Regional and annual variability in common eider nesting ecology in Labrador

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Nesting densities are often used to estimate breeding population size and with other measures of reproductive performance can be useful indicators of population status. These aspects of breeding biology often show considerable spatial and temporal variation. Between 2000 and 2003, we surveyed nesting common eiders (Somateria mollissima) on 172 islands in three archipelagos (Nain, Hopedale, Rigolet) on the Labrador coast. Rigolet was the largest archipelago (2834 km²) followed by Nain then Hopedale, and island density varied inversely with archipelago size. Overall means were: nest density 52.0 ± 141.9 (SD) nests/ha; nest initiation 12 June \pm 12 days; clutch size 3.7 \pm 1.2 eggs/nest; egg volume 98.8 \pm 10.4 cm³; and clutch volume 392.3 ± 135.0 cm³. Rigolet had the highest average egg volumes and nest densities, the highest single island nest density of 1053 nests/ha, and the earliest average nest initiation date. We found significant differences in nest densities among archipelagos and across years; significant archipelago and year interactions were detected for nest initiation date and clutch size. Significant differences were found among individual islands for all response variables except egg volume. For egg volume, within-archipelago island differences were not significant, but between-archipelago differences were significant. Thus egg volume may be a useful diagnostic to identify population affiliation.

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Understanding regional and annual variation in the breeding ecology of organisms is important for conservation and management purposes. Variable expression of ecological characteristics often occurs in response to change in the environment where organisms live. Breeding ecology can be influenced by biophysical factors at differing temporal and spatial scales (Scott et al. 2002). Aspects of avian ecology such as nest density, clutch size, and egg volume can be influenced by population dynamics, population genetics, habitat quality, and/or food availability (Lack 1967; Ryder 1970; Robertson et al. 2001; Hario & Selin

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2002). Furthermore some components of breeding ecology may be prone or resistant to annual and/or regional fluctuations in the biophysical environment (Avise & Hamrick 1996; Erikstad, Tveraa et al. 1998; Bregnballe 2002; Hario & Selin 2002).

Common eiders (*Somateria mollissima*) are an important species for many northern peoples, as a source of meat, eggs and down. These birds exhibit substantial variation in the timing of nesting, nesting density, clutch size and other aspects of their breeding ecology (Goudie et al. 2000; Robertson et al. 2001; Chaulk et al. in press). Under-

standing patterns of annual and regional variation in eider breeding performance can be very important for management and conservation purposes. Early nest initiation dates and large clutch sizes are generally indicative of favourable breeding conditions, while nesting densities are often used to estimate breeding population size. Unfortunately most eider research is limited in spatial and/or temporal scope, and the rare papers that contain multi-year and/or multi-site comparisons, are generally from temperate climes in northern Europe (Milne 1974; Swennen 1983; Coulson 1984, 1999; Hario Selin 1988; Bregnballe 2002; Hanssen et al. 2003).

The goal of this paper was to examine annual and regional variation of common eider nesting ecology at three distinct sites on the mid-Labrador coast, a subarctic region, over a four-year period (2000 to 2003). In this paper, we investigate nest density, nest initiation, clutch size, egg and clutch volume. In general we expected both regional and annual variation in most nesting characters, as these traits have been shown to vary in eiders. Of greater interest was whether annual patterns of variation were maintained across regions, or whether traits varied independently within regions across the years.

