Polar bear use of a persistent food subsidy: Insights from non-invasive genetic sampling in Alaska

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Abstract: Remains of bowhead whales (Balaena mysticetus) harvested by Inupiat whalers are deposited in bone piles along the coast of Alaska and have become persistent and reliable food sources for polar bears (Ursus maritimus). The importance of bone piles to individuals and the population, the patterns of use, and the number, sex, and age of bears using these resources are poorly understood. We implemented barbed-wire hair snaring to obtain genetic identities from bears using the Point Barrow bone pile in winter 2010–11. Eighty-three percent of genotyped samples produced individual and sex identification. We identified 97 bears from 200 samples. Using genetic mark–recapture techniques, we estimated that 228 bears used the bone pile during November to February, which would represent approximately 15% of the Southern Beaufort Sea polar bear subpopulation, if all bears were from this subpopulation. We found that polar bears of all age and sex classes simultaneously used the bone pile. More males than females used the bone pile, and males predominated in February, likely because 1/3 of adult females would be denning during this period. On average, bears spent 10 days at the bone pile (median = 5 days); the probability that an individual bear remained at the bone pile from week to week was 63% for females and 45% for males. Most bears in the sample were detected visiting the bone pile once or twice. We found some evidence of matrilineal fidelity to the bone pile, but the group of animals visiting the bone pile did not differ genetically from the Southern Beaufort Sea subpopulation, nor did patterns of relatedness. We demonstrate that bowhead whale bone piles may be an influential food subsidy for polar bears in the Barrow region in autumn and winter for all sex and age classes.

Key words: Arctic, Balaena mysticetus, Beaufort Sea, bowhead whale, capture–recapture, genetic tagging, non-invasive, population estimation, subsidy, Ursus maritimus


Polar bears (Ursus maritimus) in Alaska range in the Chukchi and southern Beaufort seas. The size and conservation status of the Chukchi Sea (CS) subpopulation, shared with the Russian Federation, is currently unknown (Obbard et al. 2010). The Southern Beaufort Sea (SB) subpopulation, shared with Canada, was estimated to be approximately 1,500 bears circa 2006 (Regehr et al. 2006); estimated survival rates at that time were associated with decreasing autumn sea ice extent, and were consistent with a declining population size (Hunter et al. 2010). The SB and CS subpopulations overlap at Barrow, Alaska (Amstrup et al. 2004) and are not genetically distinct (E. Peacock et al. 2013, unpublished report). The major prey species of polar bears from both subpopulations is the ringed seal (Pusa hispida) and bearded seal (Erginathus barbatus). In the CS, polar bears also forage on walrus (Odobenus rosmarus divergens) and beluga whale (Delphinapterus leucas; Bentzen et al. 2007, Rode et al. 2013). Polar bears are not known to prey on bowhead whales (Balaena mysticetus), and therefore their presence in polar bear diets is likely a result of scavenging on beach-cast carcasses and at bone piles created from whales harvested in subsistence hunts (Bentzen et al. 2007).

Inupiat communities of coastal Alaska have hunted bowhead whales since before written records (Stoker and Krupnik 1993, Ashjian et al. 2010). Traditionally, most hunting occurred in spring, and remains from the whales were left on the ice to sink.
into the ocean. Since the 1990s, harvesting of bowhead whales has increased during autumn, when ice has retreated from shore, as whales have become more available at this time of year. In addition, deteriorating ice conditions have reduced spring whaling (H. Brower, T. Hepa, C. George, North Slope Borough Department of Wildlife Management [NSB], Barrow, Alaska, USA, personal communications, 2011; Ashjian et al. 2010). In total, more whales are now harvested because the quota was increased to meet the subsistence needs of Inupiat communities; this harvest increase was supported by an increase in estimated abundance of the bowhead whale population (George et al. 2004). As a result, bowhead whale remains from autumn hunts are now left on-shore in greater numbers in localized bone piles, which attract polar bears and other wildlife. These bone piles now constitute reliable and persistent food resources for polar bears at 3 sites: Barrow, Cross Island, and the community of Kaktovik (Fig. 1); on average, 18 whales are deposited each autumn among these sites (Bacon et al. 2010, NSB, Barrow, Alaska, USA, unpublished data). Bone piles may consist of trimmed blubber, remnant meat, bones, and other animal carcasses including birds, caribou (*Rangifer tarandus*), and fish.

At Barrow and Kaktovik, these food subsidies occur close to villages, creating the potential for human–bear conflict (Miller et al. 2006). Moreover, polar bears have become more common on land in the autumn since the 1990s (H. Brower, C. George, personal communications, 2011; US Geological Survey [USGS], Anchorage, Alaska, USA, unpublished data), which may be due to the extent, duration, and rapidity of retreating sea ice (Schliebe et al. 2008), later freeze-up, the presence of this food subsidy, or a combination of these factors. Concom-

Fig. 1. Map of the study area, showing the approximate location of the bowhead whale bone pile at Point Barrow on the northern coast of Alaska, USA.
itantly, declines in body condition, population size, and survival of polar bears in the SB subpopulation between the 1980s and mid-2000s have been linked to a reduction in optimal sea ice habitat (Durner et al. 2009, Hunter et al. 2010, Regehr et al. 2010, Rode et al. 2010). As such, the persistent, novel, and substantial (Bentzen et al. 2007) food subsidy of bowhead whale carcasses has the potential to influence individual behavior, foraging ecology, and population dynamics.

