and the potential for more complex relationships (e.g., parabolic; Gjerdrum et al. 2003), we included all climate and oceanographic variables as covariates in subsequent analyses.

A model that included differences among islands, sampling periods, Island x period interaction, and a negative relationship to the summer (June-Juy-August) NPGO best predicted the biomass of Neocolanus in Crested Auklet bads. An identical model, but with a negative relationship to winter NPGO, also necelied considerable support. The same model (covariation with summer NPGO) best predicted explausild biomass in Least Auklet loads (Table 3-4). Both Neocolanus and explausild biomass in Least Auklet loads was best predicted by a model that included differences among islands, periods, period x island interaction, and a negative relationship with summer (June-July-August) POO (Table 3-5). A model that included only a negative relationship with winter (JDI POO best predicted the biomass of Neocolanus and explausids in Whiskered Auklet loads on bulket (Table 3-6).

Both NPGD and PDD track basin-wide SST anomalies, and as local SST was not among the highly-canked models, we investigated other possible relationships between food load composition and SST using Neocolanus in Least Auklet diet. Neocolanus accounted for most biomass in most years, and sample sizes were large on all three

Islands. We found a parabolic relationship of Neocolanus biomass to local summer SST ( $d^2 = 0.25$ ). Major et al. (2000) hypothesized that introduced Norway rats (Rattus norvegica) could affect the parabolic relationship of Neocolanus biomass to SST increased ( $d^2 = 0.71$ ), Maximal Neocolanus biomass occurred when summer (June-July) SST was ~4.5°C, was relatively constant between 3-6°C, and declined sharply outside this range flowers. S5a, b).

Correcting for False Discovery Rate, dietary overlap of Least Auklets among all three islands was correlated with winter (DII) INFO. and overlap between Kiska and Ksatacchi was related to NFGO in spring (March-April-May, MAM) and summer (June-July-August, Jul). Within islands, overlap between Least and Created Auklets was not related to any climate variable at Kiska or Kasatochi. At Buddr, Least and Whiskered Auklet overlaw searchated to shring (MAM NR).

# Discussion

Overall, our results confirm previous studies of auklet chick diet (uning adult food loads as a prony) in the Aleutian Islands (Day & Byrd 1989) which indicated that Least and Whistered auklets rely heaving on large-bodied oceanic copepode, whereas Crested Auklet adults also deliver a significant amount of explausified to their chicks. Unlike studies from SL: Paul Island in the Pribliof Islands, and on the continental shelf and shelf there are found way free Columns amountain in chick mails of theme species. The amount of C. marzhallee, a nertic copepod, in chick meak was negatively related to auklet chick survival at colonies on the continental sheef [51, Lawrence Island, Gall et al. 2006]. Unlike the larger oceanic copepods Neocolamus spop. C. marzhallee has lower lipid content, and Least Auklets are thought to seek out Neocolamus spop. J. performatily over less portfalable prey resources (Hunt Jr. & Harrison 1990; Hunt Jr. 1997). Rossell et al. 1999). Oceanic Neocolamus spo. copepods (N. plumchrus, M. flemingeri, and N. cristatua) are rich in lipids and wax esters required by auklets (Rohy et al. 1986). At St. Lawrence laland, oceanic copepods are advected by the Aualyr Current (Boringer et al. 1989), and contribute to higher chick survival (Gall et al. 2006; Sheffeld Goy et al. 2006). Soch oceanographic fronts concentrate zooplanktan, including Neocolamus spp. (Hunt Jr. 8, Harrison 1990; Hunt Jr. 897).

Around St. Lawrence Bland, Created Auklets were often associated with epibentics advected explauation wwrms (Russell et al. 1999), and their chick' diet consisted languly of *Thysonoessa* spp. explauation (Bédard 1969a; Flatt et al. 1990a). Sheffield Guy et al. 2009), inadicisto executionation, Created Auklets in the Nettania appear to also rely heavily on calanoid copepods to provision their chicks in most years (Appendix 5, Day & Byrd 1999), unlike their counterparts in the Bering Sea (Biddard 1969a; Hunt L. et al. 1981; Sheffield Guy et al. 2009). Created Auklets provisioning chicks exploid large bodied oceanic copepods when they are available (Searing 1977), in some years didwired meaks consisting where of these, commerciants with corepords

availability being generally greater in the oceanic domain of the western Aleutian Islands (Hunt Jr. 1997).

Much less is known about Whiskered Auklet chick's diet. Our results agree with previous studies of chick meaks on Buildi [Day & Byrd 1989; Hunter et al. 2002], but contrast with studies of adult stomach contents in the eastern Aleutians, where explausidis dominated diet [Troy & Bradstreet 1991], pointing to differences in the provisioning behaviour of adults depending on whether self-feeding or providing meaks to chicks, or perhaps opergraphy.

# Dietary overlap and similarity among species

In the northern Bering Sea and extern Alexitam, previous work has suggested that distributional differences among species are related to dietary differences (humt *K*. et al. 1998; Bussell et al. 1999). We found considerable dietary overlag between species and among years that could be the result of two non-actualwe hypotheses: adult authors adult and preversion different locations, or, because Miccolanus coppods are so abundunt (Muckas et al. 1998), competition for them is insignificant. Different species of aukhet may also exploit prev patches at different depths (Haney 1991); further study of individual wine details is required.

Chick diet of auklets in the Aleutian Islands exhibits high interspecific overlap compared with other seabirds. Assuming significant overlap when C  $\geq$  0.6 (Catry et al. 2009), we found nearly 80% of all C-estimates were significant. Dietary overlap among Alcultan auklets was similar to that in other subarctic and temperate seakind communities (Pearson 1968), Built & Morejoin 1977), Burrett et al. 1997; Gonzalez-Solis et al. 1997; Minich 2007), and generally higher than that in tropical communities (Achmole & Ashmole 1967), Diamond 1983; Catry et al. 2009). Overlap among auklets was also higher than among realizities at the more diverse subaratic: Cercera Hand (Bidoux 1994). While this high degree of overlap is noteworthy. It agrees with the prediction that overlap is greater in temperate and sub-polar communities where small numbers of exploited prey tasa are very abundant (Diamond 1983). Dietary overlap was greater in the Alexitans than among auklets breeding at St. Matthew Island in the northern Bernig Sea, where overlap between Least and Crested Auklets of C = 0.52 was at the lower end of the values we recorded Patrinon 1990.

To our knowledge, ours it the first study of dietary overlap over a larger range of years, and we have shown that there is considerable variation from year to year and among sites, highlighting the need for long term investigations to shucidate the ecological relationshown more globally-index members of the same foreigning and in the the context of ocenstitence. Dietary overlap at multiple sites has received little attention. We found that Least Auktet chick diet across the Alexitan Islands was nearly identical in all years and exhibited high overlap between Islands. These trends, combined with lack of interannual differences in the proportion of *Neor*abura or explausidis in most constitutions. Overlap was sover showness lands for Corela dukets but was applicant, further

supporting the interpretation that click det is similar among all three sites. Previous research at Kiska quantified near failure of Least Aukket threeding in 2001 and 2002, with productivity in those years being far lower than recorded at any Least Aukket colony in any year anywhere in Alaksa [Major *et al.* 2006; Chapter 2]. Most breeding sites failed during the chick rearing stage, with rat at chick y and hurstitoling alters being explanatory hypotheses. Our results here are relevant to this issue because we found no difference in Least Auklet chick diet (*C* = 0.99) between Buldir (where auklets had high productivity; Chapter 2] and Kiska. Food lead composition did net correlate with breeding failure at Kiska, or was it aberrant in any way in the years of failure. This adds to the evidence that introduced Norway rats were responsible for auklet breeding failure in 2001 and 2002 at Kiska.

## Food load mass

Food load mass varied across sampling periods within the breeding section in all three species. In Loast and Crested Auklets (and to a certain extent in Whiskered Auklets), food load mass increased as the breeding useson progressed, paralleling in our study than load size recorded on the Priblof Islands, (5.28g, Roby & Brink 1986a, cf. Table 8.3). If energy requirements of Least Auklet folks are the same in the Phibliof and Alexian islands, then adults in the latter location must deliver food to the chicks more frequenty (Roby Rob Rink 1995a, Roby & Rink 1996b, there are no data come frequenty (Roby Rob Rink 1995a, Roby Rob Rink 1996b, there are no data come frequenty (Roby Rob Rink 1995a, Roby Rob Rink 1997b). To our Knowledge, there are no data come frequenty (Roby Rob Rink 1995a, Roby Rob Rink 1997b).

from Crested or Whiskered auklets to make similar comparisons. The Dovekie (Alle alle), similar to auklets ecologically, adapts to changes in zooplankton abundance by increasing the rate of food load deliveries to chicks (Jakubas et al. 2007).

Load size was larger relative to adult body mass in the Whikered Auklet than in Least or Crested auklets. Whikered Auklet parents provision their chicks only once or twice per right [Conyukhov et al. 2000], much less than the other species (Roby & Brink Siges, Fraser et al. 2002). Whikered Auklets forage in tide right conserves the work other auklet species (Byrd & Williams 1993), which may allow them to carry a heavier load over a shorter distance. In addition, because most provisioning of chick's is nocturnal, adults may be under pressure to deliver larger loads to meet their chick' energy and growth formands (Verbeneg 1996; Nutter et al. 2002).

# Relationship of food loads to prey availability, and indices of ocean climate

We found no relationship between prev availability, as assessed by CPR data from near breeding colonies, and food load composition. Auklets generally forage in areas of occasi forest and upwelling cones, which have high concentrations of cooplankton (linder et al. 1983; Hust L. et al. 1999; Russell et al. 1999); CPR data are averaged over an entire tow, and would not indicate the presence of upwelling areas. The biomass of *Neocelanus* spo, copepoids in CPR tows wan also unrelated to indicate climate or occanography. CPR data were not available further south, in areas

this region, which may partially explain why we found no relationship between prey abundance and abundance in food loads in a given year.

The summer North Pacific Gyre Oscillation (NPGO) predicted food load composition of Crested Auklets, with a higher NPGO value corresponding with a decrease in the biomass of Neocalanus copepods and euphauslids. Least Auklet food load composition was related to the summer Pacific Decadal Oscillation (PDO), where a higher PDO value corresponded with a smaller biomass of both Neocalanus copepods and euphausiids. The NPGO is an index of sea-surface height and temperature anomalies over 110-180°W and 25-62°N (Di Lorenzo et al. 2008), and tends to explain more of the SST anomaly than the PDO (Bond et al. 2003). Our examination of the relationship between local summer SST and Neocolanus biomass in Least Auklet food loads showed a parabolic relationship, with copepod biomass peaking between 3-6°C. and declining significantly outside this range. Such a parabolic relationship is found when an optimum exists, such as the relationship between Tufted Puffin (Fratercula cirrhata) breeding success and summer SST (Gjerdrum et al. 2003). Neocalanus copepods require cooler temperatures to develop (ideally less than 10°C; Ikeda et al. 1990), but when temperatures are copier than 3°C, development may be delayed significantly as the copepods take longer to acquire the necessary lipids to grow and moult (Batten et al. 2003).

Interestingly, we found a decrease in Neocolanus copepods in Least Auklet chick meals at temperatures above 6°C. Such temperatures should be favourable for copepod

growth and development (Nicda et al. 1390; Batten et al. 2003), meaning that coepeod abundance would likely remain high. Warmer temperatures correspond with an earlier start of the peak abundance of Neocolanus, and a difference of 4/C in SST between southern and northern sampling stations along the west close of Nerri American is reflected in a difference of 3-4 weeks in the timing of the Neocolanus bloom (Batten et al. 2003). Despite this, the variation in local SST measured around auklet brending islands in hum and July each year remained within the themal limits of Neocolanus development ( CUC; Neak et al. 1990).

# Implications of climate-influenced diet on demography

Many studies of relationships between climite and oceanographic indices and demographic parameters in seablinds suggest bottom up control mechanism, although seldom are these tested explicitly (e.g., buratet et al. 2004; Chapter 2; Votier et al. 2005; In the Alexitan bilands, both adult survival and reproductive success are related to largescale climate indices (Chapter 2; authors' unpublished data). A more detailed analysis of the relationship between auklet demography and diet is warranted, but the similarity in food foad composition from years to year suggests that other factors (e.g., direct predation, density dependence, competition, change in abult' behaviour) are requisiting auklet oppolations in the Alexiten blands.

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T.W. Chapman and E.H. Miller, provided improvements on previous drafts of this

manuscript.

# Figures



Figure 3-1 – We studied auklet food loads at Buldir, Kiska, and Kasatochi islands,

Aleutian Islands, Alaska, from 1994-2006.



Figure 3.2 – Crested Auklet chick diet (as inferred from adult food loads) had greater proportions of euphaulish than that of Least or VM/skiered Auklets. Data are presented as the proportional biomass of *Necostabus* copepode, euphauside, and other prey types. Colour shades represent samples from early (light) mid (medium) and late (durk) chick rearing. Data are presented for Buddir (top, 1994-2006), Kiska (middle, 2001-2006), and Kastochic biotom.



Figure 3.3 — Least Auklet chick dist (as inferred from adult food loads) was almost entirely Neocolamic copepade. Data are presented as the proportional biomass of Neocolamic copepade, explausida, and other prey types. Colour shades represent samples from early [lipt); mid (medum) and late (dark) chick rearing. Data are presented for Buddir (top, 1994-2006), Kilak (middle, 2001-2006), and Kasatochi (bottom. 1995-2006).



Figure 3-4 Whiskered Auklet chick diet (as inferred from adult food loads) at Buldir (1994-2005) was very similar to that of Least Auklets, with large proportions of copepods, but are presented as the proportional biomass of Neocolanus copepods, euphausids, and other prey types. Colour shades represent samples from early (light) mid (medium) and use (rark) chick range.



Figure 3-5 — The proportion of Neocolanus copepods in Least Auklet food loads was related quadratically to summer SST around all three breeding colonies (a,  $r^2 = 0.25$ ), and at Buldir and Kasatochi only (b,  $r^2 = 0.71$ ).

# Tables

Table 5-1. The food load mass for Least, Crested, and Whiskered auklets increases as the breeding season progresses (sarly, mid, and late) at three breeding tites in the Aleutan blands, Alata, in 1994 2006. Samples were pooled among breeding sites and years. Values are estimated marginal means 1.5.E. (n) in grams (wet weight). Values sharing the same letter are not significantly different based on overlapping 95% confidence intervals.

Species	Early	Mid	Late
Crested Auklet	8.27 ± 0.83 (282)*	11.77 ± 0.74 (342) <sup>b</sup>	14.78 ± 0.73 (447) <sup>b</sup>
Least Auklet	2.94 ± 0.24 (234)*	4.23 ± 0.24 (249) <sup>b</sup>	$5.20 \pm 0.24 (200)^{b}$
Whiskered Auklet	8.02 ± 0.66 (113)*	$10.90 \pm 0.67 (108)^{b}$	9.29 ± 0.70 (95) <sup>20</sup>

Table 3-2. The overlap among Least, Crested, and Whiskered auklet food loads at Buklir, Kiska, and Kasatochi Islands in 1994-2006 is variable, but consistently high. Values are presented as Horn's (1566) modification of Morisita's (1595) overlap index (C), where C = 0 indicates completely dissimilar dates with no overlap, and C = 1 indicates identical dists (i.e. complete overlab.

	Buldir			Kasatochi	Kiska
	Crested-	Crested-	Least-	Crested-	Crested-
Year	Least	Whiskered	Whiskered	Least	Least
1994	0.697	0.981	0.803		
1995	0.634	0.746	0.985		
1996	0.412	0.460	0.998	0.813	
1997	0.513	0.496	0.999	0.949	
1998	0.499	0.469	0.999	0.952	
1999	0.873	0.897	0.995	0.836	
2000	0.977	0.976	1.000	0.553	
2001	0.324	0.328	0.998	0.822	0.010
2002	0.836	0.971	0.932	0.703	
2003	0.170	0.684	0.765	0.740	0.015
2004	0.993	0.990	0.999	0.927	
2005	0.949	0.962	0.999	0.958	
2006	0.915	0.879	0.995	0.953	0.839

Mean	0.676	0.757	0.959	0.837	0.288
95% C.I.	0.149	0.132	0.044	0.077	0.540

Table 3-3. The composition of food loads of Least and Crested auklets at Buldir, Kiska, and Kastacchi Islandsi in 1994-2006 are very similar. Values are presented as Horn's (1966) modification of Morisita's (1999) overlap index (C), where C = 0 indicates completely disimilar diets with no overlap, and C = 1 indicates identical diets (i.e. complete overlap.).

	Crested Au	ıklet		Least Auklet		
	Buldir-	Buldir-	Kasatochi-	Buldir-	Buldir-	Kasatochi-
Year	Kasatochi	Kiska	Kiska	Kasatochi	Kiska	Kiska
1996	0.944			0.959		
1997	0.702			0.929		
1998	0.743			0.992		
1999	1.000			0.877		
2000	0.626			0.969		
2001	0.806	0.815	0.527	0.995	1.000	0.996
2002	0.937			0.995	0.999	0.993
2003	0.785	0.897	0.652	0.988	1.000	0.991
2004	0.968			0.995	0.999	0.997
2005	0.998			0.993		
2006	0.945	0.981	0.990	0.973	0.934	0.964
Mean	0.859	0.898	0.723	0.969	0.986	0.988
95% C.I.	0.076	0.094	0.271	0.022	0.026	0.012

Table 3-4. Summary of the models set for predicting the proportion of Cretetd Auklet chick food load biomass represented by Wecolenus copends and euphaulids at Buldin, Kiska, and Kasatachi, Aleukan Islanda, Alaska from 1994 to 2005. Models are sorted by increasing DAC, (quasi Alaski's information criteria corrected for anile sample size) value with the model and laski's information criteria corrected for anile sample size) wale with the model of the model set Is the best approximation of the data. Model parameters: I: kland, P: sampling period, AIPI: Aleukan Low Pressare Index, NPGC: North Seciel Gyre Gocillution (DIC: Becember to February; MAM: March to May), SST: ses surface temperature. Model corrected for 2 = 1.846 (Neccolinus) and 1.811 (esphausitid), AQAIC, is the difference from the top-ranked model. Only the top 10 ranket models are shown.

