INTRODUCTION

Phocids (true seals) breed on land or ice, and the pups enter the water and start foraging either during lactation periods (4 to 90 d) or following a period of post-weaning fast (weeks or months). Because this transition from the terrestrial to the aquatic environment is critical for survival (Le Boeuf et al. 1994), knowledge of the diving behavior of pups is essential for our understanding of the ecology of seals.

Dives of pups are shallower and shorter than those of adults (Gjertz et al. 2000). This is presumably because the pups are still developing in terms of their physiology (e.g. oxygen store) and behavior (e.g. swimming skills). The physiological features with regard to diving are well studied, and pups are known to have smaller mass-specific oxygen stores than adults (Burns et al. 2005, 2007). Additionally, pups have higher mass-specific resting metabolic rates (Rea & Costa 1992, Burns et al. 2005) and poorer heart-rate control during diving than adults (Greaves et al. 2004); these likely increase the costs of diving in pups and thus limit their diving capabilities.

In contrast, little is known about the behavioral features of seal pups with regard to diving. For example, adults commonly adopt various swimming styles such as prolonged gliding and stroke-and-glide swimming (Williams et al. 2000, Davis et al. 2001, Sato et al. 2003, Watanabe et al. 2006); however, whether pups can use...
these styles is not clear. Furthermore, the swimming stroke activity of pups may be less stable and less efficient than adults, as is the case in the locomotory abilities of humans (Hausdorff et al. 1999). To test these predictions, it is necessary to monitor the stroke activity of pups using accelerometers (Sato et al. 2003), video cameras (Williams et al. 2000), or magnetic sensors (Wilson & Liebsch 2003).

Bearded seal *Erignathus barbatus* pups enter the water within a few hours of birth (Kovacs et al. 1996), possibly to avoid predators such as polar bears, and can be expected to demonstrate immature diving behavior more clearly than other species. In the present study, we deployed multi-sensor data loggers that recorded depth, swim speed, and acceleration on bearded seal pups weighing between 33 kg (newborn) and 104 kg (close to weaning). The objective of the present study was to determine the characteristics of diving behavior and swimming style of pups. By comparing these characteristics to the general pattern of adults, insights can be gained into the development of diving behavior in seals.

**MATERIALS AND METHODS**

**Study sites and seals.** Field experiments were conducted in Kongsfjorden (78.9° N, 12.5° E), Svalbard, Norway, in May 2006 and May 2007. The study periods covered the peak birthing period of bearded seals in Svalbard (the first week of May) and the subsequent lactation period that lasts about 3 wk (Lydersen & Kovacs 1999). Mother–pup pairs that were resting on ice floes or swimming were approached by boat, and 6 pups were captured using a custom-built dip net. The pups were weighed using a Salter spring scale (±0.5 kg), and their sex was determined. They were then instrumented (see next paragraph) and released. All the mothers remained in the water nearby during the entire procedure, and approached their pups and swam with them upon their release. This protocol was approved by the Governor of Svalbard (Sysselmannen) and the Norwegian Animal Research Authority. The ages (days) of 3 of the pups were known because their birth dates were known from daily censuses. The ages of the other 3 pups were estimated from their body mass, assuming a daily mass gain of 3.3 kg (Lydersen et al. 1994) and a newborn body mass of 38 kg (K. M. Kovacs & C. Lydersen unpubl. data) (see Table 1).

**Instruments.** We used 512 Mb W2000L–PD2GT data loggers (length = 127 mm, diameter = 27 mm, weight in air = 125 g; Little Leonardo), which recorded swim speed, depth, and temperature at 1 s intervals, and 2-dimensional (lateral and longitudinal) accelerations at 1/32 s intervals. The maximum range of the depth sensor was 2000 m, with a resolution of 0.5 m. Data loggers must be recovered to obtain the data, but recapture of the instrumented seals is difficult. Therefore, we used a newly developed remote-controlled release system that enabled the logger to be retrieved without recapture of the animals (Fig. 1). A mat of synthetic fibers was glued onto the fur on the back of the seals with a quick-setting epoxy resin. Then, a package composed of a data logger, a release mechanism with a semi-rigid antenna for UHF radio signals (Little Leonardo), and a float of copolymer foam (Nichiyu Giken Kogyo) was attached to the mat with a plastic cable. The cable was connected to the release mechanism. When the seals were located some days after the initial deployment, UHF radio signals were sent by a transmitter to activate the release mechanism from a distance of about 50 m. The cable was severed by an electric charge from the battery of the mechanism, and the package was released from the seal. The package also contained a VHF transmitter (Telonics) that helped in locating the seals. The total weight of the instruments was 288 g in air (0.3 to 0.9% of the body mass of the seals) and its buoyancy offset 56 g in water.