Numerous polar bears gather at these bone piles, allowing for observational studies of general patterns of use (Miller et al. 2006). Aerial surveys (Schliebe et al. 2008) can produce an estimate of the number of bears at these sites, and satellite telemetry can provide perspective on how bone piles influence movements of some individuals (Fischbach et al. 2007). However, such methods cannot provide accurate and precise estimates of the number of polar bears using the bone piles.

Understanding how these bone piles may affect population dynamics requires studies involving individually-marked animals. Physical capture or biopsy darting of bears at bone piles can be difficult due to the high number of bears in the vicinity and the location of bone piles on small spits of land or islands. Further, the number of bears that researchers could sample and identify would be small relative to the number of animals using the bone piles, rendering capture-recapture estimates imprecise. Finally, aboriginal communities, which are key partners for polar bear conservation and management across the circumpolar Arctic, often do not support such methods for polar bear research (Peacock et al. 2011). Therefore, our underlying objective was to refine the less-invasive hair sampling technique (Woods et al. 1999) for use in polar bear research at high-density sites.

We chose the bone pile at Barrow as a test case for non-invasive sampling to estimate the abundance of polar bears because of its accessibility. Previous attempts to collect samples from polar bears using hair snares have not produced enough recaptures to provide more than a minimum number of bears known alive, partly due to design of the snare and poor research access (Van Coeverden De Groot et al. 2012). Thus, we thought it appropriate to conduct our study in a location that researchers could easily access. Our primary objective in describing the bears accessing the bone piles was to use genetic capture-recapture to estimate the number of bears that used the bone pile over the 3 months following the autumn bowhead harvest (Nov–Feb). Our second objective was to characterize the demographics and temporal dynamics (frequency of visits and fidelity to the bone pile) of the bears using the resource. Finally, we examined how the group of polar bears using the bone pile compared demographically and genetically to the Southern Beaufort Sea subpopulation at large.

Methods
Field methods

Our study occurred at Point Barrow, a 7-km long gravel spit that marks the boundary between the Chukchi and Beaufort seas (Fig. 1). It is located 16 km north of Barrow, Alaska, and is the northernmost tip of the United States. Polar bears from both the CS and SB use the area (Amstrup et al. 2004). Our study was conducted from 6 November 2010 to 14 February 2011; ice reformed around the study area in early December 2010 (J.H., personal observation). Daily average temperatures during the study ranged from -27°C to -14°C with an average of -19°C (maximum, 0.5°C; minimum -43°C; http://www.arh.noaa.gov/clim/). Eight bowhead whale carcasses were deposited from 10–25 October 2010 (Suydam et al. 2010). The length of these harvested whales averaged 9.8 m, which is equivalent to approximately 3–4 tonnes of viscera and blubber per whale (George 2009).

We erected a barbed wire hair corral (Fig. 2) consisting of a single continuous strand of 4-point, 12-gauge barbed wire around the bone pile. We spaced metal t-posts approximately 10 m apart and pounded them into the gravel to a minimum depth of 40 cm. We used either bailing wire or fence clips to initially attach barbed wire to posts at a height of 0.5 m, and then adjusted the height to 0.65 m after 2 days of observation to ensure bears contacted the barbed wire and left a hair sample. We alternately attached the wire to the inside and outside of the t-posts to provide greater strength. The corral required continual maintenance because of snow drift, melt, and damage caused by bears. Therefore, we added fence posts, shoveled drifts, and adjusted snare height as necessary.

We checked the fence daily for the first 4 days of sampling and monitored polar bear use by remote cameras (Reconyx Inc., Holmen, Wisconsin, USA). After the first 4 days, we checked the snare at least weekly to ensure integrity of the wire. We collected samples by plucking hairs from a single barb and

placing them in a paper coin envelope (each envelope/barb represented one sample). In general, samples from a single barb ranged from a few hairs to a small clump. If a single barb had large quantities of hair, we did not collect it as a sample, since it could have been from more than one bear. A priori, we decided to collect only 1 sample within a 0.75 m section of fence, or approximately six barbs, to avoid collecting multiple samples from a single crossing event. All un-sampled hairs were cleaned from the fence at each visit via plucking or burning with a propane or MAPP™ gas torch. We air-dried samples and stored them at room temperature until DNA extraction.

We divided the sample collection period into eleven 1-week consecutive encounter occasions, with an exception of 4 weeks between the 8th and 9th encounter occasion when we were unable to collect samples due lack of personnel. Because of funding limitations, we sought to analyze approximately 20 samples for each occasion. Therefore, we randomly selected approximately 20 samples deemed to be high quality (at least 10 guard hairs with follicles). We excluded samples that did not include sufficient hairs to divide for subsequent stable-isotope and contaminants analysis, thus in some cases 20 samples per occasion was not attained. We submitted a total of 200 of the 407 samples collected for DNA analysis (Table 1).

