Journal of Applied Ecology 2007 **44**, 875–884

Diet reconstruction and historic population dynamics in a threatened seabird

D. RYAN NORRIS*¹, PETER ARCESE†¹, DAVID PREIKSHOT‡, DOUGLAS F. BERTRAM§ and T. KURT KYSER¶

*Department of Integrative Biology, University of Guelph, Guelph, ON, Canada N1G 2W1; †Centre for Applied Conservation Research, Department of Forest Sciences, 2424 Main Mall, University of British Columbia, Vancouver, BC, Canada V6T 1Z4; ‡Aquatic Ecosystems Research Laboratory, Fisheries Center, 2202 Main Mall, University of British Columbia, Vancouver, BC, Canada V6T 1Z4; \$Canadian Wildlife Service, clo Institute of Ocean Sciences, 9860 West Saanich Road, PO Box 6000, Sidney, BC, Canada V8L 4B2; and ¶Department of Geological Sciences and Geological Engineering, Queen's University, Kingston, ON, Canada K7L 3 N6

Summary

- 1. For the overwhelming majority of species, we lack long-term information on the dynamics of populations. As a consequence, we face considerable uncertainty about how to discriminate among competing hypotheses of population decline and design conservation plans.
- 2. The marbled murrelet *Brachyramphus marmoratus* is a small seabird that nests in coastal old-growth forest but feeds year-round in near-shore waters of the north-eastern Pacific. Although a decline in nesting habitat is the primary reason why marbled murrelets are listed as threatened in Canada, nest predation and food availability may also influence population abundance. To examine the hypothesis that murrelet populations are influenced by variation in diet quality, we analysed stable-carbon and -nitrogen isotopes in feathers of museum specimens collected in the Georgia Basin, British Columbia.
- 3. Between 1889 and 1996, we found a decline in stable isotopic signatures that was approximately equal to a 62% drop in trophic feeding level. We also found that the estimated proportion of fish in murrelet diet was related closely to murrelet abundance over the past 40 years, as estimated from volunteer surveys. Using these isotopic data, we modelled population size as a function of variation in reproductive rate due to changes in diet quality and found that our model matched closely the 40-year field estimates. We then applied our 107-year isotopic record to the model to back-cast estimates of population growth rate to 1889.
- **4.** Our results suggest that, up to the 1950s, murrelet populations in the Georgia Basin were capable of growing and were probably limited by factors other than diet quality. After this period, however, our results imply that murrelets were often, but not solely, limited by diet quality.
- **5.** Synthesis and applications. Protecting nesting habitat may not be sufficient to rebuild populations of this highly secretive and threatened seabird and recovery might also require the restoration of marine habitat quality, as well as a better understanding of how ocean climate affects prey abundance and reproductive rate. Combined with contemporary demographic data, stable isotope analysis of historic samples provides a unique opportunity to reconstruct population histories for species where we lack long-term information.

Key-words: back-cast, *Brachyramphus marmoratus*, Pacific coast, population decline, seabirds, stable isotopes, trophic level

Journal of Applied Ecology (2007) **44**, 875–884 doi: 10.1111/j.1365-2664.2007.01329.x

Introduction

Identifying factors that limit the abundance of animals is a critical step towards diagnosing the causes of population decline. For the overwhelming majority of species, however, we have little information about the historic dynamics of populations. As a consequence, we face considerable uncertainty about how to discriminate among competing hypotheses of population trend and how to design reliable conservation plans (Caughley 1994; Jackson et al. 2001; Baker & Clapham 2004). Here, we reconstruct a 107-year record of variation in diet for the marbled murrelet Brachvramphus marmoratus Gmelin 1789, a threatened seabird that nests in coastal old-growth forest but feeds year-round in near-shore waters of the north-eastern Pacific from California to Alaska (Nelson 1987), to test if diet quality influenced population growth.

Murrelets are highly secretive birds that forage individually or in small groups and nest on the limbs of large trees up to 100 km inland (Nelson 1987; Hull et al. 2001). The first nest was discovered in 1974 (Binford, Elliot & Singer 1975), and only about 200 have been described to date (Zharikov et al. 2006). Despite these impediments to research, evidence suggests that the availability of suitable nesting habitat in old-growth forest is a key factor limiting murrelet populations over much of their range, and recovery efforts have therefore focused mainly on conserving nesting habitat to promote population persistence (Ralph et al. 1995; Burger 2002). Although a decline in nesting habitat is the primary reason why marbled murrelets are listed as threatened in Canada (Burger 2002), nest predation and food availability probably also influence their population growth (Ralph et al. 1995; Burger 2002; Peery et al. 2004). So far, comprehensive tests of these potentially competing hypotheses have proved technically challenging and hard to sustain financially. Thus, it is possible that while the conservation of nesting habitat is necessary to conserve murrelet populations in future, it may not be sufficient to promote population growth.

We explored the hypothesis that murrelets are limited by feeding conditions at sea (Peery et al. 2004), specifically by the availability of forage fish relative to macro-zooplankton. Although macro-zooplankton are not necessarily of higher caloric value than fish (Vermeer & Cullen 1982; Hedd et al. 2002), forage fish are expected to be higher-quality prey for seabirds because they are much larger, which results in more energy gained per unit effort. For example, Becker & Beissinger (2006) estimated that a seabird such as the marbled murrelet would have to consume between 45 and 80 euphausiids to equal the energetic value of small forage fish such as a Pacific sardine Sardinops sadax or northern anchovy Engraulis mordax. Evidence suggests that murrelets are flexible foragers across their breeding range, taking a variety of forage fish and marine invertebrates (primarily macro-zooplankton; Burkett 1995).

To estimate diet, we analysed stable isotopes of nitrogen $(\delta^{15}N)$ and carbon $(\delta^{13}C)$ in feathers of museum specimens collected in the Georgia Basin, a highly exploited marine ecosystem in southern British Columbia, Canada (Pauly et al. 1998, 2001; Beamish et al. 2004). Both isotopes can be used to trace the structure of marine food webs (Hobson & Welch 1992; Kline & Pauly 1998), with δ^{15} N showing a stepwise enrichment between trophic levels (DeNiro & Epstein 1981; Post 2002) and δ^{13} C distinguishing between pelagic and benthic prey (Fry & Shear 1989). Feathers are metabolically inert after growth, and so provide an isotopic signature of diet during the period in which they were grown (Hobson & Wassenaar 1997). Little is known about the diets of murrelets in the Georgia Basin. Munro & Clemens (1931) reported that the stomach contents of four marbled murrelets in Departure Bay contained surfperch spp. (Cymatogaster), larval fish, euphausiids spp. and other small invertebrates. Northwest of the Georgia Basin in the Queen Charlotte Islands, Sealy (1975) found that subadult murrelet stomach contents contained 67% sandlance (Ammodytes spp.) and 27% euphausiids spp. Sealy (1975) also found that euphausiids spp. were more abundant earlier in the breeding season and presented some evidence that higher trophic level prey were taken preferentially in relation to their relative abundance, leading to the hypothesis that breeding could be influenced by the abundance of forage fish.