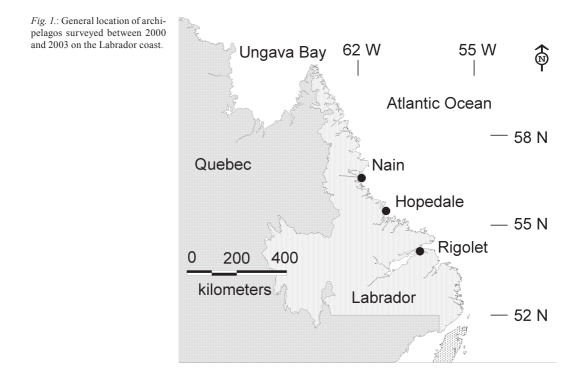
Study area

Archipelagos near the communities of Nain, Hopedale and Rigolet were surveyed from 2000 to 2003 (Fig. 1). The extent of each archipelago was determined by calculating a minimum convex polygon (Mohr 1947) containing all islands that were completely searched. The total geographic area of the three archipelagos was estimated to be 4785 km², and contained approximately of 1600 islands (Table 1). The archipelago adjacent to Rigolet covered the largest geographic area while the archipelago adjacent to Hopedale was the smallest (Table 1). Overall the average size of islands within the three archipelagos was 30.1 ha±288.8 (1 SD); on average the largest islands were found in the Rigolet archipelago (Table 1). The greatest island density occurred in the Hopedale archipelago (0.90 islands/km²) fol-

Table 1. Location, island characteristics, sampling intensity and sampling dates of archipelagos in Labrador where data were collected on nesting ecology of common eiders (Somateria mollissima), 2000–2003.

	Nain	Hopedale	Rigolet	All	
Study area location					
Mean latitude (°N)	56.36	55.33 54.18			
Mean longitude (°W)	61.06	59.81	57.41		
Study area size (km ²)	1151	800	2834	4785	
Study area perimeter (km)	145	128	28 278 551		
N (islands) in study area	497	720	335	5 1552	
Island density (islands/km ²)	0.43	0.90	0.12	0.32	
Archipelago island size					
mean±1 SD (ha)	33 ± 318	20 ± 194	47 ± 394	30 ± 289	
Range (ha)	0.01 - 6903	0.02-3875	0.02 - 5204	0.01 - 6903	
Size sampled islands					
Mean±1 SD (ha)	1.3 ± 1.5	1.6 ± 2.6	4.1 ± 10.2	2.3 ± 6.0	
Island sampling scheme					
No. one year only	13	33	17 63		
No. two years only	11	19	15	45	
No. three years only	10	14	12	36	
No. all four years	8	13 7 28		28	
Survey dates 2000	3-9 July	28-30 June	20-26 June	20 June - 9 July	
Survey dates 2001	5 - 19 July	4 - 17 July 18 - 27 June 18 June -		18 June - 19 July	
Survey dates 2002	13-22 July	3 - 17 July	17-22 June 17 June-22 July		
Survey dates 2003	11 - 13 July	3 - 7 July	14-20 June 14 June-13 Jul		

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lowed by Nain then Rigolet (Table 1).

All archipelagos shared similar environmental characteristics such as a northern maritime climate, vegetation composed primarily of mosses, lichens, forbes, grasses, and sedges. The archipelagos were typically comprised of barren islands with sparse vegetation and very limited nesting cover. Islands in the Rigolet area had denser and taller ground vegetation, and on some islands woodier cover, including stunted black spruce (Picea mariana). All three archipelagos are classified as coastal barrens (Lopoukhine et al. 1978), are considered to have a High Boreal ecoclimate (Meades 1990) and a Low Arctic oceanographic regime (Nettleship & Evans 1985). On the central and northern Labrador coast the concept of discrete archipelagos is somewhat misleading, as the island complex along this coast is for the most part continuous. The island complexes, hereafter referred to as archipelagos, selected for study were typical of this portion of the Labrador coast.

Methods

From 2000 to 2003 we surveyed three archipelagos (Nain, Hopedale, Rigolet) for nesting

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eiders. Islands were selected for study on the basis of random sampling. In all cases we limited our searches to islands that were estimated to be smaller than 30 ha. Analysis of the spatial distribution of our data set show that our samples were spatially random within the subset of islands that were less than 30 ha within each archipelago (K. G. Chaulk unpublished data). Ground survevs were conducted using standard search methods employed by the Canadian Wildlife Service (Nettleship 1980) and other researchers (Falardeau et al. 2003; Merkel 2004; Chaulk et al. in press); these consisted of two to four people walking linearly over the islands searching for signs of eider nesting. Islands in the three archipelagos were for the most part barren with limited nest-cover; hens and unattended nests were easily detected. Archipelagos were searched at approximately the same time each of the four years (Table 1). For the most part the senior author conducted all surveys, the exception being Nain and Hopedale in 2001 and 2002, where the senior author initiated the surveys but the field crew completed them.