Fig. 2. Polar bears feeding on bowhead whale carcasses inside the barbed-wire corral (a) at Point Barrow, Alaska. Polar bears were sampled while going over (b) or under (c) the barbed-wire fence. A cub negotiating the fence (d).
Laboratory methods

Hair samples were genotyped at 20 microsatellite loci by Wildlife Genetics International (Nelson, British Columbia, Canada). DNA was extracted using a Qiagen DNeasy kit (QIAGEN Inc., Valencia, California, USA) from a minimum of 10 guard hair roots to ensure sufficient quantities of DNA to minimize genotyping errors associated with low quantities of DNA (Goossens et al. 1998). When the sample was underfur, the entire clump was used in the extraction to maximize DNA quantity and quality. Initially, DNA extracts were amplified at 11 hypervariable microsatellite markers to identify individuals: G1A, G10B, G10C, CX110, G1D, G10L, G10M, MU59, G10P (Paetkau and Strobeck 1994, Taberlet et al. 1997, Proctor et al. 2002); and G10H and G10J (GenBank accession numbers U22086.1 and U22087.1, respectively). Samples that were amplified at ≤11 loci were considered unsuccessful. After individuals were identified, one sample representing each individually-identified bear was amplified at 9 additional microsatellite loci to enable co-ancestry analyses: MSUT-2, C9H9, CXX20, MU50, MU51, G10X, CXX173 (Ostrander et al. 1993, Paetkau et al. 1995, Taberlet et al. 1997, Kitahara et al. 2000, Proctor et al. 2002, An et al. 2010); and 14RENP07 and G10U (Gen Bank accession numbers AJ411284, and U22092.1, respectively). As a form of quality control, the resulting dataset was analyzed for pairs of multi-locus genotypes that differed at 1 and 2 markers (i.e., 1- and 2-mismatch pairs) to examine potential problems associated with allelic drop out (Paetkau 2003). These pairs of genotypes were scrutinized for allele size coding errors; if no errors were detected, samples were re-amplified at the specific markers to ensure correct genotype assignment. One of each sample representing each individually-identified bear was amplified at the amelogenin locus (Ennis and Gallagher 1994) to identify sex. Probability of identity for siblings ($P_{ID(sibs)}$) was calculated using the program GIMLET (Valière 2002).

Individual, sex, and family patterns of use

We tested whether the sex ratio of polar bears that visited the bone pile differed from 1:1 using a chi-square test at $\alpha = 0.05$. We sorted and matched the genotypes collected from the bone pile with genotypes of polar bears captured in the spring in the CS between 2008–2011 ($n = 185$; US Fish and Wildlife Service [USFWS], Anchorage, Alaska, USA, unpublished data) and in US portion of the SB between 1981–2011 ($n = 1,151$; USGS, unpublished data) or biopsied on the North Slope of Alaska during autumn 2010–2011 ($n = 160$; Pagano et al. 2013) to acquire ancillary data on individually-identified bears including: (1) age, (2) identities of known offspring, (3) identity of mother, (4) identity of grandmother, and (5) identities of animals the bear had been associated with in the breeding season. We specifically sought to identify matrilines to address the lay assumption that the polar bears ‘learn’ to feed at the bone pile from their mothers and subsequently return as adults. Secondly, we examined the 2010 capture (from both subpopulations) and biopsy (from SB) data to predict whether an identified bear was likely to be visiting the bone pile with a family member (mother, offspring, or sibling). This latter examination was intended to identify family groups from the bone pile genetic data (related bears that are currently in family groups). Finally, we used a sample of polar bears ($n = 75$) from the SB subpopulation that were captured on the spring sea ice in 2010 to compare sex and age class structure of the SB subpopulation at large to the group of bears identified at the bone pile. Polar bears in SB primarily occupy a small area of near-shore continental shelf sea ice during the spring, and all sex and age groups are targeted in the capture effort. Therefore, we assume the SB capture sample represents the subpopulation at large (Amstrup et al. 1986, Regehr et al. 2010) and is representative of

Table 1. Hair samples of polar bears collected at the bowhead whale bone pile at Point Barrow, Alaska, submitted for genetic analysis and successfully amplified at a minimum of 7 microsatellite loci and the sex-identification locus (amelogenin locus) during 11 sampling events between 6 November 2010 and 14 February 2011.

