Active hunting by deep-diving sperm whales: 3D dive profiles and maneuvers during bursts of speed

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ABSTRACT: Active hunting pursuit can involve high expenditures of energy and therefore requires appropriately high-energy gains from successful prey capture. Using data loggers deployed on 12 sperm whales off the Ogasawara Islands, we regularly recorded bursts of speed during deep dives. Here, we analyzed speed, acceleration, and 3D data from these maneuvers to describe the whales’ hunting behavior. Bursts occurred at depths >400 m. The median number of bursts was only 1 per dive (range: 0 to 6 per dive), and 33% of the dives did not include any bursts, suggesting prey capture may not always require bursts. Bursts of speed averaged 3.4 ± 1.0 m s⁻¹ (mean ± SD; maximum: 8 m s⁻¹), more than twice the mean speed observed during dives (1.5 ± 0.2 m s⁻¹). Bursts were generally divided into two phases: (1) rapid acceleration with active stroking, and (2) drastic deceleration and changes in body orientation. Tagged whales swam up to 405 m (mean: 120 ± 88 m) during these two phases. Such behavior suggests chasing of their prey, and should incur high energy expenditure. A large percentage (20 ± 14%) of the drag-related locomotion cost of the dives was spent in bursts. Two major types of bursts were observed: inflectional bursts with turning (87%) and linear bursts without turning (13%). Our results strongly indicate that sperm whales use an active-pursuit hunting strategy and use the bursts only to catch powerful and nutritious prey (i.e. large and/or muscular) that compensate for the energetic cost of the burst.

KEY WORDS: Diving behavior · Hunting tactic · Hunting strategy · Swim speed · Data logger · Physeter macrocephalus · Cetacean

INTRODUCTION

Two main hunting strategies have been reported for both terrestrial and aquatic predators: sit and wait and active pursuit (e.g. Huey & Bennett 1986). Active pursuit involves higher energy expenditure and requires appropriately high-energy gains from successful prey capture (e.g. Secor 1995). Several studies have reported that aquatic predators increase speed to feed on mobile and larger pelagic fishes (Davis et al. 1999) and decrease speed to feed on small inactive crustaceans (Wilson et al. 2002,
Watanabe et al. 2004). Energy expenditure and therefore oxygen consumption are predicted to increase as a cubic function of swim speeds (e.g. Kooyman 1989, Otani et al. 2001). Therefore, to maximize foraging efficiency, aquatic predators should match their pursuit speed with prey escape speed that is a function of prey size (Wilson et al. 2002). Thus, both speeds and duration of active pursuit of predators should be indicative of both the prey quality and prey character (e.g. activity and size). In the present study, we focused on pursuit speed to investigate the hunting behavior of the sperm whale *Physeter macrocephalus*.

The sperm whale is the largest toothed whale and one of the deepest- and longest-diving mammals. Sperm whales routinely dive to depths >400 m, and such dives are thought to be foraging dives (Watkins et al. 2002). There are a few hypotheses regarding the hunting strategy of sperm whales. Early hypotheses were based on the sit and wait strategy, which is energetically less costly than active pursuit (Beale 1839, Clarke 1970). The more accepted hypothesis is that sperm whales engage in an active search and approach strategy by employing echolocation clicks (e.g. Norris & Harvey 1972, Möhl et al. 2000, Madsen et al. 2002, Whitehead 2003). Using a tag with tri-axial magnetometers and accelerometers, but without a speed sensor (Dtag; Johnson & Tyack 2003), Miller et al. (2004b) showed that sperm whales exhibit a rapid decline in the click interval (buzzes) accompanied by increased body movements, suggesting that such behaviors indicate final approach and prey capture. Amano & Yoshioka (2003) reported that the movements of a tagged sperm whale had events of acceleration of up to about 7.0 m s⁻¹. Such events were termed bursts of speed and were suggested to be feeding events. These previous studies support the hypothesis that sperm whales use an active approach strategy rather than a sit and wait strategy when hunting. However, little is known about the range over which sperm whales can adjust their pursuit speed.