We then examined the relationship between predicted diet and murrelet abundance, as estimated from volunteer surveys between 1958 and 2000. Next, we tested whether reproductive rate, driven by variation in diet quality, was sufficient to explain trends in murrelet abundance by comparing a model to estimates of abundance from field observations. Finally, by applying pre-1958 stable isotope values to the model, we back-cast population growth rates to 1889 to ask what effect variation in diet quality might have had on historic population growth rate.

Methods

FEATHER SAMPLING AND STABLE ISOTOPE ANALYSIS

We sampled feathers from adult marbled murrelet specimens in alternate (breeding) plumage (n = 102) from nine museums across Canada and the United States. We analysed brown-tipped breast feathers, which are grown during the prebasic moult from March to April (Nelson 1987). Because murrelets begin breeding in late April to early May (Nelson 1987; McFarlane Tranquilla, Williams & Cooke 2003), isotope values from these feathers provide a signature of diet during the late prebreeding or the early breeding period. In the field, we have observed some murrelets that were captured in mid-April in Desolation Sound, south-western British Columbia, growing new breast

Stable isotopes and population dynamics

feathers at the same time as they were developing a brood patch (i.e. suggesting that they are beginning to breed; D. R. Norris and P. Arcese, unpublished data). All museum samples originated from the Georgia Basin (range: 50°43′-47°15′ N, 121°58′-125°29′ W) between 1889 and 1996. Stable isotope analysis was conducted at the Queen's Facility for Isotope Research, Kingston, Ontario. Isotope ratios (R) are expressed in δ units where $\delta = [(R_{\text{sample}}/R_{\text{standard}}) - 1] \times 1000$. δ^{15} N is the ratio of $^{15}N/^{14}N$ relative to air, and $\delta^{13}C$ is the ratio of ¹³C/¹²C relative to Pee Dee Belemnite. Before analysis, feathers were washed in 2:1 chloroform: methanol solution for 24 h, rinsed in distilled water, and left to air-dry for 48 h. Between 0.10 and 0.15 mg of each feather (including material from the vane and rachis) was weighed in a tin capsule, then combusted and oxidized in a TC Elemental Analyser and introduced online into a Finnegan MAT Delta Plus XL isotope ratio mass spectrometer. During analysis, we ran four standards (reported here as mean \pm SD). For carbon, these were the international standard NBS 21 graphite $(-27.97\% \pm 0.13, n = 35)$ and an in-house standard UC-1 graphite (-5.77% ± 0.22 , n = 28). For nitrogen, we used the international standard RM 8548 ammonium sulphate $(19.86\% \pm 0.40, n = 43)$. For both elements, we also used an in-house keratin-based standard: domestic chicken *Gallus gallus* feathers (δ^{13} C: -19.46% ± 0.19 ; δ^{15} N: $7.7\% \pm 0.34$, n = 25), which were prepared in the same fashion as murrelet feathers (cut up into small pieces totalling 0·10 and 0·15 mg). All four standards showed very low variability and matched values run previously in our laboratory. We also ran duplicate feather samples from the same murrelet (n = 51), which produced a mean (± SE) difference or repeatability of $\pm 0.25\%$ (± 0.04) for $\delta^{15}N$ and $\pm 0.18\%$ (± 0.03) for $\delta^{13}C$.

Some evidence suggests that individuals replace breast feathers sequentially from anterior to posterior (Nelson 1987). Using a subset of birds (n = 13), we found no difference between $\delta^{15}N$ or $\delta^{13}C$ values of breast feathers sampled from the anterior vs. posterior areas of the breast (two-tailed paired *t*-test, δ^{15} N: t = 1.28, P = 0.23; δ^{13} C: t = -0.17, P = 0.87), suggesting either that murrelets have similar diets over the prebreeding period or that they moult their breast feathers at similar times. We proceeded to use posterior breast feathers in subsequent analyses. Similar to previous work on this species (Becker & Beissinger 2006; Becker et al. 2007), we found no difference in isotopic compositions between males (δ^{15} N mean \pm SD: 14.7 ± 1.3 ; δ^{13} C mean \pm SD: -16.2 ± 1.1) and females $(\delta^{15}N: 14.8 \pm 1.3; \delta^{13}C: -16.4 \pm 1.2; \text{ two-tailed } t\text{-test},$ δ^{15} N: $t_{88} = 0.60$, P = 0.55; δ^{13} C: $t_{88} = -0.78$, P = 0.44), so sexes were pooled.

DIETARY MIXING MODEL

We used IsoError (Phillips & Gregg 2001), a dualisotope (δ^{15} N, δ^{13} C), three-source mixing model, to

estimate the relative proportion of prey groups in murrelet diet for each year between 1889 and 1996. Potential prey items (with published or unpublished source) included euphausiid spp. sandlance (Hobson, Piatt & Potocchelli 1994; E. Davies, M. Hifpner, R. C. Ydenberg, unpublished data), Pacific herring Clupea harengus, eulachon Thaleichtys pacificus, black prickleback Xiphister atropurpureus, blackbelly eelpout Lycodopsis pacifica, shiner surfperch Cymatogaster aggregata (Hobson et al. 1994) and amphipod spp. (Hobson, Drever & Kaiser 1999). All prey samples were from coastal British Columbia: Barkley Sound, south-western Vancouver Island (Hobson et al. 1994), Triangle Island, northern Vancouver Island (E. Davies, M. Hifpner, R. C. Ydenberg, unpublished data) and Queen Charlotte Islands (Hobson et al. 1999). Although there may be some seasonal variation in isotopic signatures in marine food webs (Kline 1999), most prev samples were collected in May or June, just after murrelets moulted feathers. Based on their separation along the two isotopic gradients, prey samples were grouped into three diet sources: forage fish, euphausiids and amphipods. We used a diet-tissue (- feather) fractionation factor of +3.7% for δ^{15} N and +1% δ^{13} C, the same values used for recent isotope-dietary studies on marbled murrelets (Becker & Beissinger 2006; Becker et al. 2007), which are based on diet-tissue isotopic experiments of a closely related species (common murre Uria aalge). Minor adjustments ($\pm 2\%$) to the diet-tissue fractionation factor value shifted the proportion estimated for each diet group simultaneously and did not affect rankings of prey items in the murrelet diet (as in Becker & Beissinger 2006) nor the relationship between diet and murrelet abundance.