<table>
<thead>
<tr>
<th>Sampling event</th>
<th>Collected</th>
<th>Submitted for analysis</th>
<th>Successfully amplified</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>52</td>
<td>22</td>
<td>17</td>
</tr>
<tr>
<td>2</td>
<td>37</td>
<td>15</td>
<td>9</td>
</tr>
<tr>
<td>3</td>
<td>33</td>
<td>19</td>
<td>13</td>
</tr>
<tr>
<td>4</td>
<td>47</td>
<td>20</td>
<td>17</td>
</tr>
<tr>
<td>5</td>
<td>24</td>
<td>19</td>
<td>14</td>
</tr>
<tr>
<td>6</td>
<td>48</td>
<td>20</td>
<td>18</td>
</tr>
<tr>
<td>7</td>
<td>23</td>
<td>17</td>
<td>12</td>
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<td>8</td>
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<td>12</td>
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</tr>
<tr>
<td>9</td>
<td>47</td>
<td>20</td>
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<tr>
<td>10</td>
<td>32</td>
<td>20</td>
<td>18</td>
</tr>
<tr>
<td>11</td>
<td>32</td>
<td>16</td>
<td>16</td>
</tr>
<tr>
<td>Total</td>
<td>407</td>
<td>200</td>
<td>165</td>
</tr>
</tbody>
</table>
a true demographic population of polar bears. In contrast, the capture sample in the CS subpopulation likely does not reflect the CS subpopulation at large because most bears in the subpopulation are not available for capture in the study area, and not all demographic groups are well represented (USFWS, unpublished data). Thus, we could only compare the demography of our sample of bone pile bears to the demographic groups of the SB subpopulation.

To further characterize the extent of family use of the bone pile, we examined the multi-locus genotypes from all pairs of individually-identified polar bears for relatedness using the program Coancestry (Version 1.0; Wang 2011). We used the Queller-Goodnight relatedness estimator ($r$, Queller and Goodnight 1989) to note presence of sibling–sibling or parent–offspring pairs in the dataset ($r > 0.46$).

**Genetic characterization**

To further evaluate whether polar bears at the bone pile constitute a group of more related individuals (due to cultural learning) than individuals in a demographically representative subpopulation at large (the SB 2010 capture sample), we compared patterns of relatedness within these groups of bears (we did not compare to relatedness of CS sample because the CS sample does not represent a demographic subpopulation). We tested for a difference of Queller-Goodnight $r$ metrics of all pairwise dyads between the bone pile and springtime capture groups using the bootstrapping method described in Wang (2011). We also assessed genetic differentiation between the bone pile group and the SB subpopulation using $F_{ST}$ with Arlequin version 3.11 (Excoffier et al. 2005).

**Capture–recapture analysis**

We created individual capture histories for each polar bear identified in the sample set. We used the open-population model POPAN (Schwarz and Arnason 1996) to estimate the total number of bears using the bone pile. Because bears that use the bone pile do not constitute a closed demographic population, the POPAN model assumes these bears represent a portion of a larger superpopulation (Kendall 1999). Using the POPAN model, we estimated apparent survival ($\phi$), capture probability ($p$), and the probability of entry ($pent$; probability that an animal from the superpopulation comes to the study area during the sampling interval). The apparent survival parameter ($\phi$) can be interpreted as the fidelity of a polar bear to the bone pile from one sampling occasion to the next (one minus the probability of permanently leaving the bone pile). The probability of permanently leaving the bone pile would be much higher than the likelihood that a polar bear would actually die from week to week over the course of 3 months. We allowed $p$, $pent$, $\phi$, and $N$ to vary over sampling occasion (time), because the freshness and amount of whale meat and blubber and sea-ice conditions changed during the study. We also ran models with sex as a co-variante. Because we could not differentiate age class for all bears from the hair samples, we pooled all bears with respect to age. Finally, we ran models time constant [signified by (.)]. Because there was a 4-week break of sampling between the 8th and 9th sampling occasions, we used unequal time interval in all models. All parameter combinations resulted in 54 a priori models. We implemented the POPAN model in Program MARK (White and Burnham 1999) using the RMark interface (Laake and Rexstad 2007). We used U-CARE Version 2.2 (Choquet et al. 2005) to examine fit of the models to the data due to potential over-dispersion as a result of unequal and inter-dependent capture probabilities.

We assessed differences in model fit to the data using Akaike’s Information Criterion for small sample sizes (AICc; Burnham and Anderson 2002; Arnold 2010). We model-averaged parameters from the models that constituted 95% of the weight of the model set.

**Results**

**Individual, sex, and family patterns**

Of the 200 hair samples submitted for analysis, 165 samples were successfully genotyped at a number of loci (11) sufficient to provide individual identity, representing a genotyping success rate of 82.5% (Table 1). From the 165 samples, we identified 97 individuals in the quality-controlled data set. Overall probability of identity for siblings ($P_{ID}$ (sibs)) was $3.74 \times 10^{-8}$ (see Table 2 for per locus $P_{ID}$ (sibs)), thus we are confident in our ability to distinguish individuals. Through comparison with all other datasets (i.e., all existing SB and CS genotypes from capture and biopsy efforts), 37 of these individuals (38%) had been previously caught in SB and therefore had age information (USGS, unpub-
We identified no individuals that were previously captured by research efforts in the CS (USFWS, unpublished data). The distribution of these 37 known bears among age classes was 34 adults, 1 subadult (age 4), one cub-of-the-year, and one yearling. The average age of known independent bears (subadults and adults) sampled at the bone pile was 13.6 years (SD = 4.4); the average age of independent bears in the SB subpopulation was 10.4 (SD = 4.2). The ratio of females (39) to males (58) of the 97 identified animals at the bone pile was 0.67:1 (x² = 3.72, P = 0.054). Individual female polar bears visited the bone pile less during the latter part (Jan and Feb) of the study (Fig. 3). In our sample of known animals (n = 37), adult females without dependent young (n = 3) were only caught once at the bone pile, whereas known mothers (n = 5) were caught over an average span of 19.4 days (SE = 15).