For each common eider nest observed, information was recorded on apparent clutch size, nest age, and nest status. Nests were classed as follows: incubating – current season nest containing eggs; hatching – at least one chick was breaking its shell; hatched – at least one chick was completely out of its shell; depredated – broken and bloody eggs were present in or immediately adjacent to the nest; and unknown – nest was in disrepair with no eggs or signs of depredation.

The incubation period was assumed to be 24 to 26 days (Goudie et al. 2000), with incubation generally commencing after the laying of the penultimate egg (Hanssen et al. 2002). Candling was used to age the eggs (Weller 1956) and to calculate nest initiation we added the number of eggs to the egg age and we subtracted this number plus one from the survey date. Nests with more than six eggs were not aged. Apparent clutch sizes were calculated using nests classified as incubating; nests with more than six eggs are generally considered dump nests produced by two or more females (Swennen 1983; Robertson 1995) and were omitted from the analysis. Due to lack of data, nest initiation dates were not calculated for the 2001 breeding year.

With respect to egg size measurements we randomly selected a subset of previously surveyed islands, and on these we randomly selected nests and eggs for further investigation. Each randomly selected egg was then measured using Vernier calipers: egg length was measured from pole to pole, and width was measured at the widest part of the egg; all measurements were recorded in mm. Egg volume was calculated based on the formula presented by Guild (1974) and Robertson et al. (2001); clutch volume was estimated only for nests for which we had data on egg volume and clutch size. Clutch volume was estimated as clutch size multiplied by the estimated egg volume for a given nest.

Island nest densities were calculated using islands that were completely searched. Island sizes were derived from digital 1:50000 base maps for the Labrador coast. Geodetic coordinates are reported as latitude and longitude, decimal degrees, North

American Datum 1983. Nest density, nest initiation, and clutch size were analysed using a general linear model, with the year, archipelago and its interaction as fixed factors. Island, nested within archipelago, was also included as a random factor to control for multiple measurements of islands across years. Egg volumes and clutch volumes were analysed with one-way ANOVA, with archipelago as a fixed factor and island, nested within archipelago, as a random factor. Critical alpha was set at 0.05 for all tests, which were all two-tailed.