MURRELET ABUNDANCE AND STATISTICAL ANALYSIS

As an index of abundance between 1958 and 2000, we used the number of marbled murrelets reported per hour from the National Audubon Society Christmas Bird Count (CBC; www.audubon.org). Counts were summed annually over 11 areas in the Georgia Basin: Campbell River, Comox, Lasqueti Island, Nanaimo, Nanoose Bay, Parksville/Qualicum Beach, Pender Harbour, Squamish, Sunshine coast, Vancouver and Victoria. CBC counts do not distinguish between age classes. Because the number of observers varied each year, we divided annual estimates by the number of observer hours. Annual estimates were smoothed using LOESS with a second-degree polynomial and 20-year smoothing window. The 20-year smoothing window was chosen because gaps in our isotope data ranged from 1 to 11 years over the 107 years time-series; thus larger windows risked underestimating temporal variation in annual estimates, and narrower windows introduced potential bias due to uneven sampling. Nevertheless, modest adjustments (10-30 years) to the smoothing window had little influence on the results we report

here. We first examined changes in isotopic compositions of feathers over time using linear mixed-model regression, with year as both a random factor and fixed trend effect, because there were different numbers of birds sampled in each year. Using Akaike's information criterion (AIC) (Burnham & Anderson 2002), we compared this model with one that had year as a random factor only. Differences between the models were assessed with Δ AICc and Akaike weights (w_i).

Because linear mixed models explained a relatively small portion of the variance, we fitted the isotopic time-series (δ^{15} N and δ^{13} C) using a cubic spline ($\alpha = 1$), allowing us to assess more accurately annual and decadal variation in the data. From this fit, we used predicted isotopic values for each year to estimate the proportion of prey items consumed by murrelets, as calculated by a dual-isotope, three-source mixing model (see above). We explored the relationship between isotope values (as well as predicted diet) and murrelet abundance with linear or quadratic regression.

POPULATION MODEL

We tested if variation in female reproductive rate expressed as a function of diet quality (estimated with stable isotopes) could account for annual variation in CBC estimates between 1958 and 2000. To do so, we followed the method of stock reconstruction for single species assessment using a time-series of abundance estimates outlined by Walters (1986) and Walters & Martell (2004). We modified a simple deterministic model of Beissinger & Nur (1997) by assuming that juvenile $(s_i = 0.74)$, subadult $(s_s = 0.79)$ and adult $(s_a = 0.88)$ survival rates were constant and equal to rates estimated for the Georgia Basin (Cam et al. 2003; Steventon, Sutherland & Arcese 2003, 2006). Following these authors, we also assumed that only adult females are able to breed. Thus, with N_a and N_s representing the number of adult and subadult females alive at time t, respectively, we can estimate the total number of females alive prior to breeding at time $t + 1 (N_{t+1})$ as:

$$N_{t+1} = s_a N_a + s_s N_s + s_i b_t N_a$$
 eqn 1

where b_t is the annual fraction of adult females that produce a female juvenile (murrelets produce one egg per year; Nelson 1987). If we assume further that CBC counts estimate N_t , and that adults (N_a) comprise 66% of N_t (expected percentage calculated from an agestructured model with S_a , S_s , S_t as above and assuming $b_t = 0.31$, which is the mean value based on δ^{15} N values during the period before the CBC counts), we can estimate a time-series of b_t sufficient to reproduce the CBC counts by rearranging eqn 1 as:

$$b_t = (N_{t+1} - s_a N_a - s_s N_s)/s_i N_a$$
 eqn 2

We used estimates of b_t from eqn 1 to test our prediction that b_t and $\delta^{15}N$ (at time t) were related positively

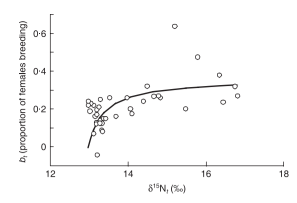


Fig. 1. Estimated variation in the fraction of adult females producing a female fledgling (b_i ; from eqn 2) in 38 years from 1958 to 1995 vs. $\delta^{15}N$ (at time t) in the breast feathers of marbled murrelets in the Georgia Basin. The solid line represents the predicted fraction of females breeding annually (RS), modelled as a saturating function of $\delta^{15}N$, where RS = [($\delta^{15}N + 12\cdot99$) × RS_{max}]/($\delta^{15}N + 12\cdot99 + \text{Half Saturation Constant}$). The line is fitted by maximum likelihood, uses a value of RS_{max} = 1·05, a Half Saturation Constant = 0·13, subtracts a constant value of 12·99 from $\delta^{15}N$ values, and scales the relationship on the x-axis. Other functions we explored such as the relationship between $\delta^{15}N$ and survival fit the data less well and modest variation in scaling had no marked effect on our final model fit to abundance estimated in the field.

(Fig. 1). After fitting this relationship, we next compared estimates of population trend from our model to CBC counts from 1958 to 2000, under the assumption that b_t varied as a function of estimated diet quality. With survival rates constant, our model population therefore grew or declined only as a function of variation in reproductive rate due to diet quality. Thus, starting population size had no influence on population growth or our comparison with CBC counts. Finally, we used isotope values from the entire time-series (including pre-1958) to estimate population growth rate (λ) from 1889 to 1996.

Results

From 1889 to 1996, we observed a decline of δ^{15} N values in marbled murrelet breast feathers (fixed effect b = -0.019, SE = 0.005, P < 0.001, r = 0.28, n = 102; mixed-model regression; Fig. 2a), suggesting that the mean trophic level of the prebreeding diet has declined over the past century. Evidence also suggested that this model was a better fit compared to a model with random effects only (mixed model: $-\log L = -165.45$, AIC = 323.96, $w_i = 0.989$; random effects model: $-\log L = -159.98$, AIC = 332.91, \triangle AIC = 8.946, w_i = 0.011; G = 5.47, d.f. = 1, P < 0.01). δ^{15} N values from pre-1900 were 15.6% compared to 13.3% between 1988 and 1996, implying a 62% drop in trophic level between these time-periods (assuming a mean difference of 3.7% between trophic levels). A non-significant trend over time was found in δ^{13} C values from the same feathers (fixed effect b = -0.01, SE = 0.005, P = 0.06, n = 102; Fig. 2b).