Of the 97 individually-identified bears, most were detected at the bone pile once (n = 54, 55%; detections include multiple detections per day) or on one day (n = 72, 74%). Twenty-seven (28%) and 23 (24%) bears were detected twice or on 2 separate days, respectively (Fig. 3, 4), 15 bears were detected 3–5 times, and 1 bear was detected 8 times. Of the bears visiting the bone pile more than once, the average duration that a bear used the bone pile was 9.6 days (SD = 15.8). The duration that individual polar bears were detected at the bone pile is from first to last detection, but does not imply continuous use of the bone pile.

Combining inferences from co-ancestry results and previously-known bears (Appendix), 21 members of 12 family groups, including 6 likely family groups with dependent young, were encountered at the bone pile. This represents 22% of the individually-identified polar bears at the bone pile. As an additional source of corroboration that family groups used the bone pile during our study, we obtained photos of mothers with 2 yearlings on 7 separate days, and one family group with 2 cubs (Fig. 2d).

### Population modeling

Of the 54 a priori models, we used the top 16 models in which all parameters were estimated, and which constituted 95% of the model weight, for model averaging (Table 3). The top model φ(sex)p(sex)psev(sex)N(.) comprised 32% of the weight and demonstrated that there were differences
in capture probability ($p$), weekly fidelity (the probability that a bear remains at the bone pile from week to week; $\phi$), and probability of arrival ($p_{em}$) to the bone pile between males and females. However, this model also suggested that similar numbers of females and males visited the bone pile. In contrast, the next-best performing model [$\phi($sex$) p(.$p_{em}($sex$)) N($sex$)]$ ($\Delta$AICc = 1.93, weight = 12%), suggested capture probability was the same for males and females, but the number of males and females visiting the bone pile differed (Fig. 5). Both models have equal number of parameters, and thus the difference in AICc is fully a result of a better fit to the data of the top model (solely the result of a change in likelihood; Arnold 2010). Although we did not detect variation over time for any parameter, the temporal sex-specific abundance estimates produced by the second model represent the approximate numbers of bears using the bone pile during any given week.

The model-averaged sex-specific weekly estimates of fidelity to the bone pile (probability of being
sampled at the bone pile from week to week) was 0.63 (SE = 0.12) for females and 0.45 (SE = 0.15) for males (Table 4). The signal-to-noise ratio (β/SE) of the sex-effect was 2.26, suggesting that sex is an important variable describing fidelity to the bone pile. The probability of a new female or male bear arriving at the bone pile in a given week was 0.07 (SE = 0.01) and 0.09 (SE = 0.004), respectively.

Table 3. Model selection results for models constituting 95% of the model weight of all examined POPAN models for capture–recapture data for polar bears visiting the bowhead whale bone pile at Point Barrow, Alaska, USA, between 6 November 2010 and 14 February 2011. In model names, (sex) is the grouping factor for males and females and (.) is a model where the parameter does not vary by sex. Parameter estimates: apparent survival (Q), which is interpreted as weekly fidelity to the bone pile; recapture (p) probability, the probability of entry (pent) or weekly arrival at the bone pile; and N, the superpopulation abundance. Estimates of parameters presented in this paper are model-averaged over all models in this table.