The mesopelagic squids found in the stomach contents of sperm whales vary greatly in activity and size, from small chiroteuthids (<100 g) to large giant squids (*Architeuthis* sp., up to 400 kg) (Clarke 1980, Kawakami 1980). The smaller squids are generally more gelatinous and slow swimming than the larger and more powerful squids preyed on by whales (Clarke et al. 1993). Therefore, large squids probably provide a challenging capture but higher gain in energy compared to the small squids that are the staple food for most sperm whales (Clarke et al. 1993, Whitehead 2003). Although large squids (e.g. *Architeuthis* sp., Ommastreohids, and *Tantingia danae*) account for a small proportion (6%) of the total number of individuals identified in sperm whales’ stomach contents, they provide a substantial proportion of the biomass consumed because of their large size (54% of total mass; Clarke et al. 1993).

In the present study, we describe bursts of speed using multi-sensor data loggers deployed on 12 sperm whales off the Ogasawara Islands of Japan. To investigate sperm whale hunting behavior, we analyzed detailed 3D dive profiles and maneuvers in relation to swim speeds. Bursts were infrequent, but they accounted for a large percentage of the drag-related locomotion cost of dives, indicating that sperm whales use the bursts only to catch powerful and nutritious prey.

### MATERIALS AND METHODS

**Tag deployment**

Tagging was conducted off the Ogasawara Islands in the western North Pacific Ocean from 2004 to 2008. Groups of female and immature sperm whales are found throughout the year off the Ogasawara Islands (Mori et al. 1995). We approached sperm whales with a fishing boat (length: 20 m) and deployed suction-cup attached tags including a data logger using a crossbow or a 6 m pole. The weight of the whole tag was from 370 to 470 g. The maximum length, width, and thickness of the whole tag was 32, 11, and 5 cm, respectively (for tag details see ‘Suction-cup attached tags’ and Fig. S1 in the supplement at www.int-res.com/articles/supp/m444p289_supp.pdf).

Three types of data loggers were used: the UWE2000-PDT (diameter: 21 mm, length: 123 mm, weight: 64 g in air; Little Leonardo), the W2000-PD2GT (diameter: 23 mm, length: 123 mm, weight: 90 g in air; Little Leonardo) and the W2000-3MPD3GT (diameter: 30 mm, length: 175 mm, weight: 140 g in air; Little Leonardo). All data loggers recorded depth, swim speed, and temperature at 1 s intervals. In addition, the 3MPD3GT recorded the 3D magnetic field at 1 s intervals and 3D accelerations at 1/16 s intervals (for detecting fluking movements and the body orientation of whales; Fig. 1). The measurement range of the depth sensor of each data logger was from 0 to 2000 m, with a resolution of 0.5 m. We attached the PDT to 2 whales, the PD2GT to 5 whales, and the 3MPD3GT to 5 whales (Table 1; details on the suction-cup attached tags of each type of data logger are in the supplement).
The start and end of dives were defined as the time when the whales descended below and ascended above a depth of 4 m, respectively. Although male sperm whales feed near the surface (Teloni et al. 2008), female and immature sperm whales routinely forage in mesopelagic waters (Amano & Yoshioka 2003, Miller et al. 2004b, Watwood et al. 2006). Off the Ogasawara Islands, female and immature sperm whales also dove to depths >400 m, and these dives were interpreted to be foraging dives (Aoki et al. 2007). According to a histogram of the maximum dive depth, a boundary was observed around 200 m between shallow and deep dives. We analyzed deep dives with a maximum depth >200 m. All deep dives were divided into 3 phases: (1) the descent phase (from the start of a dive to the first time at which the depth change rate was negative); (2) the ascent phase (from the last time at which the depth change rate was positive, to the end of the dive); and (3) the bottom phase (the time between the end of the descent phase and the beginning of the ascent phase) (see Aoki et al. 2007 for details). Dive depth was defined as the maximum depth of the dive. Post-dive surface duration was defined as the time between the end of the dive and the start of the next dive.

