



Coupling GPS tracking with dive behavior to examine the relationship between foraging strategy and fine-scale movements of northern fur seals

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ABSTRACT: The foraging strategies of diving marine species are often categorized into 3 fundamental groups (epipelagic, mesopelagic, and benthic foraging) based on diving, habitat use, and diet studies. Because these foraging strategies are influenced by the distribution and behavior of the prey being targeted, we would expect search behavior and space use to differ depending on the strategy employed. Since northern fur seals *Callorhinus ursinus* display both epipelagic and benthic foraging strategies, they were an ideal model to test the hypothesis that fine-scale movement and space-use patterns will vary when animals use markedly different foraging strategies. Dive bouts were characterized into foraging strategies based on numerous dive parameters (depth, duration, etc.). For each strategy, we compared movement patterns (e.g. transit rate and path straightness) and space use (area-restricted search [ARS] zones) around St. Paul Island, Alaska, USA. Nearly all dive parameters were significantly different between foraging strategies (epipelagic vs. benthic). In addition, epipelagic bouts were more sinuous and covered a greater total distance than benthic bouts. However, the greater distances traveled in epipelagic bouts were due to longer bout durations, as transit rates were not different between the 2 strategies. On average, <2 ARS zones were identified per trip, and the characteristics of epipelagic and benthic ARS zones were not different. By combining dive behavior with precise at-sea locations, this study has provided a greater understanding of the fine-scale foraging behavior of northern fur seals. Monitoring changes in foraging behavior over time and comparing behavior among populations with differing population trajectories may provide more clues as to why northern fur seal numbers on St. Paul Island continue to decline.

KEY WORDS: *Callorhinus ursinus* · Fractal landscape analysis · Area-restricted search

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INTRODUCTION

Understanding how animals exploit their environment is central in ecology and necessary for the protection of species and critical resources. Through a combination of information that has been gathered about diving behavior, habitat use, and diet, the foraging strategies of many air-breathing marine species have been classified into 3 fundamental patterns. Epipelagic foragers target prey within the water column at depths ranging from 0 to 200 m, whereas mesopelagic species

forage between ~200 and 1000 m. In contrast, benthic or demersal predators feed on species that are found at or near the bottom, targeting continental shelf or shelf break regions of varying depths (Tremblay & Chérel 2000, Costa & Gales 2003, Villegas-Amtmann et al. 2008).

Within marine vertebrate classes and families, species that demonstrate each of the disparate foraging strategies have been identified. For example, both gentoo *Pygoscelis papua* and Adélie penguins *P. adeliae* feed on epipelagic prey, whereas yellow-eyed

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penguins *Megadyptes antipodes* forage benthically (Croxall et al. 1988, Naito et al. 1990, Mattern et al. 2007). Leatherback sea turtles *Dermochelys coriacea* feed on epipelagic, soft-bodied invertebrates such as jellyfish, whereas mature green turtles *Chelonia mydas* feed primarily on benthic sea grasses and algae (Hochscheid et al. 1999, Hays et al. 2006). Different species of pinnipeds also exhibit each of the fundamental foraging strategies. For example, subantarctic fur seals *Arctocephalus tropicalis* feed epipelagically, adult female northern elephant seals *Mirounga angustirostris* feed mesopelagically, and walrus *Odobenus rosmarus* feed benthically (Wiig et al. 1993, Georges et al. 2000, Le Boeuf et al. 2000). A few studies have even shown that intraspecific foraging strategies can vary between sexes (e.g. in northern elephant seals; Le Boeuf et al. 2000) or individuals (e.g. in Galapagos sea lions *Zalophus wollebaeki*; Villegas-Amtmann et al. 2008).

Given that these foraging strategies are partly shaped by variability in the distribution and behavior of prey, we would expect that, as predators target different prey species, search behavior and space use will differ depending on the foraging strategy employed. However, for many air-breathing marine species, it has been difficult or impossible to examine the fine-scale movements of individuals during distinct foraging periods due to limitations in position accuracy and frequency when using traditional Argos satellite tracking (Bradshaw et al. 2004). Nevertheless, recent advances in GPS technology are helping to overcome these challenges by providing the ability to examine the detailed movement patterns of marine species (Mattern et al. 2007, Schofield et al. 2007, Kuhn et al. 2009).

Previously, for species that spend much of their time underwater, traditional GPS instruments were ineffective for obtaining at-sea locations (Sisak 1998, Jay & Garner 2002). With the development of Fastloc GPS technology (Bryant 2007), it is now possible to obtain the more accurate, consistently acquired locations that are currently available for tracking animals in the terrestrial environment (Mattern et al. 2007, Hazel 2009, Kuhn et al. 2009). Instruments that are equipped with Fastloc GPS are able to obtain satellite signals in milliseconds and can typically determine locations with an error of <50 m (Bryant 2007). During stationary tests, Fastloc GPS instruments determined positions with an average error of 40 ± 80.5 m, and 95% of locations were <102 m from the true location (Hazel 2009). In contrast, even the highest quality locations reported by Argos can have errors of up to 250 m (Argos 2006). Finally, these instruments are not limited to obtaining locations only when Argos satellites are overhead, resulting in a much higher temporal frequency of locations (Kuhn et al. 2009).

Given that fine-scale movement patterns of air-breathing marine species can now be measured, it is finally possible to examine how movement patterns vary between differing foraging strategies. Adult female northern fur seals *Callorhinus ursinus*, Linnaeus 1758 are an ideal model to examine these differences without the confounding factors of sex, location, or season because they display both benthic and epipelagic foraging behaviors (Gentry et al. 1986, Goebel 1998). In addition, identifying patterns of movement that are related to these different foraging strategies is essential to understanding northern fur seal habitat use. This species was listed as 'depleted' under the Marine Mammal Protection Act in 1988, and the largest breeding colony in Alaska (St. Paul Island) has been declining at a rate of $\sim 6.0\% \text{ yr}^{-1}$ (Towell et al. 2006, NMFS 2007). A greater understanding of foraging behavior and habitat use is vital to provide guidance for future conservation efforts and management of this species. Therefore, the objectives of this study were to (1) describe and quantify the different foraging strategies used by female northern fur seals during the breeding season, and (2) examine how fine-scale movement (e.g. transit rate, distance traveled, path straightness) and space-use (e.g. area-restricted search zones) patterns vary when animals employ differing foraging strategies.

MATERIALS AND METHODS

Animal handling. Research was conducted from September to October 2007 and August to October 2008 at St. Paul Island, Alaska (USA; Fig. 1). Forty-two adult female northern fur seals were instrumented at 2 rookeries: Reef ($n = 20$; 57.1°N , 170.3°W) and Vostochni ($n = 22$; 57.25°N , 170.1°W). Females that were observed to be nursing or calling for a pup were captured using hoop nets. Instruments were attached while animals were either physically restrained or sedated using gas anesthesia (isoflurane) that was administered with oxygen via a portable vaporizer. Each female was measured (± 1 cm) and weighed using a digital scale (± 0.1 kg). Animals were recaptured 6 to 52 d later, physically restrained for instrument removal, and reweighed prior to release.