© 2007 The Authors. Journal compilation © 2007 British Ecological Society, Journal of Applied Ecology, 44, 875–884

Stable isotopes and population dynamics

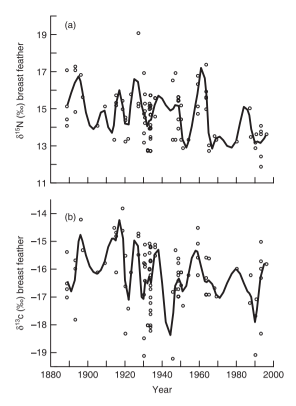


Fig. 2. Variation in the isotopic composition and diet of marbled murrelets in the Georgia Basin from 1889 to 1996. Smoothed cubic spline fit (dark line) of (a) stable-nitrogen isotopes (δ^{15} N) and (b) stable-carbon isotopes (δ^{13} C) in breast feathers (open circles). Using mixed models, we found a significant decline in δ^{15} N (fixed effect b = -0.019, SE = 0.005, P < 0.001, r = 0.28, n = 102) and a non-significant trend in δ^{13} C (b = -0.01, SE = 0.005, P = 0.06, n = 102).

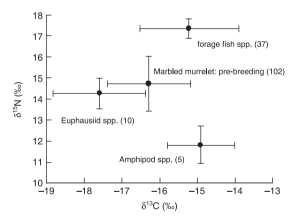


Fig. 3. Relationship between isotopic signatures of marbled murrelet feathers and potential prey. Mean (\pm SD) stable carbon (δ^{13} C) and stable-nitrogen (δ^{15} N) isotope values of marbled murrelet breast feathers sampled over a 107-year period in the Georgia Basin (subtracting an assumed diet-tissue discrimination factor of 3·7‰ for δ^{15} N and 1·0‰ for δ^{13} C), forage fish, euphausiids and amphipods (sample sizes in brackets).

After adjusting for diet-tissue discrimination, the mean prebreeding diet of all murrelets fell between the isotopic values expected if birds fed exclusively on fish or on euphausiids (Fig. 3). We used a two-isotope, three-source mixing model to reconstruct the proportion

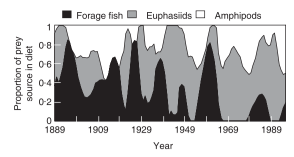


Fig. 4. Diet reconstruction of marbled murrelets over 107 years. The proportion of fish, euphuasiids and amphipods in murrelet diet predicted from a dual isotope (δ^{15} N, δ^{13} C) mixing model (see Methods). Over time, the proportion of fish showed a linear decline ($R^2 = 0.33$, P < 0.0001), euphuasiids increased ($R^2 = 0.20$, P < 0.0001) and amphipods remained relatively constant ($R^2 = 0.02$, P = 0.11).

of forage fish and zooplankton (euphausiids or amphipods) in murrelet diet over the 107-year period. We found a decline in the proportion of fish, an increase in the proportion of euphausiids, and no change in amphipods (Fig. 4). Fish comprised a mean (\pm SE) of 61% (\pm 4·4, n = 11) of the diet before 1900, but only 9% (\pm 6·3, n = 26) after 1970. In contrast, euphausiids comprised 27% (\pm 8·5) of the diet pre-1900, but rose to 61% (\pm 11·2) post-1970.

To explore the hypothesis that diet influenced murrelet population dynamics, we tested if isotope values and predicted diet composition were related to murrelet abundance in the Georgia Basin, as estimated from surveys conducted between 1958 and 2000 (see Methods). We found that both prebreeding δ^{15} N values and the proportion of fish in the murrelet diet were related closely and positively to CBC counts, with the highest variance explained in abundance when both isotopes (Fig. 5a) and predicted diet (Fig. 5b) were measured 3-4 years previous to murrelet abundance. These striking results suggest that when murrelets fed at higher trophic levels, breeding success increased due to a positive effect of diet on prebreeding condition (Peery et al. 2004). A lagged effect of diet on population size is not surprising, because juveniles do not breed until 3-5 years of age (Ralph et al. 1995); thus, annual variation in the production of juveniles should influence population size in the postbreeding period directly, but also indirectly via its effect on the total number of females that breed 3-5 years hence (e.g. Lande, Engen & Sæther 2006). Although the highest variance in murrelet abundance was explained by diet measured 3-4 years previously, it is likely that there are also smaller, more immediate effects of changes in diet on population size (as shown by R^2 values in the inset of Fig. 5).

We next compared our estimates of the annual fraction of females producing a juvenile (b_i) from eqn 2 (see Methods) to $\delta^{15}N$ in the prebreeding diet. As predicted, this relationship was positive, rising rapidly from values of $b \sim 0$ at $\delta^{15}N$ values below 13·5, then rising to an asymptote near $b \sim 0.35$ at $\delta^{15}N$ values over

© 2007 The Authors. Journal compilation © 2007 British Ecological Society, Journal of Applied Ecology, 44, 875–884

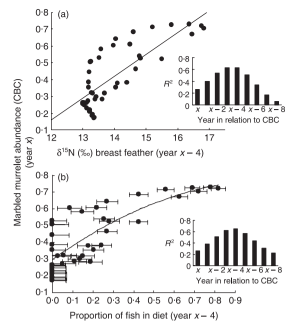
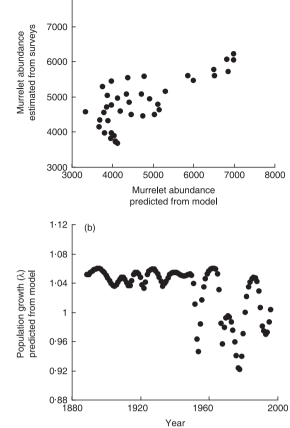


Fig. 5. Relationship between diet and marbled murrelet abundance over the past 43 years. Both (a) δ^{15} N values in breast feathers ($R^2 = 0.63$, P < 0.0001) and (b) the proportion of fish (\pm SE) predicted in the diet; ($R^2 = 0.65$, P < 0.0001) of murrelets (in year x) are significant predictors of abundance 4 years later (year x + 4), as estimated from the National Audubon Society Christmas Bird Count. Insets: in both δ^{15} N (top) and the proportion of fish (bottom), most of the variation (R^2) in murrelet abundance (CBC counts in year x) was explained when variables were measured between 3 and 4 years prior to abundance estimates.

