



Southern elephant seal foraging success in relation to temperature and light conditions: insight into prey distribution

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ABSTRACT: The distribution of southern elephant seal *Mirounga leonina* prey encounter events (PEEs) was investigated from the foraging behaviour of 29 post-breeding females simultaneously equipped with a satellite tag, a time–depth recorder and a head-mounted accelerometer. Seal diving depth and PEE were related to water temperature at 200 m (T_{200}), and light level at the surface (L_0) and at depth. Approximately half (49%) of all dives were located in waters encompassed between the southern Antarctic Circumpolar Current Front and the Polar Front. Seals dived significantly deeper during the day than at night. Diving and PEE depth increased with increasing T_{200} and for a given T_{200} according to L_0 and the percentage of surface light reaching 150 m. On average, 540 PEEs per day were recorded. Seals exhibited more PEEs per unit of time spent diving during the twilight period compared with at night, and were least successful during daylight hours. Elephant seals forage in T_{200} ranging between -1 and 13°C ; however, few PEEs were recorded at depths shallower than 400–500 m at night when the T_{200} exceeded 8°C . The diet of female Kerguelen elephant seals appears to be dominated by myctophids (lanternfish), and according to the average mass of their most likely myctophid prey (9 g, *Electrona calisbergi* and *E. antarctica*; 30 g *Gymnoscopelus nicholsi* and *G. piabilis*), we estimate that seals consumed 4.8–16.1 kg of fish daily. Despite lower catch rates in warmer waters, no relationship was found between the mean T_{200} at the scale of the foraging trip and daily or absolute mass gain, suggesting that elephant seals are compensating for lower catch rates by consuming larger/richer prey items in those waters.

KEY WORDS: Southern elephant seals · Foraging success · Light · Temperature · Diving behaviour · Prey

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INTRODUCTION

An increasing number of studies show that, along with the measurement of concomitant oceanographic data, the behaviour of top marine predators can provide valuable insights into the habitat in which they forage (e.g. Biuw et al. 2007, 2010). Accordingly, the

development of miniaturised electronic devices has opened a new era of research on the foraging behaviour of marine predators and allows the acquisition of huge quantitative data sets of behaviour from animals moving freely in their natural environment (Robert-Coudert & Wilson 2005). Behavioural (location, diving depth, prey capture) and environmental

(temperature, salinity, light, fluorescence) parameters can be continuously recorded by high-resolution multi-channel data loggers. This enables studies of the conditions under which animals live, as well as the ability to monitor their environment (Boehlert et al. 2001).

Among top marine predators, air-breathing diving species such as seals are particularly well-suited for the bio-acquisition (i.e. bio-logging) of movement, behavioural, physiological as well as environmental data using miniaturised animal-attached tags (Rutz & Hays 2009). Their large size allows them to carry electronic devices with minimal disturbance, and researchers can handle them while they are on land in their breeding colonies. Furthermore, while investigating at-sea seal ecology, measurements of oceanographic parameters within the water column have also been obtained. Consequently, foraging, diving predators have provided new and original information about physical (Charrassin et al. 2008, Costa et al. 2008, Roquet et al. 2009) and biological oceanography (Guinet et al. 2013) in often inaccessible regions.

In recent years, one of the most exciting applications of bio-logging is the determination of the distribution and availability of prey of which we have very little knowledge. Until recently, the main limitation was an *in situ* measure of direct prey ingestion. Early devices to record predation events of seabirds and marine mammals included stomach (Wilson et al. 1992) and oesophageal temperature sensors (Ancel et al. 1997, Charrassin et al. 2001), which detected temperature changes induced by prey ingestion. However, these temperature loggers have certain limits, as they are restricted to endotherms feeding on ectothermic prey and are quite invasive. Furthermore, stomach temperature recorders fail to detect multiple rapid ingestions of small prey and are often regurgitated prematurely (Ropert-Coudert et al. 2004, Liebsch et al. 2007). External loggers such as the Inter-Mandibular Angle Sensor (IMASEN) record the mouth-opening angle of predators (Wilson et al. 2002), but are sometimes difficult to apply on animals with flexible lips (Ropert-Coudert et al. 2004, Liebsch et al. 2007). Recent studies in free-ranging diving pinnipeds have overcome this methodological issue by using head- or jaw-mounted accelerometers to detect prey encounter events (PEEs) (Suzuki et al. 2009, Naito et al. 2010, 2013, Viviant et al. 2010, Gallon et al. 2013, Iwata et al. 2012). This has enabled investigations into fine-scale temporal and spatial changes in the foraging of an increasing number of pinniped species, including the southern elephant

seal *Mirounga leonina* (SES hereafter) – the largest and deepest-diving pinniped.

Both its abundance and body mass make the SES a major consumer of Southern Ocean (SO) marine resources (Guinet et al. 1996, Hindell et al. 2003). It was previously thought that SES primarily fed on squid, but a recent isotopic investigation challenged this view and, instead, indicated that female SES from the Kerguelen Islands mainly feed on myctophid fish (i.e. lanternfish) (Cherel et al. 2008). Indeed, myctophid (the abundant *Electrona antarctica*, *E. carlsbergi* and *Gymnoscopelus nicholsi*) otoliths have been found in stomachs of SES from the Kerguelen sector (Slip 1995) and elsewhere (Daneri & Carlini 2002).

Satellite tracking data show that SES are broad-ranging animals during their post-breeding and post-moult migrations, with adult females from the Kerguelen Islands foraging mainly in oceanic waters of the Antarctic and Polar Frontal Zones (Bailleul et al. 2010). While at sea, SES dive repeatedly to mesopelagic depths (300–500 m, up to 2000 m) and tend to follow the nycthemeral vertical migration of their mesopelagic prey, diving generally deeper during the day (Hindell et al. 1991, McIntyre et al. 2010). In a recent study, Jaud et al. (2012) found that light level at depth explained 70% of the variation in diving depth of the SES during daylight hours while no significant effect was detected at night. Furthermore, light level at depth in oceanic waters was explained to a large extent by phytoplankton concentration: the higher the phytoplankton concentration, the less light and the shallower the dives of SES (Jaud et al. 2012).

The objectives of this study were to investigate the variation in diving behaviour and prey encounter event rates of female SES to better characterise their prey in terms of likely size and distribution according to environmental conditions such as sea temperature and light level along a north–south gradient ranging from subtropical waters to Antarctica.

MATERIALS AND METHODS

Animal handling and instrumentation

During October/November each year from 2008 to 2012, post-breeding SES females were captured on the Kerguelen Islands (49° 20' S, 70° 20' E). Individuals were anaesthetised using a 1:1 combination of tiletamine and zolazepam (Zoletil 100), which was injected intravenously (McMahon et al. 2000). In general, post-breeding females forage at sea for durations

varying from 65 to 80 d, departing late October before returning to land throughout January to moult. A total of 34 SES females were equipped with a range of satellite tags and data loggers (see Table 1).

Eighteen seals were equipped with GPS loggers in combination with Argos satellite loggers and archival data loggers (SPLASH10-Fast-Loc GPS, Wildlife Computers). SPLASH10 devices transmitted Argos location data and collected GPS location data. The sampling interval of GPS locations was set to a minimum of 20 min, slightly shorter than the average dive duration of post-breeding females (Hindell et al. 1991, McIntyre et al. 2011a), to maximise the chance of obtaining a GPS location every time individuals surfaced between dives. An additional logger, a time–depth recorder (TDR) included in the SPLASH10 devices, collected and archived pressure, light and temperature levels at 1 or 2 s intervals.

Sixteen seals were equipped with conductivity–temperature–depth satellite-relay data loggers (CTD-SRDLs, Sea Mammal Research Unit, University of St Andrews) combined with either TDR loggers (MK9, Wildlife Computers), set to sample and archive pressure, light and temperature levels every 1 or 2 s, or TDR–accelerometer data loggers (MK10-X, Wildlife computers), sampling acceleration as well as pressure. The acceleration was measured on 3 axes: longitudinal (surge), vertical (heave) and lateral (roll) axes were used to separate dynamic accelerations from gravitational acceleration. Acceleration was sampled at 16 Hz and pressure every second.