Neocalanus	Number of parameters	QAIC	$\Delta QAIC_{c}$	w,
I+P+IP+NPGO/JJA	11	2536.183	0.000	0.57
I+P+IP+NPGO/DJF	11	2537.723	1.541	0.26
I+P+IP+NPGO/MAM	11	2540.412	4.230	0.07
I+P+IP+NPI/DJF	11	2542.858	6.675	0.02
I+P+IP+ALPI	11	2543.471	7.288	0.01
I+P+IP+SST/JJ	11	2543.980	7.797	0.01
I+P+IP+NPI/MAM	11	2544.054	7.871	0.01
I+P+IP+PDO/DJF	11	2544.230	8.048	0.01
I+P+IP+PDO/MAM	11	2544.470	8.287	0.01

Neocalanus	Number of parameters	QAIC	$\Delta \textbf{QAIC}_{c}$	W/
I+P+IP	10	4700.352	8.419	0.01

Euphausiids	Number of parameters	QAIC	∆QAIC	W/
I+P+IP+NPGO/JJA	11	2585.000	0.000	0.58
I+P+IP+NPGO/DJF	11	2586.570	1.570	0.26
I+P+IP+NPGO/MAM	11	2589.311	4.311	0.07
I+P+IP+NPI/DJF	11	2591.804	6.804	0.02
I+P+IP+ALPI	11	2592.429	7.429	0.01
I+P+IP+SST/JJ	11	2592.948	7.948	0.01
I+P+IP+NPI/MAM	11	2593.023	8.023	0.01
I+P+IP+PDO/DJF	11	2593.203	8.203	0.01
I+P+IP+PDO/MAM	11	2593.447	8.447	0.01
I+P+IP	10	2593.602	8.602	0.01

Table 3-5. Summary of the model set for predicting the proportion of Cretted Auklet chick food load biomass represented by Necolahura copepads and explausilis at Buldir, Kiska, and Kasatochi, Alexian Iahada, Alaxka from 1394 to 2006. Abbreviations are identical to those in Table 3-4. Models not corrected because E = 1.0. Only the top 10 ranket models are tooms.

Neocalanus	Number of parameters	AIC	∆AIC <sub>c</sub>	w,
I+P+IP+PDO/JJA	11	1805.820	0.000	1.00
I+P+IP+NPI/DJF	11	1817.946	12.126	0.00
I+P+IP+NPGO/DJF	11	1819.558	13.738	0.00
I+P+IP+NPGO/MAM	11	1820.426	14.606	0.00
I+P+IP+PDO/MAM	11	1822.822	17.002	0.00
I+P+IP	10	1822.864	16.989	0.00
I+P+IP+NPI/MAM	11	1824.606	18.786	0.00
I+P+IP+ALPI	11	1824.862	19.042	0.00
Ρ	4	1829.568	23.467	0.00
L	4	1856.368	50.267	0.00

Euphausiids	Number of parameters	AIC	∆AIC <sub>c</sub>	w,
I+P+IP+PDO/JJA	11	1805.820	0.000	1.00
I+P+IP+NPI/DJF	11	1817.946	12.126	0.00
I+P+IP+NPGO/DJF	11	1819.558	13.738	0.00

Euphausiids	Number of parameters	AICc	∆AIC <sub>c</sub>	W/
I+P+IP+NPGO/MAM	11	1820.426	14.606	0.00
I+P+IP+PDO/MAM	11	1822.822	17.002	0.00
I+P+IP	10	1822.864	16.989	0.00
I+P+IP+NPI/MAM	11	1824.606	18.786	0.00
I+P+IP+ALPI	11	1824.862	19.042	0.00
Ρ	4	1829.568	23.467	0.00
1	4	1856.368	50.267	0.00

Table 3-6. Summary of the model set for predicting the proportion of Whikered Auklet chick food load biomass represented by Mexcalanux copepods and euphacilia at Buldt, Kiska, and Kasatochi, Alexian Iakan, Alaxia from 1594 to 2006. Abbreviations are identical to those in Table 3-4. Model corrected for 2 = 11.673 (Mexcalanus) and 1.668 (euphauidi). Other top 10 network models are shown.

Neocalanus	Number of parameters	QAIC <sub>c</sub>	∆QAIC <sub>c</sub>	w,
PDO/DJF	2	1127.850	0.000	0.98
PDO/JJA	2	1136.976	9.126	0.01
NPGO/DJF	2	1139.201	11.351	0.00
PDO/MAM	2	1139.438	11.589	0.00
NPI/MAM	2	1140.505	12.655	0.00
NPGO/JJA	2	1141.627	13.778	0.00
NPI/DJF	2	1142.397	14.548	0.00
SST/JJ	2	1142.469	14.619	0.00
NPGO/MAM	2	1142.880	15.030	0.00
NPI/JJA	2	1142.923	15.074	0.00

Euphausiids	Number of parameters	QAIC	∆QAIC <sub>c</sub>	w <sub>i</sub>
P+PDO/DJF	5	1133.402	0.000	0.98
P+PDO/JJA	5	1143.130	9.728	0.01
P+NPGO/DJF	5	1144.466	11.064	0.00

Euphausiids	Number of parameters	QAIC	∆QAIC <sub>c</sub>	w
P+PDO/MAM	5	1145.647	12.245	0.00
P+NPI/MAM	5	1146.756	13.354	0.00
P+NPGO/JJA	5	1147.303	13.900	0.00
P+NPGO/MAM	5	1148.358	14.956	0.00
P+NPI/DJF	5	1148.435	15.032	0.00
Intercept Only	2	1148.683	15.281	0.00
P+SST/JJ	5	1148.759	15.357	0.00

# CHAPTER 4: RAPID BUT DELAYED PRIMARY MOULT BY ARCTIC-BREEDING

# LEAST AUKLETS

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## Abstract

Along with breeding and migration, moult is one of the most energetically expensive components of birds' circannual cycles. Auklets (Alcidae, tribe Aethiini) are apparently unique among the auks in that flight feather moult and breeding overlap. We investigated the degree of overlap of primary moult and breeding in Least Auklets (Aethia pusilla) at four colonies - Kiska Island (Aleutian Islands, Alaska, 52\*N), St. George Island (Pribilof Islands, Alaska, 56\*N), St. Paul Island (Pribilof Islands, 57\*N), and Cape Ulyakhpen (Chukotka Peninsula, Russia, 64"N). Flight feather moult commenced during incubation with up to four primaries replaced by the end of chick rearing. We found no difference in moult rate between adult breeders and non-breeders, or between adults and subadults at Kiska. Adult auklets replaced their first four primaries faster and initiated moult later at higher latitudes despite similarities in the length of the breeding season among colonies. The cold-water Anadyr Current, which advects lipid-rich oceanic Neocoloous son, conenads, likely contributes to the increased energetic requirements of Least Auklets breeding and moulting at higher latitudes. In Least Auklets and other members of their tribe Aethiini, sequential primary moult, rather than simultaneous moult typical of other alcids, may be possible because of auklets' lower body mass, wing loading, mass loss at hatching and foraging patterns.

## Introduction

The regular replacement of feathers through moult is costly both in energy and time (Murphy 1996; fichiwer et al. 2009). Despite the importance of moult, studies of inter-annual and inter-colony variation of moult of pelagic seabrds, and possible influencing factors are rare (Temile et al. 1990; Underhill, & Crawford 1999). Because of the cost of feather synthesis and the aerodynamic and thermoregulatory costs of missing feathers, understanding patterns of moult can be excited to investigations of anon physiology, coefficiency, and behaviour (Hoye & Buttemer 201).

Auklets (Acthini) are munuauk alchid because several species mout their primary feathers sequentially beginning during the breeding seaons (Médard & Seak) 1984; Emaile et al. 1990; Konyukhov 2001; 2009). Mots other auks mout all their flight feathers simultaneously after they leave the breeding colony, medering individuals flightless for saveral weeks (Stresemann & Stresemann 1966; Harris & Yule 1977; Seak) 1977; Bedard 1985; Euvin 1988; Harris & Wanless 1990b; Isridge 2006; Pyle 2008). Moulting patterns for some species of auk are largely unknown (e.g., spectacied Guillemot (Capphia corbo), Carvert's Murrelet (Synthiboromphus corveri) and Japanse Marrete (S. wurduszam)).

The Least Auklet (*Lething* pusiling) is endemic to the Berling and Ohbotsk seas and adjacent North Pacific Ocean, where they breed at a few denice colonies (Gaston & Jones 1998). In North America, major colonies are in the western Aleutian Islands, on the Philolis Islands, and in the northern Berling Sea (Dones 1993). In Russia, Least Auklets

breed on Ratmanov (Big Dionede) bland (Bering Starik), on the Chukokia Peninsula, and on silands in the Sea of Othotsk and Kurll Archipelago (Demet'ev et al. 1951; Kostova 1957; Konghave et al. 1998; Kondraysev et al. 2000; Artuakin et al. 2001; Zelenskaya 2009). Norkhern colonies in the Bering Sea and Sea of Othotsk are surrounded by sea ice during part of the year, whereas all Aleudian and most Kuril bland colony sites are ice-free var our off theremer et al. 2010.

Moult is often timed with bind' circannual cycles (Dawson 1998; Dawson et al. 2001), and in Least Auklets, is probably related to the timing of breeding. Auklet breeding phenology is later at higher latitudes; e.g. Least Auklets next 3-4 weeks later at 2. Learners blank (G2721) than colonies in the Aukletin blank (Sebay) 3757; Jones 1933b; Gaston & Jones 1998). We therefore predicted that the timing of moult would show a similar gradient in Least Auklets. Such latitudinal variation in timing and rate of moult has been observed in mammali. (Todorovi: 2555; King & Moody 1992) and land binds (Williamon & Emission 1371; Medwaldh & King 1376; Dawson et al. 2000), but ours is the first westgation of the phenomenon in mamine binds.

By including data from all major breeding areas (northern Bering Sea, Pribilof Islands, Akotish Islands, Chuketka Peninsuka), our goal was to Investigate how the timing of primary replacement view dwth breeding phenology, age class, breeding status (breeder us, non-breeder), and laitude of the colony. Because Least Auktest begin replacing light feathers during includation and continue through chick rearing, we predicted that addition more northerly colonies affected by sea and snowment

would mould later and faster than adults in the south. Based on previous studies we also predicted that subaduls would initiate mould later, but replace primaries faster than adults (Bédard & Seah) 1984, Emilie et al. 1990). Finally, we evaluated hypotheses to explain why the tribe Activiti is unique among the Akidae in overfapping primary replacement and tweefing.

## Methods

## Capture and moult status of wild Least Auklets

We caught Leat Aukites at Sriine Polin, Kiala landin, Alendian Ihanda, Alakak (52'08'h, 17'37'l) in June-July 2008 and 2009, Ulakais Nidge, St. George Island, Prihoff Haldnd, Aukike (56'37'k, 10'7') win June-dupat 2005; Tolebi Polin, Eata Lunding, and Zapadni Point Colosien on St. Paul Island, Prihoff Islands, Alaska (57'11'K, 13'0'16'W) in June-August 2000 and June-July 2004; and Cape Ulyakhpen on the Chukotta Perinnuka, Russia (46'37'K, 13'4'W) in June-September 1988-1900 (Figure 4-1). Summaria of data from 51; Paul St. George, and Cape Ulyakhpen are also presented by KonyAhov O'2009. Biels were caught using noose caretos with a ground heileder over hreeding crevices (Konyukhov 2009). We weighted, measured, and banded each bird, and recorded the number of replaced and mixing primarias. Adults were identified as having a black forehead with white streaks; subadults (Individuals 11.13 months).

determined the apparent breeding status of each adult by resighting marked blick and recording whether they carried a chick meal, which is evident by the distended throat pouch (Portenios 1334; Bidard 1306a). Individuals were identified as breeders in a given year if they were observed with chick meals at least twice during the chick-rearing period (late June to late July). This underestimates the number of actual breeders, as some captured birds were not seen following release and could have been breeding elsewhere in the colony, and because some breeders whose attempt failed during includion with the been seen corriging chick mell forces et al. 2002.

At 3F, Paul Island, St. George Island, and Cape Unykäheen, newk grown primaries were measured to the nearest 0.1 mm with calipters. At Icika Island, we recorded the number of newly grown and missing privarias only. We then converted feather lengths to proportion of feather mass grown (PKMG, Underhill & Summers 1993) by assuming that the mass deposition within each new feather was linear, and using masses of fully grown primaries obtained from auklets encountered dead at Itakia (all feathers) or sampled for stable-isotope analysis (P1 and P10, Table 4-1). PFMG is preferable, as it describes the energetic costs more accurately than mesured feather lengths (Underhild & Jacchini 1988), and counting replaced and missing feathers assumes that each feather is energetically identical (flohwer 2008). Statistical analyses were performed on the PFMG reliable to the feather mass of the first four primaries, as only these are replaced during the breeding season (see Results). Data from Ticks were converted to PFMG bit assuming that the outernot new primary was Nale grown, then extremot servers for the senter primaries were prevented more the senter the senter mass of the first four primaries, as only these are replaced during the breeding season (see Results). Data from Ticks were converted to PFMG bit assuming that the outernot new primary was Nale grown, then extremot were Strips was prevented to the feather mass of the first primaries, and the first primaries, and the senter outer outer strips was prevented to the primary was Nale grown, then extremot warrows the primary set and outernot new primary was Nale grown. Then extremot warrows primary was the outernot new primary was Nale grown. Then extremot warrows the primary set and outernot new primary was Nale grown. Then extremot warrows the primary set and outernot new primary was Nale grown. Then the outermot warrows the primary mass and primary the outernot the primary was Nale grown. Then extremot theorem the strips of the primary theorem prim

grown, and all others were fully-grown. This pattern is consistent with our observations based on measured lengths at other colonies (unpubl. data).

#### Museum specimens

To study (asst Aublet moult outside the hereding season, we examined specimens collected between Spetember-January in museum collections in Ganada, the United States, and Russia (Appendix 9). As with live aublets, each specimen was agred faulter students and automatic spectra and farather<sup>(1)</sup> (0), missing feather (1), new feather 125k grown (2), new feather 50% grown (3), new feather 75% grown, or new feather fully grown (3). Values were then converted into PFMG of all 30 primaries (Remisieveicz et al. 2010). Because we assumed some mixing of writter populations, and lacked any a priori knowledge about spergetion of individual Least Aublets from different breeding areas during the non breeding season, all specimens from Stembers-January were included in all and analysis.

#### Statistical methods

Using the mass of each primary (Table 4-1), we converted the measured lengths of feathers to PFMG for analysis in an Underhill Zurcchni type 5 analysis (Underhill & Zurcchni 1988; Underhill et al. 1990), hereafter "U-2 5" analysis. U-2 5 is used when individuals sampled have either or vet started, or are currently moulting primaries, and no individual has finished primary moult at the end of the study (Underhill et al. 1990). U 2 models use a maximum likelihood approach to estimate duration (and therefore, rate) of moult and the or moult initiation for a given population. Models were constructed using the package 'moult' in R.2.12.1 (B Development Core Team 2010). To test for differences in moult parameters among groups (breeders vs. non-breeders, adults vs. subadults, among coloneits, ve used paired tests: for multiple comparisons, we corrected alpha values using both the False Discovery Rate (FDR) approach (Regnaming & Horberg 1995), and Dum Sfak correction (fidat Sor).

### Results

### Moult of Least Auklets at breeding colonies

The relationship between feather length and mass was not uniform among primaries (Table 4-1); outermost primaries had greater mass per mm than did inner primaries.

On Kaka Island, we examined 214 live birds between 8 June-26 July 2008 and 115 birds between 10 June-39 July 2009. We examined 232 adult birds from 51. George Island between 30 June-5 August 2005, and 143 adults between 25 June-31 July 2007. At 51. Paul Island, we examined 31D birds between 10 June-9 August 2000, and 182 birds between 29 June-30 July 2004. At Cape Ulyakhpen, we captured 129 adults between 4 June-7 Settember 1948. Jirds July 2004. At Cape Ulyakhpen, we captured 139 adults between 4 between 2 June-31 August 1990. We found that Least Auklets begin growing up to four primary feathers sequentially during late incubation and early chick rearing.

Because of the small number of adult birds examined on Kiska in 2009 (n = 65), it was not possible to construct a meaningful U-Z model for this group, so it was removed from subsequent analysis.

Multiple comparisons were identical using both the TDR and Dunn Stäkk correction. The duration of primary moult did not differ between breeders and non-breeders (10:13 vectors (10:13 ve

There were feed differences among colonies in the duration of primary moult, with adalets at Cape Ulyakhpen moulting faster than those in the Prikiloh 12 0004 and 2005 (both p < 0.001; FB et <0.0002). The start of primary moult could be described by three groups: added to be durating the start of primary moult (could be described by three groups: added to be durating the start of primary moult (could be described by Chickota (Cape Ulyakhpen 188), 1990), with brids at Cape Ulyakhpen instanting moult the latest (Table 4.3). Variation in the start date was generally higher at solution controls (Adedgesa). Variation in the start date was generally higher at solution the latest (Table 4.3). Variation in the start date was generally higher at solution.

Start date SD was similar at Cape Ulyakhpen in 1988 & 1990 and St. Paul in 2000 (all p > 0.14), and at Cape Ulyakhpen in 1989 and St. George in 2007 (p = 0.86); all other colonies were significantly different from each other (Table 4-3).

#### Museum specimens

We examined 66 museum skins collected between September and January. Some birds had completed primary moult by the end of September, and others a month later. No birds were moulting primaries after 27 November (Table 4-4).

### Discussion

Least Aukkes replace the innermost primary feathers at the breeding colony during chick rearing, and complete primary moult by October or November. Our results contrast with moult patterns of the closely related Cassin's Aukke (Psychorramptus autoricus) an which subables tend to moult primary feathers faster, resumably because the costs of breeding reduce the amount of emergy and resources available for moulting (Emslie et al. 1990). We found no difference: adults (breeders, and non-breeders or failed breeden) moulted at the same rate as subadults. Befard and Sealy (1984) also found no difference in wing moult progress between adult breeders and failed breeders. Befard and Sealy (1984: 467) defined "subadults" as "birds in their second summer after hatching and in their Kahemate (I) (Dawnorth Sealy (Le-23-25) months old), whereas we

believe that these birds were in their first summer after hatching, hence only 11-13 months old (Pyle 2008).

One of the most distinctive features of subalditis is their extremely worn and finded flight feathers (Jones 1993; originally grown ayear before as a nestling), which are certainly in need of replacement. Subadults began moult 18 days later than adults, and individuals were much more variable in when they would start moulting than adults. 2008), and although subadult Least Auklets have a lower body mass than adults (Jones 1993b), they are no more variable (author' unpublished data). Pre-breeding birds, however, do not need to carry nutrient reserves required by breeding individuals (Gaston & Highner 2006). The moulting patterns of subladult birds require greater attention, and further study among colonies and years to determine whether the trend we observed it stypical or normaliox.