Speed through the water was measured using an external propeller on the data logger. The rotation rate (revolutions s$^{-1}$) of the propeller was converted to swim speed (m s$^{-1}$) using a calibration line for each whale. We used 2 different methods to obtain calibration lines, with the method used dependent upon the type of data logger used. A linear regression of revolutions per second against the rate of vertical depth change (m s$^{-1}$) was used to estimate a calibration line for whales tagged with the PDT or PD2GT data loggers (modified from Blackwell et al. 1999, see Aoki et al. 2007 for details; Fig. S2 in the supplement). We regressed third, fifth, and tenth percentile points of propeller rotation data to corresponding vertical depth change rates in each whale’s data and used the line with the highest regression coefficient value for each whale. Another method was employed for each whale tagged with the 3MPD3GT after analyzing acceleration data. A calibration line was obtained from a linear regression of revolutions per second against swim speed (m s$^{-1}$), which was calculated from the rate of vertical depth change divided by the sine of the pitch when the sine of the adjusted pitch (see below) was >0.9 (see Sato et al. 2003 for details and Fig. S3 in the supplement). Both methods yielded relatively high regression coefficients (range: 0.801 to 0.998; n = 12 whales). Resolutions of swim speed,
which correspond to 1 rotation of the propeller, ranged from 0.014 to 0.052 m s\(^{-1}\) (n = 12 whales). The swim speeds, directly derived from the rotation rate, were in the range of 0.1 to 3.8 m s\(^{-1}\) (n = 12 whales).

Accelerations in the tri-axis (longitudinal, lateral, and dorso-ventral axes) measured using the 3MPD3GT data logger can be broken into components related to the body orientation of the animal with respect to gravity (gravity-based components) and propulsive activities imposed by fluke thrust (dynamic components, which is also called specific or platform acceleration) (Sato et al. 2003, Watanabe et al. 2006). Lower-frequency gravity-based accelerations of all axes were extracted from raw acceleration data using the low-pass filter of the software IFDL within IGOR Pro (WaveMetrics). A frequency boundary for the filter was determined for each whale (range: 0.0933 to 0.125 Hz) by calculating the power spectral density of the raw acceleration data (see Watanabe et al. 2006 for details). For this calculation, the bottom phases of the dives were excluded because the body rotations recorded in the bottom phase seemed to occur at rates similar to those of stroking action. We therefore note that it was impossible to separate these components perfectly in the bottom phases. Lower-frequency gravity-based acceleration of the longitudinal axis was used to calculate the pitch angle of the data logger (Sato et al. 2003). The roll angle of the data logger was calculated from the gravity-based components of dorso-ventral and lateral accelerations based on Johnson & Tyack (2003) (see also Shiomi et al. 2010 for details).

Since the data logger was not always attached exactly parallel to the axes of the whale, it was necessary to correct attachment angle. Corrected values, calculated to adjust pitch, were used to compensate for possible off-axis placement on the body by using the cumulative value method for each dive per whale (see Sato et al. 2003 for details). Absolute differences in the corrected values between dives averaged 2.8 ± 2.5° (n = 65 dives), indicating good tag stability on the animal. Corrected values, calculated to adjust for roll, were obtained to compensate for possible off-axis placement on the body based on 2 criteria for each dive per whale: (1) the roll of the whale should equal zero when the whale was resting at the surface between deep dives and (2) the corrections should match the visually inspected orientation of the tag on the whale (Miller et al. 2004a). Absolute differences of the corrected values between dives averaged 4.2 ± 6.4° (n = 65 dives). Magnetism values change depending on the angles between the geomagnetic vector and each axis of the data logger. Therefore, the heading of the data logger (i.e. heading of the whale) was calculated from adjusted pitch and roll and tri-axis magnetism (see Johnson & Tyack 2003 for details) using a macro, programmed by T. Narazaki (Narazaki & Shiomi 2010), for the program IGOR Pro (WaveMetrics). We assumed that the heading of the data logger did not differ dramatically from that of the tagged whales (see ‘Suction-cup attached tags’ in the supplement for details). The tags were not initially deployed parallel to the body axes of the animals; however, they shifted to an almost parallel position after several minutes, suggesting that tags became nearly parallel to the body axes of the animals owing to the water flow (see Sakai et al. 2011 for details). However, we were unable to check the tag position during each surfacing event for all the tagged whales. Therefore, the heading of the data logger was probably not exactly parallel to body axes for all the whales. This might have caused estimates of speed to be reduced or increased to some extent (see ‘Data analysis’ in the supplement for details). However, we think that these effects did not remarkably change our main results.