Instrumentation. Each animal was equipped with an Mk10-AF tag (Wildlife Computers), which has an Argos satellite transmitter, Fastloc GPS, and time-depth recorder capabilities. Thirty of the females were also instrumented with an Mk10-L tag (Wildlife Computers), which only had time-depth recorder capabilities. The Mk10-AF tag used Fastloc technology to collect GPS locations at 15 min intervals when the animal was above the surface. Along with acquir-

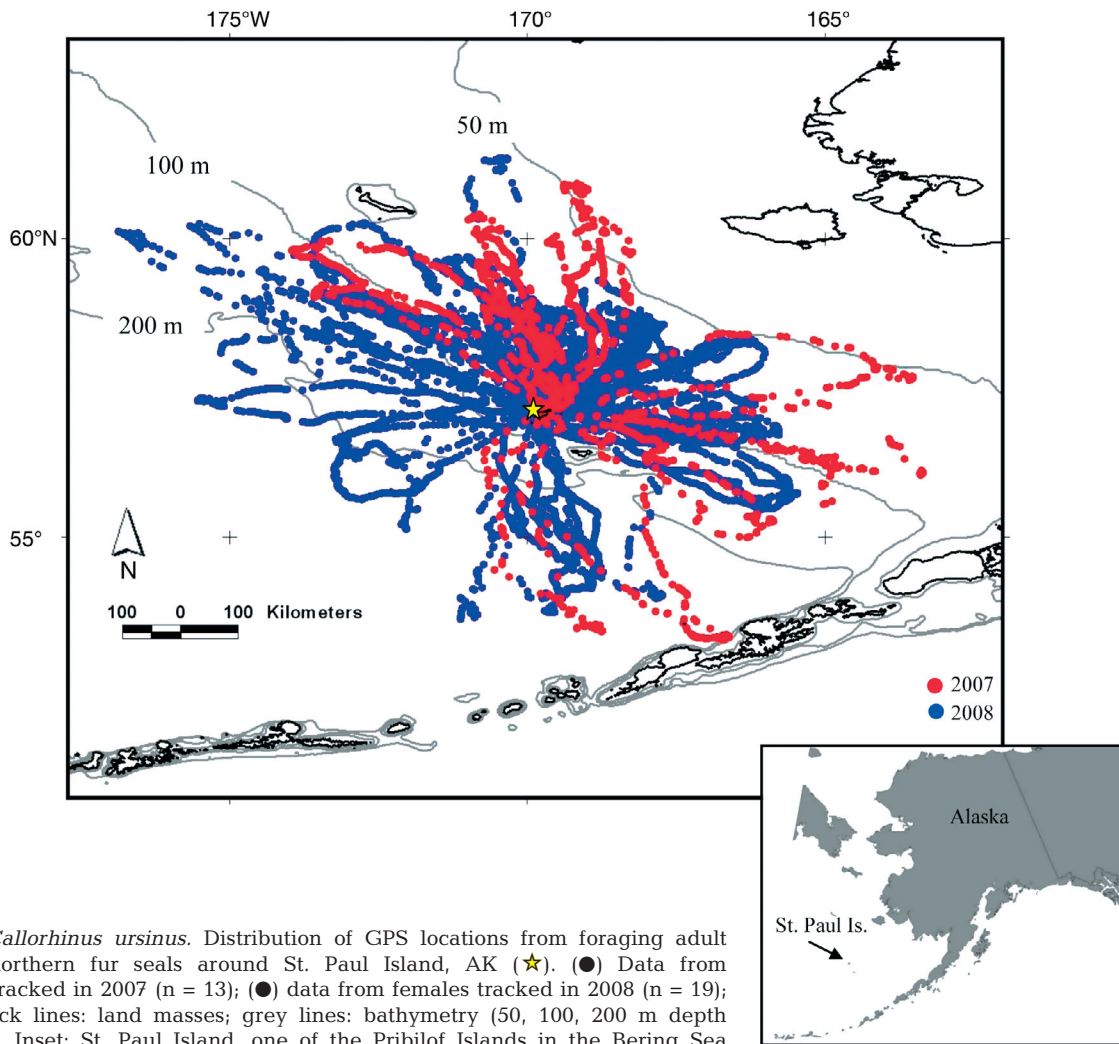


Fig. 1. *Callorhinus ursinus*. Distribution of GPS locations from foraging adult female northern fur seals around St. Paul Island, AK (★). (●) Data from females tracked in 2007 ($n = 13$); (●) data from females tracked in 2008 ($n = 19$); Solid black lines: land masses; grey lines: bathymetry (50, 100, 200 m depth contours). Inset: St. Paul Island, one of the Pribilof Islands in the Bering Sea

ing an Argos satellite position, the instrument was programmed to transmit a subset of the GPS data via the Argos system at the highest priority level. When archived GPS data were not recovered, transmitted GPS data were used.

In 2007, dive data from both Mk10-L and Mk10-AF instruments were sampled at 5 s intervals. In 2008, the Mk10-L and Mk10-AF tags sampled dive depth at 1 or 5 s intervals, respectively. Mk10-L tags were used in conjunction with stomach temperature telemeters (STT); however, only foraging trips where females no longer retained the STT were used for this study to reduce the possible influence of the STT on at-sea behavior. Finally, to facilitate instrument recovery, each female was equipped with a VHF tag (Advanced Telemetry Systems). Instruments were attached directly to the dorsal pelage using quick-set epoxy.

Dive data analysis. Dive data were processed using a purpose built zero-offset correction algorithm and analysis program written in MATLAB (IKNOS-DIVE,

IKNOS toolbox, Y. Tremblay unpubl.). A dive was defined by a minimum depth of 4 m and a minimum duration of 20 s. Maximum dive depth, dive duration, and post-dive surface interval were calculated for each dive. Bottom time was calculated as the time between the first and last inflection points at $>80\%$ of the maximum depth. Dive bouts were classified using log-survivorship curves based on the post-dive surface interval following the methods of Gentry & Kooyman (1986). An inflection point on the curve signals the duration between consecutive dives (termed the bout ending criterion, BEC) that, if surpassed was deemed to end a bout. A BEC was determined for each foraging trip. The minimum criterion for a bout was 4 dives >6 m, with the post-dive surface interval of each dive falling within the BEC. In addition, to examine movement parameters, bouts with <4 GPS locations were removed because we believe they did not provide enough movement information to test the study hypothesis.

The first goal of this study was to determine whether distinct foraging strategies are used by northern fur seals on St. Paul Island. Therefore, a hierarchical cluster analysis was used to classify dive behavior into foraging strategies using the Euclidean distance and average linking method. Bouts were independently clustered for each foraging trip. Input variables were the bout averages for dive depth (m), dive duration (s), bottom time (s), post-dive surface interval (s), descent rate (m s^{-1}), ascent rate (m s^{-1}), vertical excursions during the bottom phase (wiggles), total vertical distance covered during wiggles (m), bottom range (difference between minimum and maximum vertical distance during wiggles, m), and dive efficiency (bottom time/[dive duration + post-dive interval]). Additionally, SDs of dive duration, dive depth, bottom time, post-dive surface interval, total vertical distance, bottom range, and efficiency were used for classification.

To characterize the resulting groups of dive behavior, 2 independent variables were used: dive index (DI; dive depth/ocean depth) (Simmons et al. 2007) and intra-depth zone index (IDZ) (Tremblay & Cherel 2003). The location for the start of each dive was determined by interpolating GPS data to each dive (see details of interpolation in 'GPS tracking data analysis'). Ocean depth was calculated using 2 min gridded global relief data (ETOP02v2). The IDZ indexed the degree of differences between the maximum depths that were achieved on consecutive dives. The IDZ range was calculated as the maximum depth that was reached in the preceding dive $\pm 10\%$. An IDZ of 1 indicates that the dive fell within the IDZ range, while an IDZ of 0 indicates that a subsequent dive differed by $>10\%$ of the previous maximum depth. Based on the differences in the 2 independent variables and average dive characteristics, the resulting dive groups were classified as epipelagic, benthic, or mixed foraging strategies.