<table>
<thead>
<tr>
<th>Model</th>
<th>Number of parameters</th>
<th>AICc</th>
<th>ΔAICc</th>
<th>Model weight</th>
<th>-2 ln likelihood</th>
</tr>
</thead>
<tbody>
<tr>
<td>φ(sex)p(sex)pent(sex)N(.)</td>
<td>7</td>
<td>264.89</td>
<td>0.00</td>
<td>0.32</td>
<td>249.87</td>
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<td>φ(sex)p(sex)pent(sex)N(sex)</td>
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<td>266.82</td>
<td>1.93</td>
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<td>251.80</td>
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<td>6</td>
<td>267.03</td>
<td>2.14</td>
<td>0.11</td>
<td>254.27</td>
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<td>φ(sex)p(sex)pent(sex)N(sex)</td>
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<td>267.14</td>
<td>2.25</td>
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<td>249.82</td>
</tr>
<tr>
<td>φ(.)p(.)pent(sex)N(.)</td>
<td>5</td>
<td>267.63</td>
<td>2.74</td>
<td>0.08</td>
<td>257.09</td>
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<tr>
<td>φ(.)p(sex)pent(sex)N(sex)</td>
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<td>2.89</td>
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<tr>
<td>φ(.)(.)pent(sex)N(.)</td>
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<td>269.24</td>
<td>4.35</td>
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<tr>
<td>φ(sex)(.)pent(sex)N(.)</td>
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<td>269.77</td>
<td>4.88</td>
<td>0.03</td>
<td>257.02</td>
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<tr>
<td>φ(sex)p(sex)pent(.)N(.)</td>
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<td>270.24</td>
<td>5.35</td>
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<td>5.42</td>
<td>0.02</td>
<td>261.96</td>
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<tr>
<td>φ(.)p(pent)N(sex)</td>
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<td>270.36</td>
<td>5.47</td>
<td>0.02</td>
<td>259.82</td>
</tr>
<tr>
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<td>270.72</td>
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<td>257.96</td>
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<td>φ(.)p(.)N(.)</td>
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<td>271.33</td>
<td>6.24</td>
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<tr>
<td>φ(.)p(.)N(.)</td>
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<td>7.50</td>
<td>0.01</td>
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<td>φ(.)p(sex)N(sex)</td>
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<td>7.52</td>
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<tr>
<td>φ(sex)p(sex)pent(.)N(sex)</td>
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<td>272.47</td>
<td>7.58</td>
<td>0.01</td>
<td>257.45</td>
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</table>
Recapture rates ($p$) were 0.23 (SE = 0.10) for females and 0.34 (SE = 0.13) for males. The model-averaged superpopulation estimate for the study period was 228 (SE = 44; Table 3).

We found good fit of the data to the models using 2 groups for males and females ($\chi^2 = 1.64, 1$ df, $P = 0.99$), and no evidence of trap-dependence ($P = 0.71$) or transience ($P = 0.5$). This suggests that over-dispersion caused by inter-dependent capture probabilities of members of family groups was minimal.

**Genetic relatedness and diversity**

The group of identified polar bears visiting the hair snare ($n = 97$) did not differ genetically from the 2010 sample of animals captured in the SB ($n = 75$; pair-wise $F_{ST} = 0.001, P = 0.149$). Genetic diversity statistics were similar for both groups (Table 1). The distributions of Queller-Goodnight relatedness metric ($r$) between pairs for the groups showed little variation. The proportion of individuals related at $r \geq 0.25$ was similar; 3.0% and 3.4% percent of dyads from hair snare and capture sample respectively ($\chi^2 < 0.001, 1$ df, $P = 0.976$).

**Discussion**

Through incorporation of detection probability, we estimated that 228 polar bears used a bowhead whale bone pile on the north coast of Alaska during the 3 month study in winter 2010–11. If all bears were from SB (see discussion below regarding CS), this represents approximately 15% of this subpopulation, which was estimated at 1,526 animals in 2006 (Regehr et al. 2006). Data suggest that this subpopulation is declining (Hunter et al. 2010; USGS, unpublished data), and thus the percent of bears using the bone pile from the SB subpopulation may be higher. Given that additional bone piles are present at Cross Island and Kaktovik, which have relatively higher concentrations of bears (Schièbe et al. 2008), and that polar bears use the bone piles
throughout the year (even when ice returns; NSB
and USGS, unpublished data), it is likely that a large
proportion of the SB subpopulation uses these bone
piles. Most bears visited only one or two times, but
some made many visits, suggesting variation in the
importance of the bone pile as a food source for
polar bears.

The abundance estimates may be biased low, as
our design violated the capture–recapture assump-
tion that all individuals have equal capture proba-
bility because recapture probabilities of members of
a family group (a mother with dependent young) are
interdependent. Although we were able to determine
the age of some bears encountered by matching their
genotypes with those of bears captured previously by
other efforts, we were unable to eliminate dependent
cubs from the modeling or otherwise differentially
model recapture probability among age classes.
Nonetheless, it is likely that unequal capture pro-
babilities yielded abundance estimates that were
biased low (Boulanger et al. 2004). However, this
bias may be minimal because we sampled cubs
without sampling mothers and mothers without
sampling cubs. This conclusion is supported by a
lack of evidence for over-dispersion in the dataset.
Some evidence in other hair-snare studies of ursids
suggests that the point estimates may not be
substantially biased due to inter-dependence of
capture probabilities, but precision of parameters
may be overestimated (Boulanger et al. 2004).

The majority of the bears sampled at the bone pile
in Barrow likely came from the SB subpopulation,
although some bears may come from the CS
subpopulation. Although 38% of the individuals we
identified were previously known from other studies
in SB, this proportion is lower than the marked
proportion (54%) within a random sample of the
subpopulation (USGS, unpublished data), suggest-
ing that unmarked polar bears from CS (which has a
much lower rate of marking) may also have been
sampled. Though no bears from the 4-year capture
sample from CS were identified at the bone pile, the
capture sample was small. Further, the capture effort
for CS bears also occurs far to the south (about
530 km), north of the Bering Strait region, and it is
likely that these bears move north and west toward
Russia in the autumn (K. Rode, USGS, Anchorage,
Alaska, USA, personal communication, 2011). Thus,
if bears from the CS visit the bone pile in autumn
they may have a movement pattern that excludes the
limited area of recent spring capture effort and so
would not have been captured previously. Eighty-
five percent of bears sampled in the CS capture effort
had bowhead whale in their diet (Rode et al. 2013),
thus these bears may be visiting bone piles near Point
Hope on the Chukchi Sea coast, visiting the Barrow
bone pile at a different time of year, or exploiting
beached whale carcasses elsewhere.