The SPLASH 10, MK10-X and MK9 TDR loggers were set to sample depth (0–1500 m, ± 1 m), water temperature (-40 to $+60^\circ\text{C}$, $\pm 0.1^\circ\text{C}$) and light (5×10^{-2} to 5×10^{-12} W cm^{-2} in blue wavelength) every 1 or 2 s. Light values are converted on-board via a log transformation to compress the light measurements to a 3 digit value.

Data loggers were glued on the head of the seals, using quick-setting epoxy (Araldite AW 2101, Ciba), after cleaning the fur with acetone. Upon returning from their foraging trip, females were located via Argos data, recaptured and weighed, and loggers were retrieved.

Data processing and analysis

Female SES from the Kerguelen Island population forage mainly in pelagic waters but also to a lesser extent on the Kerguelen and Antarctic continental shelves (i.e. the main foraging habitat of Kerguelen Island sub-adult and adult males; Bailleul et al. 2010, Dragon et al. 2010, Authier et al. 2012). As part of this work we focused on the pelagic portion of the tracks by excluding data from areas where water depths were < 1000 m (Fig. 1), based on National Geophysical Data Centre ETOPO2 Global 2 Elevations (www.ngdc.noaa.gov/mgg/global/etopo2.html), to exclude all locations on the Kerguelen and Crozet plateaus where SES are known to conduct benthic dives (Bailleul et al. 2010).

Pressure sensor data were analysed from MK9 and SPLASH10 TDRs to obtain dive depth profiles. For the purpose of this study, and based on the sensor's absolute accuracy, only excursions below a depth of 15 m were analysed. Bottom depth, light and the temperature profiles associated with each dive were extracted (see Dragon et al. 2012 for details). Each dive was divided into a descent, ascent and bottom phase. The bottom phase corresponds to the period between the end of the descent and the beginning of the ascent. The different dive phases were defined according to Dragon et al. (2012) using a custom-written MATLAB code (version 7.0.1; available on request). In the present study, the diving depth is defined as the mean depth during the bottom phase of the dive.

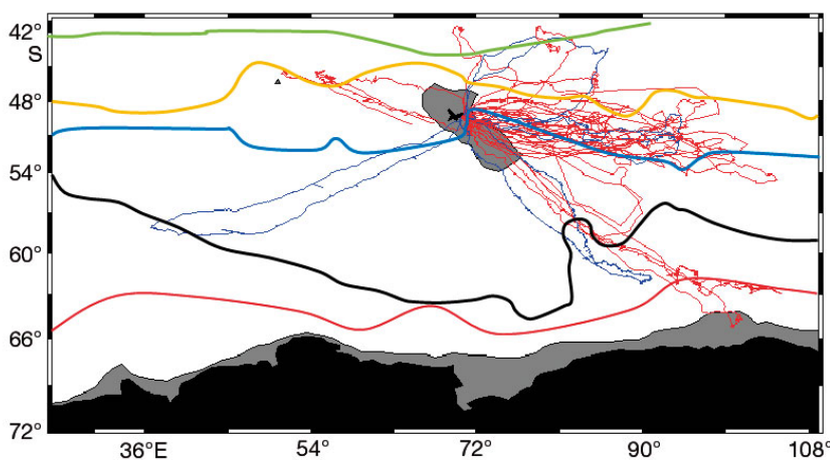


Fig. 1. Tracks of the 29 post-breeding southern elephant seal females for which high-resolution data were recovered. The blue section of track corresponds to the period when accelerometry data were collected. The main frontal structures are reported from south to north: Antarctic divergence (red), Southern Antarctic Circumpolar Current Front (black), Polar Front (blue), Subantarctic Front (orange) and the Subtropical Front (green). Shelf area from the coast to the -1000 m isobaths is indicated in grey

Data from the accelerometer were processed according to Viviant et al. (2010) and Gallon et al. (2013) using custom-written MATLAB code (available on request). Identifying individual feeding events from the accelerometry data was performed according to the following procedures. Firstly, the 3 accelerometer time series were filtered with a high-pass filter of 0.33 critical frequency for data sampled at 16 Hz to remove the remaining noise corresponding to swimming movement, highlighting the peaks in acceleration that represent head movements. Rapid head movements were used to identify possible PEEs. The 0.33 filter was defined visually on power spectral density of acceleration on the 3 axes.

Secondly, the standard deviation along a fixed 1 s window over each time series axis was calculated to sum accelerations at a 1 s scale. The standard deviation along a 5 s moving window, over each standard deviation time series, was then calculated for each axis to highlight extreme standard deviation and consequently significant accelerations. The threshold distinguishing those extreme events was defined using the *kmeans* function (MATLAB, tool box statistics) and was calculated for each seal and for each axis independently. Only head movement events that could be detected simultaneously on the 3 axes were considered as a true PEE; others were considered to be related to dives where animals did not alter their swimming behaviour whilst travelling underwater. Continuous values at 1 Hz above the threshold were considered as unique PEEs. Events separated by periods longer than 1 s below the threshold value for any given axes were considered as a different PEE. The number, timing and depth location of PEEs were determined for each dive. In the present study, a PEE does not mean that the seal was feeding with certainty, but should be considered as a relative index of a prey encounter during the dive.

From these data we then calculated PEE per unit time (PEE rate) at 2 scales: (1) the complete dive and (2) the bottom phase of the dive. The latter removes the effect of the vertical accessibility of the prey, since a seal has to undertake a longer vertical transit when it dives deeper and consequently has less time to spend at the bottom of the dive (i.e. where most PEEs are expected to occur).

For each animal, the temperature profile associated with each dive was extracted from MK9, TDR10-X and SPLASH 10 TDR data. At any given latitude, but different longitudes, water masses with very different temperature profiles can be observed depending on the location of the main frontal structures (see Fig. 1). Dives were therefore ranked re-

gardless of their latitude, in a decreasing order, according to their water temperature at 200 m (T_{200}), to reflect a general north–south gradient. Profiles were then regrouped into 1°C T_{200} classes. Each 1°C class was defined according to its lower boundary (i.e. –1°C defining the [–1°C; 0°C] temperature class). Then, for all the dives belonging to a given T_{200} class, the mean temperature for 5-m depth bins was calculated over the whole depth range. This provides a mean temperature profile for a given temperature scenario at 200 m. These mean temperature profiles were used for figure representation and statistical comparisons between the different 1°C T_{200} classes: day–night difference in the mean diving depth and comparison of PEEs per minute spent diving or spent at the bottom of the dive. Dives shallower than 200 m were allocated to a T_{200} category according to the T_{200} encountered during previous and/or following dives with a diving depth exceeding 200 m.

The SO fronts are narrow regions of sharp horizontal gradients of water properties that mark the boundaries of different water masses (Fig. 1). These structures are defined by representative values of temperature and salinity at approximately 200 m depth, where each front is generally best marked. The water mass the seal was diving in was identified using the T_{200} , as indicated by Park et al. (1991, 1998) and Orsi et al. (1995). Going north from the Antarctic continent, the Antarctic Divergence (AD) has been identified with the most shoreward penetration of the 0.8°C isotherm at depths below the Winter Water. The Southern Antarctic Circumpolar Current Front (SACCF) is defined by a temperature of 1.6°C and a salinity of 34.6 PSU at 200 m depth (Park et al. 1998). The Polar Front (PF) is approximated by the northern limit of the subsurface temperature minimum of 2.8°C at the 100–300 m depth layer (Park et al. 1993, Belkin & Gordon 1996). Although the Subantarctic Front (SAF) is less well defined, values of 6.8°C are considered as subsurface expressions of this front in the Indian sector (Park et al. 1993) and, therefore, we used 7°C at 200 m as the SAF limit. Values of 10.8°C at 200 m depth are adopted to define the Subtropical Front (STF) (Park et al. 1993); therefore, we used 11°C at 200 m as the STF limit in this study. We defined the Polar Frontal Zone as the area located between the STF (northern limit) and the PF (southern limit), and the Antarctic Circumpolar Current (ACC) zone is defined as the area between the SAF and the SACCF.