14.5 (Fig. 1). We then fitted this relationship to a saturating curve (Fig. 1). Predicted values of female reproductive rate from this fitted relationship in the last 5 years of our $\delta^{15}N$ time-series (1992–96) reached a high of 23% of females breeding successfully with $\delta^{15}N=13.61$, to a low of 15% with $\delta^{15}N=13.17$ (Fig. 1). These values closely matched empirical estimates of reproductive rates from radio-tracked murrelets in the Georgia Basin between 1998 and 2001 (0·19–0·23 female offspring/female; Bradley *et al.* 2004).

We then asked if our model, using the fitted relationship between reproductive rate and δ^{15} N, predicted murrelet abundance similar to CBC estimates from the field between 1958 and 1996. We modified a simple, three age-class deterministic model of Beissinger & Nur (1997) and assumed constant rates of adult, subadult and juvenile survival as estimated in the Georgia Basin by Cam *et al.* (2003), such that variation in λ occurred only as a consequence of variation in female reproductive rate. We found that model predictions and CBC estimates of variation in abundance were correlated closely and positively (Spearman's correlation, $r_s = 0.74$, P < 0.0001, n = 39; Fig. 6a), consistent with the hypothesis that murrelet abundance is influenced by the effects of prebreeding diet quality on reproductive rate.

To explore the potential population consequences of long-term declines of $\delta^{15}N$ values in the murrelet diet,



8000

(a)

Fig. 6. Historical population dynamics of marbled murrelets. (a) Murrelet abundance estimates from field surveys (as in Fig. 5 but scaled) was correlated highly with abundance predicted from a population model, which is based on the assumption that the proportion of females breeding is a saturating function of δ^{15} N (Fig. 1). (b) Back-cast time-series of population growth (λ) derived from the population model.

we used our model to back-cast estimated population growth rates (λ) to 1889, when field surveys of abundance were unavailable. Our results suggest that population growth was relatively constant prior to 1950 but fluctuated greatly thereafter (Fig. 6b). These results also imply that murrelet populations in the Georgia Basin were limited by feeding conditions at sea more often after 1950, perhaps as a consequence of over-fishing or regional changes in climate (Pauly *et al.* 1998). In contrast, the relatively high and invariant growth rates prior to 1950 suggest that marine conditions probably played a smaller role in limiting population growth during this period.

Discussion

Our results demonstrate that stable isotopes can be a powerful tool for identifying the effect of trophic feeding level on population dynamics in a species that is both difficult to study and threatened. In the absence of historic data on population trend, there is likely to be considerable uncertainty about the key mechanisms driving population change (Jackson *et al.* 2001). Failing to identify these mechanisms increases the possibility

© 2007 The Authors. Journal compilation © 2007 British Ecological Society, Journal of Applied Ecology, 44, 875–884

Stable isotopes and population dynamics

that current conservation actions will be insufficient to prevent future population declines (Baker & Clapham 2004).

The decline in δ^{15} N values in murrelets reported here are even larger than declines recorded in California. In Monterey Bay, δ^{15} N values in murrelet breast feathers sampled from 1895 to 1911 were 1.4% higher (38% higher trophic level) during cold conditions and 0.5% higher (14% higher trophic level) during warm conditions compared to feathers sampled during equivalent conditions between 1998 and 2002 (Becker & Beissinger 2006). From 1889 to 1996, we estimated that δ^{15} N values declined by 2.3\% in the Georgia Basin, BC, implying a 62% drop in trophic level. High variation in isotopic signatures over this period (Fig. 2) also led us to ask if diet might influence murrelet numbers via its effect on the annual fraction of females that reproduce successfully. We found a strong correlation between diet and the abundance of murrelets 3-4 years later (Fig. 5). Interestingly, our simple model, which assumed constant survival rates and variation in reproduction linked to isotopic signatures, produced a lag similar to that observed between isotopic signatures in museum specimens and an independent index of abundance from field surveys (Fig. 6a). Overall, our results suggest that variation in diet quality may influence the dynamics of murrelet populations in the Georgia Basin, as has already been shown for colonial breeding alcids in British Columbia (Gjerdrum et al. 2003; Hedd et al. 2006) and marbled murrelets breeding in California (Peery et al. 2004; Becker et al. 2007).

Isotopic signatures of archived museum specimens can provide a unique wealth of information for reconstructing historical diets and habitat use (Dalerum & Angerbjörn 2005; Hilton et al. 2006). For example, δ^{15} N and δ^{13} C signatures in feathers of northern fulmars Fulmarus glacialis in the north-east Atlantic were higher before 1940 compared to 1993, suggesting that this species was feeding on whale offal in the early half of the 20th century (Thompson, Furness & Lewis 1995). Although δ¹⁵N values of North Pacific pinnipeds have shown little change over the last half of the 20th century, δ^{13} C values have declined over the same timeperiod, suggesting a shift in foraging locations or a change in isotopic signatures at the base of the food web (Hirons, Schell & Finney 2001; Hobson et al. 2004). Over a longer time-period, changes in δ^{15} N and δ^{13} C signatures in feathers and bone collagen of California condors Gymnogyps californianus demonstrated a diet switch from a heavy reliance of marine-derived nutrients during the Pleistocene to exclusively terrestrialderived nutrients over the past century (Chamberlain et al. 2005).

Our results demonstrate a new approach to understanding the dynamics of a little-known species where traditional demographic studies are technically too difficult or costly to advance. Potentially, our method can be applied to a wide range of species in different ecosystems. For example, Levin *et al.* (2006) showed

recently that variation in stable-oxygen isotopes ($\delta^{18}O$) in the tooth enamel of east African mammals tracked long-term variation in environmental aridity, suggesting that one might use such data to reconstruct population histories for species limited in their abundance by rainfall (Mduma, Sinclair & Hilborn 1999). In the Chesapeake Bay (USA) an increase in $\delta^{13}C$ values in striped bass *Morone saxatillis*, related to shifts in diet or foraging location, have occurred with declines in physical condition (Pruell, Taplin & Cicchelli 2003), raising the possibility that archived tissues, such as scales, might help reconstruct estimates of stock size and population dynamics in this species.