Finally, for each foraging strategy, we examined whether there were significant differences in bout start time in relation to time of day. Due to the significant change in daylight hours over the tracking period, bouts were classified as starting either during the day or during the night based on local sunrise and sunset times for each day. Based on previous descriptions of pinniped dive behavior, we predicted that epipelagic bouts would start more often during night time hours when diel vertically migrating prey move to shallow waters, whereas benthic dive bouts would occur throughout the day when animals target non-migrating prey on the bottom (Costa & Gales 2003, Ream et al. 2005, Villegas-Amtmann et al. 2008).

GPS tracking data analysis. Erroneous GPS locations were filtered based on a maximum transit rate of 2.5 m s^{-1} (Tremblay et al. 2006, Kuhn et al. 2009).

Tracks were interpolated using a hermite curve to associate each dive with a location based on the methods described by Tremblay et al. (2006). The sum of distances between locations was used to calculate total distance travelled (km). Average bout transit rate (km h^{-1}) was calculated by dividing the total distance travelled by the bout duration. A straightness index was used to determine the sinuosity of the path by dividing the straight-line distance between the bout start and end locations by the total distance travelled (Weimerskirch et al. 2002).

In addition to changes in movement patterns, periods of area-restricted search (ARS) are often used to identify foraging behavior and important feeding habitats (Kareiva & Odell 1987, Fauchald & Tveraa 2003, Tremblay et al. 2007, Weimerskirch et al. 2007). Periods of ARS were determined using the fractal landscape method (Tremblay et al. 2007). First, speeds between successive interpolated locations were ranked. The distance traveled between locations was determined for the lowest third of the ranked speeds, and the speed that was associated with the longest distance was selected. This distance was then used to segment the seal's track around each location. The fractal dimension (fractal D) for each segment was calculated based on the area covered by the seal in relation to the segment size. An automatically detected threshold value for each track was used to distinguish between fractal peaks and minor oscillations. Finally, the characteristics of each peak were determined, including the ARS duration and search area (see Tremblay et al. 2007 for more details). An index of search intensity (area peak) was calculated based on both time in ARS and distance covered. An index of track sinuosity was determined based on mean fractal D for each ARS period. Because ARS zones can include one or more bouts, these zones were compared based on the dominant strategy that was used during the ARS period.

Statistical analysis. Summary data are reported as mean \pm SE, unless otherwise noted. Statistical analysis was conducted using R (R Foundation; www.r-project.org). Bout classification and movement parameters were compared using linear mixed models. Random factors for the models included animal ID and foraging trip within each animal. Year was included as a fixed factor in the models. Sampling frequency ('fast' = 1 s, 'slow' = 5 s) was also included as a fixed factor to test the influence of sampling rate on dive data. The most appropriate model was chosen from a combination of all random and fixed factors based on the lowest Bayesian information criterion (BIC; Burnham & Anderson 2002). Because data for path straightness, IDZ, and DI were proportion data that did not show a normal distribution after

transformation, models were fitted using Markov Chain Monte Carlo (MCMC) simulations using the R2jags package (Smithson & Verkuilen 2006). A total of 10 000 iterations were used with a burn-in of 2000. The mean and 95% CIs were calculated for the random and fixed factors to determine differences between bout types. Ranges presented are the minimum and maximum of the female averages. Data that were non-normal were \log_{10} or square-root transformed. All contrasts were considered significantly different at $p < 0.05$.

RESULTS

In 2007, 5 fur seals were instrumented with an Mk10-AF tag and 8 were instrumented with both an Mk10-AF and an Mk10-L tag. One seal lost the Mk10-L tag prior to recapture. In 2008, 7 seals were equipped with an Mk10-AF and 22 seals were equipped with both an Mk10-AF and an Mk10-L tag. In 2008, there was significant instrument loss and it was discovered at recapture that the tracking instruments had separated from the glue base. This resulted in recaptures of 12 seals with both Mk10-AF and Mk10-L tags, 6 seals with only Mk10-AF tags, 3 seals with only Mk10-L tags, and 8 seals with no tags. Of the 8 seals that did not retain their instruments, 4 were relocated with the VHF transmitter still attached. Because no markers were used to identify individuals, we were unable to determine whether seals with glue patches that were resighted on the rookery were individuals that lost all instruments or were previously recaptured seals.

For 1 seal that was recaptured with only an Mk10-L (2008), no GPS data were transmitted and the dive data were therefore excluded from this analysis. An additional seal from 2008 was removed from the analysis as an outlier: she made 2 trips that were >19 d, and because she was not seen with a pup at recapture, it was presumed that she lost her pup. Although the Mk10-AF instruments were programmed to transmit dive data, these data were received in a histogram format and therefore could not be used for the analysis of fine-scale behavior.

In total, 71 foraging trips with matching GPS and dive data were obtained from 32 ind. At instrumentation, mean fur seal mass was 39.3 ± 0.6 kg, and females gained an average of 1.1 ± 0.7 kg over the tracking period (24.3 ± 2.8 d). In cases where duplicate dive data was acquired from an individual (both Mk10-AF and Mk10-L instruments), the highest resolution data was used for analysis (1 s sampling rate). As a result, dive data were analyzed from 21 Mk10-L tags (2007: 7, 2008: 14) and 11 Mk10-AF tags (2007: 6, 2008: 5).

Dive data

Average BEC was 31.1 ± 0.5 min with a range of 21.7 to 41.2 min; if a subsequent dive did not occur within this amount of time, a dive bout was considered over. On average, 14.7 ± 0.9 bouts were recorded per trip (range: 6 – 25) and these bouts lasted for 5.4 ± 0.1 h. Cluster analysis of the dive bouts resulted in 3 distinct groups of dive behavior that were classified as epipelagic bouts ($n = 613$), benthic bouts ($n = 212$), and mixed dive bouts ($n = 27$; Fig. 2). Visual examination of the mixed group showed that these dive bouts began as either epipelagic or benthic and then switched part way through (Fig. 2). Therefore, these mixed dive bouts were not included in subsequent analysis. Six females each displayed only 1 foraging strategy (100% epipelagic, $n = 5$; 100% benthic, $n = 1$), whereas the rest showed some combination of the two, both within and among trips (average proportion of epipelagic bouts for each female: 0.79 ± 0.02). After selecting bouts that met the minimum GPS location criterion, 412 bouts remained for the analysis of movement patterns (342 epipelagic bouts, 70 benthic bouts).

All of the dive parameters were significantly different between epipelagic and benthic bouts except for bottom time and the number of wiggles in the bottom phase (Table 1); for all parameters there was an effect of one or more of the random variables examined (Table 2). During benthic bouts, dives were over 3× deeper, nearly 2× longer, and had longer post-dive surface intervals than during epipelagic bouts (Table 1). Dive efficiency in epipelagic bouts was also 3× higher than in benthic bouts. Although the number of wiggles per dive did not differ, females covered a greater overall bottom range during the bottom phase of epipelagic bouts (Table 1). A detectable effect was found for both foraging trip and animal ID for dive depth, duration, bottom time, post-dive interval, wiggles in the bottom phase, dive efficiency, and dives per bout (Table 2). Animal ID showed a measurable effect for total vertical distance and foraging trip showed an effect for bottom range (Table 2). Sampling rate showed a detectable effect for the number of dive wiggles and the total vertical distance. Year was not a significant factor for any of the dive variables examined.