Light levels within the water column vary greatly according to a number of factors, such as sun angle, cloud cover and phytoplankton concentration within oceanic waters. This latter factor explains approxi-

mately 50% of the variation of light attenuation within the water column (Jaud et al. 2012). For each dive record from the MK9, MK10 X and SPLASH10 TDR data, we calculated the level of light available at each depth measurement along the entire dive. Surface light level (L_0) was calculated as the mean light reading during the surface interval after the dive. The percentage of surface light reaching 150 m (L_{150}/L_0), corresponding in general to the euphotic depth in the SO (Knox 2007), was calculated during the ascent phase of the dive prior to surfacing. To assess the effect of phytoplankton and particle concentration on light level available at depth, light at depth was expressed as a percentage of L_0 .

Each dive was attributed to one of 3 periods: day, night or twilight. This was defined according to the solar angle, taking into account the geographical location of the seal provided by the Argos/GPS data and the time at the beginning of the dive. Day was defined when the sun was above the horizon (0°); twilight, when the sun was between 0° and 6° below the horizon; and night, when the sun was more than 6° below the horizon. The twilight period used in our study corresponds to civil dawn and dusk, i.e. the time at which there is enough light for objects to be distinguishable in the air and the light is therefore likely to influence the vertical distribution of biological organisms within the water column. The sun angle according to the geographical location and time was calculated using the *suncycle* function in MATLAB.

Statistical analyses

The relationships between (1) bottom duration according to dive duration and diving depth, (2) dive duration according to T_{200} , diving depth, the number of days elapsed since departure from the colony, L_0 , L_{150}/L_0 for day, night and twilight periods and (3) the mean depth at the bottom of the dive and PEE depth according to T_{200} , L_0 and L_{150}/L_0 for day and night (the same analysis was performed north and south of the SAF, i.e. for T_{200} higher and lower than 7°C) were analysed using a linear mixed model (package nlme in R; R Development Core Team 2009). Light level analyses were performed only during day and night. Indeed during twilight, light level varies to a great extent and is known to induce very large variation in elephant seal diving behaviour. Instead, this study aimed to assess whether female SES diving behaviour responds to small changes in L_0 and L_{150}/L_0 during either the day or the night. The variation in PEE according to diving depth, with dive duration or bot-

tom time accounted for in the model, was analysed using a generalised linear mixed model with multivariate normal random effects, using penalized quasi-likelihood (package GlmmPQL in R; R Development Core Team 2009) on night, day and twilight separately. Individual seal was included as a random factor to take into account inter-individual variability. An autoregressive variance-covariance matrix (cor AR1 in R; R Development Core Team 2009) representing first-order autocorrelation structure was used to model the serial correlation among observations for each individual (Zuur et al. 2009). All variables were standardised (centred and scaled) to facilitate model convergence and to be able to compare the respective contribution of the predictors (Zuur et al. 2009). Using GLmmPQL, model selection can only be performed according to p-values. We implemented a stepwise procedure, with the threshold set at $p = 0.05$. Data are presented as means \pm SD. The variation in PEE rate throughout the dive and at the bottom of the dive and according to T_{200} classes was compared between day, night and twilight using a Wilcoxon test.

RESULTS

The mean mass and length of the 34 post-breeding females prior to their departure were 296 ± 54 kg and 242 ± 15 cm, respectively. Among these 34 equipped seals, all but 3 were recaptured and high-resolution diving data were successfully recovered for 29 of 31 remaining females. The mean foraging trip duration was 82 ± 9 d ($n = 31$; Table 1), the mean recovery mass was 363 ± 55 kg ($n = 25$); therefore, the mean overall mass gain was 75 ± 36 kg while the daily mass gain was 0.9 ± 0.4 kg per day spent at sea ($n = 25$; Table 1).

The tracks of the 29 SES females for which we had high-frequency measurements of pressure, temperature and light are shown in Fig. 1. These tracks extended from the subtropical zone to the Antarctic shelf. On average, SES performed 66 ± 9 dives daily. Among those, PEE data were obtained for 12 SES along their track (see Fig. 1, Table 1). An example of data obtained for one complete foraging trip is presented in Fig. 2.

Dive, light and temperature distribution

Seals ranged from the AD to the STF front (Figs. 1 & 3). Among the 106 313 dives performed by the 29 post breeding SES females, 62.2, 29.9 and 7.9%

Table 1. Female southern elephant seal identity, deployment and recovery dates, foraging trip duration, mass, standard body length, mean 200 m temperature (T_{200}) and equipment type deployed

Seal ID	Deployment	Recovery	Trip (d)	Departure mass (kg)	Return mass (kg)	Mass gain (kg)	Daily mass gain (kg)	Length (cm)	T_{200} (°C)	Argos-CTD	Splash-10-F	TDR-MK9	TDR-MK 10-X
2008-8	13.10.2008	05.01.2009	84	231.6				225.0	3.0		X		
2009-1	15.10.2009	12.01.2010	89	273.5	294.0	20.5	0.2	225.0	3.5	X		X	
2009-2	15.10.2009	30.12.2009	76	267.5	310.0	42.5	0.6	235.0	5.9	X		X	
2009-3	16.10.2009	11.01.2010	87	268.0	300.0	32.0	0.4	228.0	3.0	X		X	
2009-4	16.10.2009	06.01.2010	82	269.5	370.0	100.5	1.2	230.0	3.1	X		X	
2009-5	17.10.2009	12.01.2010	87	397.5				276.0	0.4		X		
2009-6	19.10.2009	05.01.2010	78	319.5	440.0	120.5	1.5	258.0	5.6		X		
2009-8	20.10.2009	20.01.2010	92	314.5	395.0	80.5	0.9	250.0	1.3	X		X	
2009-9	24.10.2009	10.01.2010	78	272.5	332.5	60.0	0.8	237.0	2.8		X		
2009-10	25.10.2009	20.01.2010	87	321.5	370.0	48.5	0.6	249.0	2.8		X		
2009-11	25.10.2009	12.01.2010	79	300.0	375.0	75.0	0.9	219.0	2.8		X		
2010-8	15.10.2010	29.01.2011	106	237.5	295.0	57.5	0.5	238.0	1.1	X		X	
2010-9	26.10.2010	11.01.2011	77	379.5	430.0	50.5	0.7	255.0	3.4	X		X	
2010-10	16.10.2010	15.01.2011	91	214.0	321.5	107.5	1.2	231.0		X		X	
2010-11	18.10.2010	03.01.2011	77	314.5	392.0	77.5	1.0	260.0	3.3	X		X	
2010-12	16.10.2010			427.0				280.0		X		X	
2010-13	15.10.2010	23.01.2011	100	384.8	448.0	63.2	0.6	261.0	1.6	X		X	
2010-14	18.10.2010	15.01.2011	89	283.5				236.0		X		X	
2010-15	24.10.2010	02.01.2011	70	330.5				270.0	1.2	X	X	X	X
2010-18	26.10.2010	03.01.2011	69	331.0	352.5	21.5	0.3	246.0	7.5		X		X
2010-19	31.10.2010	20.01.2011	81	395.0				266.0	1.2		X		X
2010-20	01.11.2010	15.01.2011	75	342.8	481.0	138.2	1.8	255.0	5.3		X		X
2010-21	18.11.2010	02.01.2011	45	376.5	425.0	48.5	1.1	218.0	1.0		X		X
2011-14	25.10.2011	26.01.2012	93	252.0	305.0	53.0	0.6	240.0	3.2	X			X
2011-16	26.10.2011	20.01.2012	86	255.0	425.0	170.0	2.0	254.0	3.0		X		X
2011-17	26.10.2011	19.01.2012	85	225.0	329.0	104.0	1.2	225.0	1.8	X			X
2011-18	26.10.2011	17.01.2012	83	245.0	327.0	82.0	1.0	238.0	2.8		X		X
2011-21	28.10.2011	31.01.2012	95	245.0				225.0	4.5		X		X
2011-22	28.10.2011	20.01.2012	84	296.0	367.0	71.0	0.8	256.0	2.0	X			X
2011-26	30.10.2011	26.01.2012	88	255.0				232.0	2.8		X		X
2011-27	30.10.2011	16.01.2012	78	236.0	307.0	71.0	0.9	235.0	8.1		X		X
2011-28	30.10.2011	14.01.2012	76	249.0	328.0	79.0	1.0	240.0	3.6		X		X
2011-29	03.11.2011			262.0				230.0			X		X
2011-30	27.10.2011			300.0				240.0		X			X
Mean			83	296.2	366.4	75.4	0.9	243.0	3.2				
SD			11	56.5	56.3	35.9	0.4	16.4	1.9				