We found that bereding status had no relationships to moult progress, as Beldard and Seaky (1984) also noted for wing moult in three Aethio auklets. Two points must be considered however, fins, our method understimates the number of actual treeders in our sample, as individuals that failed during incubation were not counted, and some bereders may only be seen once with a chick meal. Some livits captured using noos carpets were non-breeders not teld to a specific part of the colony or were breeding to that the red device on the colony counter on the colony counter of the colony or were breeding to that the red device on the colony counter on the colony counter.

moult differs among colonies and years. There is annual variation in the timing and rate of primary moult in other auklets (Emslie et al. 1990).

Birds breeding at high latitudes often experience a briefer summer breeding period (e.g., Summers et al. 2004; Coppack 2007), and moult is often faster at more northerly locations, presumably to minimize overlap of two life-history stages (moult and breeding, Wingfield 2008). Delays in moult can result in an increase in moult rate and decrease in feather guality (Dawson 2004). Least Auklets do not show a compressed breeding season or compression in moult duration at higher latitudes (cf. Cooper et al. 2011), where incubation (30 d) and fledging periods (29 d: Piatt et al. 1990b) from St. Lawrence Island (63°24'N) are similar in duration to those in the Aleutian Islands (29 d: Renner 2005). There is little information on auklet moult outside the breeding season (i.e., in migration, or on wintering grounds) and few museum specimens have been collected between October and May. Auklets from more northerly locations face longer migrations, as colonies in the northern Bering Sea and the Sea of Okhotsk are covered by sea ice, unlike the Aleutian Islands (Fetterer et al. 2010), so may have to moult faster (de la Hera et al. 2009).

Least Auklets breeding in Chaketka likely exploit the cold-water Anadyr Current, which advects oceanic zooplankton, like Nercationuc copepods (Springer et al. 1987; Platt & Springer 2003). Sea-surface temperature of the Anadyr Current is colder than around the Pribliod and Alexitan Islands (Platt & Springer 2003). Combined with the increased at ear of mout a Cape Urghthem compared with other colonies in our study.

we conclude that northerly-breeding Least Auklets must consume more calories per unit time than their southern conspecifics – a situation made possible by the Anadyr Current's advection of oceanic copepods.

Our estimate of the complete duration of growth of Least Auklets' P1 feather is around 20-30 days (Konyukhov 2009; Rohwer et al. 2009). Our modelled values for primary initiation indicated that moult starts during incubation and that the first primary is often replaced by the mean hatch date. The rate of PFMG increase per day is not likely linear among feathers, so estimating a moult completion date in the absence of a large sample of birds is difficult. Based on observations of museum specimens, primary moult concludes before the end of November. This is similar to estimates of moult duration for the congeneric and similar-sized (118 g) Whiskered Auklet (Aethia pygmaea) at Buldir Island, Aleutian Islands, where moult was predicted to end around October (Konyukhov 2001). Essential to understanding and interpreting stable-isotope data from feathers, however, is knowledge of the range of dates over which a feather was grown, as feathers usually incorporate exogenous nutrients from the time of synthesis (Hohson & Clark 1992a) and, in species with moult migrations, endogenously catabolised nutrients that were laid down prior to migration (Fox et al. 2009). In adult auklets, it is therefore possible to select feathers representing three different times in the annual cycle - the last (outermost) primary grown the previous winter after young have fledged, body feathers grown during the spring (Pyle 2008), and the first primary grown during incubation.

#### Primary moult in the Alcidae

Least Auklets are the smallest of the auks, and therefore have smaller feathers to replace. Dovekies (Alle alle) are not closely related to the auklets, but are ecologically similar (small planktivorous auks), breed at higher latitudes than auklets and thus experience briefer summers, and have short breeding periods (Stempniewicz 1981). Dovekies are thought to moult their primaries synchronously (Salomonsen 1944; Dement'ev et al. 1951; Storer 1960; Stresemann & Stresemann 1966; Bédard 1985; Stempniewicz 2001), although recent evidence suggests that, like Common Murres [Uria aglae), primary moult originates at a single foci between P4 and P7, proceeding in two concurrent waves towards P1 and P10 (Thompson et al. 1998); further investigation is required on the precise moulting pattern. Nevertheless, the difference between Dovekies and the Aethiini is clear: Dovekie's primaries are moulted rapidly and moult is accompanied by a flightless period, whereas auklets have a relatively protracted moult and maintain flying ability. This difference could be because of Dovekie's long-distance post-breeding migration (Thompson et al. 1998), or a phylogenetic effect. Unlike auklets, Dovekies travel several thousands of km (e.g., from NW Greenland to the North Atlantic) and must moult before migration (Stempniewicz 2001). Auklets move on the order of hundreds of km after the breeding season (e.g., Diomede islands to southern Bering Sea: see below), and are therefore not under as much pressure to complete moult as Dovekies. Dovekies (subfamily Alcinae, tribe Alcini) are closely related to

murres, Razorbills, and the Great Auk (Pinguinus impennis), all of which moult primaries simultaneously, while auklets (subfamily Fraterculinae, tribe Aethiini) moult primaries synchronously (Friesen *et al.* 1996; Pereira & Baker 2008).

The auklets have among the lawest wing loading (body mass per unit wing area) of the aicki (Stempniewicz 1982; Minami et al. 1991; Spear & Aniley 1997; Ortega-Jiménese et al. 2010) and thus are able to fly with missing primaries. The larger auka (mirrelets, pulling, murres, guillemots, and Rasorbid (Ariz trand) would experience a significant decrease in wing area were they to mouth flight feathers sequentially. This would result in greater wing loading (Bridge 2004), likely imdering them flightess for a largthy period of time (Levery 1988), in contrast, the flightless of eart Auk likely and a primary moult similar to that of Common Murres (Salomonsen 1945; Thompson et al. 1998). Least Auklet sequences and smarks loads when the Arearing commerces, and may obtain their own flood from more instore areas, which reduces wing loading and flight costs (Lones 1994). This may be a mechanism to accommodate flight feather replacement while maintaining fliping ability. Other auto also decrease in mass during chick rearing, bot this is apparently insufficient to allow for a reduction in wing area via moult (Cost et al. 1921; Charter 3).

Moult-breeding overlap is also associated with species that have limited postbreeding migration or disperal (liridge 2006). Least Auklets' wintering areas are largely unknown, but they are more abundant in the western Pacific during the winter and sping (lyvakin 1831; Sydeman et al. 2020), and some winter in the Sea of Japan off the

coast of Primoye, Rusia (Shuttor 1965; Velihainin 1977a; Kondrahve et al. 2000). Sea ice cover in the Bering-Sea and Sea d'Ohitchi, however, requires some auklest to disperse up to 1000 km from their breeding colony (Fetterer et al. 2010). Bridge (2006) cusclified least Auklets as "dispersive", or "Tristis that leave the breeding area after fieldgring chick but that speed the non-breeding period in the general area or region of the breeding she" (Bridge 2006; 9). It is clear, however, that Least Auklets breeding in the northern Bering Sea (Chuketla Peninsula, Ratmanov, Little Diomede, 9: Lawernce, 9:. Matthew ilandi) and the Sea of Ohibetki (Yamakaye tislandi, and Iona (5: Lonah) baland, for example) must disperse at least hundreds of km during winter (Fetterer et al. 2010.

Our results indicate that primary moult in Least Auklets responds to local occanographic conditions that differ with latitude, and hat, like land birdt, auklets' rote of moult is gratest at high latitudes. Physiological and behavioural factors, including mass loss and a change in foraging area, allow Least Auklets to moult during chick resing, presumably the time of higher tood availability.

### Acknowledgements

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Figure 4-1 – We investigated Least Auklets' primary moult at four colonies throughout their breeding range in the Bering Sea.



Figure 4.2 – Duration and initiation date of primary moult in Least Aukites varied among years and colonies, with adults from the northern colony (Dape Ulyakhper) moulting later (higher x-intercept) and faster jahorter duration; greater slope). The solid lines are the mean estimates and 50% confidence intervals, and dots represent individual observations. Researchers arrived after the start of primary moult on St. Paul in 2004, and on St. Genergin 2005.

# Tables

Table 4.1 – The muss and length of Least Auklet primary feathers used to calculate the proportion of feather mass grown for Underhill-Succhini moult models (n × 5 individuals, except for P1 and P10, where n = 5.13 is dividuals). Primary lengths are from Konyukhov (2009). Total feather mass ± 5.01 signers for the five individuals for whom all 30 primaries were weighted.

Primary Number	Mass ± S.D. (mg)	Length ± S.D. (mm)	Mean mg mm <sup>-1</sup>
1	11.1 ± 1.3	43.1 ± 1.7	0.26
2	$12.9\pm2.3$	46.0 ± 1.9	0.28
3	$14.3 \pm 2.7$	48.7 ± 1.5	0.29
4	$16.5 \pm 2.6$	51.1 ± 2.0	0.32
5	18.3 ± 2.8	$54.5 \pm 1.4$	0.34
6	19.1 ± 3.2	55.9 ± 1.7	0.34
7	$19.8 \pm 3.6$	$57.1 \pm 1.6$	0.35
8	$20.8 \pm 3.7$	57.8 ± 2.1	0.36
9	22.8 ± 3.4	58.3 ± 1.7	0.39
10	24.5 ± 2.8	56.5±1.8	0.43
Total	176.6 ± 30.0		0.33
Table 4.2 - At Kiska Island in 2008, adult breeding status, and age class had no effect on the duration or initiation date of primary moult in Least Aukkts. Subadults' initiation date was significanty more variable than that of adults. Groups sharing the same letter are not considered statistically different.

Group	n	Initiation Date	Initiation Date	Duration ± S.E.
		± S.E. (days)	S.D. ± S.E. (days)	(days)
Adult breeders	84	167 ± 5.9 <sup>a</sup>	$18.3\pm0.2^3$	79.5 ± 24.8°
Adult non-	71	$162 \pm 3.7^{a}$	$18.7\pm0.2^{\rm s}$	$104.5 \pm 16.8^{\circ}$
breeders				
All adults	155	164 ± 3.0*	$19.5\pm0.1^{a}$	93.0 ± 12.4 <sup>a</sup>
Subadults	56	182 ± 7.7*	$24.3\pm0.5^{\rm b}$	100.7 ± 52.6 <sup>a</sup>

Table 4-3 - Adult Least Auklets from the most northern colony (Cape Ulyakhpen) tended to moult their primaries faster, and to start their primary moult later than consectlics in the Pribiols (St. Paul, St. George), or the Aleutians (Kiska). Groups sharing the same letter are not considered statistically different.

Colony	Year	n	Initiation Date	Initiation Date	Duration ± S.E.
			± S.E. (days)	S.D. ± S.E.	(days)
				(days)	
Cape	1988	129	28 Jul ± 1.8°	$11.3\pm0.1^{\rm b}$	$54.4\pm6.6^{\circ}$
Ulyakhpen					
Cape	1989	195	6 Jul ± 2.5 <sup>b</sup>	$11.3\pm0.1^{\rm b}$	$75.2\pm8.3^{ab}$
Ulyakhpen					
Cape	1990	386	12 Jul ± 2.2 <sup>b</sup>	$10.2\pm0.1^{a}$	60.5 ± 7.2*
Ulyakhpen					
St. Paul	2000	310	9 Jul ± 2.3 <sup>b</sup>	$11.5\pm0.1^{\rm b}$	$66.1\pm8.6^{*}$
St. Paul	2004	182	10 Jun ± 5.3ª	$17.9\pm0.2^{\circ}$	$114.0\pm18.b^6$
St. George	2005	232	14 Jun ± 6.3ª	$23.0\pm0.2^{e}$	133.2 ± 23.1 <sup>b</sup>
St. George	2007	143	7 Jul ± 2.2 <sup>b</sup>	$10.3\pm0.1^{\rm a}$	76.6 ± 9.1 <sup>ab</sup>
Kiska	2008	214	13 Jun ± 3.5°	$19.6\pm0.1^d$	$93.6\pm14.3^{ab}$

Table 4-4 - The majority of Least Auklets examined from museum collections had completed primary moult by the end of October, and all had completed primary moult by the end of November. Data are pooled among sampling locations, age classes, and years. PFMG = proportion of feather mass grown (P1 through P10).

Month	n	PFMG ± S.D.	Range
September	25	0.32 ± 0.27	0.03-1.00
October	13	$0.91 \pm 0.15$	0.62-1.00
November	7	$0.99 \pm 0.03$	0.93-1.00
December	11	$1.00 \pm 0.00$	1.00-1.00
January	10	$1.00 \pm 0.00$	1.00-1.00

# CHAPTER 5: STABLE ISOTOPES REVEAL VARIATION IN FORAGING ACTIVITY OF

# INCUBATING LEAST AUKLETS (AETHIA PUSILLA)

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# Abstract

Central-place foraging birds, such as breeding seabirds, face a trade-off between satisfying the energetic demands of growing nestlings and meeting their own nutritional requirements. We investigated the foraging patterns of breeding adult Least Auklets (Aethia pusilla) at Kiska Island, Aleutian Islands, Alaska, in 2009. Because of Least Auklets' small size (85 g), suitable attached data recorders were not available at the time of our study, so we used within-feather variation in \delta13C and 615N stable isotopes of the first primary feather (P1), which is grown during the incubation period. Controlling for individual differences, both 812 C and 815 N increased as the breeding season progressed, indicating that Least Auklet adults shifted their foraging patterns significantly. These results could be explained by adults gradually switching to different prev taxa, a seasonal change in isotopic ratios in a constant prev taxon, or a shift to a slightly more productive foraging location, such as an oceanographic front. Other work has shown that no seasonal change in chick diet prev taxa occurs, perhaps pointing to the latter hypothesis, which could involve a mechanism to balance adults' own energetic requirements with the energy required for the successful survival of their single egg and chick. Together, these results indicate that adult Least Auklets may balance their own demands with those of their egg and chick by a shift in foraging activity.

# Introduction

Stable-isotope ratios in seabird feathers, which are grown incrementally, provide information about foraging patterns, distribution, and environmental variability experienced by individuals during periods when they are inaccessible outside the breeding season (Cherel & Hobson 2007: Bond et al. 2010a). Hidden aspects of breeding birds' foraging activity can be revealed by stable-isotope ratios in feathers that are grown during the breeding season. Apart from advances in laboratory techniques and equipment that have reduced the cost of analyses, many recent advances in the application of stable isotopes to ecological questions have been because tissues such as feathers, blood, or claws can be sampled non-destructively (Hobson & Clark 1992a; Bearhop et al. 2004). Because only a small amount of tissue is required for analysis (approximately 0.25 mg), large feathers can be sub-sampled several times to gain insight into species' foraging ecology with greater temporal resolution. Variation in foraging patterns during feather growth will result in within-feather isotopic heterogeneity because exogenous nutrients are incorporated immediately into growing feathers (Murphy 1996), except in species that fast during moult (Cherel et al. 2005a; Fox et al. 2009).

We examined within-feather variation in  $\delta^{3/2}$  and  $\delta^{3/3}$  Ni in the first primary feather (P1) of Least Auklets (Aethyla pushillo), a small planktivorous auk endemic to the Bering and Okhotik seas. Adults eat calanoid coopspods, mainly Neocolonus plumchrus and A. (Jemingero (Rédard 1969a; Hum Jr. et al. 1998), which are advected by

oceanographic fronts (huter, Jr. & Harrison 1909, Huut Jr. 1907; Hustell et al. 1999). Previous studies of adult auklet diet were based on stomach contents (Bédard 1969a; Harrison 1990; Huut Jr. et al. 1998), which often represent the previous meal only, and do not account for temporal variation (DUF) & Jacksten 1980). Auklett foraging distribution has also been studied in relation to oceanographic features (Kinder et al. 1983; Harrison et al. 1990; Russell et al. 1999), but vessel surveys are indicative of population-level patterns, do not address individual afferences, and are a to coarse a resolution to infer differences at a fine temporal scale (e.g., within the incubation preford). Recause colonies (= 1 million individuality; Sowis et al. 1978), attached data recorders are not yet practical. The stable-isotope analysis approach therefore offers a unique opportunity to investigate individual addut' foraging activity with a fine temporal resolution, in one ato examine potential changes in adult self-feeding as chicks' demands increase.

Least Aukket' P1, the intermost primary is approximately 43 mm lone. The old P1 feather is shed near the beginning of incubation, and the new replacement feather takes 20 days to grow approximately 23 mm day<sup>10</sup> (Heary EJL comytaley 2000; Rohwer et al. 2000). Least Auhlets incubate their egg for approximately 30 d (Seal) 1984; Furt et ol. 1990b, and at Kiska, peak hatching date is around 30 June 13 d (Chapter 2). With a man moult initiation date at Kiska of 13 June 13.5 d (Harey 1991), the majoric of the feather's incubated entity incubation, when datils must balance

their own nutritional requirements with the successful incubation of their egg. Our objectives were to 1] quantify within-feather heterogeneity in  $\delta^{10}$ C and  $\delta^{10}$  M in Least Aulide 1P inmaries replaced during the incubation period; 2] use these isotopic signatures to elucidate possible changes in adult foraging patterns, and 3] to identify important areas for future investigation of questions related to the trade-off between set maintenance and parental area in this patkitrixous adult.

# Methods

We sampled the right P1 from ten adult teast Addets at Sirus Point, Kiaka Island, Aleutian Islands, Alaska (32'08'N, 177'37'E), in July 2009 (Rond & Jones 2009a). Heathers were 75-100% grown. We cleaned feathers by washing in 0.25M NAOH (Beacherp et al 2000; Bond & Diamod 2009). Feathers were divided into four regions (A-D), from distal to proximal ends of feathers, each representing approximately 5-8 mm of the feather vance. Approximately 0.25 mg of feather harb was sampled from each region, representing tissue that was grown during days 1-3 (A), 4-8 (B), 9-12 (C), and 13-13 (D) (for each feather. You fail vance areas were sampled (ag., days 15-20), because one feather were not fully grown. Each sample was placed in at nou, cruthed, and combusted at 1800°C in a Carlo Erba NA1500 Series II elemental analyzer for analysis of compounds for instoper ratios. Resultant gases were segarated by a packed G column and analyzed by a Dela V Pius continuous-flow isotope ratio mass a 50 of repeated

measurements; MEA-H1 &<sup>3</sup>Nc 0.52 ± 0.13, IAEA-N2 &<sup>3</sup>Nc 20.32 ± 0.04, IAEA-CH6 &<sup>13</sup>C · -10.45 ± 0.06, U5G5-24 Graphite &<sup>13</sup>C · -16.04 ± 0.14, U5G5-25 &<sup>55</sup>Nc · -30.25 ± 0.04) and lab standards (MUN Sulphanilamide &<sup>13</sup>C · -28.23 ± 0.22, &<sup>13</sup>Nc · -1.51 ± 0.09, MUN-CO-2 &<sup>13</sup>C · -40.11 ± 0.09), which cover the range of isotopic values in our samples.