**Data analysis.** Based on the absolute accuracy of the sensor, a dive was defined as any excursion below the sea surface to a depth of >2 m. The diving capability of
the pups was first examined by their dive depths and durations. To examine their swimming behavior in greater detail, each dive was subdivided into a descent phase (from the beginning of a dive to the time of the first ascent), an ascent phase (from the time of the last descent to the end of the dive), and a bottom phase (the time between the end of descent and the beginning of ascent). We excluded dives in which the depth at the start or end of the bottom phase was < 5 m, because swimming parameters (e.g. speed and number of strokes) measured over very short descent or ascent are not reliable.

The relative swim speed through water was recorded as the number of rotations per second (rps) of an external propeller mounted at the anterior end of the loggers. This value was converted to the actual swimming speed (m s⁻¹) using the calibration method of Sato et al. (2003).

From lateral acceleration records, we extracted information on stroke activity of the seals by filtering out the low-frequency signals with IGOR Pro (WaveMetrics) (Sato et al. 2003). A single stroke cycle was defined as the period taken by the flipper to move from one extreme lateral position back to the original position. Gliding was defined as the period when the seals stop stroking for more than 2 s, which corresponded to approximately twice the mean duration of the stroke cycle for all pups. Distance covered by a stroke cycle (m) was calculated as the swim speed (m s⁻¹) integrated over a period (s) divided by the number of strokes during the period.

Modulation of the stroke activity was first visualized by creating the Gabor spectrograms for filtered lateral acceleration with IGOR Pro (Sato et al. 2008). Then, the variability of the stroke cycle frequency within individuals was quantified by determining bandwidth, which represents the deviations around the centroid frequency and is usually used for acoustic data (e.g. Madsen et al. 2004). To obtain bandwidth, power spectral density (PSD) of lateral acceleration was calculated by fast Fourier transformation with IGOR Pro. PSD showed the peak frequency that represented the dominant stroke cycle frequency of each seal (Sato et al. 2007). Bandwidth of the signals was calculated as the range of frequencies where the power of the fast Fourier transformation was above 95% of the maximum value. This value (95%) was chosen because it gave the maximum bandwidth excluding the low-frequency signals, which are assumed to be the result of turning and rolling movements of the seals.

A single-factor ANOVA followed by Scheffe’s post-hoc test was performed with StatView (SAS Institute) to compare diving (e.g. dive depth) and swimming parameters (e.g. speed) among individuals. Because successive dives made by 1 individual can be auto-correlated, we also randomly selected 110 and 50 dives (i.e. the lowest number of dives recorded for 1 individual) from each seal for the analyses of diving and swimming parameters, respectively, and performed the same tests. The value for statistical significance was set at p < 0.05. Mean ± SD values are reported.

RESULTS

Behavior was recorded for approximately 3 d (63 to 78 h) for each pup, except for the largest pup, in which the data logger was accidentally lost 3 h after deployment and recovered from the sea surface the following day (Table 1). The pups spent an average of 58 ± 12% (n = 5) of their time hauled out on the ice and the rest of their time in the water, excluding the largest pup, which was in the water throughout the 3 h recording period. The smallest pup showed the highest haul-out time (78%), and its daily haul-out time decreased continuously from 84 to 73% during the 3 d. A total of 5518 dives were recorded for the 6 pups. The dives of the smallest pup were the shallowest (ANOVA, \( F_{5,5512} = 27.5, \ p < 0.0001 \); Scheffe’s post-hoc test, \( p < 0.0001 \) for all 5 combinations) and the shortest if excluding the largest pup (\( F_{5,5512} = 40.8, \ p < 0.0001 \); \( p < 0.0001 \) for 4 combinations), a result probably affected by the short recording period for the largest pup. Tests using 660 randomly-selected dives (110 dives for each seal) gave the same results, except that dive depth of the smallest pup did not differ significantly from 2 medium-sized pups (Eb-07-26 and Eb-06-02), and dive duration of the smallest pup did not differ significantly from one of the medium-sized pups (Eb-07-26). All the diving parameters (i.e. daily mean and maximum dive depth and duration) increased continuously only in the smallest pup. All pups showed dives with a bottom phase during which the swim speed dropped below the stall speed of the sensor. Such dives constituted 26 ± 14% (n = 6) of the total number of dives (Table 1).