It is possible that long-term isotopic signatures are also influenced by factors unrelated to changes in the diet of marbled murrelets. For example, anthropogenic inputs of atmospheric CO₂ may lower δ¹³C values in marine food webs (the 'Suess effect': Gruber et al. 1999: Hilton et al. 2006). However, we found little (nonsignificant) trend in δ^{13} C values in breast feathers over time, consistent with our assumption that variation in isotopic ratios was driven mainly by temporal changes in diet. Adjusting δ^{13} C values to account for the Suess effect (estimated as 0.16% decline per decade; Quay et al. 2003) had the effect of decreasing the predicted proportion of fish estimated in the murrelet diet (Figs 3 and 4), supporting further our conclusion that the trophic feeding level of murrelets has declined over the last century.

The prevalence of nitrates or the degree of vertical or horizontal advection might also cause variation in δ^{15} N values at the base of the food web. However, Rau, Ohman & Pierrot-Bults (2003) showed that δ^{15} N values of zooplankton sampled off the California coast varied only \pm 1–2‰ over a 50-year period with no significant trend over time. Schell, Barnett & Vinette (1998) also found that $\delta^{15}N$ (and $\delta^{13}C$) values of zooplankton in high-latitude Pacific waters did not vary significantly over 10 years. Despite low variation of isotopic signatures, Rau et al. (2003) also provided evidence that zooplankton were enriched in ¹⁵N during El Niño years. In contrast, we have found no evidence that $\delta^{15}N$ values were correlated with large-scale ocean climate variables, such as the Pacific Decadal Oscillation or Southern Oscillation Index (D. R. Norris, P. Arcese, unpublished data), suggesting either that climatic events had little impact on isotopic signatures at the base of the food web, or that annual variation in $\delta^{15}N$ values of zooplankton due to climate was small compared to annual variation in the murrelet diet.

Our interpretation of the positive correlation between murrelet abundance estimated by the CBC and our model assumes that individuals counted each winter in the Georgia Basin represent mainly birds that also breed in the region. Although it is possible that some murrelets wintering in the Georgia Basin breed in northern British Columbia or Alaska, Becker & Beissinger (2006) and Peery *et al.* (2004) found that most marked birds in northern California over-wintered within

300 km of their breeding areas. Although our model suggests that murrelets are influenced by marine conditions our results should be interpreted with caution, given our assumptions of limited movements between the breeding and non-breeding seasons. A priority for future research is to obtain an independent validation of our general model by comparison with other trend estimators, such as more recent radar and at sea surveys. We are also working to test whether the fraction of birds nesting varies annually with diet quality.

It is also possible that our isotopic 'landscape' of prey was not representative of actual murrelet diets as our prey samples were from only three locations. However, the δ^{13} C and δ^{15} N values of euphasiid and forage fish did not differ significantly between these locations or between these locations and areas further north (see values in Hobson *et al.* 1994; Kline 1999). This suggests that the isotopic signatures of the prey items that we utilized show relatively low variation along the northwest Pacific coast. Nevertheless, we recognize that additional prey sampling is needed to confirm these relationships, including sampling other fish species and over multiple years and seasons.

Our results emphasize the potential importance of conserving multiple habitats throughout the annual cycle in highly vagile species that are difficult to track year-round (Janzen 1986; Marra, Hobson & Holmes 1998: Cherel et al. 2006: Norris & Taylor 2006). For murrelets, our work is consistent with Peery et al.'s (2004) conclusion that protecting nesting habitat may be insufficient to promote population recovery. In this case, population recovery might require the restoration of marine habitat quality, as well as a better understanding of how long-term trends in ocean climate affect prey abundance and reproductive rate (Becker et al. 2007). Quantifying long-term changes in the amount and quality of habitats used over the entire life history of species is likely to be critical for correctly diagnosing the causes of population decline and designing reliable recovery plans in many species.

Acknowledgements

We thank the Burke Museum (University of Washington), Slater Museum (University of Puget Sound), Royal BC Museum, Cowan Museum (University of British Columbia), Canadian Museum Nature, Field Museum, Conner Museum (University of Washington State), Carnegie Museum, Royal Ontario Museum and the Academy of Natural Sciences-Philadelphia for their generous donation of feather samples. The study was supported generously by H. and W. Hesse, Endangered Species Recovery Fund (World Wildlife Fund Canada and Environment Canada), the Natural Sciences and Engineering Research Council and the Izaak Walton Killam Memorial Trust. M. Longstaffe and K. Klassen kindly assisted with stable isotope analysis. Gary Shugart and Carl Walters provided essential advice on sampling protocol and parameter estimation. Jennifer Gill, Amy Newman and two anonymous reviewers provided excellent comments.