## Statistical Analysis

We analysed trends within individuals using a repeated measures MOVIA in 5955 16 (5955 (nc., Chicago, IL, USA), and tested the assumption of sphericity using Mauchly S W (Mauchly 1540) and homogeneity of variance using Levent's test (Levene 1960). We also included a post-hoc analysis to detect any linear or quadratic trends over the feature length of the feather.

# Results

Within individuals, the range of values of  $\delta^{10}$ C was between 0.70-2.43%, and differences within feathers in  $\delta^{20}$ N were between 0.30-2.42%, isotoper ratios met the assumptions of the linear model (Levens's test, all p = 0.10), but net sphericity for  $\delta^{10}$ C (Mauchy's W,  $\delta^{10}$ C, p < 0.033,  $\delta^{20}$ Hz p = 0.473). We therefore used the Greenhouse-Geisser (1995) correction for subsequent analyses of  $\delta^{10}$ C, as  $\varepsilon < 0.375$  (Girden 1992). Controlling for individual differences, there were significant within feather differences in  $\delta^{10}$ C is c < 0.013 at  $\delta^{10}$ C is c < 0.013 at  $\delta^{10}$ C is c < 0.013 at  $\delta^{10}$ C.

individuals, with both  $\delta^{10}$ C and  $\delta^{15}$ N increasing through the incubation period (mean difference between sections A and D ± SD;  $\delta^{10}$ C: +1.5 ± 0.7%, p < 0.001,  $\delta^{13}$ N: + 0.8 ± 0.7%, p = 0.022; Figure 5-1).

#### Discussion

Seabrids, as central-place foragers, are expected to minimize transit time between their breeding colony and food resources (Baind 1991), and to balance the trade-off between distance to foraging grounds and the energy and nutrients sequired (Okset et al. 1995; Effect et al. 2009; Differences in time allocated cost eff-anitematic and parental cue should balance the life-history trade off between current reproduction and survival in adults (Clark & Ydenberg 1990s; Stearns 1992; Dobon & Jouventin 2020b), Auks that forage by wing-propetied pursuit-diving, such as teart Auklets, have high wing lossdings compared with other brids, and the energetic costs of flying between foragine areas and the horeform tide are Will food at al. 1993.

In Least Auklets, we found a consistent shift towards higher 8<sup>10</sup>C in feasther tissue deposited as incluation progressed, with a similar but weaker trend for 8<sup>10</sup>C. In feasther results could be explained by several processes: 1) adults gradually switched to different prey taxs; 2) a seasonal change in isotopic ratios in prey; 3) a shift in forraging location, or 4) metabolism of endermous nutrients.

Heterogeneity in feather 3<sup>15</sup>C and 6<sup>18</sup>C an reflect changes in the isotopic content of diet (and hence in foraging patterns) at a fine temporal resolution. We did not sample adult stomachs to determine diet, but there were no within-season differences in chick diet during 1993-2006 (Chapter 3), so we have no reason to suppert a diet shift in adults. Least Addets' primary previ n the Alextical (Veccalarous coopeo)d are the most abundant zooplankter in the North Pacific Ocean (Mackas et al. 1998), and their peak abundance coincides with Least Auklets' nesting period June and July: Batten & Mackas 2009). We therefore have no a priori reason to suppert that adult diet composition changes throughout the breeding season, although independent verification is regulared to support this are for.

A seasonal change in prev isotopic ratios (hypothesis 2) would explain the change in feather instope ratios we observed (assuming taxonomic constancy in diet). Such a phenomenon has not been documented in the Bering Sea (but see Rau et al. 2005 for discussions on inter-annual variability in plankton stable isotope ratios in the California Current). Because no such data exists for the Bering Sea, we cannot confirm or refute this hypothesis.

Least Addets lose up to 7% of their body mass around the time the egg hatches, while maintaining a relatively constant mass during incubation. This mass loss is presumably a mechanism to decrease flight costs while adults provision next-bound chicks (lones 1994). Mass loss may be related to replacement of up to four flight feathers at the same ine (linen; 1993). Metaboling endgement antireit reserves

(hypothesis 4) could influence the isotopic ratios in feathers during this time (Fox et al. 2009), and would be represented in the most proximal feather section (section D) in our study.

A change in foraging behaviour during incubation, such as foraging closer to the colony site, might also explain the results (hypothesis 3), "Isoscapes," or geographic patterns in stable-isotope ratios, have been described for calanoid copepods in the Bering Sea (Schell et al. 1998). These isoscapes show relatively constant 813C and 815N values in copepods in the western Aleutian Islands, but at a coarse resolution (areas pooled in 2% contours), and there is no information on temporal variability. The consistent shift among individuals along a geographic isotope gradient suggests a geographic shift. Higher 813C is associated to increase phytoplankton growth as occurs in nutrient-rich waters, such as inshore systems or oceanic fronts (Goericke & Fry 1994; Popp et al. 1998; Graham et al. 2010). Least Auklets preferred prey, Neocalanus copeopds, are often concentrated in such fronts in the Bering Sea (Hunt Jr. et al. 1998: Russell et al. 1999). A shift towards more productive systems at the end of incubation and beginning of chick rearing could be related to adults' increased energetic demands as chicks hatch, and require their parents to provision them several times per day, with increasing amounts of food (Roby & Brink 1986a; Chapter 3).

During incubation, adults expend energy caring for their egg, and have developed several strategies to accommodate chicks' demands. Incubation shifts average 23.6 ± 5.6 h (Roby & Brink 1986a), and during this time, off-duty parents can range wider, and

exploit more distant resources. Once the chick hatches, it is broaded continuously for the first five days (lones 1993b), and fed 2.6 ± 0.7 times/day (Roby & Brink 1986a), restricting the parents' ability to travel far. By foraging in productive areas, such as coasile forst, close to be breeding colony, adults reduce their costs of selfmaintenance, and can increase the time allotted to chick provisioning. Adults may not exploit these productive areas during incubation because they are less selective in prey choice, or temporal effects on the distribution and abundance of zooplankton, and development of upwellings.

There was also a trend in increasing 6<sup>19</sup>N through the incubation period. This change was small in 7 of 10 cases (range < 1.0%), but might also reflect a distary change based on shift in foraging area. Quantitative estimates of adult auktet diet using stable inotope analysis are not possible because discrimination factors, the change in 8<sup>10</sup>C or 8<sup>10</sup>N between prey and consumer, are specific to each consumer (Caut et al. 2008), and can influence mixing model output greatly (Bond & Diamond 2011); no such discrimination factors are available for advects.

Continued miniaturization of data loggers will permit telemetric studies in the near future that would provide much more detailed insight into adult auklets' foraging patterns. Jocotopic investigations of differences in adult and chick diet are also possible provided that the differences in metabolism between growing chicks and adults can be accounted for (Promote & Averbach, 1992): Chern et al. 2005.

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Figure 5-1 – Controlling for individual differences, both 5<sup>10</sup>C (A, p < 0.001) and 5<sup>10</sup>N (B, p = 0.022) in individual Least Auklets' first primary feather (P1) increased from incubation (feather section A) to early chick rearing (feather section D). Each line represents totope values from a single individual.

# CHAPTER 6: STABLE ISOTOPES INDICATE SEASONAL AND AGE-RELATED

# VARIATION IN FORAGING BY LEAST AND CRESTED AUKLETS IN THE ALEUTIAN

# ISLANDS, ALASKA

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## Abstract

Least and Crested Auklets (Aethia pusilla and A. cristatella) replace breast feathers during the pre-breeding period, and primaries form during the incubation and post-breeding periods. Thus, by sampling different feathers from a single individual on one occasion, we can obtain foraging information from the majority of the annual cycle using stable-isotope analysis. (SIA). We sampled adult and subadult Least Auklets and adult Crested Auklets at Buldir, Kiska, and Gareloi Islands, in the western Aleutian Islands, Alaska, from 2007-2010, and Least Auklet chicks at Kiska in 2009. There were no differences in isotope ratios among islands or years, but significant differences between species, among seasons (feather types), and within species between age classes. Adult Least Auklets had higher &<sup>15</sup>N than subadults during the pre-breeding period, and chicks during chick rearing. Crested Auklets had higher 615N than Least Auklets at Gareloi throughout the year. Both species showed a trophic shift during the post-breeding season, characterised by an increase in \delta<sup>15</sup>N, and for Least Auklets, in \delta<sup>13</sup>C as well, Species' isotopic differences were likely caused by differences in diet composition of Crested (euphausiids) and Least Auklets (copepods). Subadults are inherently less experienced than adults, probably still learning to forage during their first winter and thus taking different food items than adults. Least Auklet adults' higher  $\delta^{15} N$  relative to chicks likely reflects the abundance of zooplankton during the breeding season, and changes in adult physiology and self-feeding patterns during chick rearing. Our results

provide hypotheses for testing in future studies using attached data loggers to measure foraging locations and movement outside of the breeding season.

## Introduction

Prev availability and variability in the diet of marine animals have profound influences on demography (Gall et al. 2006: Sprensen et al. 2009), are influenced by oceanographic and climatic patterns at the landscape level (Durant et al. 2003; Hobson et al. 2004) and determine individual health and breeding condition (Sorensen et al. 2010). During the breeding season, adult seabirds provision nest-bound young, and often face a trade-off between their own energetic demands, and the requirements of their eee or young (Clark & Ydenberg 1990b). This has resulted in a diversification of strategies to balance current reproductive effort with future survival, including alternative strategies when adults self-provision. An often-implicit assumption of seahird dietary studies is that adult and chick diet are similar, but this is often not the case (Wehle 1982, 1983; Wilson et al. 2004). A good example is the bimodal foraging strategy of some species, where adults make frequent brief trips to provision young, but make longer less frequent foraging trips to replenish their stores with high energycontent prev (e.e., Chaurand & Weimerskirch 1994; Weimerskirch et al. 1994; Welcker et al. 2009a). To gain a full understanding of the foraging behaviour and requirements of adults, quantifying the diet provided to nestlings is insufficient.

Aukkets (Aerkhir spp.) are the most abundant seabirds in the North Pacific Ocean and Bering Sea (Baston & Jones 1998). Aukkets breed in dense colonies in the Aleutian Islands, Alaska, and winter in the North Pacific Ocean (Jones 1993a, b). Aukket diet during the breeding season is relatively well studied, although most information

concerns nesting diel. In the Alexitain Islands, adults provision chicks with calanoid coopeops (Detectionus spc) and exphassids (Thystoneous spc). [Day & Byrd 1999; Chapter 3). Limited information suggests that, in the Alexitains, adults consume coopeopds, exphassids, and smaller numbers of amphipods and cephalopods during the breeding season (Day & Byrd 1999; Troy & Braditeret 1991; Hunt J. *et al.* 1998). The winter ecology and behaviour of auklets is largely unknown, and consequently there are few good quantitative descriptions of winter diet (Stejineger 1885; Troy & Braditeret 1991). The diet of subadult Least Auklets (birds in their second year of life) is unknown. These individuals attend breeding coionies sporadically as non-breedert and, not bundened with the constraints or bundened with the contraints of bundened bundened with the contraints of bundened with the contraints of bundened bundened with the contraints of bundened with the contraints of bundened bundened with the contraints of bundened with the contraints of bundened bundened bundened bundened bundened with the contraints of bundened bund

In the past 25 years, biochemical techniques to evaluate avian det indirectly have been developed (Holson 1987; herone et al. 2004). Using these forensic techniques has enabled researchers to study seabird diet during periods when birds are away from breeding outlooks, which for many species, covers the majority of the annual cycle. Stable-isotope ratios of carbon and nitrogen are now used frequently to characterise seabird foraging behaviour (Barrett et al. 2007). Values are expressed as the parts-gene-thousand ratio of the havier ("N or "C) to the lighter isotope ("N or "C) as compared with an international standard (Pee De Belennite for C, atmospheric air for N),  $\delta^{10}N$ , or the ratio of  ${}^{10}N'^{10}N_{1}$  is a continuous measure of trephic position, and it increases between 276 with their throtics the because" N is exceeded arected areforemating to the share of  ${}^{10}N'^{10}N_{1}$  is a container of the peince position, and it increases between 276 with their throtics the because "N is exceeded preferentially and the located preferentially to the specific of the section of the because of the careted preferentially to the specific of the specific of the careted preferentially of the lighter position, and it increases between 276 with their throtics the because "N is accented preferentially to the lighter position, and it increases between 276 with their throtics the because "N is accented preferential to the lighter baseling".

Introgenoou water (Skeele & Daniel 1978; Minagawa & Wada 1984; Keiny 2000). There is list troppic enrichment of  $\delta^{12}$ C at higher troppic levels (Bau *et al.* 1983; Hobona & Welch 1992), but marine (Bicardonate-based) and terrestrial carbon sources (CO<sub>2</sub> based) differ in their  $\delta^{12}$ C values (Pitersona & Fry 1987).  $\delta^{12}$  C an also provide information on foraging area in the marine environment because of spatial "isoccapes" in the environment (Hobosn 1999; Cherel & Hobon 2007; Graham *et al.* 2010). Variation in  $\delta^{12}$ C results from varying levels of primary production, with areas of grater phytophatom growth such as upwellings and inshore areas being enriched in  $^{12}$ C relative to unimed perface; rothers (Socciae & Fry 1994; Proper of 1. 1996).

Ferthers are very useful for trable-isotope analysis because they can be sampled non-lethally, stored easily in remote field settings, and (because they are here none growth their totope calces represend et al. the time of synthesis (blobon & Clark 1992a; but see Fox et al. 2009). Availets moult primary feathers sequentially, beginning around the time when chicks start to hatch, and continuing from P1 to P10 (innermost to outermost primary) until like October or early Nevember (Kornykahov 2000; Claptier 4); the same is likely true for Crested Availets (Bédard & Sew) 1984; Pole 2000; and those of subadult birds between December-April (Pyle 2000; blue by sampling three feathers from each individual, one can investigate stable lostopes from late incubation/early clack rearing (P1), autumn post-breeding (P10), and spring pre-breeding (Doby; Pyle 2009).

We had several objectives: Inst, because the diet and foraging ecology of subaduit Least Auklets is entirely unknown, we compared the spring pre-breeding isotopic ratios of Least Auklet adults and subadults at Kiska Island between 2007 2010. We also compared <sup>34</sup>° and <sup>34</sup><sup>15</sup> between adults and chicks. Second, we compared the diet of Least Auklet adults from three breeding colonies throughout the annual cycle, as chick diet is relatively consistent among islands [Chapter 3]. Finally, we compared the isotopic ratios of adult Least and Crestel Auklets from Gareloi Island throughout the annual cycle to gain insight into the isotopic overlap between these ecologically similar species.

### Methods

#### Auklet capture & field sampling

We captured Least Auklets on Main Talus, Build Island (52/2374, 177397; adults in 2008 and 2009), Sirius Point, Kiska Island (52/08/N, 177397; adults and Subadatis in 2007 2010, clicks in 2008), and Southeast Colory, Garelei Island (51/44/N, 178/45/W, adults in 2009). Crested Auklets were captured at Carelei in 2009 only. We captured birds using noose carpets set in the breeding colony during the auklets' activity period between 0000 1600 Hawiii Aveutian Daylight Time (Byrd et al. 1988); Jones 1992). Adult Least Auklets were those in definitive alternate plumage, indicated by extensive white stranks on the forehead, a redder bill gain dies wom grunteris compared with subadults. Subadults (12 months doll in July) have a brown forthead with no streaking, worn primaries that appear brown, and a darber bill and often have dark spotted or entriety dark throast feathering (Bédard & Sealy 1994, Jones 1993b). Created Auklet adults were identified by their large, orange lith hamphotheca. grey forthead (x. brown in subadults), and fresher primaries (Jones 1993a). Birlds 24 months and older are indistinguishable. Once we determined a bird's app, we measured and banded birds, and took feather samples (P1 and P12 from the right wing, breast feathers from the upper breast). We placed feathers in individual paper envelopes, kept cool in the field, and freasest a 20° cmo our return from the field.

#### Laboratory analysis

Prior to analysis, feathers were washed with 0.25M NaOH and the main-drift of remove external contamination (Bearloo et al. 2000); Bond & Diamond 2009). Individual aukket feathers are heterogeneous isotopically (Chapter 5), so we sampled each feather in the same way. Approximately 0.2.0.3m (from the distalt for the wave was cut, placed in this cape, and crushed using sterilized forreise). Samples were combusted at 1800°C in a Carlo Erba NAI500 Series II elemental analyser for continuous flow analysis of compounds for intoge ratios. The resultant games were separated by a packed GC column and analysed by a Delta Y Plus isotope ratio mass spectrometre though a continuous flow interface (CF-IRMS), isotope values were calibrated uning method blanks and stander derevence matrixel MRAP4 (men s Sb., 5<sup>4</sup>% 20.32 z ; 0.09%a), MEA-CHE (0<sup>3</sup>C: -1.0.3 5: 0.09%a), USGS-24 (graphite, 0<sup>13</sup>C: -16.11: 0.15%a) and USGS-35 (0<sup>13</sup>C: -30.25: 0.05%a) and Internal lab standards MUP CO 2 (0<sup>13</sup>C: -40.11: 1 0.10%a) and MUN Sulphamilamide (0<sup>13</sup>C: -38.36: 1.0.11; 8<sup>-1</sup>N: -1.42: 1.0.09). Standards covered the range of isotopic values in our samples. Mean recovery of all standards ranged from 99-101%. Within-run duplicate samples (9% of total samples) yielded mean standard deviations of 0.02%, for 8<sup>13</sup> can do 405%e (for 6<sup>13</sup> %

### Statistical methods

Quantitative lootope mixing models are sensitive to variation in species-specific discrimination factors (the change in 631c or 635M between prev and consumer; Bord & Diamond 2011). We have restricted our analysis to a comparative description because in discrimination factors have been established for availates.