Horizontal swim speed of each whale was calculated by cosine (pitch of the whale) × swim speed. Thereafter, the 3D dive profiles were estimated by depth, heading, and horizontal swim speed of the whale (i.e. dead reckoning; Wilson et al. 1991). To identify strokes, the higher frequency dynamic acceleration of dorso-ventral axis was obtained by subtracting low-frequency components from the raw acceleration data. The peaks and troughs of the signal with absolute amplitudes greater than a specified threshold were considered to represent individual strokes (Sato et al. 2003). To select the threshold, we made a histogram of absolute amplitudes from all peaks and troughs for each whale. This analysis showed a clear boundary between small noises and strokes. We defined the boundary as the threshold for each whale (range: 0.25 to 1.67 m s\(^{-2}\)).

A burst (Fig. 2) was defined based on the following 2 conditions: (1) forward acceleration was >0.15 m s\(^{-2}\) calculated from swim speed at 5 s intervals and (2) swim speed reached more than the mean + 1.5 SD of the bottom phase of each dive (see also Fig. S4 in the supplement). The start of a burst was defined as the time when swim speed began to increase. The end of a burst was defined as the time when the tagged whales began to swim constantly again, or the swim speed decreased to less than the mean speed of the bottom phase. Bursts were expressed as events that included relatively high speed with strong forward accelerations (Fig. S4 in the supplement). We therefore focused on the tails of swim speed histograms.
Aoki et al.: Active hunting of sperm whales during deep dives as bursts (see Fig. 3a). To investigate maneuvers of the whales, changes in body orientation (i.e. pitch, roll, and heading) (° s−1) were calculated at 1 s intervals. High rates of change in body orientation (>mean + SD during bursts which corresponded to means + 10 SD during deep dives: pitch: 7° s−1, roll: 15° s−1, heading: 21° s−1) were picked up as rapid changes in body orientation during bursts. We investigated temporal occurrence of rapid changes in body orientation, changes in depths, and the presence of turns during bursts (see ‘Data analysis’ in the supplement for details).

The lack of data on the metabolic rate of sperm whales prevents precise calculation of the locomotion cost of bursts. The mechanistic model used to describe locomotion cost is based on the hydrodynamic drag experienced by the animal (Hind & Gurney 1997). The drag force scales with the square of swim speed, U (Vogel 1981). The power required to overcome this drag force can be approximately scaled with \( U^3 \) (Hind & Gurney 1997), if we assume that the propeller efficiency and drag coefficient are constant over a range of swim speeds. Therefore, the ratio of the cost of bursts to the whole dive locomotion cost \( (CL_{b/d}) \) can be estimated by:

\[
CL_{b/d} = \left( \int_{t_b}^{t_d} U^3 \, dt \right) / \left( \int_{t_b}^{t_d} U^3 \, dt \right)
\]

where \( t_b \) is the duration of bursts and \( t_d \) is the duration of the whole dive (Aguilar Soto et al. 2008).

All statistical analyses were performed using the R 2.6.2 package (The R Foundation for Statistical Computing, Vienna) (see the supplement for details of statistical analyses). Values for statistical significance were set at \( p < 0.05 \). We report means ± SD.

RESULTS

A total of 116.8 h of data, including 126 complete dives (maximum depth: >200 m), were obtained from 12 tagged sperm whales Physeter macrocephalus off the Ogasawara Islands. All tagged whales were determined to be mature females or immature individuals on the basis of their size and external appearance (estimated body length by sight: 6 to 10 m). Maximum dive depths and duration were 1422 m and 53 min, respectively (mean dive depth: 695 ± 247 m, mean dive duration: 33 ± 6 min; n = 126 dives). The whales swam continuously during deep dives (Fig. 2). The mean swim speed during a whole dive was 1.5 ± 0.2 m s−1 (n = 126 dives; Table 2, Fig. 3a). The mean speed of each phase was 1.4 ± 0.3 m s −1 during descent, 1.6 ± 0.3 m s−1 at the bottom of the dive, and 1.6 ± 0.3 m s−1 during ascent (n = 126 dives). The tagged whales spent most of their time in deep
dives that exceeded a maximum depth of 200 m (81.4 ± 24.5%; n = 12 whales). All whales tagged with the 3MPD3GT glided more during ascent than in descent (gliding rate — descent: 29.1 ± 5.0%, ascent: 57.1 ± 19.8%; n = 5 whales).