The IDZ of 51% (Table 1) that was obtained for epipelagic bouts was indicative of extensive variation in dive depths, consistent with fur seals tracking the movement of vertically migrating prey (Kooyman & Goebel 1986, Goebel 1998). In contrast, benthic bouts had significantly higher IDZ (77%; Table 1), showing a high association in dive depth between sequential dives. The differences between DIs show an even stronger distinction between the 2 bout types, with epipelagic bout mean values being only 0.28 ± 0.01

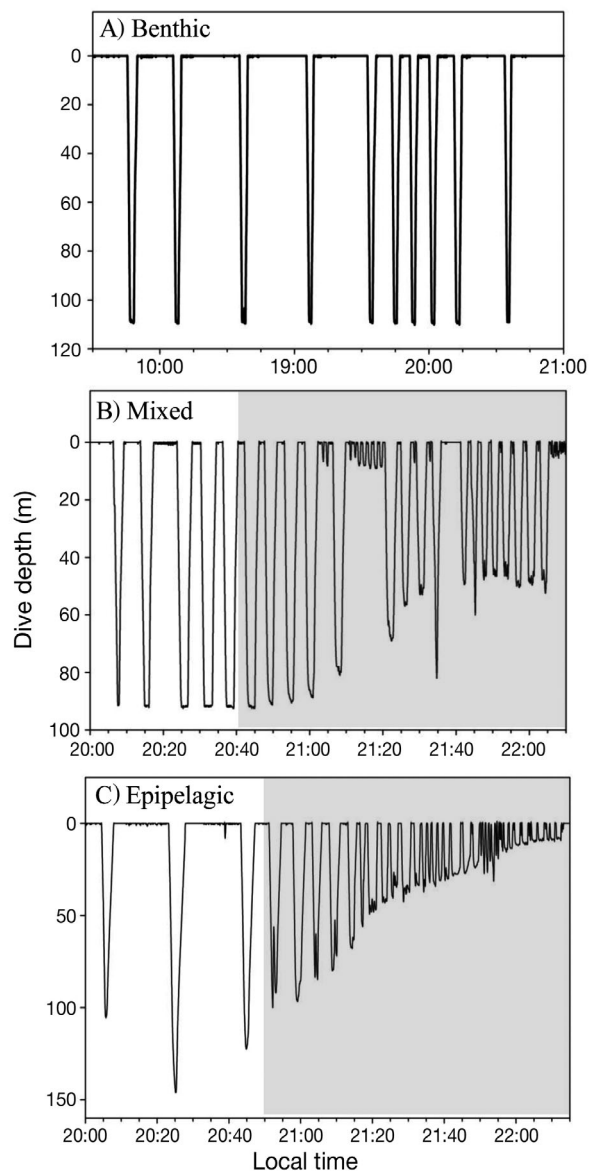


Fig. 2. *Callorhinus ursinus*. Examples of the 3 foraging strategies that were observed in distinct dive bouts. (A) Benthic diving was identified by repeated dives to the same depth, usually occurring during daylight hours. This is supported by the high dive index (DI) of 0.80 observed for this bout, which means dive depths were close to the ocean floor at this location. (B) Mixed diving was identified by a switch in diving strategy during a single bout. The mixed diving strategy occurred in <4% of all bouts. The switch regularly occurred just after sunset (shaded area). The average DI for mixed dive bouts was 0.47 ± 0.03 , which was midway between the DI for benthic and epipelagic dive bouts (see Table 1). (C) Epipelagic dive bouts were identified by shallow diving that had distinct variation in dive depths as measured by a lower intra-depth zone index (IDZ; see Table 2). This epipelagic dive bout began just prior to sunset (shaded area) and had a DI of 0.28, indicating that the dives were not near the ocean floor. Note the decreasing trend in dive depth just after sunset, which likely resulted from this female tracking prey as it vertically ascended with decreasing light levels

and benthic bout mean values being significantly higher at 0.83 ± 0.02 (Table 1). As was found with the previously described dive parameters, DI showed a detectable effect of animal ID, whereas IDZ showed an effect of year (Table 3). Finally, nearly equal numbers of epipelagic bouts occurred during the day and at night (45 and 55%, respectively), whereas 78.6% of the benthic bouts began during the daytime hours.

Tracking data

Archived GPS tracks were acquired from 30 females (68 foraging trips), and transmitted GPS tracks (GPS-t) were used for the remaining 2 females (3 foraging trips). A total of 14 169 GPS locations were acquired and 13 488 remained after filtering (Fig. 1). This resulted in 192.2 ± 13.5 locations trip⁻¹ and 26.9 ± 1.8 locations d⁻¹ at sea. The average time between GPS locations was 1.3 ± 0.1 h, with an average maximum for all females of 10.6 ± 1.1 h between locations. Although GPS-t data are generally acquired at a lower frequency than archived GPS data (Kuhn et al. 2009), removing bouts with <4 GPS locations rendered the temporal resolution of the GPS-t data (average time between locations: 1.6 ± 0.04 h) within the range acquired by archived GPS data.

Trip duration averaged 7.4 ± 0.2 d and ranged from 3.7 to 11.4 d. Trip durations were significantly shorter in 2008, and there was a detectable effect of individual (difference estimate = -1.7 , SE = 0.61, $Z = -2.8$; animal ID: variance = 1.9, SD = 1.4). These foraging trips covered areas around St. Paul Island that were within the 50 m depth contour to off-shelf regions over 1000 m deep (Fig. 1).

Females travelled almost 2× farther during epipelagic bouts than during benthic bouts (Tables 1 & 2); however, epipelagic bouts were over 2× longer in duration (Table 1). When transit rates during a bout were compared, the difference was not significant. Both foraging strategies had high straightness indices; nevertheless, seals travelled in a slightly straighter path during benthic foraging (Table 1, Fig. 3). For all movement parameters except path straightness, there was a detectable effect of one or more of the random factors (Tables 2 & 3).

The number of ARS periods detected was low, averaging 1.3 ± 0.1 per foraging trip; during 14 trips, no ARS zones were detected ($n = 74$, range: 0 to 3 trip⁻¹; Fig. 4). Time in ARS zones had a wide range of 0.08 to 3.3 d and females travelled an average of 36.8 ± 3.1 km while in these ARS zones (range: 5.8 to 128.5 km; Table 4). Sixty-five ARS periods were identified with predominantly epipelagic bouts and 8 with predominantly benthic bouts. An additional ARS period had no

Table 1. *Callorhinus ursinus*. Average (\pm SE) dive and movement parameters for each foraging strategy. Comparisons were made between the 2 foraging strategies using a linear mixed model or Markov Chain Monte Carlo (MCMC) simulations (path straightness, IDZ and DI, see 'Materials and methods' for details). Parameters with CIs that do not include 0 denote significant differences (*). For calculation of dive and movement parameters see 'Materials and methods'