Analysies were conducted in 5455 16 (5455 km, Chicago, IL, USA). Becaute 0<sup>13</sup>C and 0<sup>34</sup>N from the same sample are not statistically independent, we used multivariate analysis of variance (MANOVA) to assess differences among species and age classes in stabilise instoperatios based analysis of variance (RAOVA) to examine each totope ratio separately in cases where the MANOVA revealed a significant difference. To test for differences among feather types within individuals, we used repeatedmeasures ANOVA, and tested for sphericity using Mauchy's W (Mauchy 1940). When the sphericity assumption was not me, we used the Greenbource Gostes (1959)

correction when e < 0.75, and the Huynh-Feldt (1976) correction when e > 0.75 (Girden 1992). We assessed homogeneity of variance using Levene's test (Levene 1960). We used 95% confidence intervals for post hoc comparisons. All tests were considered significant when p > 0.05. biotope data are presented as mean \$ 5.0.

### Results

## Differences among Least Auklet age classes

We tested age class differences using treast features from Kika collected in 2007-2010. Variances of 3<sup>3</sup>°C were homogenous, but those of 3<sup>3</sup><sup>3</sup>N were not [Levent's test, b<sup>3</sup>°C<sub>2</sub> = 1 0.00, b<sup>3</sup>N<sub>1</sub> = 0.03). AMAGVA detected significant differences between exploses in isotoper ratios (Kika adults usidadits. With's  $\lambda = 0.66$ ,  $F_{1,m} = 17.73$ , p <0.001), but not years (2007-2010, Wilks'  $\lambda = 0.97$ ,  $F_{1,1m} = 0.42$ ,  $\mu = 0.87$ ) or the year\*age interaction (Wilks'  $\lambda = 0.91$ ,  $F_{1,1m} = 0.82$  p = 0.56). Adults had higher b<sup>3</sup>°C (-18.81 ± 1.8% vs. -1.395 ± 1.8%,  $F_{1,2m} = 6.88$ ,  $\mu = 0.013$ ) and b<sup>3</sup>N (13.21 ± 1.6% +vs. 1.10 ± 1.4%4,  $F_{1,2m} = 34.94$ ,  $\mu < 0.003$ ) isotope ratios than subadults during the pre-breeding period (Figure 6-1).

To compare adults and chicks from Kiska, we used chick breast feathers (collected in 3008) and adult P1 feathers, which were grown at the same time. Because we found no differences among years in adult P1 feathers (see "Differences among colonies and seasons" below, dast form Kiska in 2008 and 2009 were polede. Neither

 $\Delta^{10}$  C no  $\Delta^{10}$  En or  $\Delta^{10}$  En had homogenous variances (Leven's test, both p < 0.001). Isotope ratios differed between age classes (WIIS'  $\lambda = 0.103$ ,  $F_{1,23} = 10.83$ , p < 0.001). Adults had higher  $\Delta^{10}$ C (18.98 ± 1.85% vs. -20.67 ± 0.35%,  $F_{1,24} = 6.43$ , p = 0.018) and  $\Delta^{10}$ N (12.30 ± 0.95% vs. 8.88 ± 0.95%  $F_{1,24} = 8.23$ . p = 0.001) than chick in Figure 6-2).

### Differences among colonies and seasons

We used P1, P10, and breast feathers to test for differences in  $\delta^{13}C$  and  $\delta^{15}N$ among the three colonies (Buldir, Kiska, Gareloi) in 2009. Neither  $\delta^{13}$ C nor  $\delta^{15}$ N had homogenous variances (Levene's test, both p < 0.001). Isotope ratios differed among islands, feather types, years, and in some of their interactions, so we used repeatedmeasures univariate analyses to examine differences in detail.  $\delta^{13}$ C did not meet the assumptions of sphericity (Mauchly's W = 0.80,  $\gamma^2_2$  = 9.44, p = 0.009), and we used the Huynh-Feldt correction ( $\epsilon = 0.94$ ). Isotope ratios among feather types varied within individuals (F<sub>1.83,83,01</sub> = 11.09, p < 0.001): P10 had higher  $\delta^{13}C$  values than P1 or breast feathers (Figure 6-3). Among individuals, there were no differences in 813C among the three islands (F144 = 1.59, p = 0.22), or years (F144 = 3.48, p = 0.07), and the island x year interaction was not significant ( $F_{1,44} = 0.86$ , p = 0.36).  $\delta^{15}N$  also did not meet the assumption of sphericity (Mauchly's W = 0.78,  $\gamma^2_2$  = 10.87, p = 0.004), and we again applied the Huynh-Feldt correction ( $\epsilon = 0.92$ ). Isotope ratios in feather types differed within individuals (F1843112 = 13.54, p < 0.001). P10 also was enriched in <sup>15</sup>N compared

with P1 and breast feathers (Figure 6-3). Among individuals, there was no difference in  $\delta^{15}N$  among islands ( $F_{2.44} = 0.77$ , p = 0.47), or years ( $F_{2.44} = 1.82$ , p = 0.19), and the island x vear interaction was not significant ( $F_{1.44} = 1.76$ , p = 0.19).

## Differences among species

We tested differences between Lesst and Crested Auklets using P. P. P.D, and breast features from sarrolin 2009. Our data met the assumptions of Inear models, and variances of both isotopes were equal (Lewen's test, both p > 0.20). Isotope ratios differed among feature types (Wilk' > 0.78, p = 0.01) and species (Wilk' > 0.72, p < 0.00), but the feather types (Wilk' > 0.78, p = 0.01) and species (Wilk' > 0.70, p = 0.42). Both  $h^{12}$  and  $h^{13}$  met the assumptions of sphericity (Masuley's W.  $h^{12}$ Co. B.7,  $\chi^2$ , h = 2.7, p = 0.31,  $h^{11}$ L 0.79,  $\chi^2$ , h = 2.7, p = 0.11,  $h^{12}$ C did not differ between species at Garolo (F<sub>1,10</sub> = 1.40, p = 0.25) or among feather types (F<sub>1,10</sub> = 0.70, p = 0.41), and the species a relater type interaction was not significant (F<sub>1,10</sub> = 0.01, p = 0.41), and the species a feather type interaction was entitively (F<sub>1,10</sub> = 5.15, p = 0.036, Figure 6.44), and for both species, P10 was enriched in <sup>10</sup> (Compared with P1 or breast feathers (F<sub>1,10</sub> = 1.306, p = 0.002; Figure 6.41, dasin, the feather type species interaction was not species fractively (Figure 6.4), dasin, the feather type species interaction was not species fractively (Figure 6.4), dasin, the feather type species interaction was not species fractively (Figure 6.4), dasin, the feather type species interaction was not species fractively (Figure 6.4), dasin, the feather type species interaction was not species fractively (Figure 6.4), dasin, the feather type species interaction was not species fractively (Figure 6.4), dasin, the feather type species interaction was not species fractively (Figure 6.4), dasin, the feather type species interaction was not species fractively (Figure 6.4), dasin, the feather type species interaction was not species fractively (Figure 6.4), dasin, the feather type species interaction was not species fractively (Figure 6.4), dasin, the feather type species interaction was not species fractively (Figure 6.4), dasin, the feather type species interaction was not species fe

# Discussion

The use of stable intolese to lifer foraging patterns can provide valuable insight into the ecology of difficult-to-study groups, such as pre-bereding subadults, or cionial birds away from breeding colonias (Barter et al. 2007; Inger & Bearhop 2008). Quantitative die etterstates uning stable sotope ratios in commune alto require inotope ratios in prey, and die-tissue discrimination factors. Information on audiest' winter diet is lacking (lones 1933b), and discrimination factors are unknown (Bond & Jones 2009b), and simultaneous prey sampling to monitor spatial and temporal variability in zooplanktom stable-istope ratios was lacking (Cline Jr 1999), which limits our interpretation to a qualitative comparison.

### Differences among age classes

Stable lostopes revealed both differences and similarities among proves of aukiets. We found age-related differences in both  $\delta^{12}$ C and  $\delta^{10}$  N of breast feathers of adult and subadult Lasst Aukiets at Kiska. Breast feathers are replaced during the prebreeding period in both age classes around March-May (Vye 2008). The difference in  $\delta^{12}$ C between adults (at least 3 years old) and subadults (approximately 12 months old, Pyle 2008) wave about 5%, within the error estimate of adultets' proy (Schell et al. 1998), with locides defined 12 kinetravia, although three can be both report and anglatal There is a horown losscape of zooplankton  $\delta^{12}$ C in the being Sea (Schell et al. 1998), with locides defined 12 kinetravia, although three can be both report and adpatial variation in zooplankton incoope ratios (IIIne Jr 1939). Least Auklets begin attending breeding colonies in April. Differences in  $\delta^{3/2}$  are therefore likely the result of variation in the precise time and location of feather replacement among individuals rather than spatial segregation of age classes.

There was a 2% difference between adults and subadults in  $\delta^{15}$ N. There are three possible causes for this difference. First, adults and subadults may forage in separate areas that differ in prev &<sup>15</sup>N. Very small differences among age classes in &<sup>13</sup>C make this hypothesis unlikely. Second, there may be metabolic differences that cause differences in isotopic discrimination between age classes, as growing seabirds' discrimination factors differ from adults (Sears et al. 2009). While subadult Least Auklets are smaller than adults (Jones 1993b, authors' unpubl. data), Sears et al. (2009) found that once juveniles reached a stable mass, isotopic discrimination was similar to that of fully-grown adults. When Least Auklet chicks depart from Kiska at the end of the breeding season, they are at 90% of adult mass (73 g vs. 81 g, Major et al, 2006). The mass of subadults we sampled (79 ± 4 g) was similar to that of adults sampled (82 ± 3 g). so differences in isotopic discrimination were not likely the cause of differences in  $\delta^{15}N$ . The difference in δ<sup>15</sup>N between age classes may reflect true dietary differences. Quantitative data on the winter diet of Least Auklets are lacking (Steineger 1885; Troy & Bradstreet 1991), subadults' diet is completely unknown. In general, diet data for prebreeding-aged seabirds are sparse because the birds attend breeding colonies irregularly and are seldom captured (Hamer et al. 2002). Previous studies have found differences

between diets of adult and subadult Tufted Putlins (*Protercula cirbata*) in the Aleutian Islands (Wehle 1982) and in the North Pacific (Tanaka 1989). Our results suggest that subadult Least Auklets forage at a lower trophic position than adults during the prebreeding period. This is most likely the result of subadult birds learning to forage for themselves, as younger birds are often less proficient at finding and capturing highsuplicy regretates. Rechers 1996, Cherol et al. 2008).

Adult Least Auklets had higher 513C and 515N than chicks during the breeding season. As mentioned, growing chicks' metabolism likely contributed to some of these differences, but the magnitude of the differences we found (1.5% for  $\delta^{13}$ C, 3.5% for  $\delta^{15}$ N) are larger than growth-related effects (Sears et al. 2009). There may also be a small effect of feather type (contour vs. primary) caused by differences in the concentration of the pigment melanin (Michalik et al. 2010; Wiley et al. 2010), breast feathers being mostly white, and primaries mostly dark (Jones 1993b). We believe that our qualitative comparisons of adult and chick stable-isotope ratios remain valid. however (Ponsard & Averbuch 1999: Vanderklift & Ponsard 2003: Cherel 2008). Least Auklet chick diet in the Aleutian Islands is well documented, and consists mainly of Neocalanus spp. copepods (often > 90% of biomass), and Thysanoessa spp. euphauslids (Day & Byrd 1989: Chapter 3). Adult diet during the breeding season is less well known. but the few studies have indicated similar prev in adult stomachs (Bédard 1969a: Harrison 1990; Hunt Jr. & Harrison 1990; Hunt Jr. et al. 1998; Russell et al. 1999).

Adult Least Auklets experienced a shift in diet through the breeding season. increasing both their  $\delta^{15}N$  and  $\delta^{13}C$  values from incubation to early chick rearing (Chapter 5), while the composition of food delivered to chicks remained constant during the same time (Chapter 3). Adults therefore forage for prey from different trophic positions ( $\delta^{15}N$ ) and in different areas ( $\delta^{13}C$ ) depending on whether the prey is meant for self-feeding or delivery to their chick. Increased  $\delta^{13}$ C is associated with increased phytoplankton photosynthetic activity and growth (Goericke & Fry 1994; Popp et al. 1998), and is characteristic of inshore and upwelling systems (Graham et al. 2010; Jaeger et al. 2010). Least Auklets forage in upwellings and areas where zooplankton is advected, where Neocalanus copepods are concentrated (Springer & Roseneau 1985; Hunt Jr. 1997; Hunt Jr. et al. 1998; Russell et al. 1999; Platt & Springer 2003; Springer et al. 2007). Nevertheless, the magnitude of the difference in stable-isotope ratios between adults in chicks remains puzzling. Our results suggest that adult Least Auklets are foraging on different prey from their chicks, or in areas with vastly different isotopic ratios. Least auklet individuals are very difficult to observe at sea, so direct observations of feeding are challenging logistically, and this species is too small (85 g mean mass) to carry telemetric equipment that would measure foraging behaviour in detail. Further experiments using existing captive populations should quantify Least Auklets' stableisotope discrimination factors, and use these to construct quantitative isotope mixing models for all age classes.

Our results contrast with bimodal foraging strategies, reported mainly in tropical and subtropical Procellariiformes (Cherel et al. 2005b; Congdon et al. 2005; Kojadinovic et al. 2008; Bond et al. 2010a), and recently described for Dovekies (Alle alle), a polar non-Procellariiform, and ecological counterpart to Least Auklets in the Atlantic Ocean (Welcker et al. 2009b). Bimodal-foraging species alternate between frequent, short trips to provision chicks, and less frequent longer foraging trips to replenish their own nutrient reserves (Chaurand & Weimerskirch 1994: Weimerskirch et al. 1994) - Least Auklets do not apparently make use of this strategy. During the breeding season, auklets forage heavily on Neocalanus spp. copepods, the most abundant zooplankter in the North Pacific (Mackas et al. 1998), and copepods are advected by oceanic fronts. where auklets are found often in vessel surveys (Kinder et al. 1983; Harrison et al. 1990; Troy & Bradstreet 1991; Hunt Jr. 1997; Hunt Jr. et al. 1998). There is also little annual variation in the proportion of Neocolanus spp. copepods in food loads brought by adult Least Auklets to their chick (Chapter 3). Unlike less productive tropical and subtropical systems, or more variable polar systems. Least Auklets' preferred prev appears to be readily available, and in large supply. Adults' response to increased energetic demand by their chick is likely therefore reflected in a geographic shift and a shortening of their own foraging distance (Chapter 3).

Differences between adults and chicks in 8<sup>15</sup>N have been studied in many seabirds, but there is no consistent pattern in the differences. In some studies, chicks' 8<sup>15</sup>N is higher, and is explained by chicks' increased nutritional demands, and the (often

unverified) assumption that higher-\$<sup>10</sup>N foods are of higher quality (e.g., Bother et al. 2000; Hodum & Hobson 2000; Cherrel 2008; Richoux et al. 2010). In other studies, adults' higher \$<sup>10</sup>N is explained by a bimodal forarging strategy (e.g., Cherel et al. 2002; Harding et al. 2008; Bond et al. 2010a), and in other studies still, there is no difference (Hedd & Montevecchi 2006; Cherrel et al. 2008; Davies et al. 2009). The difference we found in mean \$<sup>10</sup>N between adults and chick Lossf Auldes (3.42%) is the greatest difference reported were adults have higher \$<sup>10</sup>N.

## Differences among colonies and seasons

We found no differences in 8<sup>1</sup>° cor 8<sup>1</sup>% among Least Auklets among tires or seasons (Resher types), suggesting that the composition of adult Least Auklets' prov is uniform during the breading season, and that adults moult in the same locations during the fail and apring. The Minis attached geolocitors were too large for Least Auklets during our study, our results suggest that telemetric results from birds from one colony may be generalized to those from other western Akuluin breeding sites. There were also no differences between years in any intospic comparison, which implies that birds moult in the same areas year year (borne) et al. 2000).

We documented seasonal shifts in  $\delta^{11}C$  and  $\delta^{12}N$  within individual Least and Crested Auklets. Both species showed a trophic shift following breeding; as P10 feathers (autumn, post-breeding) had higher  $\delta^{12}N$  than breast (spring, pre-breeding) or P1

(summer, chick rearing) feathers. The relatively rapid and distinct change in  $\delta^{13}$ C in Least Auklets likely resulted from migration to wintering grounds. Unlike the congeneric Whiskered Auklet (Aethia avamaea), which is resident near breeding colonies yearround (Byrd & Williams 1993; Zubakin & Konyukhov 1994), Least Auklets likely disperse to the northwestern Pacific Ocean to areas around Japan (Shuntov 1965; Vyatkin 1981; Kondratyev et al. 2000; Sydeman et al. 2010). Some Crested Auklets winter close to the Aleutian Islands (Renner et al. 2008), and therefore a less significant seasonal change in Crested Auklets' & 33C was expected. The similarity between breast and P1 feathers' & 33C and  $\delta^{15}N$  suggests that body contour feathers may be moulted near breeding colonies. The timing of auklets' return to Aleutian breeding colonies is not known - adults have been observed at colony sites in April (Jones 1993a, b), and may perhaps arrive earlier. Both species showed a dietary shift during the post-breeding season, and withinindividual changes ranged from increases of 0-5.64% (mean ± S.D.: 2.05 ± 1.40%). It is possible that this shift is related to adults improving their condition following chick rearing and concurrent primary moult (Chapter 4). A portion of this increase could also be an artefact of protein metabolism resulting in putritive stress, which can alter &<sup>15</sup>N (Hobson et al. 1993; Sears et al. 2009). Within species, however, there was a large amount of variation in 513C and 515N during each season among individuals (Appendix 10). This suggests that there could be considerable individual specialization on certain prey types, or that individuals forage in different areas (Bearhop et al. 2006; Cherel et al. 2007).

## Differences among species

There were differences in stable-sidope ratios between species sampled at Garelsi. Created Auklets had higher  $\delta^{10}$  than Least Auklets, but we found no difference in  $\delta^{10}$ C. In the southern Bering Sea, explavatids have higher  $\delta^{10}$  than copeopds [Schell et al. 1990; Remains of higher trophic prey (squid beaks, gaid otolith) have also been found in adult Created Auklet stomachts (Braditeret 1985; Harrison 1990; Hunt Jr. et al. 1990); have only rarely been found in adult Least Auklet stomachts. Three may be some degree of differentiation between Least and Crested Auklets in their foranging patterns, specifically, their food composition. Both auklet species had similar  $\delta^{10}$ C, suggesting that both species forage in areas with similar oceanographic conditions. This contrast with previous dulles near Gareloi than how shown peographic sagregation between the two species, but at much finer spatial resolution (at the level of deldies and upwellings) rather than the more come resolution of 70° (butter, et al. 1995) Stoff et al. 1990.