Bursts of speed (Fig. 2) were observed from all tagged whales. A total of 181 bursts were recorded (median: 1 per dive, range: 0 to 6 per dive; n = 126 dives; Table 2). However, 33% of dives did not include any bursts, and the remaining 67% of dives included 1 burst at least. The mean and maximum durations of the bursts were 58 ± 37 s and 176 s, respectively (n = 181 bursts). During a burst, maximum swim speed was generally between 2 and 5 m s⁻¹ (3.4 ± 1.0 m s⁻¹; n = 161 bursts), but reached 8.0 m s⁻¹ (Fig. 3b). The total swimming distance of the bursts had a range of 8 to 405 m (120 ± 88 m; n = 161 bursts). The ratio of the cost of bursts to the whole dive locomotion cost averaged 20 ± 14% (range: 1 to 70%) at dives where at least 1 burst was observed (n = 73 dives). We note that 12 dives including bursts were not used for this calculation (see supplement for details). Burst rates were significantly higher during the bottom phase (4.2 bursts h⁻¹) than during descent (1.2 bursts h⁻¹) or ascent phases (0.7 bursts h⁻¹) (Kruskal-Wallis test, χ² = 15.1, p < 0.001). More than 90% of bursts occurred at depths >400 m. Bursts were most frequent between 800 and 900 m (Fig. 3c). In contrast, the tagged whales most frequently stayed between 400 and 500 m, and time spent at depths beyond 400 m decreased with increasing depth (Fig. 3c).

Maneuvers and 3D dive profiles during bursts

A total of 69.5 h of data, including 106 bursts, were obtained from 5 whales tagged with the 3MPD3GT. Tagged whales swam steadily before the start of 61% of bursts, but gliding was recorded prior to 39% of bursts (duration of gliding: 57 ± 42 s, maximum: 177 s; n = 41). Change in body orientation was slow during gliding, with no active stroking (mean change rate: pitch: 1 ± 1° s⁻¹, roll: 2 ± 1° s⁻¹, heading: 2 ± 1° s⁻¹; n = 41), and the mean swim speed was 1.2 ± 0.4 m s⁻¹ (n = 41).

During bursts, whales exhibited rapid acceleration and drastic deceleration of swim speed (maximum acceleration during each burst: 0.28 ± 0.11 m s⁻², maximum deceleration during each burst: −0.55 ± 0.36 m s⁻²; n = 106). To investigate whether deceleration during bursts was passive or actively generated by the animal, we compared deceleration rates dur-
ing bursts with those recorded during glides. We extracted deceleration rates during horizontal swimming for both bursts and glides because the buoyancy of the animals causes acceleration/deceleration during descent and ascent glides (Miller et al. 2004a, Watanabe et al. 2006). Additionally, we used only the deceleration rates during bursts that had the same range of swim speeds as the glides (0.8 to 2.2 m s$^{-1}$), because faster speed causes larger drag. Our statistical model included decelerating rates as a dependent factor, mean speed of decelerating periods and phase (glide/burst) as fixed independent factors, and whale ID as a random factor. Deceleration rates during bursts were substantially larger than those recorded during glides (deceleration during burst: $-0.55 \pm 0.51$ m s$^{-2}$, $n = 21$; deceleration during horizontal glide: $-0.026 \pm 0.015$ m s$^{-2}$, $n = 34$; GLMM, $\chi^2_{1} = 105.7$, p < 0.0001), indicating that deceleration during bursts was not passively attained. Rapid decelerations coincided with larger changes in pitch and heading (GLMM — pitch: $\chi^2_{1} = 15.8$, p < 0.0001; heading: $\chi^2_{1} = 25.9$, p < 0.0001; swim speed: $\chi^2_{1} = 65.5$, p < 0.0001). Such changes in heading or pitch are expected during horizontal or vertical turns. 

Bursts exhibited a diverse combination of the whale’s maneuvers. According to 3D scatter plots of the principal components that were extracted from maximum speed, maximum deceleration, change rate of heading, and rapid and simultaneous changes in body orientation during bursts (Table S1 in the supplement, Fig. 4), the bursts were divided into 2 types: inflectional bursts (n = 87 bursts, 82%) and linear bursts (n = 19 bursts, 18%). Inflectional bursts had more rapid changes in body orientation and more drastic deceleration than linear bursts (Table 3). The details of each type are shown below and in Fig. 5.