References

- Baker, C.S. & Clapham, P.J. (2004) Modeling the past and future of whales and whaling. *Trends in Ecology and Evolution*, **19**, 365–371.
- Beamish, R.J., Benson, A.J., Sweeting, R.M. & Neville, C.M. (2004) Regimes and the history of the major fisheries off Canada's west coast. *Progress in Oceanography*, **60**, 355–385
- Becker, B.H. & Beissinger, S.R. (2006) Centennial decline in the trophic level of an endangered seabird after a fisheries decline. *Conservation Biology*, **20**, 470–479.
- Becker, B.H., Peery, M.Z. & Beissinger, S.R. (2007) Ocean climate affects the trophic level and reproductive success of the marbled murrelet, an endangered seabird. *Marine Ecology Progress Series*, **329**, 267–279.
- Beissinger, S.R. & Nur, N. (1997) Appendix B: Population trends of the marbled murrelet projected from demographic analysis. *Recovery Plan for the Threatened Marbled Murrelet* (Brachyramphus marmoratus) *in Washington, Oregon and California*, pp. B1–B35. US Fish and Wildlife Service, Portland, OR.
- Bindford, L.C., Elliot, B.W. & Singer, S.W. (1975) Discovery of a nest and the downy young of the marbled murrelet. *Wilson Bulletin*, **87**, 303–306.
- Bradley, R.W., Cooke, F., Lougheed, L.W. & Boyd, S. (2004) Inferring breeding success through radiotelemetry in the marbled murrelet. *Journal of Wildlife Management*, **68**, 318–331
- Burger, A.E. (2002) Conservation Assessment of Marbled Murrelets in British Columbia: Review of the Biology, Populations, Habitat Associations, and Conservation. Technical Report Series no. 387. Canadian Wildlife Service, Delta, BC.
- Burkett, E.E. (1995) Marbled murrelet food habits and prey ecology. *Ecology and Conservation of the Marbled Murrelet* (eds C.J. Ralph, G.L. Hunt, M.G. Raphael Jr & J.F. Piatt), pp. 223–246. General Technical Report PSW-GTR 152. US Department of Agriculture Forest Service, Albany, CA.
- Burnham, K.P. & Anderson, D.R. (2002) Model Selection and Multimodel Inference: a Practical Information-Theoretic Approach, 2nd edn. Springer-Verlag, New York.
- Cam, E., Lougheed, L., Bradley, R. & Cooke, F. (2003) Demographic assessment of a marbled murrelet population from capture–recapture data. *Conservation Biology*, 17, 1118–1126.
- Caughley, G. (1994) Directions in conservation biology. *Journal of Animal Ecology*, **63**, 215–244.
- Chamberlain, C.P., Waldbauer, J.R., Fox-Dobbs, K., Newsome, S.D., Koch, P.L., Smith, D.R., Church, M.E., Chamberlain, S.D., Sorensen, K.J. & Risebrough, R. (2005) Pleistocene to recent dietary shifts in California Condors. Proceedings of the National Academy of Sciences USA, 46, 16707–16711.
- Cherel, Y., Phillips, R.A., Hobson, K.A. & McGill, R. (2006) Stable isotope evidence of diverse species-specific and individual wintering strategies in seabirds. *Biology Letters*, 2, 301–303
- Dalerum, F. & Angerbjörn, A. (2005) Resolving temporal variation in vertebrate diets using naturally occurring stable isotopes. *Oecologia*, 144, 647–658.
- DeNiro, M.J. & Epstein, S. (1981) Influence of diet on the distribution of nitrogen isotopes in animals. *Geochimica et Cosmochimica Acta*, 45, 341–351.
- Fry, B. & Shear, E.B. (1989) δ^{13} C Measurements as indicators of carbon flow in marine and freshwater ecosystems. *Stable Isotopes in Ecological Research* (eds P.W. Rundel, J.R.

Stable isotopes and population dynamics

- Ehleringer & K.A. Nagy), pp. 196–229. Springer-Verlag, New York.
- Gjerdrum, C.A., Vallee, A.M.J., St Clair, C.C., Bertram, D.F., Ryder, J.L. & Blackburn, G.S. (2003) Tufted puffin reproduction reveals ocean climate variability. *Proceedings of the National Academy of Sciences USA*, 100, 9377–9382.
- Gruber, N., Keeling, C.D., Bacastow, R.B., Guenther, P.R., Lucker, T.J., Wahlen, M., Meijer, A.J., Mook, W.G. & Stocker, T.F. (1999) Spatiotemporal patterns of carbon-13 in global surface oceans and the oceanic Suess effect. *Global Biogeochemical Cycles*, 13, 307–335.
- Hedd, A., Bertram, D.F., Ryder, J.L. & Jones, I.L. (2006) Effects of interdecadal climate variability on marine trophic interactions: rhinoceros auklets and their fish prey. *Marine Ecology Progress Series*, 309, 263–278.
- Hedd, A., Ryder, J.L., Cowen, L.L. & Bertram, D.F. (2002) Inter-annual variation in the diet, provisioning and growth of Cassin's auklet at Triangle Island, British Columbia: responses to variation in ocean climate. *Marine Ecology Progress Series*, 229, 221–232.
- Hilton, G.M., Thompson, D.R., Sagar, P.M., Cuthbert, R.J., Cherel, Y. & Bury, S.J. (2006) A stable isotopic investigation into the causes of decline in a sub-Antarctic predator, the rockhopper penguin *Eudyptes chrysocome*. *Global Change Biology*, 12, 611–625.
- Hirons, A.C., Schell, D.M. & Finney, B.P. (2001) Temporal records of δ¹³C and δ¹⁵N in North Pacific pinnipeds: inferences regarding environmental change and diet. *Oecologia*, 129, 591–601.
- Hobson, K.A., Drever, M.C. & Kaiser, G.W. (1999) Norway rats as predators of burrow-nesting seabirds: insights from stable isotope analyses. *Journal of Wildlife Management*, 63, 14–25.
- Hobson, K.A., Piatt, J.F. & Potocchelli, J. (1994) Using stable isotopes to determine seabird trophic relationships. *Journal* of *Animal Ecology*, 63, 786–798.
- Hobson, K.A., Sinclair, E.H., York, A.E., Thomason, J.R. & Merrick, R.E. (2004) Retrospective analysis of Stellar sea lion tooth annuli and sea-bird feathers: a cross taxa approach to investigating regime and dietary shifts in the Gulf of Alaska. *Marine Mammal Science*, 20, 621–638.
- Hobson, K.A. & Wassenaar, L.I. (1997) Linking breeding and wintering grounds of neotropical migrant songbirds using stable hydrogen isotopic analysis of feathers. *Oecologia*, 109, 142–148.
- Hobson, K.A. & Welch, H.E. (1992) Determination of trophic relationships within a high Arctic marine food web using δ¹³C and δ¹⁵N analysis. *Marine Ecology Progress* Series, 84, 9–18.
- Hull, C.L., Kaiser, G.W., Lougheed, C., Lougheed, L., Boyd, S. & Cooke, F. (2001) Intraspecific variation in commuting distance of marbled murrelets (*Brachyramphus marmoratus*): ecological and energetic consequences of nesting further inland. *Auk*, 118, 1036–1046.
- Jackson, J.B.C., Kirby, M.X., Berger, W.H., Bjorndal, K.A., Botsford, L.W., Bourque, B.J., Bradbury, R.H., Cooke, R., Erlandson, J., Estes, J.A. *et al.* (2001) Historical overfishing and the recent collapse of coastal ecosystems. *Science*, 293, 629–638.
- Janzen, D.H. (1986) The enternal external threat. Conservation Biology: the Science of Scarcity and Diversity (ed. M.E. Soule), pp. 286–303. Sinauer, Sunderland, MA.
- Kline, T.J. (1999) Temporal and spatial variability of ¹³C/¹²C and ¹⁵N/¹⁴N in pelagic biota of Prince William Sound, Alaska. *Canadian Journal of Fisheries and Aquatic Sciences*, **56** (Suppl. 1), 94–117.
- Kline, T.C. Jr & Pauly. D. (1998) Cross-validation of trophic level estimates from a mass-balance model of Prince William Sound using ¹⁵N/¹⁴N data. Proceedings of the International Symposium on Fishery Stock Assessment Models (eds T. Quinn, F.I.I. Funk, J. Heifetz, J.N. Ianelli,