# Conclusions

Combined with knowledge of moulting patterns, stable isotopes offer insight into the foraging patterns of small seabirds, including audies, where attached data loggers remain too large. We found significant age differences in foraging of Least Auklets, and present the first information on the foraging of subadults. There was no availation

among breeding colonies spanning over 250 km in the western Aleutian Islands, which agrees with inter-island similarities in chck diet (Chapter 3). We also found a significant trophic shift during the post-breeding season for both Least and Crested Auklets. Further research should focus on the wintering ecology of these seabids, and the temporal and spatial variability in acoglation stable-intologe ratios.

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Figures



Figure 6-1 – Adult Least Auklets had significantly higher  $\delta^{15}N$  during the pre-breeding season than subadults sampled in 2007-2010 from Kiska Island, Aleutian Islands, Alaska.













# CHAPTER 7: GENERAL DISCUSSION

Seabards can be ureful ecosystem sentinel species for investigating the directs of climate variability on populations over time, and over large spatial extent; (Harris & Manless 1990a; Frederiksen et al. 2007). Furthermore, by studying ecological fators and population responses at multiple sites, local effects and population-specific factors can often be identified (Weise et al. 2004; Lavers et al. 2009). In this series of studies, I sought to investigate the relationships between large-scale indices of climate in the marine emissionment, and the responses of plankthorous auklets at multiple sites throughout their tweeling range.

An initial hypothesis was that, because auklets are planktivores, changes in occanographic conditions would affect their pirmary prey, Netschanus coepeoids and Thypanoesian euphausids, which could influence reproductive success (Chapter 2). This study was one of the first to investigate the relationships between occanography and reproductive success in multiple species of auklet at multiple sites over multiple among species and sland, but productivity on all species on all islands was negatively correlated with the Akustain low Pressure Index, a measure of the intensity of the Aleutain low pressure system between December and March (Chapter 2). We hypothesized that this relationship was the result of bottom-up control by Imitigation Doplankton abudance dwing the following bereding season.

We then addressed this hypothesis by examining the composition of chick meals brought by adults to nest-bound chicks over the same period. We found that breeding-

seano accomprophic conditions influenced the proportion of main prey taxa in auxilists' diet, with the exception of Whinkered Auklets, whose diet composition was related to wither conditions. We need very title charge among years in the proportion of copepeds and explausidis in one of the first studies to relate seabind diet to cocanography (Durant et al. 2009). Based on this analysis, we concluded that auklets are able to buffer bottem-up foreign of demographic parameters, like productivity, because the composition of chick meals has remaind relatively constant over 14 years of environmental availability in the Bering Sea (Chapter 3). This highlights the realinee of auklets to withstand considerable environmental change without immediate negative response, at least to date (Folke et al. 2004). It also suggests that direct effects on adults during the winter are realisely than food limitation during the breeding season to influence amart exproductive success.

In addition to breeding, moult is an important, but expensive, life history stage, especially for smaller brids [Holmgren & Hendenström 1995]; Schleitz & Murphy 1997; Barta et al. 2008; Winglied 2008; Newton 2009). Unlike many other species, the Arthio addiets moult during the breeding seasos (Idead & Sashy 1984; Konyukhov 2001; 2000; Chapter 4). We used data collected on the progress of primary feather moult in Lesst Audets at four colonies from 52-64°W to test the hypothesis that brids at higher latitudes institute moult later, and moult faster than southern conspectives (like Mould & King 1976; Hennborg et al. 2001; Ryder & Rimmer 2003; Davison 2004). We found that task tudeis take that a hyber a Ritidae donuclitary, and start than those weak tudeis the set of the progress of a donuclitary, and start than those to start. Audets breeding a hyber a Ritidae donuclitary, and start than those to start. Audets breeding a hyber a Ritidae donuclitary and start than those to start. Audets breeding a hyber a Ritidae donuclitary and start than those

breeding at more southerly latitudes, even though the length of the breeding season is similar among locations (Chapter 4). The cold-water Anadyr Current sweeps south of Cape Ulyakhpan, our most northerly study site, bringing with it oceanic copepods that can fuel adults' mouth-breeding overlap (Pavlov & Pavlov 1996; Stabeno et al. 1999; Platt & Springer 2003; Geinrich 2005). Detailed descriptions of moulting patterns are required to interpret results of stable isotope analysis from feathers properly (Inger & Benchro 2008; Bond & Jones 2009b).

We then used our knowledge of moutine patterns in sublets to examine two aspects of their foraging patterns using stable isotope analysis of feathers: seasonal and species differences between Least and Crested Aublets sampled throughout the Aloutan tablets, and within-individual differences during the Least Aublets' including and their of Ususe synthesis, stable isotoper ratios in Least Aublets' memory primary (P1) will indicate the bird' foraging patterns during includation (Chapter 4). We found that both  $\Delta^{10}$  can  $\Delta^{10}$  th increased throughout the 20 days of feather growth, likely indicating a abilit in foraging location towards more productive exempraphic from stab table bills concentrations of Necealanus copepods (Harrison et al. 1990; Russell et al. 1999). This shift, accompanied by the relatively unchanging composition of meals brought to nest bound shick (Chapter 1), may indicate that adults balance the trade-off batween self maintenance and chick provisioning by shifting their own foraging location tome productive occanorapposite features in the second (Chapter 4).

Feathers moulted at different times of the annual cycle can provide information on birds' foraging away from the breeding colony. Auklets' outermost primary (P10) is moulted in the autumn following breeding (Chapter 3), and breast feathers are replaced during the spring pre-alternate moult (Pyle 2008). We found similar  $\delta^{13}$ C and  $\delta^{15}$ N values in each year and at each site within groups. Adult and subadult (one year-old) Least Auklets differed significantly in their foraging patterns during the pre-breeding period as did adults and chicks during chick rearing. Crested Auklets fed at a consistently higher trophic position than Least Auklets throughout the annual cycle (Chapter 6). These results differ from our observations of chick meal composition (Chanter 3) in that they indicate a lesser degree of overlap among species between adult Least and Crested Auklets. Similar to chick meals, however, there were no differences among islands or years. They expand on Chapter 3, however, in elucidating the trophic relationships among species and age classes, showing differences outside the breeding season, and among age classes whose diets are not sampled easily.

### Introduced Norway rats at Kiska Island

Since the 1940s, introduced Norway rats have been present at Kiska (Murie 1959). We found significant differences in reproductive success between Kiska and ratfree Buldir and Kasatochi Islands, driven primarily by reproductive failure in 2001 and 2002 (Chapter 2). In an initial assessment (Major et al. 2006), two hypotheses were put forward to explain the poor reproductive success: 1) poor food resources around Kiska, 555 and 2) predation and disturbance by rats. We found no significant differences in chick meal composition among years at (Kiska, or between Kiska and rat-free islands in the Aleutan Islands (Chapter 3). Furthermore, we found no significant differences in the pre-breeding dilutes a inferred by stable isotopes between Kiska and rat-free Buldr and Gareloi lislands (Chapter 6). These results strongly implicate introduced rats in the breeding failures of 2001 and 2002.

Least Auklet reproductive success was significantly lower on Riska than on Budit or Kasatochi, and a paucity of consistent correlations with basin-wide oceanographic or large-scale climatic indices in the North Pacific further suggests that colony-specific actions play an important role in influencing auklet demographics (Chupter 2). The East Auklet population on Riska is declining (HL Major et al., unpublished data), and the Crested Auklet population is also likely declining as well (Bond et al. unpublished). Longliked species, such as seabirds, hypically have high survival and low feerindity, and survival is often the most influential demographic parameter in determining population growth (Humr et al. 2002). In any given year, text and Crested Aukles at Kisla had aignificantly different survival than those on Budit ar Kasatochi (cometimes higher, other times lower, but never in phase) and was often less-related to large-scale indices of climate than their conspecifics on rat-free Islands (Jones et al., submitted). Together, these results suggest that introduced rats have had a negative effect on the auklet population at Kiska Island.

### Novel techniques and findings

We were the first to incorporate a spatial approach to examining the relationships between oceanographic and climatic variables and the demography of top marine predators. Spatially variable parameters, such as SST or SLP are used often in studies of climate-linked demography (Durant et al. 2009), but often over a restricted area. By examining landscape-wide variation in SST and SLP, and its relationship to reproductive success, we moved closer to a mechanistic approach that provided hypotheses for future studies.

In addition, we were also the first to demonstrate a latitudinal gradient in the intensity of primary moult in a marine bird. Such detailed study of moult, and investigation of the moult/breeding overlap in auklets, made detailed study of foraging using stable isotopes available. This included an analysis of auklet foraging during the pre- and post-breeding periods, and of subadults and chicks. Prior to this study, no information analysis for target systematic for these periods or age classes.

#### Future directions

Future auklet research should focus on three areas: winter ecology, metapopulation dynamics, and quantification of isotopic discrimination factors. The first two of these overlap to some degree, since the degree of mixture of breeding populations during the non-breeding period is unknown. The migratory patterns and wintering areas of aukkets are known mostly through needodial observations and ship-based surveys (Gizenka 1555; Veithanin 1977;; Vyakin 1981; Zubakin & Konyakhor 1994; Renner et al. 2008). Much of the new information presented here is interpreted in the light of limited data on the winter distribution, behaviour, and ecology of aukkets. Less and Whiskered Aukkets (85 and 118 g body mass respectively) are, at present, too small for attached delemetry equipment, such as geolocations or satellite transmitter; Crested Aukkets (260 g) are more suitable. Such telemetric studies should use a multi-colony approach. There are fundamental differences in the environmental pressures on aukkets beredling in the Aukutan Islands, the Problef Islands, and the northern Bering Sex, primary among these is the degree of winter sea ice and spring snow cover. Aukkets beredling in the Aukutans, and the problef Islands, and the northern Bering Sex, primary among these is the degree of winter sea ice and spring snow cover. Aukkets beredling in the Aukutans, and the present framer than those breedling in the Alkutans, and may therefore differ in their nutritional requirements, moult characteristics, and demography.

Somewhat related is the degree of inter-colony movement, especially in light of possible population declines throughout the Aleutian Islands. Reliable, repeatable survey methods for chebia auklets are cuarterity lacking, so dedecarable indefinition provide the only information on populator relation. In the Aleutian Islands, Least and Crested Auklets currently breed at site main sites (Buddin, Kiska, Segula, Semisopochnoi, Garelio, Chagulak), and formerly at Eastarchi, prior to its volcanic eruption in August 2008 (Smith et al. 2010; Williams et al. 2010). Of these, populations at the two colonies with any monitoring fuller, Kiska, Segula have be defining (Brann & Rendol 2002).

Bond, Jones, Najoe et al. unpublished data). Populations in the Berling Sea may also be declining (Roby & Brink 1986); Renner & Renner 2010], and the status of Russian populations is poorly homon (Gaston & Najoes 1998; Kondrayev et al. 2000). Introduced readation, (Rochardsev et al. 2000; Najoet et al. 2000; Junico aching vegetation (Roby & Brink 1986); Renner & Reynolds 2006], and volcanism (Williams et al. 2010; Bond et al. submitted) can threaten aukkt population, and population declines observed at some colonies may also include a cartain degree of Inter-Colony movement. Colonies of millions of auklets and irregular monitoring at most colonies make measuring immigration or emigration challenging. Indeed, as of 2011; there is only one Alaskan aukkt colony where demographic information will be collected (Bukidri, There is considerably less information on the distribution of nocturnal Whiskered Auklets, despite their listing as a species of Conservation Concern by the U.S. Fina and Wildlife served 2008].

Second, lostopic studies would be greatly improved by determining stable lostope discrimination factors for auklett. Discrimination factors, the change in 8<sup>-10</sup>C and 8<sup>-10</sup>R between prevail consumer, are tissue- species- and prev-specific [Caut et al. 2008, 2009). Cagdive populations are often required for the accurate determination discrimination factors, and few exist for Activa auklets. By determining auklet discrimination factors, quantifactive lostopic mining models (e.g., MicRif, SURF, Moore & Semmers 2008, Parnell et al. 2010) can be used to estimate the proportional contribution of gree you to auklet of discriminate the proportional to prostingbioting of gree to auklet of discrimination factors.

occanographic sampling during the auklets' pre- and post-breeding season to establish possible pery abundance, and composition, and data from telemetry studies, we can gain further insight into the ecology of these abundant plankthores during the 9 months when they are away from their breeding colonies, and relatively inaccessible to researchers.

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## APPENDICES

## Appendix 1. The number of breeding pairs of least (LEAU), crested (CRAU) and

whiskered auklets (!	WHAU)	studied at Bu	Idir, Kiska, and	Kasatochi during	1990-2008.
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	Buldir			Kiska		Kasatoc	hi
Year	LEAU	CRAU	WHAU	LEAU	CRAU	LEAU	CRAU
1990	61	67		•		•	
1991	81	74	46				
1992	89	79	55	-			
1993	44	49	54	-			
1994	64	67	57	-			
1995	64	66	67				
1996	57	66	57	-		54	43
1997	84	82	90	-		91	76
1998	76	70	78			95	104
1999						100	110
2000	69	78	70			89	110
2001	65	75	75	210	31	85	109
2002	50	81	100	205	23	94	109
2003	83	45	44	208	20	110	136
2004	81	67	66	197	31	91	107

	Buldir		Kiska	Kiska		Kasatochi	
2005	73	79	70		-	93	103
2006	84	73	68	180	34	77	88
2007	68	66	67	173	36	124	143
2008	67	75	68	157	29	98	109

Appendix 2. Reproductive success (number of chicks fledged/number of eggs laid) of least, crested and whiskered auklets at Buldir, Kiska, and Kasatochi from 1990-2008. Abbreviations and sample sizes are in Appendix 1.

Year	Buldir			Kiska		Kasatoc	hi
	LEAU	CRAU	WHAU	LEAU	CRAU	LEAU	CRAU
1990	0.38	0.49	•				
1991	0.62	0.58	0.50				
1992	0.54	0.72	0.60				
1993	0.50	0.78	0.57	-			
1994	0.41	0.69	0.74				
1995	0.70	0.77	0.67				
1996	0.60	0.61	0.70			0.69	0.74
1997	0.60	0.76	0.66			0.55	0.55
1998	0.45	0.76	0.53			0.49	0.66
1999						0.46	0.69
2000	0.48	0.62	0.46			0.66	0.75
2001	0.55	0.64	0.36	0.13	0.39	0.55	0.45
2002	0.60	0.60	0.48	0.10	0.43	0.53	0.59
2003	0.34	0.11	0.55	0.50	0.45	0.63	0.63
2004	0.53	0.45	0.53	0.52	0.68	0.53	0.70
2005	0.60	0.78	0.76			0.39	0.61

Year	Buldir			Kiska		Kasato	chi
2006	0.75	0.64	0.62	0.54	0.85	0.44	0.66
2007	0.66	0.85	0.85	0.58	0.58	0.61	0.76
2008	0.70	0.83	0.84	0.64	0.59	0.61	0.77

Appendix 3. Timini	z of breeding	(mean egg	hatch date) of	least, crested	and whiskered
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auklets at Buldir, Kiska, and Kasatochi from 1990-2008. Abbreviations are in Appendix 1.

Year	Buldir			Kiska		Kasatochi	
	LEAU	CRAU	WHAU	LEAU	CRAU	LEAU	CRAU
1990	27 June	25 June	24 June			-	-
1991	30 June	29 June	27 June	-	-	-	-
1992	29 June	26 June	18 June	-	-	-	-
1993	26 June	27 June	22 June	-	-	-	-
1994	24 June	25 June	19 June	-	-	-	-
1995	29 June	29 June	25 June	-	-	-	-
1996	25 June	26 June	22 June	-	-	25 June	30 June
1997	27 June	28 June	24 June	-	-	28 June	01 July
1998	30 June	05 July	23 June	-	-	03 July	03 July
1999	-	-	-	-		30 June	29 June
2000	25 June	29 June	17 June	-	-	28 June	29 June
2001	26 June	29 June	22 June	28 June	-	28 June	30 June
2002	25 June	25 June	24 June	5 July	2 July	27 June	27 June
2003	27 June	02 July	25 June	2 July	-	25 June	29 June
2004	28 June	05 July	21 June	27 June	4 July	30 June	28 June
2005	25 June	25 June	18 June	-	-	29 June	29 June
2006	30 June	ylut e0	27 June	1 July	7 July	03 July	03 July

 2007
 27 June
 23 June
 17 June
 28 June
 28 June
 30 June
 01 July

 2008
 24 June
 26 June
 21 June
 28 June
 29 June
 30 June
 01 July

Prey taxa	Estimated mass (g)
Cnidaria	
Scyphozoa	
Jellyfish	N/A
Nemata	
Nematode	N/A
Arthropoda	
Insecta	
Unidentified insect	0.00100
Ostracoda	
Unidentified ostracod	0.10000
Maxillopoda	
Euchaetidae	
Paraeuchaeta	
Paraeuchaeta birostrata	0.00200
Paraeuchaeta elongata	0.00440
Calanidae	
Calanus	
Calanus marshallae	0.00126
	0.00010

Appendix 4. Taxonomic list of prey taxa and values used to calculate biomass.