The group of polar bears that used the bone pile
was dynamic, with an average stay at the bone pile of
10 days (median = 5 days). However, overall the
fidelity parameter suggested an approximate turn-
over of 50% of the bears from week to week.
Females showed greater weekly fidelity to the
bone pile than males, perhaps reflective of shorter
movements when adult females are accompanied by
cubs. Our data suggested that mothers, relative to
unencumbered females, may drive the higher fidelity

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Group</th>
<th>Estimate</th>
<th>SE</th>
<th>$\beta/SE^a$</th>
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<td>0.01</td>
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<tr>
<td>total</td>
<td>228</td>
<td>44</td>
<td></td>
<td></td>
</tr>
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</table>

$^a$For $\varphi$, $\rho_{ent}$, and $\rho$, $\beta/SE$ is from the top model [$\varphi$(sex)$\rho$(sex)$\rho_{ent}$(sex)$N$(.)]. For $N$, the ratio is from the model [$\varphi$(sex)$\rho$(.)$\rho_{ent}$(sex)$N$(sex)]
with $\Delta AICc = 1.93$.
(estimated in the models) of females to the bone pile. Nevertheless, the median number of days that mothers visited the bone pile was one. Pregnant female polar bears enter dens between October and December and do not emerge until March or April (Lentfer 1975), which may have resulted in the observation that fewer females used the bone pile later in the study. Overall, our results suggest more males than females used the bone pile.

We determined that at least 6 family groups with dependent young used the bone pile, further corroborating that these food resources are used by all sex and age classes of polar bears (Miller et al. 2006). Ovsyanikov et al. (1994) also found that polar bear family groups with dependent young will use aggregations of walrus carcasses, despite the presence of other bears including potentially infanticidal (Amstrup et al. 2006) adult males. Though all ursids tend to be solitary, mixed sex and age groups are common at carcasses, salmon (*Oncorhynchus* spp.)-spawning streams, or garbage where food is abundant (Craighead et al. 1995). However, it is also known that mixed sex and age aggregations of ursids are not without risk of injury (Hessing and Aumiller 1994), and sexual segregation in ursids occurs (Derocher and Stirling 1990). In 2012, USGS researchers found the carcass of a yearling polar bear with puncture wounds in the skull at the bone pile at Cross Island (USGS, unpublished data), and brown bears (*U. arctos*) are known to displace polar bears at the bone pile at Kaktovik (Miller et al. 2006). Further, there seems to be a general tendency for family groups to use the bone piles at a different time of day than adult males (Miller et al. 2006). As with other ursids, polar bears will likely behave such that they optimally balance risk with reward (Gende and Quinn 2004).

Through our co-ancestry analysis and information from previously-known bears, we identified 11 adult-aged sibling–sibling and parent–offspring relationships among the bears visiting the Barrow bone pile. However, the comparisons of genetic diversity and distribution of the relatedness metric suggest the genetic diversity within the group of animals using the bone pile is not lower than within the SB subpopulation as a whole. Thus, though we detected some matriline fidelity, the assemblage of bears at the bone pile did not constitute a group of highly-related individuals.

The inferences from our study benefited from the accompanying information from long-term capture and genetic databases. Without these accompanying data, data derived from hair snares preclude knowledge of age. Hair snaring, however, can provide accurate characterization of differential sex use, and detection-probabilities enabling estimation of un-observable individuals. Further, hair snare samples provide data during day and night and during periods of the year when capture operations are impossible (e.g., the sun was below the horizon at Point Barrow from 20 Nov to 23 Jan). Given that long-term marking and genetic data exist in various regions of the Arctic (e.g., in the eastern Arctic of Canada), hair snaring can be used as supplementary data. The methods we used can be employed at bowhead whale bone piles across the North Slope of Alaska to further characterize the influence of these food subsidies on the status of the SB subpopulation. Similarly, the method could be used across the Arctic to provide samples to study polar bear abundance and behavior at localized remains of marine mammals on shore, whether they are bone piles from traditional harvest or beach cast remains. For example, walrus carcasses resulting from trampling events at large terrestrial haulouts on Wrangel Island are frequently used by polar bears (Ovsyanikov et al. 1994, Kochnev 2002). Samples collected at such sites could be used to understand behavior and local abundance of polar bears, and could supplement larger-scale multi-source genetic capture–recapture studies (Boulanger et al. 2008). However, we do not envision the use of non-invasive hair snaring as the sole method for estimating abundance of an entire subpopulation as has been used for terrestrial ursids because polar bears are far-ranging and many subpopulations occur in remote, inaccessible areas. In addition, the sea ice habitat used by most polar bears is mobile. Nevertheless, non-invasive hair sampling can be conducted by local people (Van Coeverden De Groot et al. 2012), and the method has the potential to increase public participation in polar bear research and management in a community-based-monitoring framework (Peacock et al. 2011, Vongraven et al. 2012).