Parameters	Epipelagic	Benthic	95 % CI
Dive depth (m) ^a	22.0 (0.62)	85.4 (2.92)	-0.58, -0.48*
Dive duration (s) ^a	97.7 (1.99)	187.5 (4.01)	-0.30, -0.22*
Bottom time (s) ^a	49.9 (1.23)	54.9 (2.13)	-0.09, 0.02
Post-dive interval (s) ^a	100.5 (5.0)	768.8 (68.8)	-1.4, -1.8*
Wiggles in bottom phase ^a	4.7 (0.14)	5.9 (.036)	-0.03, 0.05
Total vertical distance (m) ^a	7.7 (0.23)	6.7 (0.39)	0.04, 0.15*
Bottom range (m) ^a	3.0 (0.070)	2.9 (0.24)	0.02, 0.13*
Dive efficiency ^a	0.32 (0.0058)	0.10 (0.0096)	0.16, 0.22*
Dives per bout	201.0 (8.0)	21.8 (2.5)	5.9, 8.7*
Intra-depth zone index (IDZ)	0.51 (0.01)	0.77 (0.02)	0.8, 1.4*
Dive index (DI)	0.28 (0.01)	0.83 (0.02)	1.3, 2.1*
Bout duration (h)	8.2 (0.2)	3.8 (0.3)	0.24, 0.38*
Total distance traveled (km)	31.4 (1.1)	15.8 (1.3)	0.78, 1.68*
Transit rate (km h ⁻¹)	4.02 (0.10)	4.26 (0.27)	-0.22, 0.06
Straightness	0.82 (0.012)	0.93 (0.013)	0.1, 0.7*

^aDive parameters used for cluster analysis

bouts that matched our diving criteria although diving occurred. There were no significant differences for the number of bouts between epipelagic and benthic ARS periods ($F_{1,71} = 2.2$, $p = 0.14$; Table 4). In addition, there were no significant differences for many of the ARS period characteristics measured (duration: $F_{1,71} = 0.006$, $p = 0.93$; area of circle: $F_{1,84} = 0.002$, $p = 0.89$; area peak: $F_{1,84} = 0.07$, $p = 0.40$). However, epipelagic ARS periods had a significantly higher track sinuosity (mean fractal D: $F_{2,70} = 3.9$, $p = 0.02$) and there was an effect of year, with the trend of 2008 being higher than that of 2007 ($p = 0.05$).

DISCUSSION

Dive behavior

Cluster analysis of the dive behavior of northern fur seals revealed flexibility in foraging strategies within the St. Paul Island population. Although 3 strategies were identified, >96% of the bouts were characterized as epipelagic or benthic diving and most females employed both of these strategies within a foraging trip. Although flexibility in foraging behavior was observed, foraging predominantly occurred in the epipelagic zone (>75% of all bouts). The ability to alternate between foraging strategies is not unique to northern fur seals and has been demonstrated in a variety of marine predators, including seabirds (Croxall et al. 1988, Tremblay & Cherel 2000), marine mammals (Mattlin et al. 1998, Burns et al. 2004), and sea turtles (Hawkes et al. 2006). In some species, these

changes are a response to aging/growth or seasonal variation. For example, loggerhead turtles *Caretta caretta* show a transition from epipelagic to benthic foraging with increased age (Hawkes et al. 2006, Hatase et al. 2007). Seasonal changes from summer epipelagic foraging to winter benthic foraging were also documented in New Zealand fur seals *Arctocephalus forsteri* (Mattlin et al. 1998) and crabeater seals *Lobodon carcinophagus* (Burns et al. 2004). The change in behavior for the New Zealand fur seals was a result of targeting different prey species, whereas Burns et al. (2004) suggested that crabeater seals were following the movements of a single prey species, krill *Euphausia superba*, toward the seafloor in winter.

For a limited number of marine species, flexibility in foraging behavior within a single trip has also been described. Both rockhopper *Eudyptes chrysolophus* and gentoo penguins show periods of epipelagic and benthic diving within a foraging trip, and the proportion of each strategy varied among individuals (Croxall et al. 1988, Tremblay & Cherel 2000). As in New Zealand fur seals, the shift in foraging strategy for gentoo penguins was likely a result of transitioning from shallow-water, vertically migrating prey to benthic fish species (Croxall et al. 1988). The flexible behavior of northern fur seals may result from a shift in the targeted age class of the primary prey species, and/or targeting different prey species all together.

On St. Paul Island, the primary prey for adult female fur seals is walleye pollock *Theragra chalcogramma*, although other epipelagic and benthic prey

Table 2. *Callorhinus ursinus*. Linear mixed model comparisons for dive and movement parameters. For each selected model, both the fixed and random effects results are presented. ID is a unique value for each tracked animal and TripID refers to each foraging trip. Year and dive sampling rate (fast = 1 s, slow = 5 s) were tested as a fixed effect for all models

	Fixed effect			Random effect			
	Estimate	SE	Z	TripID		ID	
				Variance	SD	Variance	SD
Dive depth (log, m)							
Benthic	1.81	0.040	45.3	0.012	0.11	0.022	0.15
Epipelagic	-0.53	0.027	-19.8				
Dive duration (log, s)							
Benthic	2.21	0.030	70.4	0.0081	0.090	0.014	0.12
Epipelagic	-0.26	0.021	-12.2				
Bottom time (log, s)							
Benthic	1.63	0.041	40.4	0.012	0.11	0.019	0.14
Epipelagic	-0.036	0.030	-1.2				
Post-dive interval (log, s)							
Benthic	6.0	0.12	50.2	0.080	0.28	0.16	0.40
Epipelagic	-1.6	0.091	-17.5				
Wiggles in bottom phase (log)							
Benthic	0.41	0.04	9.7	0.0038	0.061	0.018	0.13
Epipelagic	0.009	0.021	0.4				
Fast sampling rate	0.31	0.055	5.7				
Total vertical distance (log, m)							
Benthic	0.65	0.039	17.0			0.018	0.14
Epipelagic	0.096	0.029	3.3				
Fast sampling rate	0.17	0.038	4.4				
Bottom range (log, m)							
Benthic	0.37	0.030	12.7	0.012	0.11		
Epipelagic	0.073	0.029	2.5				
Dive efficiency							
Benthic	0.11	0.018	6.3	0.0022	0.047	0.0037	0.061
Epipelagic	0.19	0.014	14.1				
Dives per bout (sqrt)							
Benthic	6.3	0.79	1.8	3.4	1.8	4.2	2.1
Epipelagic	7.3	0.69	10.7				
Bout duration (log, h)							
Benthic	-0.83	0.035	-23.7	0.009	0.09		
Epipelagic	0.31	0.036	8.5				
Total distance (sqrt, km)							
Benthic	4.18	0.25	16.8			0.44	0.67
Epipelagic	1.23	0.23	5.4				
Transit rate (sqrt, km h⁻¹)							
Benthic	1.36	0.07	20.4	0.032	0.18	0.20	0.44
Epipelagic	-0.08	0.069	-1.1				

Table 3. *Callorhinus ursinus*. Results of Markov Chain Monte Carlo (MCMC) comparisons for dive and movement parameters. Comparisons were made between epipelagic and benthic dive bouts, and both fixed and random effects were included. ID is a unique value for each tracked animal and TripID refers to each foraging trip. In addition, year was tested as a fixed effect. Parameters with CIs that do not include 0 denote significant differences (*)

	Type			Year			ID			TripID		
	Mean	SD	95% CI	Mean	SD	95% CI	Mean	SD	95% CI	Mean	SD	95% CI
Intra-depth zone index (IDZ)	1.07	0.16	0.8, 1.4*	0.38	0.19	0.02, 0.8*	0.076	0.059	0.0, 0.2	0.073	0.054	0.0, 0.2
Dive index (DI)	1.7	0.18	1.3, 2.1*	-0.04	0.24	-0.5, 0.4	0.37	0.11	0.2, 0.6*	0.09	0.07	0.0, 0.3
Straightness	0.34	0.16	0.1, 0.7*	-0.13	0.19	-0.5, 0.3	0.085	0.06	0.0, 0.24	0.065	0.05	0.0, 0.18