Prey taxa	Estimated mass (g
Neocalanus	
Neocalanus cristatus	0.01390
Neocalanus plumchrus/flemingeri	0.00276
Neocolanus spp.	0.00500
Malacostraca	
Amphipoda	
Unidentified amphipod	0.00220
Ischyroceridae	
Erichthonius	
Erichthonius spp.	0.00220
lschyrocerus	
Ischyrocerus spp.	0.00220
Eusiridae	
Pontogeneia	
Pontogeneia spp.	0.05000
Gammaridae	
Unidentified gammarid amphipod	
Calliopiidae	
Halirages	
Halirages bungei	0.05000
Prey taxa	Estimated mass (g
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Lysianassidae	
Unidentified lysianassid amphipod	0.00400
Stenotholdae	
Unidentified stenothold amphipod	0.10000
Talitridae	
Unidentified talitrid amphipod	0.00220
Hyperiidae	
Hyperia	
Hyperia spp.	0.00200
Hyperoche	
Hyperoche medusarum	0.00385
Primno	
Primno macropa	0.00300
Themisto	
Themisto pacifica (<4mm)	0.00368
Themisto spp. (<4mm)	0.00385
Tanaidacea	
Unidentified Tanaid crustacean	0.05000
Euphausiacea	
Euphausiidae	

Prey taxa	Estimated mass (g
Euphausiid furcila	0.00600
Euphausiid spp. (<7mm)	0.00600
Euphausiid spp. (lg, > 7mm)	0.08400
Thysanoessa	
Thysanoessa inermis (<7mm)	0.02000
Thysanoessa inermis (>12mm)	0.07500
Thysanoessa raschii (<7mm)	0.03050
Thysanoessa raschii (>12mm)	0.09780
Thysanoessa spp. (>12mm)	0.07895
Decapoda	
Unidentified shrimp	0.05000
Unidentified shrimp larvae (<7mm)	0.01200
Shrimp zoae	0.01000
Crab zoae	0.01000
Hippolytidae	
Hippolytidae juvenile	0.03700
Pandalidae	
Pandalid shrimp (>12mm)	0.04865
Pandalid shrimp larvae	0.01200
Paguridae	

Prey taxa	Estimated mass (g)
Paguridae glaucothoe	0.00500
Paguridae megalopa	0.01500
Paguridae zoea	0.00162
Atelecyclidae	
Atelecyclidae megalopa	0.01500
Mollusca	
Gastropoda	
Unidentified gastropod	0.00500
Thecosomata	
Unidentified pteropod	0.00100
Limacinidae	
Limacina	
Limacina helicinia	0.00200
Limacina spp.	0.00350
Cephalopoda	
Teuthida	
Unidentified squid	0.06000
Gonatidae	
Unidentified armhook squid	0.06000
Chordata	

Prey taxa	Estimated mass (g)
Actinopterygii	
Unidentified fish	N/A
Unidentified fish larvae	0.48500
Scorpaeniformes	
Hexagrammidae	
Hexagrammos	
Hexagrammos spp. (1+ yr)	11.00000

Appendix 5. Summary of food load composition of Crested Auklet chick meals.

Table A5-1. Summary of food load composition of Crested Auklets at Buldir Island, 1994-2006. Data are presented as annual values of per cent biomass.

Year	1994	1995	1996	1997	1998	1999
n	37	46	78	82	103	88
Copepoda	37.13%	26.83%	75.28%	24.75%	28.32%	83.81%
Calanoid spp.	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Calonus marshallae	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Neocolanus Total	37.13%	26.83%	75.28%	24.75%	28.32%	83.81%
Neocalanus spp.	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Neocalanus cristatus	37.10%	26.79%	73.45%	21.71%	28.01%	83.13%
Neocalanus plumchrus/flemingeri	0.03%	0.04%	1.83%	3.04%	0.31%	0.68%
Amphipoda	0.93%	0.80%	0.58%	0.79%	0.14%	1.57%
Hyperoche medusarum	0.00%	0.00%	0.01%	0.00%	0.00%	0.00%
Primno macropa	0.04%	0.00%	0.00%	0.00%	0.00%	0.01%
Themisto pacifica (<4mm)	0.88%	0.80%	0.57%	0.79%	0.14%	1.56%
Themisto spp. (<4mm)	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Euphauslacea	61.79%	71.73%	23.75%	74.38%	71.52%	13.81%
Thysanoessa inermis (<7mm)	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Thysonoessa inermis (>12mm)	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Euphausiid furcilia	0.00%	0.00%	0.00%	0.00%	0.00%	0.01%

1999
% 13.80%
0.00%
0.81%
0.74%
0.00%
0.00%
0.00%
0.01%
0.00%
0.05%
0.00%

# Table A5-1 continued.

Year	2000	2001	2002	2003	2004	2005	2005
n	29	45	54	23	33	39	36
Copepoda	82.74%	49.33%	71.94%	4.37%	86.72%	81.76%	76.92%
Calanoid spp.	0.00%	0.53%	0.00%	0.00%	0.00%	0.00%	0.00%
Colanus morshollae	0.00%	0.00%	0.00%	0.00%	0.00%	0.07%	0.00%
Neocalanus Total	82.74%	48.80%	71.94%	4.37%	86.72%	81.69%	76.92%
Neocolanus spp.	2.58%	0.00%	0.00%	3.67%	0.00%	0.00%	0.00%
Neocolanus cristatus	79.97%	48.74%	70.21%	0.70%	86.66%	81.13%	74.58%
Neocalanus							
plamchrus/flemingeri	0.19%	0.05%	1.73%	0.00%	0.05%	0.56%	2.34%
Amphipoda	0.00%	3.03%	5.32%	12.87%	0.17%	1.25%	0.05%
Hyperoche medusarum	0.00%	0.00%	0.01%	0.00%	0.00%	0.00%	0.00%
Primno macropa	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Themisto pacifica (<4mm)	0.00%	2.83%	4.89%	0.00%	0.17%	0.01%	0.03%
Themisto spp. (<4mm)	0.00%	0.20%	0.42%	12.87%	0.00%	1.24%	0.02%
Euphausiacea	16.56%	42.58%	20.28%	82.43%	13.00%	16.30%	23.03%
Thysanoessa inermis (<7mm)	3.47%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Thysanoessa inermis (>12mm)	8.95%	0.00%	0.00%	0.00%	6.64%	0.00%	0.00%
Euphausiid furcilia	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Euphausiid spp. (<7mm)	4.14%	6.13%	8.94%	3.76%	2.41%	3.29%	23.03%

Year	2000	2001	2002	2003	2004	2005	2006
Thysanoessa spp. (>12mm)	0.00%	36.45%	11.34%	78.67%	3.95%	13.00%	0.00%
Decapoda	0.69%	5.06%	2.46%	0.33%	0.11%	0.69%	0.00%
Hippolytidae juvenile	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Pandalid shrimp (>12mm)	0.00%	0.11%	2.44%	0.00%	0.11%	0.69%	0.00%
Pandalid shrimp larvae	0.69%	4.96%	0.00%	0.33%	0.00%	0.00%	0.00%
Shrimp zoea	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Paguridae megalopa	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Atelecyclidae megalopa	0.00%	0.00%	0.02%	0.00%	0.00%	0.00%	0.00%
Crab zoea	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Unid. squid	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%

Table A5-2. Summary of food load composition of Crested Auklets at Kiska Island, 2001-

Year	2001	2003	2006
n	7	3	24
Copepoda	0.00%	0.00%	87.39%
Neocolanus Total	0.00%	0.00%	87.39%
Neocalanus cristatus	0.00%	0.00%	87.21%
Neocalanus plumchrus/flemingeri	0.00%	0.00%	0.18%
Amphipoda	0.00%	0.00%	0.06%
Hyperache medusarum	0.00%	0.00%	0.04%
Themisto pacifica (c4mm)	0.00%	0.00%	0.02%
Euphausiacea	100.00%	100.00%	12.10%
Euphausiid spp. (<7mm)	99.63%	0.00%	0.00%
Euphausiid spp. (lg, > 7mm)	0.37%	100.00%	12.10%
Decapoda	0.00%	0.00%	0.38%
Larval shrimp (<7mm)	0.00%	0.00%	0.38%
Unid. squid	0.00%	0.00%	0.07%

2006. Data are presented as annual values of per cent biomass.

Table A5-3. Summary of food load composition of Crested Auklets at Kasatochi Island,

Year	1996	1997	1998	1999	2000
n	36	39	35	36	34
Ostracoda	0.00%	0.19%	0.00%	0.00%	0.00%
Copepoda	36.58%	90.35%	73.02%	67.04%	39.76%
Unid. copepod	0.04%	0.00%	0.00%	0.00%	0.00%
Calanus marshallae	0.00%	0.00%	0.00%	0.00%	0.03%
Neocologue Total	36 54%	90.35%	73.02%	67.04%	39.73%
Neocolanus cristatus	33.49%	73.14%	67.57%	62.21%	37.72%
Neocolanus plumchrus/flemingeri	3.05%	17.21%	5.44%	4.83%	2.02%
Amphipoda	0.14%	0.33%	2.83%	1.27%	0.09%
Gammaridae	0.00%	0.05%	0.00%	0.00%	0.00%
Themisto pacifica (<4mm)	0.14%	0.27%	2.10%	1.27%	0.09%
Pterepod spp.	0.00%	0.01%	0.00%	0.00%	0.00%
Shrimp zoea	0.00%	0.00%	0.74%	0.00%	0.00%
Themisto spp. (<4mm)	0.00%	0.00%	0.00%	0.00%	0.00%
Euphausiacea	63.04%	6.75%	23.37%	30.63%	60.07%
Euphausiid furcilia	0.00%	0.00%	0.00%	0.05%	0.00%

1996-2006. Data are presented as annual values of per cent biomass.

Year	1996	1997	1998	1999	2000
Euphausiid spp. (<7mm)	0.00%	6.75%	0.00%	0.00%	16.89%
Thysanoessa spp. (>12mm)	63.04%	0.00%	23.37%	30.58%	43.18%
Decapoda	0.24%	2.26%	0.18%	1.07%	0.08%
Hippolytidae juvenile	0.00%	0.00%	0.00%	0.74%	0.00%
Larval shrimp (<7mm)	0.17%	1.86%	0.00%	0.00%	0.08%
Pandalid shrimp (>12mm)	0.00%	0.00%	0.00%	0.00%	0.00%
Primno macropa	0.00%	0.03%	0.00%	0.00%	0.00%
Paguridae glaucothoe	0.00%	0.00%	0.00%	0.09%	0.00%
Paguridae megalopa	0.00%	0.00%	0.05%	0.00%	0.00%
Atelecyclidae megalopa	0.04%	0.29%	0.11%	0.00%	0.00%
Crab zoea	0.03%	0.09%	0.02%	0.24%	0.00%
Unid. squid	0.00%	0.11%	0.06%	0.00%	0.00%
Unid. Fish	0.00%	0.00%	0.53%	0.00%	0.00%

# Table A5-3 continued.

Year	2001	2002	2003	2004	2005	2006
n	31	39	30	34	35	34
Ostracoda	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Copepoda	90.63%	65.07%	65.11%	74.01%	88.48%	82.58%
Unid. copepod	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Colanus marshallae	0.00%	0.21%	0.00%	0.00%	0.00%	0.00%
Neocolanus Total	90.63%	65.86%	66.11%	74.01%	88.48%	82.58%
Neocolanus cristatus	90.29%	62.39%	65.81%	74.01%	86.55%	81.65%
Neocolanus plumchrus/flemingeri	0.34%	3.47%	0.29%	0.00%	1.93%	0.93%
Amphipoda	0.10%	1.02%	0.03%	20.50%	0.18%	0.13%
Gammaridae	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Themisto pacifica (<4mm)	0.02%	0.19%	0.00%	0.00%	0.00%	0.05%
Pterepod spp.	0.00%	0.00%	0.00%	0.00%	0.00%	0.03%
Shrimp zoea	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Themisto spp. (<4mm)	0.08%	0.83%	0.03%	20.50%	0.18%	0.05%
Euphausiacea	8.34%	32.22%	32.03%	5.49%	10.97%	16.46%
Euphausiid furcilia	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Euphausiid spp. (<7mm)	6.98%	27.30%	11.81%	5.49%	8.07%	11.26%
Thysanoessa spp. (>12mm)	1.36%	4.92%	20.22%	0.00%	2.90%	5.20%

Year	2001	2002	2003	2004	2005	2006
Decapoda	0.92%	0.70%	1.82%	0.00%	0.37%	0.75%
Hippolytidae juvenile	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Larval shrimp (<7mm)	0.65%	0.14%	0.68%	0.00%	0.30%	0.61%
Pandalid shrimp (>12mm)	0.26%	0.56%	1.07%	0.00%	0.00%	0.00%
Primno macropa	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Paguridae glaucothoe	0.00%	0.00%	0.00%	0.00%	0.01%	0.00%
Paguridae megalopa	0.00%	0.00%	0.05%	0.00%	0.00%	0.00%
Atelecyclidae megalopa	0.01%	0.00%	0.03%	0.00%	0.07%	0.14%
Crab zoea	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Unid. squid	0.00%	0.00%	0.00%	0.00%	0.00%	0.07%
Unid. Fish	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%

Appendix 6. Summary of food load composition of Least Auklet chick meals.

Table A6-1. Summary of food load composition of Least Auklets at Buldir Island, 1994-

Year	1994	1995	1996	1997	1998	1999
n	4	8	15	31	25	32
Copepoda	84.54%	94.58%	96.05%	93.87%	94.37%	95.59%
Paraeuchaeta birostrata	0.00%	0.00%	0.00%	0.00%	0.00%	0.01%
Calanus marshallae	0.00%	0.00%	0.00%	0.00%	0.04%	0.00%
Neocolonus Total	84.54%	94.58%	96.06%	93.87%	94.33%	95.58%
Neocolanus spp.	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Neocolanus cristatus	13.95%	1.85%	10.02%	15.56%	25.77%	42.44%
Neocalanus plumchrus/flemingeri	70.59%	92.73%	86.04%	78.31%	68.55%	53.15%
Amphipoda	1.48%	1.17%	0.81%	0.88%	0.47%	0.36%
Erichithanius spp.	0.00%	0.00%	0.00%	0.00%	0.40%	0.00%
Hyperoche medusorum	0.00%	0.74%	0.07%	0.00%	0.00%	0.00%
Primno macropa	0.39%	0.00%	0.00%	0.37%	0.00%	0.14%
Themisto pacifica (<4mm)	1.09%	0.43%	0.75%	0.51%	0.07%	0.22%
Euphausiacea	0.00%	0.18%	2.36%	1.46%	3.96%	0.61%
Thysanoessa raschii (>12mm)	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Thysanoessa spp. (>12mm)	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Euphausiid furcilla	0.00%	0.00%	0.00%	0.00%	2.54%	0.46%

2006. Data are presented as annual values of per cent biomass.

Year	1994	1995	1996	1997	1998	1999
Euphausiid spp. (sm, <7mm)	0.00%	0.18%	2.36%	1.46%	1.42%	0.15%
Decapoda	4.93%	2.65%	0.62%	2.35%	0.66%	3.16%
Pandalid shrimp (>12mm)	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Pandalid shrimp larvae	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Larval shrimp (<7mm)	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Hippolytidae juvenile	0.00%	0.00%	0.00%	0.00%	0.00%	2.66%
Shrimp zoea	4.93%	2.65%	0.45%	1.18%	0.56%	0.35%
Paguridae megalopa	0.00%	0.00%	0.00%	0.00%	0.00%	0.10%
Crab zoea	0.00%	0.00%	0.00%	0.20%	0.00%	0.01%
Atelecyclidae/crab megalopa	0.00%	0.00%	0.17%	0.96%	0.10%	0.04%
Gastropoda	1.07%	1.42%	0.15%	1.44%	0.55%	0.10%
Limacina spp.	0.00%	0.00%	0.00%	0.00%	0.55%	0.00%
Unid. snail	1.07%	0.00%	0.00%	0.00%	0.00%	0.00%
Pterepod spp.	0.00%	1.42%	0.15%	1.44%	0.00%	0.10%
Other	7.98%	0.00%	0.00%	0.00%	0.00%	0.16%
Gonatidae	0.00%	0.00%	0.00%	0.00%	0.00%	0.16%
Unid. fish larvae	7.98%	0.00%	0.00%	0.00%	0.00%	0.00%

# Table A6-1 continued.

Year	2000	2001	2002	2003	2004	2005	2006
n	12	32	31	21	30	24	29
Copepoda	98.20%	96.82%	97.45%	99.06%	92.52%	96.43%	75.83%
Paraeuchaeta birostrata	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Calanus marshallae	0.35%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%

Neocalanus Total	97.85%	96.82%	97.45%	99.06%	92.52%	96.43%	75.83%
Neocolanus spp.	3.16%	0.00%	0.00%	84.75%	0.00%	0.00%	0.00%
Neocalanus cristatus	80.95%	20.27%	1.70%	14.31%	3.06%	71.51%	17.87%
Neocolonus plumchrus/flemingeri	13.74%	76.55%	95.75%	0.00%	89.47%	24.91%	57.96%

Amphipoda	0.02%	2.68%	1.42%	0.42%	0.00%	0.01%	0.06%	
Erichithanius spp.	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	
Hyperache medusarum	0.00%	0.00%	0.02%	0.00%	0.00%	0.00%	0.00%	
Primno macropa	0.00%	0.00%	0.02%	0.01%	0.00%	0.00%	0.00%	
Themisto pacifica (<4mm)	0.02%	2.68%	1.38%	0.41%	0.00%	0.01%	0.06%	

Euphausiacea	1.66%	0.37%	0.76%	0.47%	5.10%	1.13%	23.00%
Thysanoessa raschii (>12mm)	0.00%	0.00%	0.00%	0.00%	0.00%	1.05%	0.00%
Thysanoessa spp. (>12mm)	0.00%	0.00%	0.00%	0.14%	0.00%	0.00%	0.65%
Euphausiid furcilla	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Euphausiid spp. (sm, <7mm)	1.66%	0.37%	0.76%	0.33%	5.10%	0.09%	22.35%
-	0.08%	0.128	0.38%	0.05%	0.00%	2.210	1.110

Year	2000	2001	2002	2003	2004	2005	2006
Pandalid shrimp (>12mm)	0.00%	0.00%	0.00%	0.00%	0.00%	0.35%	0.00%
Pandalid shrimp larvae	0.00%	0.13%	0.38%	0.00%	0.00%	0.00%	0.00%
Larval shrimp (<7mm)	0.00%	0.00%	0.00%	0.00%	0.00%	1.91%	0.00%
Hippolytidae juvenile	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Shrimp zoea	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Paguridae megalopa	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Crab zoea	0.00%	0.00%	0.00%	0.02%	0.00%	0.00%	0.00%
Atelecyclidae/crab megalopa	0.08%	0.00%	0.00%	0.03%	0.00%	0.05%	1.11%
Gastropoda	0.04%	0.00%	0.00%	0.00%	0.00%	0.13%	0.00%
Limacina spp.	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Unid. snail	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Pterepod spp.	0.04%	0.00%	0.00%	0.00%	0.00%	0.13%	0.00%
Other	0.00%	0.00%	0.00%	0.00%	2.37%	0.00%	0.00%
Gonatidae	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Unid. fish larvae	0.00%	0.00%	0.00%	0.00%	2.37%	0.00%	0.00%

Table A6-2. Summary of food load composition of Least Auklets at Kiska Island, 2001-