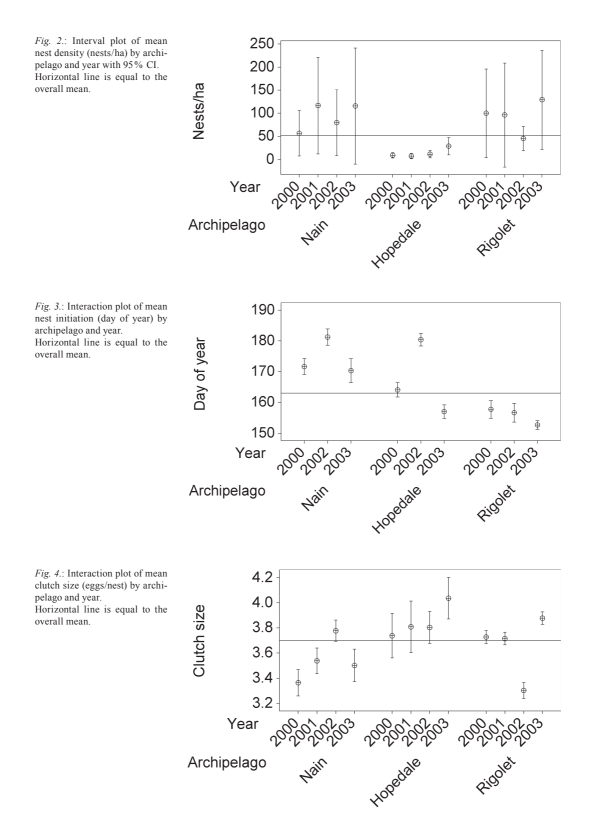
Results

We sampled 172 islands in the three archipelagos with the greatest island sampling effort expended in Hopedale (Table 1). The average size of sampled islands was 2.3 ± 6.0 ha (Table 1). On the islands that were completely searched we counted 10962 eider nests and these contained 35401 eggs. Overall, on average there were 52 nests/ha or 172 eggs/ha (Table 2). The highest nest counts were observed in Rigolet; one island had 654 nests, and another island, 0.18 ha in size, had the equivalent of 1053 nests/ha. For island nest-density both archipelago and year were significant factors, although they did not show statistically significant interaction (Table 3). There appeared to be greater between-archipelago variation than between-year variation in nest density (Fig. 2).

The overall average nest initiation date was 12 June, the earliest average nest initiation dates occurred in the south at Rigolet (3 June) and the latest in the north at Nain (23 June) (Fig. 3). However, we detected a significant interaction between archipelago and year for nest initiation date (Table 3, Fig. 4). The overall average clutch size was 3.7 ± 1.2 eggs/nest (Table 2). The largest average clutch size was observed in Hopedale at 3.8 ± 1.2 eggs, followed by Rigolet at 3.7 ± 1.2 eggs, and Nain at 3.6 ± 1.2 eggs; howev-

Table 2. Mean (\pm 1 SD) and ranges of nesting traits of common eiders breeding in Labrador, 2000–2003.

	Period	Ν	Mean
Nest density (nests/ha)	2000-03	331	52.0 ± 141.9 (range = 0 - 1053)
Nest initiation	2000, 02-03	272	12 June \pm 12 d (range = 21 May to 9 July)
Clutch size	2000-03	10137	3.7 ± 1.2 (range = 0-6)
Egg volume (cm ³)	2003	415	98.8 ± 10.4 (range = 61.0 - 160.0)
Clutch volume (cm ³)	2003	415	392.3±135 (range = 77.2 - 1008.4)



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er, we detected a significant interaction between archipelago and year for clutch size (Table 3, Fig. 4). Egg volume varied significantly among archipelagoes but not across individual islands (Table 3). The largest egg volume was observed in Rigolet and the smallest in Nain (Fig. 5). The largest clutch volume was observed in Hopedale (Table 2, Fig. 5) although the differences among archipelagos were not significant (Table 3).

Discussion

The highest nest density that we observed occurred in Rigolet (1053 nest/ha). This observation was made on a small island approximately 0.18 ha, and is high relative to nest densities reported elsewhere. For example, *S. m. borealis* typically nest at low nest densities (0.8 to 4.5 nests/ha; Cooch 1986), though there are some exceptions, such as East Bay, Southampton Island (Abraham & Finney 1986). Nest densities for *S. m. dresseri* in the St. Lawrence River averaged 3 nests/ha, but reached as high as 741.5 nests/ha (Chapdelaine et al. 1986). Nesting densities are probably influenced by numerous interacting local factors, including—but not limited to—available nesting islands and brood rearing areas, predators, disturbance, and overall population size.

Previously, we also observed significant differences in nest densities among archipelagos across a larger geographic range in Labrador, but were not able to examine annual variation (Chaulk et al. in press). The present analysis suggests that annual variation is important, and suggests increasing nest densities over the 2000–2003 study period.

Nest initiation

Spring ice break-up was late in both the Hopedale and Nain archipelagos during 2002. We found a significant interaction between archipelago and year with respect to nest initiation date, with Rigolet in the south showing relatively consistent timing, and the two northern archipelagos showing greater annual variation.