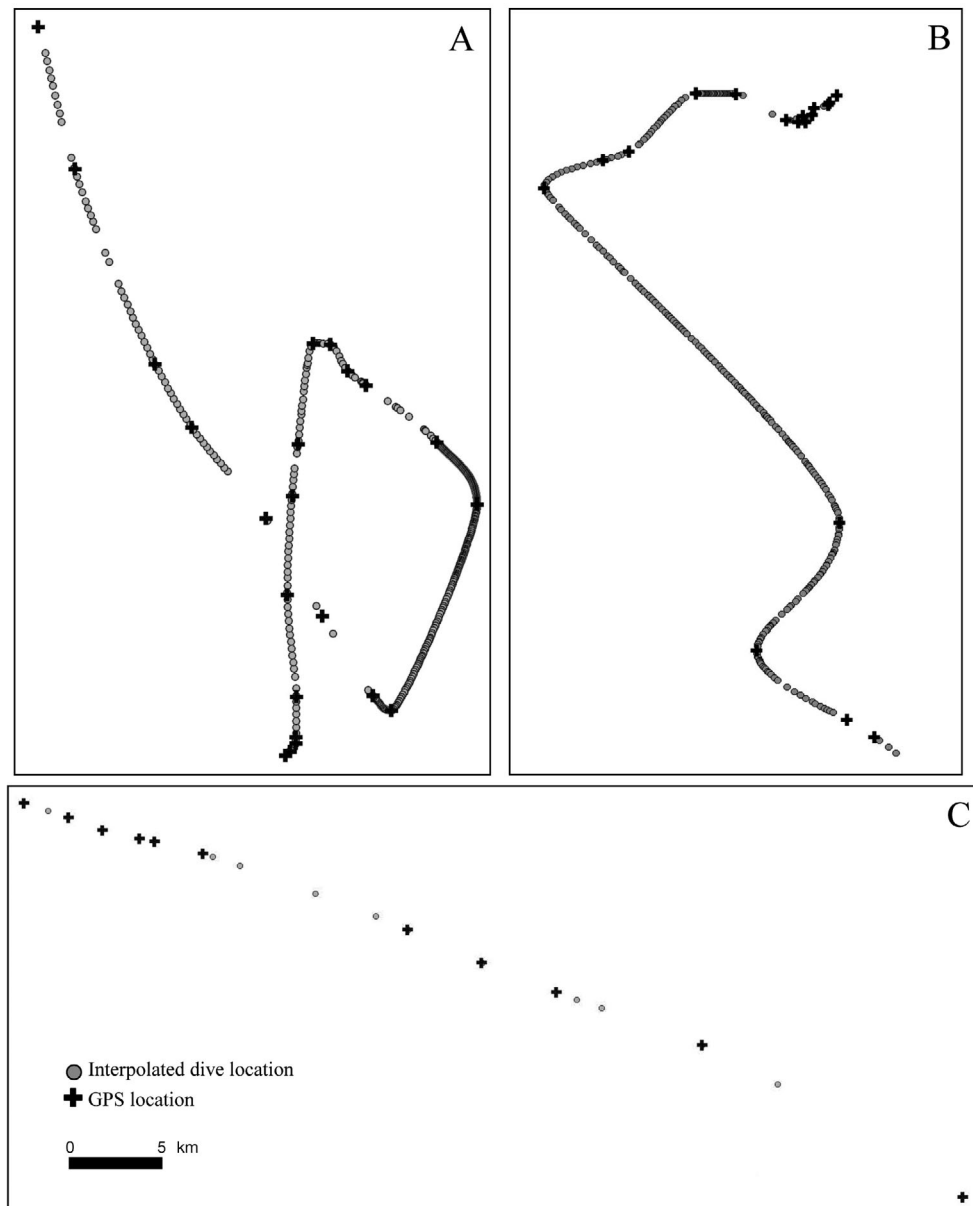


Fig. 3. *Callorhinus ursinus*. Examples of variation in path straightness for females displaying (A,B) epipelagic and (C) benthic dive bouts

species are also important (Antonelis et al. 1997, Gudmundson et al. 2006, Zeppelin & Ream 2006). The proportion of the pollock in the diet varies among rookeries, ranging from 64 to 84% frequency of occurrence (Zeppelin & Ream 2006). At the 2 rookeries used during this study, pollock frequency of occurrence was 66 and 64% (Vostochni and Reef, respectively; Zeppelin & Ream 2006) and the next highest ranked species at both sites had just over 11% frequency of occurrence (Vostochni: Pacific sand lance *Ammodytes hexapterus*; Reef: squid, *Gonatopsis borealis* and/or *Berryteuthis magister*). Previous

work has shown that in addition to differences in diet, females from these rookeries also tend to travel in different directions when foraging, resulting in habitat segregation between rookeries (Robson et al. 2004). Interestingly, although these patterns of habitat segregation appeared to be maintained by the females in this study, equal proportions of benthic and epipelagic dive bouts were recorded from females instrumented at each location (epipelagic bouts: Reef = 0.84, Vostochni = 0.83). This similarity in dive behavior may be due to the equally high proportions of pollock in the diet at each rookery.