We selected 2146 dives for the analysis of swimming behavior (Table 1). Swim speed during transit phases (i.e. descents and ascents) was the lowest in the smallest pup (\( F_{5,2140} = 83.4, \ p < 0.0001 \); \( p < 0.0001 \) for all 5 combinations) and highest in the largest pup (Scheffe’s post-hoc test, \( p < 0.0001 \) for all 5 combinations). Distance covered by a stroke cycle during transit phases was the lowest in the smallest pup (\( F_{5,2140} = 227, \ p < 0.0001 \); Scheffe’s post-hoc test, \( p < 0.0001 \) for all 5 combinations) and the highest in the largest pup (Scheffe’s post-hoc test, \( p < 0.0001 \) for all 5 combinations). Tests using 300 randomly-selected dives (50 dives for each seal) gave the same results. The 2 smallest pups stroked continuously throughout both descent and ascent (Fig. 2a).
Fig. 2. *Erignathus barbatus*. Single dives made by pups (a) Eb-07-16, (b) Eb-06-02, and (c) Eb-07-04 with depth, swim speed, filtered lateral acceleration (indicative of stroke activity), and a spectrogram of filtered lateral acceleration where magnitude was expressed by color. Red vertical dashed lines delimit 3 dive phases: descent, bottom, and ascent; \(\text{J} \): continuous strokes; \(\text{JJ} \): prolonged gliding. (d–f) Power spectral density calculated from filtered lateral accelerations during the dives.
Three of the medium-sized pups (Eb-07-26, Eb-06-02, and Eb-06-04) glided during the last part of the descent of deep dives (>40 m) (Fig. 2b). However, gliding was not the main swimming style for any of the medium-sized pups; mean glide time during transit phases constituted only 1 to 5% of the transit time (Table 1). In contrast, the largest pup glided during most of the ascent (Fig. 2c), and its gliding time during transit phases was by far the highest ($F_{5,2140} = 257, p < 0.0001$; $p < 0.0001$ for all 5 combinations) (Table 1). A test using 300 randomly-selected dives (50 dives for each seal) gave the same result.

Stroke-and-glide swimming was not observed in any pups.

Spectrograms showed different modulation patterns of the stroke activity among the pups (Fig. 2a–c). The stroke cycle frequency of the largest pup during descent was stable at 1.2 Hz, while that of the smallest pup was more variable. The 4 medium-sized pups showed intermediate patterns between the smallest and largest pups. These differences in the spectrograms corresponded to the differences in the PSD profiles calculated for each dive (Fig. 2d–f). The slopes around the peak, which corresponded to the dominant stroke cycle frequency of the dive, were steep in the largest pup and gentle in the smallest pup, indicating higher stability in stroke cycle frequency in the largest pup. The examples in Fig. 2 show that the width of the peak band on the PSD profiles is a good indicator of the variability in the stroke cycle frequency; these findings support our application of bandwidth to acceleration data. The PSD profiles of the entire lateral acceleration records differed qualitatively among the pups (Fig. 3).

### DISCUSSION

The pups spent approximately half of their time hauled out, and the other half in water; this was in agreement with Lydersen et al. (1994), who studied bearded seal pups weighing between 41 and 54 kg — a

---

**Table 1. Erignathus barbatus. Diving and swimming behavior of bearded seal pups. Individuals are shown in order of weight. Means ± SD are presented.**

<table>
<thead>
<tr>
<th>Attribute</th>
<th>Pup identification</th>
<th>Sex</th>
<th>Date of capture</th>
<th>Body mass at capture (kg)</th>
<th>Age at capture (d)</th>
<th>No. of dives</th>
<th>Dives with bottom phase where speed drops (%)</th>
<th>Dive depth (m)</th>
<th>Haul-out time (%)</th>
<th>No. of dives selected</th>
<th>Swim speed (m s$^{-1}$)</th>
<th>Dominant stroke cycle frequency (Hz)</th>
<th>Dominant stroke cycle duration (min)</th>
<th>Distance covered by a stroke cycle (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Descriptive information</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sex</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Date of capture</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Body mass at capture (kg)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Age at capture (d)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Diving behavior</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Data length (h)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Haul-out time (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No. of dives</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dives with bottom phase where</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>speed drops (%)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dive depth (m)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>mean</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>maximum</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Swim speed (m s$^{-1}$)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>mean</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>maximum</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Swimming behavior</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>No. of dives selected</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>mean</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>maximum</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*Age estimated from weight*
narrower range than in the present study. However, the newborn in this study spent 84% of its time hauled out on the first day of life, and gradually decreased its daily haul-out time over the next 2 d. Adult females of this species spend only 8% of their time hauled out during the nursing period, spending the majority of their time diving and remaining at the sea surface adjacent to the ice floes where their pups are resting (Krafft et al. 2000). Our results indicate that bearded seals begin to adopt a more aquatic lifestyle (i.e. less time on floating ice) during the early stages of life. Unlike most phocids, pups of this species start foraging before weaning (Lydersen et al. 1996). Because bearded seals prey on fishes, crustaceans, mollusks, and worms at the sea floor (Hjelset et al. 1999) using their sensitive whiskers (Marshall et al. 2006), the bottom phases where swim speeds drop suggest they are foraging. In the present study, all pups including the newborn demonstrated such dives. This suggests that the pups imitated the swimming behavior of the accompanying mothers, who frequently dive shallowly (<10 m) and forage during the nursing period (Krafft et al. 2000). Independent prey acquisition probably does not begin until after 1 or 2 wk of life (K. M. Kovacs & C. Lydersen unpubl. data).