dives were performed during day, night and twilight, respectively. The mean dive duration and mean diving depth were 21.4 ± 4.8 min and 519.2 ± 208.3 m for day; 16.7 ± 4.2 min and 384.4 ± 199.2 m for night; and 18.6 ± 3.8 min and 391.5 ± 196.3 m for twilight, respectively. The bottom duration of a dive was found to increase with dive duration but to decrease with increasing diving depth (Table 2). Dive duration increased with the number of days elapsed since departure from land. T_{200} exercised a different effect between day and night. During the day, for a given diving depth, seals increased their dive duration with increasing T_{200} while a negative effect was found at night and no relationship for twilight (Table 3).

The vast majority of dives (88.8%) were concentrated in water masses having a T_{200} encompassed between the AD and the SAF (Fig. 3). However, it is worth noting that 49.0% of the dives were distributed within water masses with a T_{200} ranging between 1 and 3°C, revealing that while foraging within the ACC zone, post-breeding female SES favoured water masses located primarily south of the PF. Approximately one-third (35.5%) of the dives were located in water masses encompassed between the PF and SAF. Only 6.0% of the dives were between the SAF and STF, with just 3.5% just north of the STF and 1.8% south of the AD (Fig. 3).

Diving depth increased with increasing L_0 , L_{150}/L_0 , as well as T_{200} both during day and night. Daytime

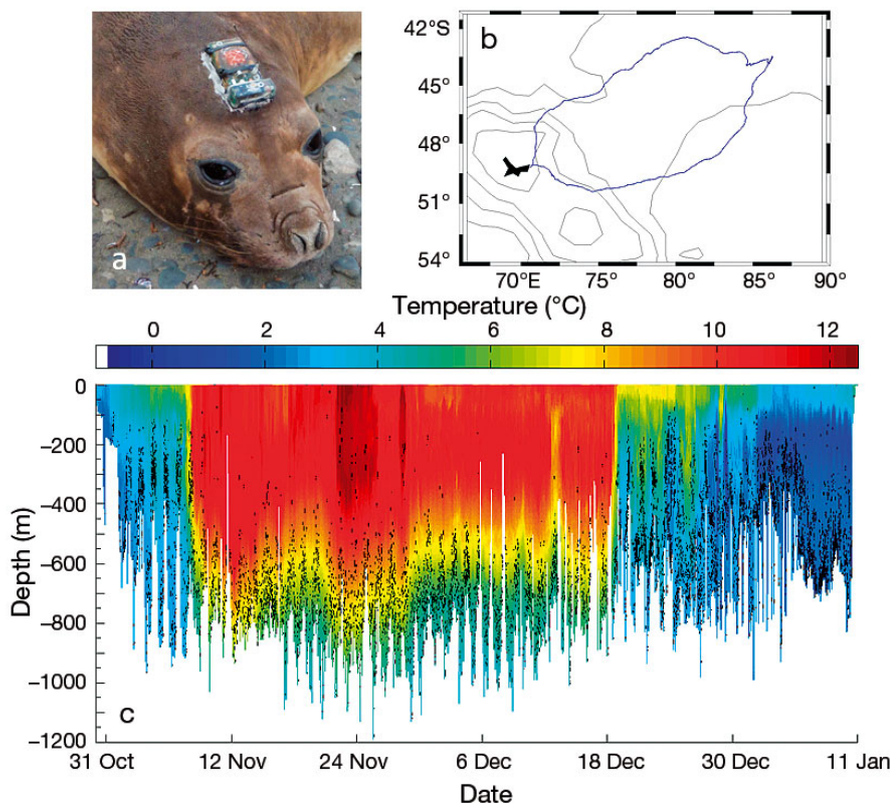


Fig. 2. An example of one female southern elephant seal (a) equipped with a SPLASH 10 tag combined with a head-mounted accelerometer enabling the reconstruction of the (b) seal track as well as (c) the distribution of prey encounter events (PEEs; black circles) according to depth and temperature monitored along the track of the animal

PEE depth increased with increasing L_0 , L_{150}/L_0 and T_{200} , while no effect of T_{200} was detected at night (Table 4).

The difference in mean diving depth between day and night varied more to the south than to the north of the SAF (Fig. 4). Consequently, there was less difference between the depths of day and night dives as the individuals headed towards the STF (Fig. 4). This was particularly noticeable for T_{200} exceeding 7°C (i.e. north of the SAF), for which PEE depth tended to increase rapidly with increasing T_{200} , during both the day and the night (Fig. 4). Indeed, mean diving depth increased significantly with increasing T_{200} during the day but not at night south of the SAF ($T_{200} < 7^\circ\text{C}$). The same significant effect of T_{200} on diving depth was found north of the SAF ($T_{200} > 7^\circ\text{C}$; Table 5), but during both day and night. Furthermore, diving depth was positively related to increasing L_0 north and south of the SAF during both day and night. In addition, diving depth was also related to L_{150}/L_0 during day and night south of the SAF and only at night north of the SAF (Table 5).

Variation in foraging success

For the 12 SES equipped with head-mounted accelerometers, the PEE

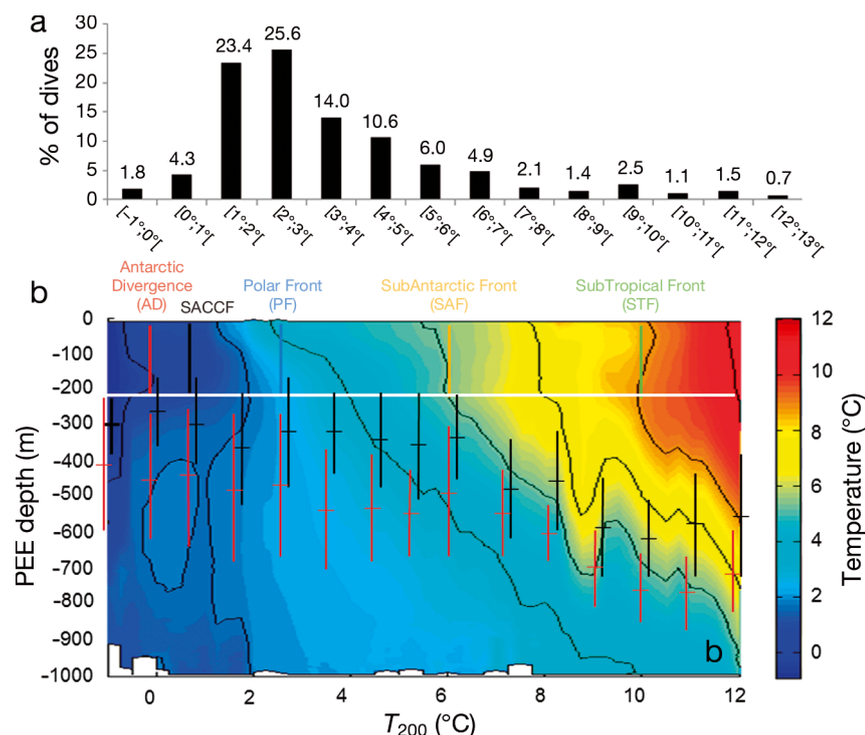


Fig. 3. A temperature transect reconstructed according to the temperature profiles ranked according to temperature at 200 m (T_{200}). (a) The percentage of dives for each 1°C temperature class. Number on top of bar is the % of dives observed for each class. (b) The main frontal structures are located along this transect according to their T_{200} signature (white line, see Fig. 1 for details). Mean (\pm SD) prey encounter event (PEE) depth during day (red) and night (black) obtained from the 12 seals equipped with head-mounted accelerometers for each 1°C temperature class overlaid on the temperature transect constructed from the data collected by the 29 seals. SACCf: Southern Antarctic Circumpolar Current Front