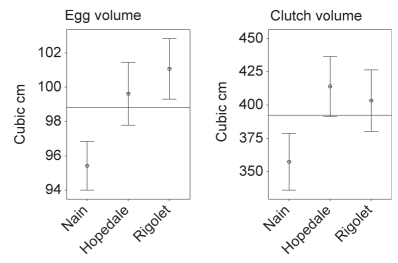
Earlier, we documented that nest initiation dates in Labrador were positively related to lat-

Table 3. Summary of general linear model analysis by variable for nesting characteristics of common eiders breeding in Labrador, 2000–2003.

Variable	e Model fit Factors		df	F	р
Nest density	90.3	Year	3	4.2	< 0.01
		Archipelago	2	3.9	0.02
		Year-archipelago interaction	6	1.1	0.36
		Island nested in archipelago	165	17.9	< 0.01
Nest initiation	72.6	Year	2	42.7	< 0.01
		Archipelago	2	62.1	< 0.01
		Year-archipelago interaction	4	7.1	< 0.01
		Island nested in archipelago	63	2.4	< 0.01
Clutch size	6.6	Year	3	3.5	< 0.01
		Archipelago	2	9.4	0.01
		Year-archipelago interaction	6	10.1	< 0.01
		Island nested in archipelago	116	4.7	< 0.01
Egg volume	6.1	Archipelago	3	5.5	< 0.01
		Island nested in archipelago	43	1.1	0.27
Clutch volume	11.5	Archipelago	3	0.8	0.44
		Island nested in archipelago	43	2.0	< 0.01

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Fig. 5.: Interval plot of mean egg and clutch volume (cm³) by archipelago with 95 % CI. Horizontal lines are equal to the overall means.



itude, but we were not able to test year effects (Chaulk et al. in press). This latitudinal pattern still holds over multiple years, but there can be substantial between-year variation in the timing of nest initiation. This highlights the importance of multi-year studies, especially for those aspects of breeding ecology that can be easily influenced by biophysical factors, such as sea ice break-up in the spring (Laurila & Hario 1988; Goudie et al. 2000). Given that we found an interaction between archipelago and year in nest initiation date, it is likely that nest initiation date is influenced by ice conditions at the local level. It is important to recognize that common eider nest initiation can be influenced by annual variation in the timing of spring break-up. This could mean that common eiders are a potentially good species for monitoring the effects of climate change in this region of North America, but our results show that more than one site would need to be monitored

Clutch size

Clutch size is often used as a comparative measure between different populations (Lewis 1939; Milne 1974; Swennen 1983; Coulson 1984; Robertson et al. 2001; Bregnballe 2002; Chaulk et al. in press). Common eider clutch size is influenced by female body condition, food availability, disease, body parasites, severity of winter, timing of spring, predation and nest parasitism (Rohwer 1992, Erikstad, Bustnes et al. 1993).

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Some researchers have suggested that average clutch size may increase during population growth (Hario & Selin 1988), whereas other have found no such trend (Swennen 2002).

Our analysis revealed a significant interaction of archipelago and year on clutch size, indicating that any annual patterns were not matched across regions. However, our model had a low R^2 value; therefore a significant amount of variability remains unexplained by either archipelago, year or their interaction. Interestingly, Bregnballe (2002) did find that clutch size varied across years in a similar way across five colonies in Denmark. At a larger spatial scale, Coulson (1999) found that clutch size varied independently across years between Scottish and Dutch colonies. Clearly, the geographic scale of the analysis appears to be important. Previously, we found significant archipelago differences in clutch size but this finding was based on analysis of one year of data (Chaulk et al. in press). Annual and regional variation in clutch size is not surprising. Our new findings suggest that average clutch size varies by archipelago and year at the scale of coastal Labrador, and we do not recommend the use of single measures (i.e., one archipelago and/ or year) of clutch size as a basis to assess population productivity. Instead long-term measurements over several archipelagos are needed for robust comparisons between different populations.