Our work suggests important refinements of the hair-snaring technique for polar bears in the Arctic. We suggest that our sampling success was improved by frequent (daily to multiple times a week) snare maintenance. During periods of high bear use (>2 samples/day), wire was often stretched, snapped,
or dislodged, and occasionally poles were bent. Wire became more brittle as temperatures declined; particularly when temperatures fell below -40°C. The need for regular maintenance suggests hair snaring may have limited application in regions where snares cannot be checked frequently. Also, the MAPP™ gas torch was more useful as propane became unusable at approximately -10°C despite attempts to keep the cylinders warm.

Our data suggest the bowhead whale bone pile at Point Barrow, and by extension, bone piles on the North Slope of Alaska, are used by a large proportion of polar bears from the surrounding region. Although most detected bears used the bone pile once or twice, some individuals spent much more time at the bone pile, suggesting variation in how individual bears use the bone pile. Bentzen et al. (2007) also showed variation in polar bear use of the bone pile: winter diets varied from 0–26% bowhead whale. Although these bone piles may benefit some polar bears, they may also have a negative influence. For example, bone piles affect the distribution of polar bears while on land (Schliebe et al. 2008) and likely attract bears to land as they constitute a reliable food source. Thus, increased use of these areas by bears near communities may increase the likelihood of being killed for subsistence harvest or in defense of life and property by Native hunters, residents, tourists, or industry workers. Changes in bowhead whale migration or in autumn whaling conditions as a result of climate change in the future (Ashjian et al. 2010) have the potential to exacerbate such conflicts in the case of reduced whale harvest. It will be important for wildlife managers and the communities that rely on bowhead whales and polar bears for subsistence to consider both the positive and negative aspects of the bone piles to polar bear ecology and management.

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Literature cited


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Appendix

Within the dataset of 37 previously-known individuals sampled at the bone pile, members belonging
to at least 5 and possibly 6 family groups with dependent young were identified (Fig. 3).

- In one case (a known mother with a known yearling), both the yearling and mother were sampled at the bone pile (each once on two different days).
- In the second case, only a known cub-of-the-year was sampled (on one day), but his known mother and male sibling were not sampled.
- In the third case, a 21-year old known mother was sampled, and likely also one of her two yearlings (i.e., an unknown female that was related to the mother at \( r = 0.5 \) was also sampled): yearlings were observed in October 2010 when the mother was biopsied, but the yearlings were not biopsied and so were not ‘known’ genetically.
- In the three other instances, the known mothers were encountered, but two cubs-of-the-year, two yearlings, or one yearling (all known genetically), respectively, which presumably were accompanying their mothers (i.e., recently observed together), were not sampled.

To further characterize the relatedness of bears that used the bone pile, we looked at all individuals identified by coancestry as related (\( r \geq 0.46 \)) which we used as evidence of sibling–sibling or parent–offspring relationships (13 dyads from 97 individuals, a–m).

- Four dyads (a–d) contained previously unknown animals, and thus no additional insight on family relationships can be drawn.
- Two dyads (e, f) involved pairs of previously-known individuals; both were pairs of different-aged adult males, which were only captured as adults, and therefore their relationships are unknown. These two pairs may be pairs of brothers; however, the age difference between one of the pairs (8 years) is large enough that the pair might be father–offspring.

The 7 remaining dyads (g–m) consisted of one previously-known polar bear and one previously-unknown polar bear.

- One dyad (g) could represent a father–offspring or brother–brother relationship. Another adult male polar bear is in two other pairings (h, i), each with a female polar bear (these relationships could be offspring–mother or sibling–sibling).
- One dyad (j) of an adult female which, based on typical polar bear reproduction cycles, could have had a cub-of-the-year in the autumn of 2010 and was paired with a female; this relationship is not necessarily mother–offspring, but could also be a sibling–sibling relationship.
- One dyad (k), also mentioned above is likely a mother–cub relationship; one 21-year old female was seen two months prior to the bone pile study with two yearlings, thus her pair in this dataset likely represents the capture of one of her yearlings (a female). A 14-year-old mother (member of dyad j) was paired with a previously unknown male in our data set; however, this unknown male is not her cub that was captured (and genetically identified) with her in 2009. Thus this dyad (l) may represent an adult family member (i.e., the unknown bear is an older cub or her father) or a sibling–sibling relationship.
- A final dyad (m) relates a 19-year old male with this same unknown male bear (in dyad l) The only relationship possible for this pair (m) is of a father–offspring relationship (not sibling), as the 19-year old adult male was too old to be an offspring of the cub’s mother (14 years old; from dyad l).