- J.E. Powers, J.F. Schweigert, P.J. Sullivan & C.-I. Zhang), pp. 693–702. Alaska Sea Grant College Program Report no. 98–01. Alaska Sea Grant, Fairbanks, AK.
- Lande, R., Engen, E. & Sæther, B.-E. (2006) Estimating the strength of density dependence from time series of population age structure. *American Naturalist*, **168**, 76–87.
- Levin, N., Cerling, T.E., Passet, B.H., Harris, J.M. & Ehleringer, J.R. (2006) A stable isotope aridity index for terrestrial environments. *Proceedings of the National Academy of Sciences USA*, 103, 11201–11205.
- Marra, P.P., Hobson, K.A. & Holmes, R.T. (1998) Linking winter and summer events in a migratory bird by using stable carbon isotopes. *Science*, **282**, 1884–1886.
- McFarlane Tranquilla, L., Williams, T. & Cooke, F. (2003) Using Vitellogenin to identify interannual variation in breeding chronology of marbled murrelets (*Brachyramphus marmoratus*). *Auk*, **120**, 512–521.
- Mduma, S., Sinclair, A.R.E. & Hilborn, R. (1999) Food regulates the Serengeti wildebeest: a forty-year record. *Journal of Animal Ecology*, 68, 1101–1122.
- Munro, J.A. & Clemens, W.A. (1931) Water fowl in relation to the spawning of herring in British Columbia. *Biological Board of Canada Bulletin*, **17**, 1–46.
- Nelson, S.K. (1987) The marbled murrelet (*Brachyramphus Marmoratus*). The Birds of North America, no. 276 (eds A. Poole & F. Gill). Academy of Natural Sciences and American Ornithologists' Union, Washington, DC.
- Norris, D.R. & Taylor, C.M. (2006) Predicting the consequences of carry-over effects for migratory populations. *Biology Letters*, **2**, 148–151.
- Pauly, D., Palmores, Ma.L., Froese, R., Pascualita, Sa-a., Vakily, M., Preikshot, D. & Wallace, S. (2001) Fishing down Canadian aquatic food webs. *Canadian Journal of Fisheries and Aquatic Sciences*, 58, 51–62.
- Pauly, D., Pitcher, T., Preikshot, D. & Hearne, J., eds (1998)
 Back to the Future: Reconstructing the Strait of Georgia
 Ecosystem. University of British Columbia Fisheries
 Center Research Report, Vancouver, BC.
- Peery, M.Z., Beissinger, S.R., Newman, S.H., Burkett, E.B. & Williams, T.D. (2004) Applying the declining population paradigm: diagnosing causes of poor reproduction in the marbled murrelet. *Conservation Biology*, 18, 1088–1098.
- Phillips, D.L. & Gregg, J.W. (2001) Uncertainty in source partitioning using stable isotopes. *Oecologia*, **127**, 171–179.
- Post, D.M. (2002) Using stable isotopes to estimate trophic position models, methods, and assumptions. *Ecology*, 83, 703–718.
- Pruell, R.J., Taplin, B.K. & Cicchelli, K. (2003) Stable isotope ratios in archived striped bass scales suggest changes in trophic structure. *Fisheries Management and Ecology*, 10, 329–336
- Quay, P., Sonnerut, R., Westby, T., Stutsman, J. & McNichol, A. (2003) Changes in the ¹³C/¹²C of dissolved inorganic carbon in the ocean as a tracer of anthropogenic CO₂ uptake. *Global Biogeochemical Cycles*, **17**, 1–20.
- Ralph, C.J., Hunt, G.L., Raphael, M.G. & Piatt, J.F. (1995) Ecology and conservation of the marbled murrelet in North America. General Technical Report. USDA Forest Service, Albany, CA.
- Rau, G.H., Ohman, M.D. & Pierrot-Bults, A. (2003) Linking nitrogen dynamics to climate variability off central California: a 51 year record based on ¹⁵N/¹⁴N in CalCOFI zooplankton. *Deep-Sea Research*, **50**, 2431–2447.
- Schell, D.M., Barnett, B.A. & Vinette, K.A. (1998) Carbon and nitrogen isotope ratios in zooplankton of the Bering, Chukchi and Beaufort seas. *Marine Ecology Progress* Series, 162, 11–23.
- Sealy, S.G. (1975) Feeding ecology of the ancient and marbled murrelets near Langara Island, British Columbia. *Canadian Journal of Zoology*, 53, 418–433.
- Steventon, J.D., Sutherland, G.D. & Arcese, P. (2003)

884

D. Ryan Norris et al.

- Long-Term Risks to Marbled Murrelet (Brachyramphus marmoratus) Populations: Assessing Alternative Forest Management Policies in Coastal British Columbia. Technical Report 012. Research Branch, BC, Ministry of Forests, Victoria, BC.
- Steventon, J.D., Sutherland, G.D. & Arcese, P. (2006) Case study: a population-viability based risk analysis of marbled murrelet nesting habitat policy in British Columbia. *Canadian Journal of Forest Research*, **36**, 3075–3086.
- Thompson, D.R., Furness, R.W. & Lewis, S.A. (1995) Diets and long-term changes in δ^{13} C and δ^{15} N values in northern fulmars *Fulmarus glacialis* from two northeast Atlantic colonies. *Marine Ecology Progress Series*, **125**, 3–11.
- Vermeer, K. & Cullen, L. (1982) Growth comparison of a plankton- and a fish-feeding alcid. *Murrelet*, **63**, 34–39.

- Walters, C.J. (1986) Adaptive Management of Renewable Resources. Macmillan, New York, NY.
- Walters, C.J. & Martell, S.J.D. (2004) Fisheries Ecology and Management. Princeton University Press, Princeton, NJ.
- Zharikov, Y., Lank, D., Huettmann, F., Bradley, R.W., Parker, N., Yen, P.-W., Mcfarlane-Tranquilla, L. & Cooke, F. (2006) Habitat selection and breeding success in a forestnesting Alcid, the marbled murrelet, in two landscapes with different degrees of forest fragmentation. *Landscape Ecology*, 21, 107–120.

Received 6 November 2006; final copy received 12 March 2007 Editor: Morten Frederiksen