Egg and clutch volume

Some studies have shown that annual variation in eider egg volume is limited (Swennen & van der Meer 1992; Robertson 1995; Laurila & Hario 1998; Hanssen et al. 2002). As such, egg volume may hold value as a comparative measure between populations at large geographic scales (Robertson et al. 2001). In 2003, egg volume differed significantly by archipelago, but not by island, whereas clutch volume differed by island but not by archipelago. Clutch volume is largely influenced by clutch size, and likely has limited value as a comparative measure between archipelagos (see above). Our data support the idea that average egg volume has merit to assess population structure at moderate geographic scales (i.e., 100s of km). But with only one year of data and low R² values for our egg and clutch volume models additional data are required to assess spatial and temporal interactions.

Using values presented in Goudie et al. (2000) we calculated average egg volumes for two subspecies of common eiders in North America (bore $alis = 96.4 \pm 4.4 \text{ cm}^3$, range = 93.8 - 102.8; dresseri $= 108.0 \pm 5.5$, range = 100.6 - 115.8). Based on an examination of these values it appears that egg volumes from Nain were most similar to those of borealis, while the values for Hopedale and Rigolet were on the high end for borealis and on the low end for dresseri. This pattern is likely the result of intergradation between borealis and dresseri in the zone of overlap that is considered to occur in both Hopedale and Rigolet (Mendall 1980; Chaulk et al. in press). Egg volume followed a latitudinal pattern with eiders from Rigolet in the south having the largest egg volumes and eiders from Nain in the north having the smallest.

Finally, we feel the need to comment on our sampling scheme, as the average size of sampled islands was significantly lower than the average size of islands in each archipelago. We actively excluded islands larger than 30 ha from our surveys. We did this for logistical reasons. Large islands require significant effort to search, so instead we focused on smaller islands that could be easily censused by small field crews over restricted time periods. Goudie et al. (2000) reported that common eiders preferred nesting on islands <75 - 100 ha. Other researchers have used island size thresholds to help identify islands for investigation during eider breeding research (Nakashima & Murray 1988; Robertson & Gil-

christ 1998; Merkel 2004) or focused on small islands during breeding surveys (Korschgen 1977; Götmark & Åhlund 1984). It is possible that omitting islands > 30 ha has impacted our results, most likely with respect to our estimates of nest density and percentage of occupied islands. In addition, larger islands might have greater vegetative coverage, and cover has been shown to increase nest success (Choate 1967; Milne 1974; Schmutz et al. 1983), and nest success could in turn impact some of the breeding characteristics that we discuss in this paper. Since we lack data from islands > 30 ha we have no way of knowing the magnitude or direction of these differences. However, our personal observations and discussions with local aboriginal people support the idea that eiders breeding on the mid-Labrador coast rarely use large islands (>30 ha) for nesting. We recognize that the spatial structuring and the biophysical characteristics of eider breeding islands are important, but these variables are beyond the scope of this paper. We hope to investigate habitat requirements of common eiders breeding in Labrador in future research.

Conclusions

To summarize our findings, when annual effects were detected, they often interacted with regional effects. Based on the inter-annual and interregional variation, as well as their interactions. long-term surveys over wide geographic regions are needed for comprehensive understanding of population dynamics and responses to environmental changes. Such understanding is needed for sound management decisions. For example, large clutch sizes and early breeding could lead to a liberalization of hunting regulations given the expected large number of young birds in the fall flight. Conversely, reduced clutches and late breeding might be used as rationale to reduce harvest quotas and limit hunting seasons. However, if only one site is assessed, our results suggest that these indicators of breeding conditions would not be representative for an entire breeding range. Finally, egg volume does not appear to vary between islands at small geographic scales (i.e., within an archipelago), but does vary at moderate geographic scales (i.e., 100s of km) and could hold promise as an indicator of population genetic differences (Robertson et al. 2001).

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