events were extracted along the track (time) and related to the corresponding pressure and T_{200} measurements (see Fig. 3). A total of 204 045 PEEs were detected, with 63.3, 26.1 and 10.6% occurring during day, night and twilight, respectively. A total of 73, 14 and 13% of the PEEs took place at the bottom, during the descent and during the ascent of the dive, respectively. When considering the PEE per unit of time spent diving or spent at the bottom of the dive, seals were found to be significantly more efficient during twilight compared with at night (dive: $W = 7\,075\,193$, $p < 0.001$; bottom: $W = 7\,680\,996$, $p < 0.001$) or during the day (dive: $W = 44\,313\,536$, $p < 0.001$; bottom: $W =$

49 752 533, $p < 0.001$), and they were also more efficient during the night compared with during the day (dive: $W = 19\,339\,935$, $p < 0.001$; bottom: $W = 19\,852\,229$, $p < 0.001$; see Fig. 5).

Drift dives, which are considered essentially as non-foraging dives and function as periods of recovery (Crocker et al. 1997), had very few prey captures (0.39 ± 1.93 PEEs per drift dive, $n = 501$). Excluding these dives, an average of 8.2 ± 6.6 PEEs were recorded per dive ($n = 25\,761$). The mean PEE depth was 467 ± 211 , 304 ± 138 and 308 ± 151 m for day, night and twilight, respectively.

A visual examination of PEE depth distribution reveals that it varied between day and night and according to T_{200} (Fig. 6). For T_{200} ranging between 0 and 7°C , the PEE tended to exhibit a bimodal distribution both during day and night. However, both modes are shallower at night compared with the daylight hours. In contrast, for T_{200} higher than 7°C , individuals were capturing prey much deeper both at night and during the day. Furthermore, PEE depth distribution was unimodal and very deep (500–800 m), which tended to be slightly shallower at night compared with during daylight hours.

The number of PEEs for a complete diving event was negatively related to both dive duration and diving depth regardless of the time period (day, night or twilight). The number of PEEs at the bottom of the dive was negatively related to diving depth during both day and

Table 2. Bottom duration according to dive duration and diving depth. *** $p < 0.001$

Model	Parameter	Estimate	<i>t</i>
Bottom duration	Dive duration	307.65	858.06***
	Diving depth	-271.94	-800.29***

Table 3. Dive duration of female southern elephant seals according to 200 m temperature (T_{200}), diving depth and number of days elapsed since departure for day, night and twilight. ** $p < 0.01$; *** $p < 0.001$

Model	Parameter	Day		Night		Twilight	
		Estimate	<i>t</i>	Estimate	<i>t</i>	Estimate	<i>t</i>
Dive duration	T_{200}	18.73	7.31**	-31.42	-7.30**	-	ns
	Diving depth	35.01	32.27***	40.71	26.53***	45.13	15.67***
	Days since departure	67.49	31.62***	61.89	24.75***	63.84	14.50***

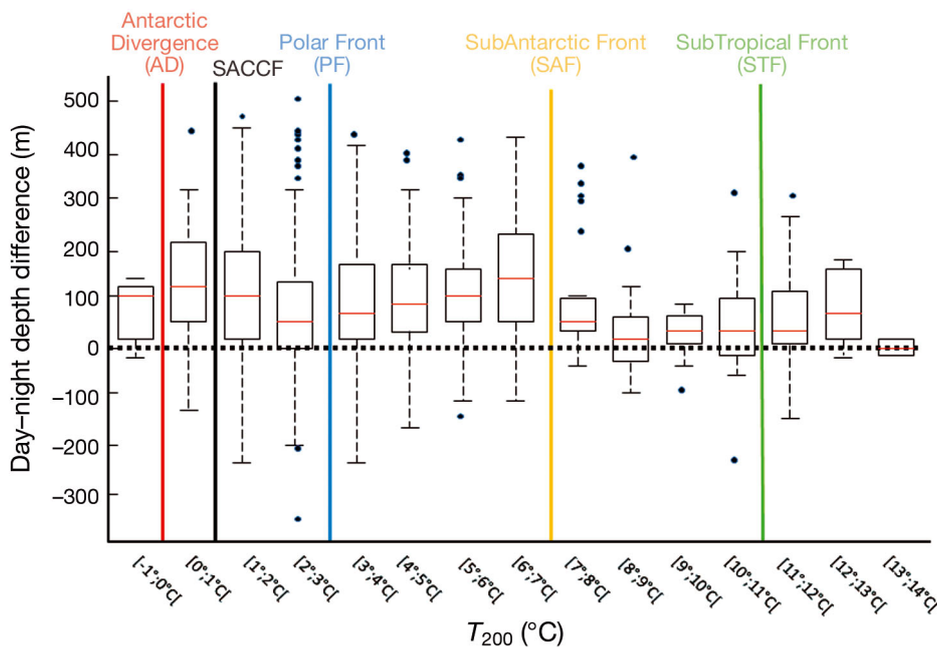


Fig. 4. Box plot of the mean depth differences between day and night dive according to each 1°C temperature at 200 m (T_{200}) class. Horizontal lines represent the median, the box represents the 0.75 to 0.25 quartiles, and whiskers represent 1.5 times the interquartile range (IQR). Small circles represent outliers (i.e. values exceeding 1.5 times IQR). SACC: Southern Antarctic Circumpolar Current Front.

Table 4. Variation in diving depth and prey encounter event (PEE) depth of female southern elephant seals according to surface light (L_0), the percentage of surface light reaching 150 m (L_{150}/L_0) and 200 m temperature (T_{200}), for day and night. *** $p < 0.001$

Model	Parameter	Day		Night	
		Estimate	<i>t</i>	Estimate	<i>t</i>
Diving depth	L_0	36.81	25.66***	25.57	10.54***
	L_{150}/L_0	30.80	19.64***	32.25	15.97***
	T_{200}	32.83	16.81***	29.98	11.68***
PEE depth	L_0	36.76	28.32***	–	ns
	L_{150}/L_0	38.94	26.55***	16.16	9.87***
	T_{200}	37.49	18.90***	31.43	14.85***