There was high variability in (1) changes in swim speed and (2) changes in depth recorded during inflectional bursts. The majority of inflectional bursts could be categorized into 2 different types of changes in swim speed (Table 3): a single acceleration phase or 2 acceleration phases. Changes in depth during inflectional bursts occurred mainly as one of 3 types: descent (n = 35 bursts, 40%), horizontal (n = 28 bursts, 32%), or ascent (n = 17 bursts, 20%). Although inflectional bursts showed similar patterns, including rapid changes in body orientation, drastic deceleration, and turns (n = 73 bursts, 87%), there were variable combinations of these factors (Table 3).
Linear bursts showed more gradual deceleration rates and slower changes in body angle compared with inflectional bursts (Table 3). Linear bursts did not include any turns. For some linear bursts, no rapid changes in body orientation were recorded (n = 6 bursts, 32%). Almost all linear bursts had a single acceleration event (Table 3). As with inflectional bursts, depth change during linear bursts occurred in 3 main forms: descent (n = 5 bursts, 26%), horizontal (n = 6 bursts, 32%), and ascent (n = 5 bursts, 26%).

**DISCUSSION**

Efficient locomotion during breath-hold dives provides the advantage of increased available time at depth for locating and catching prey (Williams et al. 1996). Watwood et al. (2006) concluded that the high diving efficiencies of sperm whales *Physeter macrocephalus* may be related to efficient locomotion. Deep-diving mammals may conserve oxygen and increase dive durations by employing steady swimming or gliding, when possible, and avoiding energetically costly rapid accelerations. However, all tagged whales in the present study showed short events of highly active swimming at foraging depths, including bursts of speed (Fig. 2). During such bursts, they accelerated rapidly, turned with drastic deceleration, and swam up to 405 m. Here we discuss (1) bursts of speed as part of sperm whale feeding behavior and (2) the hunting process and strategy of sperm whales.

**Bursts of speed as part of sperm whale feeding behavior**

Toothed whales employ echolocation clicks to detect their prey and emit buzzes (i.e. a rapid decline in the click interval) to approach prey. Thus, emissions of buzzes have been used as an indicator of feeding events (e.g. Miller et al. 2004b, Watwood et al. 2006, Aguilar Soto et al. 2008, Johnson et al. 2008). By using acoustic tag data, the movements of toothed whales in relation to buzz emissions have previously been investigated. In the case of pilot whales, an increase in vertical speed concurrently with buzz emissions has been shown to reflect prey capture attempts (Aguilar Soto et al. 2008). Sperm whales exhibit buzzes in association with increased changes in body orientation, and such behaviors are thought to indicate final approaches and prey capture (Miller et al. 2004b). Both studies noted that buzz emissions correspond with increasing changes in body orientation or swim speed. Unfortunately, acoustic data were not available in our study. However, our results also showed that 87% of bursts (i.e. inflectional bursts) included rapid changes in body orientation. Additionally, bursts of speed occurred at depths >400 m, corresponding with depths where female and immature sperm whales were found to emit buzzes in the Atlantic Ocean, the Gulf of Mexico, and the Ligurian Sea (Watwood et al. 2006). Thus, our results strongly suggest that bursts of speed by the tagged whales recorded in our study are related to feeding events. However, we could not exclude other potential explanations; for example, that the bursts of speed indicate inter- or intra-specific competition. Whales may detect a prey over a wide range and sense another approaching whale. They then may increase their speed to reach the prey before the other whale does. If this behavior occurs, bursts of speed could also be interpreted in relation to feeding events. However, sperm whales probably do not compete against each other. Female sperm whales live in a long-term stable association with each other (Whitehead 2003). Female sperm whales disperse and form a ranked pattern while foraging, thus, possibly avoiding interference by other foraging animals in the same area (Whitehead 2003). Off the Ogasawara Islands, the encounter rate of other deep-
diving toothed whales (e.g. ziphiid whales) is considerably lower than that of sperm whales (authors’ unpubl. data). Therefore, all bursts observed in the present study do not seem to be caused exclusively by inter-specific competition. Another explanation for bursts could be predator avoidance. However, we consider the probability of this explanation to be quite low. This is because killer whales *Orcinus orca* are the main predators of sperm whales (Whitehead 2003) and usually dive to depths <200 m (Baird et al. 2005). However, the majority of bursts in the present study occurred at depths >400 m. Even if deep-water predators were distributed in our study area, the deceleration behavior observed during the bursts contradicts escape behavior.