Year	2001	2002	2003	2004	2006
•	59	17	31	23	30
Insect	0.00%	0.01%	0.00%	0.00%	0.00%
Copepoda	97.30%	97.86%	95.91%	80.65%	98.29%
Calanus marshallae	0.03%	0.00%	0.00%	0.00%	0.00%
Neocalanus Total	97.27%	97,86%	95.91%	80.65%	98.29%
Neocalanus spp.	0.00%	95.98%	0.00%	0.00%	0.00%
Neocalanus cristatus	8.09%	1.88%	0.00%	3.47%	3.19%
Neocalonus plumchrus/flemingeri	89.18%	0.00%	95.91%	77.18%	95.10%
Amphipoda	0.26%	0.05%	0.15%	0.32%	0.02%
Hyperoche medusarum	0.00%	0.00%	0.00%	0.13%	0.01%
Primno macropa	0.01%	0.01%	0.00%	0.00%	0.01%
Themista nacifica (c4mm)	0.59%	0.00%	0.09%	1.035	0.02%
Theories are defend		0.315	0.516	0.106	0.05%
inemisto spp. («+mm)	0.43%	0.21%	0.31.6	0.10/4	0.000
Euphausiacea	1.06%	0.05%	2.02%	9.06%	0.67%
Euphausiid spp. (lg, > 7mm)	0.87%	0.00%	1.80%	0.00%	0.00%
Euphausiid spp. (sm, <7mm)	0.19%	0.05%	0.21%	9.06%	0.67%

2006. Data are presented as annual values of per cent biomass.

Year	2001	2002	2003	2004	2006
Decapoda	0.14%	1.67%	1.36%	0.07%	0.92%
Larval shrimp (<7mm)	0.12%	0.00%	0.00%	0.07%	0.85%
Pandalid shrimp (>12mm)	0.00%	1.65%	0.00%	0.00%	0.00%
Paguridae zoea	0.01%	0.00%	0.00%	0.00%	0.00%
Atelecyclidae/crab megalopa	0.00%	0.00%	0.00%	0.00%	0.04%
Crab zoea	0.00%	0.03%	1.36%	0.00%	0.03%
Gastropoda	0.45%	0.19%	0.11%	0.51%	0.03%
Limacina helicinia	0.00%	0.19%	0.00%	0.00%	0.00%
Pterepod spp.	0.45%	0.00%	0.11%	0.51%	0.03%
Unid. fish larvae	0.00%	0.00%	0.00%	8.45%	0.00%

Table A6-3. Summary of food load composition of Least Auklets at Kasatochi Island,

Year	1996	1997	1998	1999	2000
	19	51	36	33	33
Unid. crustacean	0.10%	0.00%	0.00%	0.00%	0.00%
Copepoda	62.53%	57.09%	84.00%	56.46%	81.85%
Calanus marshallae	0.00%	0.00%	0.00%	0.00%	0.00%
Neocalanus Total	62.53%	57.09%	84.00%	56.46%	81.85%
Neocolanus cristatus	4.23%	5.42%	11.96%	18.92%	9.33%
Neocolanus plumchrus/flemingeri	58.31%	51.67%	72.04%	37.54%	72.52%
Amphipoda	0.51%	10.91%	1.22%	1.74%	4.74%
Erichithonius spp.	0.00%	0.00%	0.48%	0.00%	0.00%
Eusiridae	0.00%	0.00%	0.00%	0.00%	2.92%
Gammaridae	0.00%	10.11%	0.00%	0.00%	0.00%
Halirages bungel	0.00%	0.00%	0.00%	1.42%	0.00%
Ischyrocerus spp.	0.00%	0.00%	0.00%	0.00%	0.26%
Pontogeneia spp.	0.00%	0.00%	0.00%	0.07%	0.00%
Stenothoidae	0.00%	0.36%	0.00%	0.00%	0.00%
Talitridae	0.02%	0.00%	0.00%	0.00%	0.00%
Hyperoche medusarum	0.46%	0.00%	0.00%	0.00%	0.00%

1996-2006. Data are presented as annual values of per cent biomass.

Year	1996	1997	1998	1999	2000
Primno macropa	0.00%	0.30%	0.00%	0.03%	0.00%
Themisto pacifica (<4mm)	0.04%	0.13%	0.75%	0.20%	1.56%
Themisto spp. (<4mm)	0.00%	0.00%	0.00%	0.00%	0.00%
Tanaidacea	0.00%	0.00%	0.00%	0.00%	0.71%
Euphausiacea	31.30%	11.07%	5.53%	1.70%	4.09%
Euphausiid furcilla	0.00%	0.00%	5.10%	0.64%	0.00%
Euphausiid spp. (sm, <7mm)	0.00%	0.00%	0.00%	0.00%	3.95%
Thysanoessa spp. (>12mm)	31.30%	11.07%	0.43%	1.06%	0.14%
Decapoda	3.68%	18.89%	5.56%	38.51%	5.10%
Hippolytidae juvenile	0.00%	0.00%	0.00%	38.22%	0.00%
Larval shrimp (<7mm)	2.38%	17.03%	0.00%	0.00%	4.76%
Pandalid shrimp (>12mm)	0.00%	0.00%	0.00%	0.00%	0.00%
Shrimp zoea	0.00%	0.00%	5.10%	0.00%	0.00%
Paguridae glaucothoe	0.00%	0.00%	0.15%	0.16%	0.02%
Paguridae zoea	0.00%	0.00%	0.00%	0.00%	0.00%
Atelecyclidae/crab megalopa	1.13%	1.42%	0.16%	0.09%	0.32%
Crab zoea	0.17%	0.44%	0.15%	0.04%	0.00%
Gastropoda	1.88%	2.04%	3.68%	1.59%	3.51%
Limacina spp.	0.00%	0.00%	3.68%	1.59%	3.51%
Pterepod spp.	1.88%	2.04%	0.00%	0.00%	0.00%

# Table A6-3 continued

Year	2001	2002	2003	2004	2005	2006
n	33	37	24	33	29	28
Unid. crustacean	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Copepoda	90.12%	92.55%	84.39%	88.53%	88.63%	77.02%
Colanus marshallae	0.00%	0.34%	0.09%	0.00%	0.00%	0.00%
Neocolonus Total	90.12%	92.21%	84.30%	88.53%	88.63%	77.02%
Neocolonus cristotus	0.27%	2.58%	1.86%	1.55%	42.96%	22.88%
Neocolanus plumchrus/flemingeri	89.84%	89.63%	82.44%	86.97%	45.67%	54.14%
Amphipoda	0.00%	1.13%	0.85%	0.23%	0.48%	0.00%
Erichithonius spp.	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Eusiridae	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Gammaridae	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Hailrages bungel	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
ischyrocerus spp.	0.00%	0.44%	0.19%	0.16%	0.06%	0.00%
Pontogeneia spp.	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Stenothoidae	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Talitridae	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Hyperoche medusarum	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Primno mocropo	0.00%	0.00%	0.01%	0.01%	0.00%	0.00%
Themisto pacifica (<4mm)	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Themisto spp. (<4mm)	0.00%	0.70%	0.65%	0.06%	0.42%	0.00%

Year	2001	2002	2003	2004	2005	2006
Tanaidacea	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
	4.775		10 3/1		1.269	
Luphausiacea	4.73%	3.12%	10.36%	3.04%	1.20%	0.02.1
Euphausiid furcilla	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Euphausiid spp. (sm, <7mm)	4.73%	2.43%	10.36%	3.84%	1.26%	8.82%
Thysanoessa spp. (>12mm)	0.00%	0.68%	0.00%	0.00%	0.00%	0.00%
Decapoda	4.49%	2.86%	4.41%	5.10%	8.74%	13.09%
Hippolytidae juvenile	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Larval shrimp (<7mm)	2.44%	2.31%	2.02%	4.60%	8.70%	12.95%
Pandalid shrimp (>12mm)	1.92%	0.42%	1.93%	0.00%	0.00%	0.00%
Shrimp zoea	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Paguridae glaucothoe	0.00%	0.00%	0.30%	0.00%	0.00%	0.00%
Paguridae zoea	0.00%	0.01%	0.02%	0.00%	0.00%	0.01%
Atelecyclidae/crab megalopa	0.12%	0.13%	0.15%	0.50%	0.03%	0.14%
Crab zoea	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Gastropoda	0.67%	0.33%	0.00%	2.30%	0.89%	1.07%
Limacina spp.	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Pterepod spp.	0.67%	0.33%	0.00%	2.30%	0.89%	1.07%

Appendix 7. Summary of food load composition of Whiskered Auklet chick meals.

Table A7-1. Summary of food load composition of Whiskered Auklets at Buldir Island,

1993-2006. Data ar	e presented as annu	al values of per cent biomass.
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Year	1993	1994	1995	1996	1997	1998	1999
n	23	16	48	71	36	26	38
Unid. crustacean	7.40%	0.00%	0.26%	0.00%	0.00%	0.00%	0.00%
Copepoda	19.14%	43.88%	76.67%	93.29%	93.39%	96.95%	93.54%
Calanoid spp.	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Paraeuchaeta birostrata	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.18%
Paraeuchaeta elongata	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Pseudhalaptilus pacificus	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.25%
Calonus marshallae	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Calanus pacificus	0.00%	0.00%	0.06%	0.00%	0.00%	0.00%	0.00%
Neocolonus Total	19.14%	43.88%	76.61%	93.29%	93.39%	96.95%	93.11%
Neocolanus spp.	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Neocolanus cristatus	11.48%	36.80%	36.09%	36.19%	15.50%	75.57%	92.49%
Neocolanus plumchrus/flemingeri	7.66%	7.08%	40.53%	57.10%	77.88%	21.38%	0.61%
Amphipoda	0.09%	4.37%	1.28%	0.26%	0.25%	0.09%	0.33%
Hyalidae	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Lysianassidae	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Hyperoche medusarum	0.00%	0.00%	1.11%	0.26%	0.00%	0.00%	0.00%

Year	1993	1994	1995	1996	1997	1998	1999
Primno macropa	0.01%	3.82%	0.00%	0.00%	0.24%	0.09%	0.33%
Themista pacifica (<4mm)	0.00%	0.55%	0.17%	0.00%	0.01%	0.00%	0.00%
Themisto spp. (<4mm)	0.08%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Euphausiacea	60.22%	48.80%	20.07%	5.99%	0.23%	1.81%	1.37%
Euphausiid furcilia	0.00%	0.00%	0.00%	0.00%	0.00%	0.35%	0.00%
Euphausiid spp. (lg, > 7mm)	0.00%	0.00%	20.07%	5.99%	0.23%	1.46%	1.37%
Thysanoessa spp. (>12mm)	60.22%	48.80%	0.00%	0.00%	0.00%	0.00%	0.00%
Decapoda	0.00%	2.79%	1.50%	0.34%	5.53%	0.59%	2.89%
Hippolytidae juvenile	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.98%
Larval shrimp (<7mm)	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Pandalid shrimp (>12mm)	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Pandalid shrimp larvae	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Shrimp zoea	0.00%	2.79%	1.50%	0.14%	4.43%	0.59%	1.92%
Atelecyclidae megalopa	0.00%	0.00%	0.00%	0.16%	0.71%	0.00%	0.00%
Crab zoea	0.00%	0.00%	0.00%	0.04%	0.39%	0.00%	0.00%
Gastropoda	0.02%	0.00%	0.21%	0.11%	0.61%	0.55%	1.87%
Limacina helicinia	0.02%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Limacina spp.	0.00%	0.00%	0.00%	0.00%	0.00%	0.55%	0.00%
Pterepod spp.	0.00%	0.00%	0.21%	0.11%	0.61%	0.00%	1.87%
Unid. snail	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%

Year	1993	1994	1995	1996	1997	1998	1999
Unid. squid	0.00%	0.16%	0.00%	0.00%	0.00%	0.00%	0.00%
Haranness on Paul	12 126	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Nexogrammas spp. (1+ yr)	13.13%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%

# Table A7-1 continued.

Year	2000	2001	2002	2003	2004	2005	2006
n	33	28	32	31	37	31	36
Unid. crustacean	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Copepoda	96.70%	99.68%	85.39%	39.55%	98.32%	96.19%	91.42%
Calanoid spp.	0.00%	2.82%	0.00%	0.00%	0.00%	0.00%	0.00%
Paraeuchaeta birostrata	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Parasuchasta elangata	0.00%	0.13%	0.12%	0.00%	0.00%	0.00%	0.00%
Pseudhaloptilus pacificus	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Colanus marshallae	0.46%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Calanus pacificus	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Neocaionus Total	96.24%	96.73%	85.27%	39.55N	98.32%	96.19%	91.42%
Neocolonus spp.	5.08%	49.31%	0.00%	0.00%	0.00%	0.00%	0.00%
Neocolanus cristatus	70.04%	47.42%	71.96%	8.36%	19.01%	17.71%	58.35%
Neocolanus plumchrus/fiemingeri	21.12%	0.00%	13.32%	31.19%	79.32%	78.49%	33.07%
Amphipoda	0.00%	0.00%	0.00%	0.91%	0.00%	0.01%	0.00%
Hyalidae	0.00%	0.00%	0.00%	0.90%	0.00%	0.00%	0.00%
Lysianassidae	0.00%	0.00%	0.00%	0.01%	0.00%	0.00%	0.00%
Hyperoche medusarum	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Primno mocropo	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Themisto pacifica (<4mm)	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Themisto son (cdmm)	0.00%	0.00%	0.00%	0.00%	0.00%	0.01%	0.00%

Year	2000	2001	2002	2003	2004	2005	2005
Euphausiacea	1.57%	0.30%	13.52%	52.03%	1.61%	3.14%	8.45%
Euphausiid furcilia	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Euphausiid spp. (lg, > 7mm)	1.57%	0.30%	11.73%	4.23%	0.51%	3.03%	8.45%
Thysanoessa spp. (>12mm)	0.00%	0.00%	1.79%	47.80%	1.09%	0.11%	0.00%
Decapoda	1.70%	0.02%	0.69%	7.52%	0.07%	0.66%	0.12%
Hippolytidae juvenile	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Larval shrimp (<7mm)	0.00%	0.00%	0.00%	0.00%	0.00%	0.59%	0.12%
Pandalid shrimp (>12mm)	0.00%	0.00%	0.58%	7.49%	0.00%	0.00%	0.00%
Pandalid shrimp larvae	0.00%	0.02%	0.10%	0.03%	0.00%	0.00%	0.00%
Shrimp zoea	1.70%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Atelecyclidae megalopa	0.00%	0.00%	0.00%	0.00%	0.04%	0.04%	0.00%
Crab zoea	0.00%	0.00%	0.00%	0.00%	0.03%	0.03%	0.00%
Gastropoda	0.03%	0.00%	0.40%	0.00%	0.00%	0.00%	0.00%
Limacina helicinia	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Limacina spp.	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Pterepod spp.	0.03%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Unid. snail	0.00%	0.00%	0.40%	0.00%	0.00%	0.00%	0.00%
Unid. squid	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Hexagrammos spp. (1+ yr)	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%

Appendix 8. Biomass (g) of Neocolanus spp. copepods and euphausilds estimated from Continuous Plankton Recorder tows in our study area during the auklet breeding season (May-August), 2000-2006.

	Year	N. plumchrus/flemingeri	N. cristatus	Euphausiacea
-	2000	3.061	5.296	0.078
	2001	0.279	0.0834	0.174
	2002	3.130	1.529	0.048
	2003	0.298	0.042	0.084
	2004	3.853	1.154	0.366
	2005	1.322	0.056	0.048
	2006	5.501	11.468	2.586

Agenetia S. Catalogen numbers of Lasat Auklet specimene examined at the American Museum of Natural Hostory (New York, New York, USA, MANHS), Academy of Natural Sciences (Philadelphia, Pennylvania, USA, MSR), Cowan Vertebrate Museum (Vancover, Britis Columbia, Canada; CVM), Delaware Museum of Natural History (Weinington, Delaware, USA; DMNHG), Denver Museum of Natura end Science (Denver, Colorado, USA; DMNG), Field Museum of Natural History (Weinington, Delaware, USA; DMNHG), Denver Museum of Natural end Science (Denver, Colorado, USA; DMNG), Field Museum (Moscow, Russia, MSUZM), Harvard Museum of Comparative Zoologic (Zemhörgh, Russiachustets, USA WCL), University of Michigan Museum of Zoology (Ann Arbor, Michigan, USA; UMMK2), National Museum of Natural History–Smithsonian Institution (Washington, C., USA; USMM), an Diego Natural History Museum (San Diego, California, USA; 2004), and the Yamashina Institute for Onthology (Chiba, Japar; Vo).

AMMHE 30040, 30041, 30043, 59350, 498234, APA215, 748241, 749286, 748240, 7 738406, 738407, 753408, 753409, 753501, 753501, 753503, 753504, AM55-55379, 148115, 148112; CVM-31290; DMMH-55060; DMME-15168; PMMH-19171, 591118, 159119; MUXDA-41221, 4724, 64650, 70053, 70777, 79193, 81118, 68668, 85669, 98259, 98259, 119172, 14724, 64650, 70053, 70777, 79193, 81118, 68668, 85669, 98259, 20059, 317567, 13989, 337386; UMMZ-150027, USMM-32080, 237505, 460386, 270939, 317567, 13989, 337386; UMMZ-150027, USMM-32980, 237505, 460386, 460337, 500MH+16003, 16064, 15065, 15066, 39723, VG-12766, 12769, 12771, 1277, 12778, 12778, 2778, 2788, 2788, 2788, 2789, 27605, 20059,

Appendix 10. Means, standard deviations, and sample sizes from stable-isotope analysis of Least and Crestef Auklets at Biddler, Kinka, and Garetlei Islanda, Akeutan Islanda, Alaska, from 2007-2010. There were no significant differences within species among islands, or among years, to data were pooled. Data are from breast (pre-breeding), P1 (chick rearing), and P1 (Doo threeding) externs.

Species	Age class	Season	n	$\delta^{13}C \pm S.D.$	$\delta^{15}N \pm S.D.$
Crested Auklet	Adult	Pre-breeding	10	-18.78 ± 1.74	14.73 ± 2.27
		Chick rearing	10	$-18.70 \pm 0.97$	13.70 ± 1.78
		Post-breeding	10	-18.54 ± 1.24	$16.71\pm0.90$
Least Auklet	Adult	Pre-breeding	50	$-18.65 \pm 1.63$	$12.84 \pm 1.63$
		Chick rearing	50	$-19.02 \pm 1.71$	12.30 ± 1.37
		Post-breeding	50	-17.54 ± 1.03	$14.37 \pm 1.04$
Least Auklet	Subadult	Pre-breeding	39	-19.95 ± 1.88	$11.10 \pm 1.44$
Least Auklet	Chick	Chick rearing	8	-20.67 ± 0.35	8.88 ± 0.23