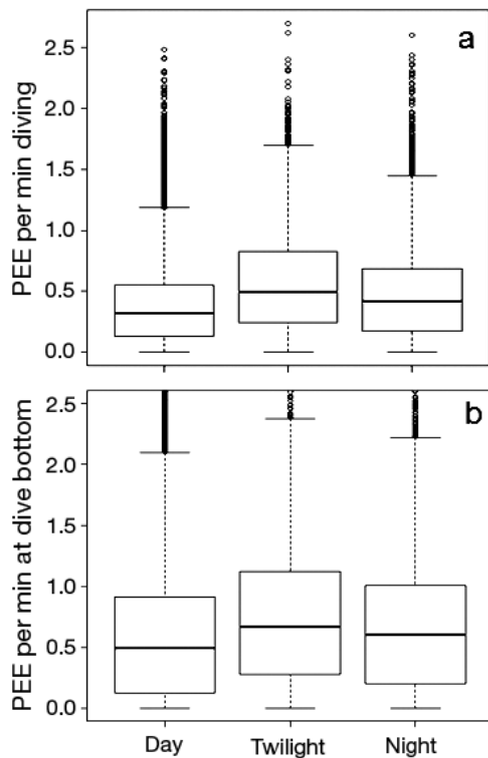


Fig. 5. (a) Box plot of prey encounter events (PEE) per minute spent diving for day, night and twilight periods. (b) Box plot of PEE per minute spent at the bottom of the dive for day, night and twilight periods (see text for details). See Fig. 4 for further details

night, but was positively related to the duration of the bottom phase of the dive during day and twilight periods (no effect was detected at night). Furthermore, PEEs for a complete dive and at the bottom of the dive were negatively related to the L_{150}/L_0 , with no effect of L_0 , during the day but positively related to both L_0 and L_{150}/L_0 during night and twilight (Table 6).

PEEs per unit of time spent diving or at the bottom of the dive were also found to vary significantly according to the water masses visited by the SES. Compared to what is expected from an even distribution of PEE rate according to T_{200} , individuals were performing more PEEs per unit of time for the following T_{200} water classes: -1°C , 0°C , 2°C , 3°C , 4°C , 5°C , 6°C and 7°C (Fig. 7). However, no relationship was found between either the absolute or the daily mass gain and the mean T_{200} of the water masses visited by the seal ($r^2 < 0.01$, $p > 0.9$ in both cases).

DISCUSSION

The results of the present study clearly demonstrate that during late spring and early summer the diving behaviour and foraging success of SES females were profoundly affected by the physical properties of the water masses visited. Indeed, these results show that at depth, water temperature, surface light level and the percentage of surface light reaching 150 m (used as an indicator of phytoplankton concentration) influence the SES foraging during day, night and twilight periods. Correspondingly, these data reveal the vertical distribution of the SES prey guild in relation to environmental variables.

Firstly, SES females dive deeper as T_{200} increases, indicating that the vertical accessibility of the prey decreases from the AD to subtropical waters. This finding is consistent with previous studies indicating that SES are diving deeper in warmer waters (Biuw et al. 2007, McIntyre et al. 2011b). Interestingly, when controlling for T_{200} , the deeper the SES dive,

Table 5. Variation in diving depth of female southern elephant seals according to surface light (L_0), the percentage of surface light reaching 150 m (L_{150}/L_0) and 200 m temperature (T_{200}) south and north of the Subantarctic Front for day and night. * $p < 0.05$; ** $p < 0.01$; *** $p < 0.001$; ns: not significant

Model	Parameter	$T_{200} < 7^{\circ}\text{C}$				$T_{200} > 7^{\circ}\text{C}$			
		Day		Night		Day		Night	
		Estimate	<i>t</i>	Estimate	<i>t</i>	Estimate	<i>t</i>	Estimate	<i>t</i>
Diving depth	L_0	38.69	25.51***	24.97	8.67**	24.41	6.09**	23.07	8.27**
	L_{150}/L_0	33.58	20.73***	29.05	14.65***	–	ns	29.45	7.33**
	T_{200}	18.64	10.46***	–	ns	233.5	3.59*	21.02	4.54**

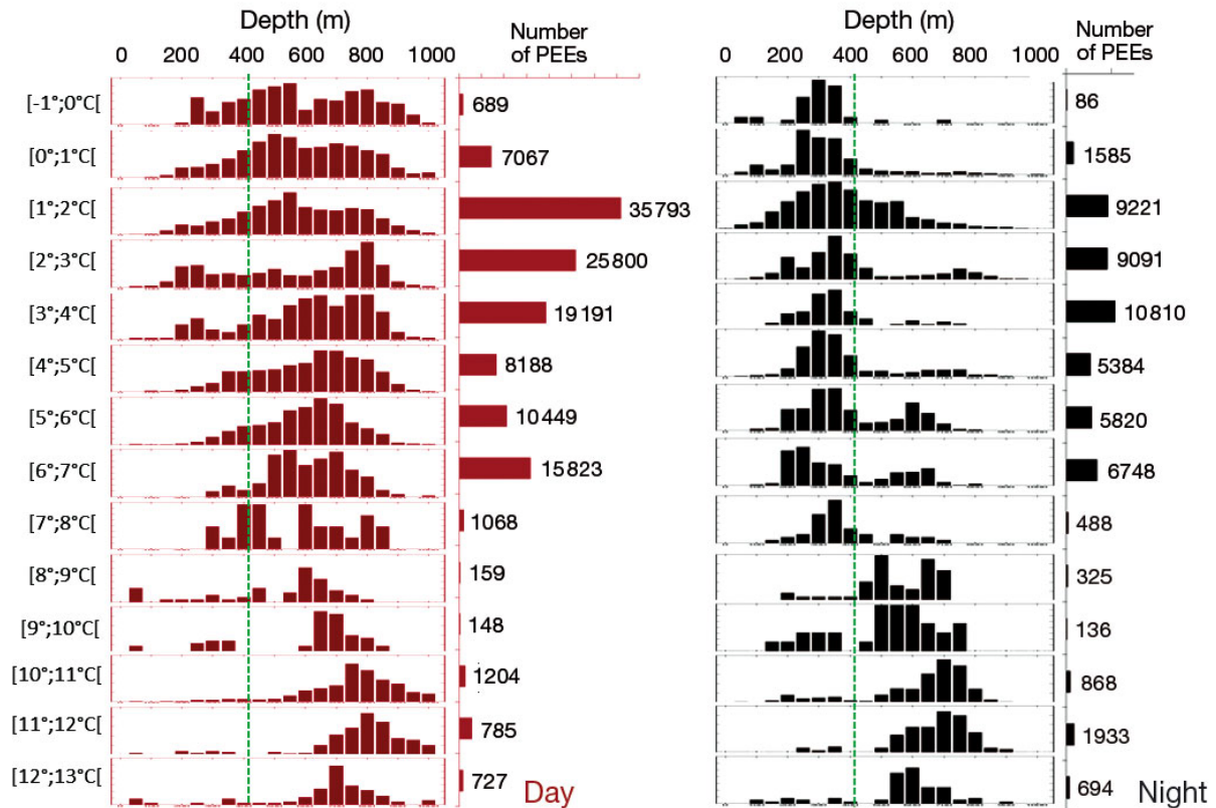


Fig. 6. Depth distribution of prey encounter events (PEEs) according to 50 m depth class during day (red) and night (black) according to each 1°C temperature at 200 m (T_{200}) class. The number of PEEs during day and night for each T_{200} class is indicated

the fewer prey they capture for a given bottom duration, suggesting a general decrease in prey density with increasing depth (Table 6). For waters with a T_{200} warmer than 7°C, a significant decrease in PEEs per unit of time spent diving or at the bottom of the dive was found, suggesting that lower prey densities and at greater depth were found north of the SAF.