Although our results indicate that the bursts of speed are feeding events, sperm whales may not always increase speed to capture their prey. The median number of bursts was 1 per dive (range: 0 to 6 per dive), and 33% of dives did not include any bursts.
In contrast, Watwood et al. (2006) reported that the number of clicks (buzzes) in sperm whales averaged 18 per dive in the Atlantic Ocean, the Gulf of Mexico, and the Ligurian Sea. Whitehead (2003) estimated that a female sperm whale consumes, on average, 87 squid per dive, based on stomach content analyses conducted by Clarke (1980). The ratio of bursts to total feeding events is unclear, because we were unable to measure all feeding events. However, the number of bursts is lower than these values. Therefore, prey capture may not always require bursts.

During bursts, tagged whales accelerated with stroking, and usually reached a speed of between 2 and 5 m s\(^{-1}\) (Fig. 3b). When a dive included 1 or 4 bursts, the energetic cost of the bursts accounted for 13 and 52% of the total cost of the dive, respectively (Fig. 2). We roughly estimated the locomotion costs of the burst based on the hydrodynamic drag that the whale experienced (Hind & Gurney 1997; see ‘Locomotion cost’ in the supplement for details), and the cost reaches a maximum of 5830 kJ. The mean calorific value is 4.3 \(\text{kJ g}^{-1}\) in wet weight for muscular squids and 2.3 \(\text{kJ g}^{-1}\) in wet weight for ammoniacal squids (Clarke et al. 1985). Using these values, we calculated the calories from a small ammoniacal squid weighing 200 g (cranchids), a muscular squid weighing 8 kg (ommastrephids), and an ammoniacal squid weighing 24 kg (architeuthids) to be 460, 34 400, and 55 200 kJ, respectively (typical weight of each squid preyed upon by sperm whales according to Whitehead 2003). The calorific value of the squids might be overestimated, because we did not consider digestibility. However, costly bursts should be compensated for by the consumption of relatively nutritious prey, such as large and/or muscular squids, to balance the energy budget. The variable maximum speeds of the bursts (Fig. 3b) suggest that sperm whales can adjust their pursuit speed according to the escape capabilities of their prey, which is most probably related to quality (in size and/or muscular content).

Some large and/or muscular squids were found in the stomach contents of sperm whales off Japan (Kawakami 1980). These squids were observed to be at approximately the diving depths of the whales in our field study, at the same time of year and in the same area. For example, an underwater camera recorded flying squids Ommastrephes bartrami (Lesueur) with muscular mantles between 400 and 800 m (Kubodera & Mori 2003), giant squids at 900 m, and Taningia danae from 240 to 940 m (Kubodera & Mori 2005, Kubodera et al. 2007). The maximum swim speeds of flying squids and T. danae were estimated from video data at 5.3 m s\(^{-1}\) (T. Kubodera unpubl. data) and 2.5 m s\(^{-1}\) (Kubodera et al. 2007), respectively. As pursuit speeds of predators are predicted to be a function of prey escape speeds (Wilson et al. 2002), the range of maximum speeds of bursts should correspond to the escape speeds of such actively swimming, large and/or muscular squids.