The dive depths and durations recorded in the present study were similar to those previously reported for bearded seal pups (Lydersen et al. 1994) and were less than those of adults (Krafft et al. 2000). The dives of the newborn, in particular, were shallow and short. However, the diving capability of this pup increased on a daily basis, suggesting a rapid development of diving capabilities during early post-natal life.

The mean swim speed was the lowest (0.5 m s⁻¹) in the smallest pup and the highest (0.9 m s⁻¹) in the largest pup. The speed of the largest pup was similar to that of adult bearded seals (0.9 to 1.4 m s⁻¹, Krafft et al. 2000). The increase in swim speed during the nursing period cannot be attributed to the increase in body length, which changes little during the first year of life (K. M. Kovacs & C. Lydersen unpubl. data). Furthermore, neither muscle mass (as a percent of body mass) nor mass-specific oxygen stores increase during the nursing periods in seals (Burns et al. 2007). Therefore, the increase in speeds may have been achieved by experience or the development of neuromuscular coordination.

Fig. 3. *Erignathus barbatus*. Power spectral density calculated from the entire lateral acceleration records for each pup. Pups are shown in order of weight. −−: Bandwidth; arrows indicate dominant stroke cycle frequency.
Only the largest pup showed substantial periods of prolonged gliding, an efficient means of locomotion through water (Williams et al. 2000). The effectiveness of gliding was evident by the largest distance covered by a stroke cycle (0.6 m) of this pup. This pup, which was close to weaning, glided throughout ascent, indicating that gliding was assisted by positive buoyancy caused by the increase in the subcutaneous blubber layer during nursing (Biuw et al. 2003, Sato et al. 2003, Watanabe et al. 2006). In contrast, 3 of the medium-sized pups glided only during the last parts of descent in deep dives, when they were presumably negatively buoyant. The 2 smallest pups did not glide, suggesting that pups develop the ability to glide during the nursing period. No pups in this study showed stroke-and-glide swimming, which is an efficient locomotory mode, at least in theory (Weihs 1974). Our results suggest that the swimming style of pre-weaned pups was immature, but it developed over the short nursing period.

Spectrograms and PSD profiles of lateral acceleration indicate that the stroke cycle frequency of the smallest pup was highly variable and that of the largest pup was relatively stable. Steady strokes are physiologically preferable for cruising because muscle contraction is most efficient over a narrow range of contraction speeds and loads (Hill 1950, 1964). In addition, steady swimming is physically advantageous because it decreases body inertia and acceleration reaction, which animals must work against (Webb 1988). Therefore, a decrease in the variability of the stroke cycle frequency presumably represents an improvement in the swimming performance. The shortest distance covered by a stroke cycle in the smallest pup (0.3 m) might be attributed to the largest variability in its stroke activity. A similar phenomenon is seen in human children with respect to the variability of stride duration, which decreases with age (Hausdorff et al. 1999). Until now, ontogeny of locomotor performance was studied only in humans (Hausdorff et al. 1999), captive animals (Noren et al. 2006), and wild animals under controlled conditions (Irschick 2000). Accelerometers provide a new tool for quantifying the developmental changes in locomotory performance in wild animals in their natural habitats.

In conclusion, the swimming performance of pre-weaned bearded seal pups was immature, but it developed during the short nursing period. The mean swim speed and the distance covered by a stroke cycle was lowest in the smallest pup and highest in the largest pup. Only the largest pup, which was close to weaning, showed substantial periods of gliding during ascent. Stroke-and-glide swimming was not observed in any pups. The stroke cycle frequency was highly variable in the smallest pup but relatively stable in the largest pup.

Acknowledgements. We thank M. Fedak, C. Freitas, M. Haupt, M. Hindell, and C. Marshall for their assistance with the field-work, T. Morisaka and K. Yoda for advice on data analysis, and P. Miller and 4 anonymous referees for helpful comments on the manuscript. This work was funded by the Norwegian Research Council, the Norwegian Polar Institute, the Japan Society for the Promotion of Science (A19255001), JSPS Research Fellowships for Young Scientists (Y.W.), and the program Bio-logging Science of The University of Tokyo (UTBLS).

LITERATURE CITED


Editorial responsibility: Rory Wilson, Swansea, UK

Submitted: August 4, 2008; Accepted: October 29, 2008
Proofs received from author(s): March 28, 2009