Furthermore, as SES dive deeper, swimming costs associated with the bottom-surface transit increase and, as a consequence, the amount of time the seal is able to allocate to foraging at the bottom of the dive (i.e. the efficient foraging part of the dive) decreases with increasing diving depth. Furthermore, when considering a complete dive, and contrary to our

Table 6. Variation of the number of prey encounter events (PEE) by female southern elephant seals during the dive and at the bottom of the dive scale according to dive metrics and environmental conditions (surface light [L_0], the percentage of surface light reaching 150 m [L_{150}/L_0] and 200 m temperature [T_{200}]). * $p < 0.05$; ** $p < 0.01$; *** $p < 0.001$; ns: not significant

Model	Parameter	Day		Night		Twilight	
		Estimate	<i>t</i>	Estimate	<i>t</i>	Estimate	<i>t</i>
PEE dive	T_{200}	–	ns	–	ns	–	ns
	L_0	–	ns	0.220	11.75***	0.090	6.05**
	L_{150}/L_0	–0.110	–10.96***	0.210	10.42***	0.052	2.65*
	Dive duration	–0.070	–7.79**	–0.171	–12.21***	–0.600	–3.31**
	Diving depth	–0.150	–20.33***	–0.029	–2.67*	–0.210	–11.55***
PEE bottom	T_{200}	–	ns	–	ns	–	ns
	L_0	–	ns	0.289	13.69***	0.114	6.82**
	L_{150}/L_0	–0.080	–6.56**	0.280	12.87***	0.115	5.47**
	Bottom duration	0.079	8.06**	–	ns	0.060	2.66*
	Diving depth	–0.38	–38.60***	–0.360	–25.97***	–0.51	–19.61***

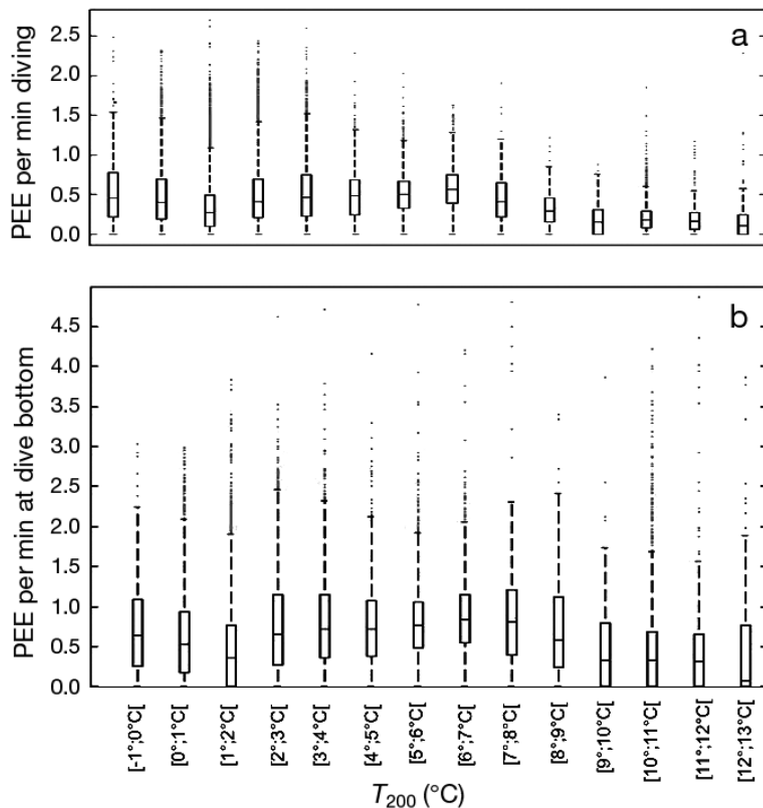


Fig. 7. (a) Box plot of prey encounter events (PEEs) per minute spent diving according to each 1°C temperature at 200 m (T_{200}) class. (b) Box plot of PEEs per minute spent at the bottom of the dive according to each 1°C T_{200} class. See Fig. 4 for further details

expectation, the number of PEEs decreases with increasing dive duration, suggesting that successful foraging dives were energetically more costly (i.e. higher swimming effort) than unsuccessful ones, forcing the seals to come back earlier to the surface. An alternative and non-exclusive hypothesis is that elephant seals modify their dive duration by changing the ascent or descent behaviour (i.e. by descending and ascending more rapidly to the surface) when they encounter high-quality prey patches, as reported in king penguins (Hanuise et al. 2013). Furthermore, when controlling for diving depth, the number of PEEs at the bottom of the dive increases with bottom duration during the day and twilight periods, but not at night, indicating that seals adjust the amount of time allocated to the efficient phase of their dive (bottom) in relation to prey density when generally diving deeper during the day and twilight compared with shallower night dives. Future studies should investigate how these observed variations match with the predictions of the optimal diving theory (Carbone & Houston 1994).

PEEs both during the whole dive and at the bottom of the dive were not related to T_{200} , suggesting that T_{200} has a general effect on prey vertical accessibility but not on foraging success when controlling for dive duration and diving depth. Correspondingly, in the present study, very few seals foraged in water with T_{200} exceeding 7°C (i.e. north of the SAF). Such a behavioural change associated with the SAF is consistent with previous studies indicating that the SAF represents an important biogeographic boundary regarding the latitudinal distribution of warm-water zooplankton (Ansorge et al. 1999) and myctophid species (Koubbi 1993, Koubbi et al. 2011), as well as top marine predators such as seabirds (Bost et al. 2009).

Individuals were found to exhibit lower PEE rates in water masses with T_{200} ranging between 1 and 2°C, corresponding to the zone encompassed between the PF and the SACCF. This is consistent with the finding of Biuw et al. (2007) that SES exhibited negative changes in body condition while migrating between the PF and the SACCF. In the Indian Ocean sector, this zone broadens latitudinally in the Kerguelen

region (Orsi et al. 1995) and this sector seems to represent a large region of unfavourable foraging conditions for SES. Despite our results supporting previous findings that dive capacity increases throughout the duration of the foraging trips (Bennett et al. 2001, Hassrick et al. 2010, McIntyre et al. 2011b), both T_{200} and diving depth were also found to influence dive duration in the present study.

For T_{200} lower than 7°C, T_{200} , L_0 and L_{150}/L_0 had a positive effect on the diving depth of SES during the day. No effect of T_{200} was found at night (Table 5). As L_0 is mostly related to both cloud cover and sun angle during the day and cloud cover and moon light at night, the present study emphasises the critical effect of light, mediated through its effect on the distribution of SES prey, on the diving behaviour of SES both during the day, as previously found by Jaud et al. (2012), and at night.

Light level at depth depends on the amount of light reaching the surface of the ocean and the intensity of light attenuation within the water column. In oceanic waters, phytoplankton concentra-

tion is a critical factor controlling most of the surface light attenuation within the water column and, therefore, light levels below the euphotic layer (Morel 1988). Light level at 150 m, combining the effect of surface light and light attenuation within the first 150 m of the water column, has been found to explain nearly 70% of the variation in the daylight diving depth of seals foraging between the PF and the SAF (Jaud et al. 2012). In the present study, with a larger data set we were able to show that SES also modified their diving behaviour at night according to both L_0 and L_{150}/L_0 , diving deeper when more light was available at the surface and when that light was less attenuated within the water column, suggesting that in clearer waters with lower phytoplankton concentrations seals had to dive deeper to reach their prey. This finding was consistent with the results obtained by Dragon et al. (2010) showing that the diving depth of SES was negatively related to remotely sensed surface chlorophyll *a* concentration.

The fact that light levels at the surface had an effect on the diving depth of SES at night suggests that variation of moonlight intensity in relation to moon cycle controls the vertical extent of the nycthemeral migration of SES prey and, therefore, their vertical accessibility, with prey items migrating closer to the ocean surface during the darkest nights. A similar effect of moonlight on diving behaviour has previously been observed in Galapagos fur seals *Arctocephalus galapagoensis* (Horning & Trillmich 1999) and northern fur seals *Callorhinus ursinus* (Lea et al. 2010). Several studies have shown that the extent of the vertical, nocturnal migration of zooplankton is related to changes in L_0 and water transparency, and that L_0 is dependent on the moon phase and/or cloud cover (Rudstam et al. 1989, Dodson 1990). Indeed, light is an essential factor for the vertical distribution of a broad range of mesopelagic marine animals, ranging from zooplankton (Liu et al. 2003) to fish (Batty et al. 1990), which distribute themselves according to precise light isolines, with threshold light intensity ('barrier isolume') found to initiate an avoidance reaction by these organisms (Boden & Kampa 1967, Kampa 1976, Gal et al. 1999, Cotté & Simard 2005).