**Hunting process and strategy of sperm whales**

For large carnivores the process of hunting usually consists of locating, stalking, chasing, and capturing (Curio 1976). The swimming distances of almost all the bursts were less than about 200 m (mean: 120 m; Table 3). The maximum detection range of the usual clicks is 325 m for an individual squid with a mantle length of 25 cm (Madsen et al. 2007), therefore, the distance of the bursts was shorter than the estimated echolocation detection range. Thus, we infer that the bursts did not begin immediately when the whales detected their prey. During the approaching phases of some carnivores, an advance is made toward the unaware prey until the predator is within attacking range (Curio 1976, Bothma & le Riche 1989). Such an approach is termed stalking (Curio 1976). Stalking predators avoid noisy body movements that may alert their prey, and their stealthy advance is slow (Curio 1976, MacFarland 1981). Cephalopods are the main prey of sperm whales and are quite sensitive to low-frequency vibrations (Hanlon & Messenger 1997, Mooney et al. 2010). Furthermore, they respond to local water movements of up to several hundred Hz (Budelmann & Bleckmann 1988). Although the typical range of echolocation clicks in sperm whales is between 5 and 25 kHz (see Madsen et al. 2002 for details), the squid Loligo pealei, which is heavily preyed upon by toothed whales, does not respond to intense echolocation clicks from squid-eating toothed whales (Wilson et al. 2007). Thus, cephalopods probably cannot hear echolocation clicks, but detect the water movements caused by predators. Field observations have shown that squids detect and avoid the swimming-generated low-frequency cues of certain fish predators (Hanlon & Budelmann 1987). Our results show that gliding (duration: 57 ± 42 s), slow speed (1.2 ± 0.4 m s\(^{-1}\)), and inactive body movements were employed before bursts began, suggesting that the whales avoided body movements that might have signaled their presence to the prey. We suspect that the bursts occur within a certain distance during the approach and are sometimes preceded by stalking, which may increase the whale’s chances of prey capture (Mac-
Farland 1981) and reduce the cost of chasing. An alternative explanation for the gliding behavior just prior to bursts is that whales may be selecting a certain prey over other potential prey. Our data, however, are not sufficient to confirm either of these hypotheses. Increasing speed is a costly event; therefore, whales may limit this behavior until they are within target of a specific prey.

Tagged whales swam up to about 405 m during bursts, suggesting that the whales were chasing fleeing prey. At the end of a successful chase, hunting predators intersect a moving path of their prey to capture it. The prey can escape, even if the speed of predators intersect a moving path of their prey to capture it. The prey can escape, even if the speed of the prey (U_{prey}) is lower than the speed of the predator (U_{pred}), if its turning radius (R_{pred}) is smaller than that of the predator (i.e. U_{pred}^2/R_{pred} < U_{prey}^2/R_{prey}; Alexander 2003). Therefore, if the prey turns at the last possible moment, the predator cannot intercept the prey even if the predator turns. However, the predator can intercept the prey by turning if turning of the prey occurs too soon (see Fig. 1.2 in Alexander 2003 for details). Thus, predators should adjust their distance to prey and the time of turning. The majority of bursts were inflectional bursts with turning (87%), suggesting that sperm whales may make sharp turns to intercept their prey. The remaining 13% of bursts were linear bursts without turning. For this type, the whales did not need to make sharp turns probably because the prey was not as maneuverable. The two different major types of bursts (inflectional bursts with turning and linear bursts without turning) suggest that sperm whales change their hunting behavior depending on the maneuverability of their prey.

Predator–prey interactions shape the hunting techniques of predators (Curio 1976). Ziphiids whales forage in a similar depth range to sperm whales (Hooker & Baird 1999, Tyack et al. 2006), and they consume mesopelagic squids (Clarke 1996). Whitehead (2003) suggested that sperm whales have wider feeding niche breadths than the northern bottlenose whale {Hyperoodon ampullatus} (Forster) and Cuvier’s beaked whale {Ziphius cavirostris} (G. Cuvier), because the ratio of the number of genera to the total number of lower beaks in stomach contents is higher in sperm whales than in the other 2 species. In mammalian carnivores, the ability to hunt a large variety of prey types is influenced by the predator’s flexibility in their killing methods (Rowe-Rowe 1978, Ben-David et al. 1991). Therefore, it is likely that sperm whales have a variety of hunting methods to catch a wider variety of prey than other deep-diving species. Our results show that 2 major types of bursts exist, and there are various patterns in 3D dive profiles, swim speed, and body movement within and between types (Table 3, Fig. 5). Therefore, these results suggest that sperm whales change their hunting behavior according to prey type and prey response, which likely differs by prey type. Teloni et al. (2008) showed that the duration and occurrence of buzzes of male sperm whales varied between foraging depths, suggesting that the whales were foraging on several types of prey with different patterns of mobility. We suspect that sperm whales are capable of using a diverse range of hunting techniques, which makes it possible for them to have wider feeding-niche breadths. The large aerobic capacity of sperm whales allows them to utilize variable hunting methods, including active pursuit, during long-duration dives in order to ensure a broad dietary range.

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LITERATURE CITED


Ben-David M, Pellis SM, Pellis VC (1991) Feeding habits and predatory behavior in the marbled polecat {Vormela


Teloni V, Johnson MP, Miller PJO, Madsen PT (2008) Shal-


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