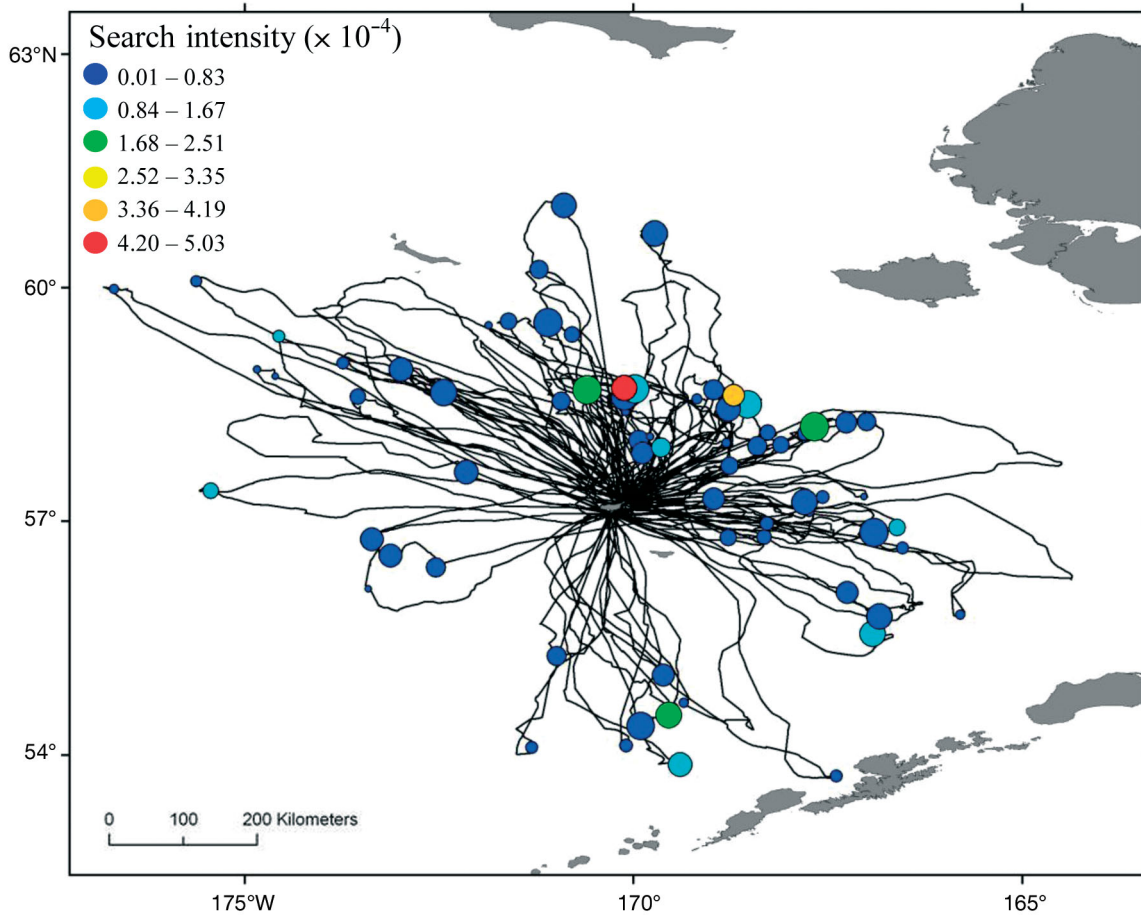


Fig. 4. *Callorhinus ursinus*. Foraging tracks (black lines) around St. Paul Island, AK, with periods of area-restricted search (ARS) behavior highlighted ($n = 32$ females, 71 trips). The size of each circle represents the area of the circle containing the ARS period. Colors signify an index of the search intensity (see Tremblay et al. 2007 for more details). ARS periods occurred both along the foraging track and at the distal end and varied in both size and intensity throughout the foraging range

A combination of scat and regurgitate analyses from both rookeries showed that the age classes of pollock that were consumed ranged between 0 and 5 (or greater, Gudmundson et al. 2006). Because the vertical distribution and behavior of pollock vary with age (Bailey 1989), it may be possible to link fine-scale foraging behavior with the targeting of differing pollock age classes. For example, juvenile pollock tend to remain above the thermocline (<40 m) and migrate into shallow surface waters during the night as they follow vertically migrating zooplankton (Incze et al. 1988). This closely matches the shallow night diving that was observed in female fur seals during epipelagic bouts. Foraging on juvenile pollock could also explain the high percentage of epipelagic bouts during the day, as females may be targeting concentrated groups of fish at the thermocline. In contrast, adult pollock remain at or near the sea floor (Francis & Bailey 1983, Bailey 1989), which would explain the deeper benthic diving.

At Vostochni rookery, Pacific sand lance is also an important prey species that uses both midwater and benthic habitats (Eschmeyer et al. 1983, Zeppelin & Ream 2006), similar to pollock, which could also explain the alternative foraging strategies.

Future studies linking individual diet with at-sea behavior will be necessary to directly relate differences in fine-scale foraging behavior to prey selection in this species. A recent study of Antarctic fur seals *Arctocephalus gazella* found that the accuracy of foraging habitat models increased significantly when individual diet was incorporated in the models (Casper et al. 2010). For the declining northern fur seal population on St. Paul Island, a similar study integrating diet and fine-scale movements could aid in the interpretation of at-sea behavior and help define critical foraging habitat. In addition, by creating foraging habitat models that take into account rookery-level differences in diet, it may be possible to make unique predictions

Table 4. *Callorhinus ursinus*. Average (\pm SE) characteristics for northern fur seal area-restricted search (ARS) periods consisting primarily of epipelagic and benthic dive bouts. Because ARS zones included >1 bouts, these zones were classified based on the dominant foraging strategy used during each ARS period. Count: no. of bouts per ARS period, ranging from 0–9; Area above peak: an index to measure ARS intensity (see Fig. 4). Mean fractal D (an index of the sinuosity of the track within the ARS zone) was the only characteristic that was significantly different (*)

	Epipelagic bouts	Benthic bouts	All ARS periods
Count	2.0 \pm 0.16	2.8 \pm 0.56	2.1 \pm 0.15
Duration (d)	1.1 \pm 0.10	1.0 \pm 0.19	1.1 \pm 0.09
Area above peak ($\times 10^{-5}$)	5.8 \pm 1.1	2.3 \pm 0.6	5.4 \pm 1.0
Mean fractal D*	1.1 \pm 0.01	1.0 \pm 0.02	1.1 \pm 0.009
Area of circle (km ²)	208.9 \pm 22.9	206.6 \pm 50.1	207.0 \pm 20.8
Dist. traveled in circle (km)	37.1 \pm 3.4	35.1 \pm 7.1	36.8 \pm 3.1

about how each rookery will be impacted by changes to the environment or prey resources, which could guide important conservation decisions.

Movement patterns

Although the dive behavior of northern fur seals has been described previously (Gentry et al. 1986, Goebel et al. 1991, Goebel 1998), this study was unique in that we were able to examine how space use varied between alternative foraging strategies by employing highly accurate GPS tracking technology. In contrast to the differences that were found in the dive data between epipelagic and benthic foraging bouts, only total distance traveled and path straightness were significantly different for movement patterns. However, the significantly higher total distance traveled can be explained by the longer bout durations during epipelagic foraging, since transit rates were not different between the 2 strategies. Therefore, the only remaining difference in movement patterns between the 2 foraging patterns was the difference in path straightness.

The overall straight nature of travel during both epipelagic and benthic bouts was surprising, as we expected females to concentrate their fine-scale movements in areas of high prey density or recent prey capture (Kareiva & Odell 1987, Fauchald & Tveraa 2003). Yet, even with the high level of path straightness, the most sinuous tracks were identified during epipelagic bouts, which is consistent with encircling or following schools of patchy prey. Similar movements have also been described for basking sharks *Cetorhinus maximus*, whose swimming paths become more convoluted while feeding on high densities of patchy zooplankton (Sims & Quayle 1998). It is also important to keep in mind that other factors, both physical and biological, may influence

fine-scale movements during these epipelagic bouts, including water and prey movement patterns. For example, various currents flow along the shelf domains and across the Bering shelf (Stabeno et al. 2001), which could impact the movement direction and speed of the foraging fur seals regardless of the type of foraging strategy being used. In addition, juvenile pollock can be persistent swimmers and are able to maintain swimming speeds of up to 2.5 km h⁻¹ for >3 h (Olla et al. 1997). If a school of juvenile pollock that is being targeted maintains a generally straight swimming path, this could also explain the high path straightness during epipelagic foraging bouts.

In contrast, the high level of path straightness during benthic dive bouts might have resulted from the seals hunting single large adult pollock; hence, concentrating movements in a small area might not be as effective. In fact, studies using underwater cameras on both monk *Monachus schauinslandi* and harbor seals *Phoca vitulina* found that animals tend to cruise rapidly along the bottom instead of focusing their efforts in a single area (Bowen et al. 2002, Parrish et al. 2000). For harbor seals, this type of movement occurred most often when densities of cryptic bottom prey were low (Bowen et al. 2002). Parrish et al. (2000) proposed that this rapid movement along the bottom was a method to minimize the amount of time prey had to detect the seal, and could result in increased foraging success.

Another possible explanation for the higher path straightness during benthic foraging could be related to the increased surface intervals that were observed with benthic diving. These increased surface intervals may be required to recover from the longer dive durations or could be periods when the seal is handling and consuming the larger adult pollock. The increased time at the surface along with the travel time to and from the sea floor may give benthic prey a chance to move or disperse, which would require the seal to transit to locate more prey. In contrast, the short surface intervals and shallow depths of epipelagic dives mean that the seal could maintain contact with the school of prey during the entire foraging bout.

Area-restricted search behavior

When foraging in areas with patchily distributed resources, predators can increase foraging success by modifying movements in response to prey encounters (Kareiva & Odell 1987, Fauchald & Tveraa 2003).

These modifications lead to ARS patterns that are often characterized by decreased transit rates and increased turning rates. Due to the patchy distribution of prey in the marine environment, multiple studies have used ARS behavior to identify foraging periods and locate important foraging habitat of marine predators (Weimerskirch et al. 2008, Weng et al. 2008, Bailey et al. 2009). For example, using ARS locations, a recent study identified the North Pacific Transition Zone as an important foraging area for Laysan albatross *Phoebastria immutabilis* (Tremblay et al. 2007). Bailey & Thompson (2006) used similar indices of increased search effort (first-passage time) to identify potential foraging sites for bottlenose dolphins *Tursiops truncatus*. These authors found that visually identified foraging behaviors (e.g. fish tossing) were more likely to occur in areas with the highest search effort.

As central place foragers during the breeding season, female fur seals focus foraging efforts on the areas surrounding St. Paul Island (Robson et al. 2004), and as expected, ARS behavior was found throughout this region. However, only a few ARS periods were identified per trip and these were generally low in overall intensity. The limited or low intensity ARS periods that were observed in the present study could result from a variety of factors including uniformly distributed prey, low density prey patches, or decreased ability to locate prey patches. For example, if pollock were evenly distributed within the foraging habitat, then foraging success would increase with distance covered, leading to more straight than sinuous paths (Krakauer & Rodríguez-Gironés 1995, Austin et al. 2004, Scharf et al. 2009). Using simulation modeling, Scharf et al. (2009) showed that after prey were encountered, searching was more directional when resources were regularly distributed. However, studies of pollock distribution in the Bering Sea do not support the hypothesis that this species is uniformly distributed (Wespestad et al. 2000); hence this does not explain the low number of ARS zones detected.

If prey patches were of poor quality (i.e. fewer or smaller prey), females should also show more long-distance movements with low turning frequency while searching for new prey patches, leading to minimal time in ARS behavior (Zach & Falls 1976, Fauchald 1999, Nolet & Mooij 2002). Similarly, if females were struggling to find prey resources in general, then there would also be a lack of ARS periods along the tracks. For example, female northern elephant seals spent proportionally more time in directed transit during years of low prey availability (Crocker et al. 2006). However, on average, the fur seals tracked in this study gained mass over the tracking period while continuing to nurse a young pup, suggesting that both of these hypotheses are not supported. In 2007, we found

some evidence that females had difficulty locating prey resources, as trip durations were significantly longer and females travelled greater distances from the rookery (Kuhn et al. 2009). Nevertheless, these differences in at-sea movements did not result in differences in ARS characteristics between years. Thus, there may be another explanation for the lack of ARS periods identified for female northern fur seals, including the possibility that ARS analysis does not reliably identify foraging effort in this species.

This lack of association between ARS periods and foraging effort may also explain why there were no differences in the characteristics of epipelagic and benthic ARS periods. One potential limitation could be the method used to categorize ARS periods, as each ARS period included one or more bouts and many also included extended periods between dive bouts. The lack of association between ARS behavior and foraging was also observed in the wandering albatross *Diomedea exulans*, where researchers showed that prey capture usually did not result in increased track sinuosity (Weimerskirch et al. 2007). ARS periods with little or no bout diving were also identified, further supporting the conclusion that ARS analysis may not be the most appropriate method to describe the at-sea movements of, or identify key foraging areas for, northern fur seals during the breeding season. This could result from the generally straight nature of fur seal travel or the extensive time this species spends in resting and grooming at the surface (Insley et al. 2008). These resting and grooming periods result in increased residence time in an area, which may mistakenly be identified as ARS behavior.

Finally, it is important to consider the resolution of the tracking data in relation to both the temporal and spatial scale of ARS patterns. The GPS locations in this study were obtained at a rate of just over 1 location h^{-1} , with an average distance between locations of ~4 km (Kuhn et al. 2009). If the ARS behavior of northern fur seals occurred on shorter temporal or smaller spatial scales, it would be impossible to detect using this dataset. This was the case for both masked boobies *Sula dactylatra* and bottlenose dolphins where increased sampling rate or visual observations allowed the identification of ARS zones that were <4 km (Bailey & Thompson 2006, Weimerskirch et al. 2008). An additional study using GPS tracking with a higher sampling rate is one way to determine whether northern fur seals in fact show ARS behavior on a much smaller scale.

Ecological implications

While GPS tracking has provided a greater understanding of the fine-scale movements of the northern

fur seal, GPS technology can also be used to gain insight into prey distributions and the impacts of environmental change. When comparing the findings of this study to previous tracking studies, it appears that the foraging ranges of female northern fur seals breeding on St. Paul are generally fixed around the island (Robson et al. 2004, Call et al. 2008). Therefore, differences in use patterns among years within this larger foraging range could provide insight into changes in prey distribution. For example, previous studies in other marine species have shown that locations of increased foraging effort are directly linked to high prey abundance in an area (e.g. Guinet et al. 2001). Additionally, changes in bout diving characteristics could be used to understand changes in prey abundance (Harcourt et al. 2002, Mori & Boyd 2004, Austin et al. 2006). Mori & Boyd (2004) showed that both bout length and time between bouts were correlated with krill abundance in foraging Antarctic fur seals. In fact, these authors suggested that examining changes in fur seal behavior has advantages over traditional hydro-acoustic surveys for estimating krill abundance. In northern fur seals, because females have a variety of foraging options (benthic, epipelagic, or mixed), changes in the proportion of each foraging strategy used may also provide insight into changes in prey resources. In Weddell seals *Leptonychotes weddellii*, temporal changes in prey abundance were shown to impact the foraging strategy employed (either benthic or epipelagic), and it was suggested that seals would not switch to benthic foraging when prey were available in the water column (Plötz et al. 2001). Due to the increased challenges of benthic foraging (Costa & Gales 2003), we also predict that female northern fur seals would only transition to benthic foraging when epipelagic resources in an area decline or are scarce.

Given that northern fur seals are currently listed as depleted and pup production on St. Paul Island has shown no changes from the recent pattern of decline (Towell et al. 2006, NMFS 2007), scientists and managers continue to search for causes. Among the list of potential concerns are changes in the foraging environment, depleted prey resources, and competition with the commercial fisheries that occur throughout the fur seal's foraging range (NMFS 2007, Hiatt et al. 2008). Consequently, the northern fur seal conservation plan listed among its highest-level priorities the tasks of describing essential fur seal foraging habitat and monitoring changes in foraging behavior and habitat use over time (NMFS 2007). The present study directly contributes to this conservation objective by linking dive behavior with fine-scale movement patterns to describe essential foraging habitat both geographically and within the water column. By monitoring fine-scale habitat use and at-sea behavior over

time and making comparisons between St. Paul Island and other islands (including both stable and recovering islands), it may be possible to gain a greater understanding of the interactions between this species and local environmental conditions, fish resources, and commercial fisheries, all of which are essential for effective management and conservation.

In conclusion, the combination of highly accurate GPS locations with dive data resulted in the first description of fine-scale foraging behavior in this species, making it possible to examine whether movement patterns differ when fur seals employ alternative foraging strategies. Although northern fur seals appear to employ 2 distinct foraging strategies during the breeding season, the predominant use of epipelagic dive bouts suggests that this may be a more profitable foraging strategy. Interestingly, while dive characteristics were strikingly different between the 2 foraging strategies, most movement patterns were not significantly different. Future studies integrating analysis of fine-scale foraging behavior with data about individual diet and environmental conditions within the foraging habitat will be essential to add to our understanding of northern fur seal ecology, increase the effectiveness of conservation efforts, and aid in developing predictions about the impacts of environmental change on this depleted species.

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