The strong influence of light on the diving behaviour of SES leads us to hypothesise that the large variability in diving depth observed in water for a given T_{200} south of the SAF is likely to be explained by high variability in the amount of light available at depth in response to variations in cloud cover, phytoplankton concentration and sunlight or moonlight.

Therefore, while the change in phytoplankton concentration in summer is likely to have an effect on the overall SO productivity and the amount of prey available to SES several months later, it also exerts substantial influence on the vertical accessibility of SES prey. This has direct consequences on the foraging efficiency of these predators, as the deeper they dive, the less efficient (in terms of PEE events per unit of time) they are.

Interestingly, the influence of light on PEEs differed temporally. Surface light (L_0) had no effect on PEEs during the day, but had a positive effect during the night and twilight periods (see Table 6). Similarly, L_{150}/L_0 had a negative effect on PEEs during the day but a positive effect at night and twilight. L_{150}/L_0 is related to the concentration of particles (including phytoplankton) present in the water (Jaud et al. 2012). Therefore, the negative relationship found between L_{150}/L_0 and PEEs would suggest that SES are more successful during the day in areas where higher concentration of particles such as phytoplankton but also possibly zooplankton are encountered. In contrast, both L_0 and L_{150}/L_0 were found to have a positive effect on PEEs during twilight and at night. This suggests that light level within the water column could indirectly impact the foraging success of SES, potentially by influencing the distribution, availability and/or detectability of prey on clear nights and in clear waters.

SES dive generally to shallower depths at night than during daylight hours (Hindell et al. 1991, Biuw et al. 2007); in some instances they have been observed to dive to greater depths at night compared with the day, suggesting that they might be foraging on different prey resources on those occasions (McIntyre et al. 2011a). Consistent with Biuw et al. (2007), the greatest differences between day and night diving depths observed in the present study were between the SAF and the SACCF (i.e. the ACC zone). A noticeable exception was in both Antarctic waters and waters north of the SAF, where the diurnal variation diminished. Reduced differences in diving depth between day and night were observed for high and low T_{200} . However, the factors responsible for these differences between Antarctic and subtropical waters are likely to be different. For the subantarctic–subtropical zone, the contribution of L_0 and L_{150}/L_0 to the variation in diving depth was found to be lower to the north than to the south of the SAF. We interpret this result as the consequence of the deepening of SES females dives in search of prey found in deeper, colder waters. The 8°C isotherm appears to represent the upper limit of the vertical distribution

of most of the SES's prey (see Figs. 2, 3) and the diving depth of SES deepens with this isotherm from the SAF to the STF (Fig. 3). Indeed, as light levels decrease exponentially with depth, the influence of light levels should decrease north of the SAF at all times due to the increased diving depth required to reach the SES's prey; as a consequence, the diurnal differences in dive depth should decrease, as observed. A reduced diurnal difference in the diving depths of SES was also observed in Antarctic waters. This is consistent with a previous report (Biuw et al. 2007) that interpreted this as indicating animals foraging benthically on the Antarctic Shelf. However, all SES females in our study foraged in oceanic waters associated with the AD, well off the Antarctic Shelf. Therefore, the reduced diurnal difference in diving depth (Fig. 4) suggests that SES females were feeding on prey remaining close to the surface during daylight hours.

The SO is characterised by broad concentric bands of water masses around the Antarctic continent, with each zone maintaining its unique physical properties (e.g. Orsi et al. 1995, Belkin & Gordon 1996). This has 2 implications. Firstly, the concentric nature of the current and water masses of the SO ensures circum-polar continuity of its ecosystems and zooplankton and fish species (Baker 1954). Therefore, our findings regarding the vertical distribution of SES prey could be extrapolated to other parts of the SO. Secondly, the T_{200} gradient used in our study largely reflects the latitudinal structuring of the SO from subtropical to Antarctic waters.

The first assumption of our study is that female SES diet is dominated by myctophid fish. Recent studies suggest that myctophids are likely to represent a major proportion of the SES female diet (Cherel et al. 2008, Bailleul et al. 2010) and 3 of the dominant myctophid species in the SO (*Electrona antarctica*, *E. calbergi* and *Gymnoscopelus nicholsi*) have previously been found in stomach contents of SES (Slip 1995, Daneri & Carlini 2002). Furthermore, a recent study using a head-mounted camera on northern elephant seals (Naito et al. 2013) identified 2 small mesopelagic fishes as prey items, a myctophid and a bathypelagic, suggesting that elephant seals, despite their large size, are feeding on abundant but small prey items. Indeed, myctophid fish are a high-calorific prey (9.3 kJ g^{-1} wet mass) when compared with icefish (5.4 kJ g^{-1} wet mass) and prey items such as squid ($1.7\text{--}4.5 \text{ kJ g}^{-1}$ wet mass; Lea et al. 2002a,b). However, we acknowledge that other prey items such as squids are also likely consumed by SES.

The second assumption of our study is that a PEE provides a robust proxy for actual prey encounters. According to the calculated mean catch rate of 8.2 prey items per dive, we estimate that on average a SES female performing approximately 66 dives per day will catch approximately 540 prey items daily. Assuming that *Electrona calbergi*, *E. antarctica*, *Gymnoscopelus nicholsi* and *G. piabilis* represent the major prey items within the ACC zone and that their mass generally ranges between 9 g (*Electrona calbergi*, *E. antarctica*) and 30 g (*G. nicholsi*, *G. piabilis*; C. A. Bost & Y. Cherel pers. comm.), we estimate that a range of 4.8–16.1 kg of fish are ingested daily per seal. These values are not inconsistent with the daily mass gains (mean $0.9 \pm 0.4 \text{ kg}$, range: 0.2–2 kg) observed in the post-breeding females in our study using a mean 10% energy transfer efficiency to body mass growth (see Ashwell Erickson & Elsner 1981, Rosen & Trites 1999, Jeanniard du Dot et al. 2008). These estimates are consistent with those based on calculations using at-sea field metabolic rates and photographs obtained by head-mounted cameras (Naito et al. 2013), revealing that northern elephant seal females feed on small mesopelagic prey (10–20 g), catching 430 to 860 prey items and an estimated ingested mass ranging between 4.3 and 17.2 kg of prey daily.

The SO mesopelagic ichthyofauna is dominated by myctophids (Kock 1992), with approximately 35 species (Hulley 1990) and a crude estimated biomass ranging between 70 and 340 million tonnes (Lubimova et al. 1987, Sabourenkov 1991). Previous studies suggest that temperature, light and oxygen content of the water are key environmental factors controlling their distribution (Hulley & Lutjeharms 1995).

Four species of myctophids contribute the bulk of the SO fish biomass: *Krefflichthys anderssoni*, *Electrona antarctica*, *E. calbergi* and *Gymnoscopelus nicholsi* (Sabourenkov 1991). Despite their ecological significance and considerable biomass, remarkably little is known about the basic biology of myctophids (Collins et al. 2008). While they are known to perform daily vertical migrations in the water column, remaining at deep depths during the day and feeding in shallower depths at night (Gjosæter & Kawaguchi 1980), there is a major lack of knowledge regarding myctophid habitats and environmental factors controlling their distribution (Gjosæter & Kawaguchi 1980, Catul et al. 2011). Assuming that myctophids represent the main component of SES female diet, the fine-scale monitoring of SES foraging behaviour according to environmental factors provides indirect

information on the myctophid vertical distribution in relation to temperature and light conditions along a north–south latitudinal gradient.

Our results suggest that SES may target different communities of myctophids at different depths north of the SAF. Indeed, as part of the ICHTYOKER transect run between Kerguelen and St. Paul Island, data from trawls at night to depths ranging between 0 and 330 m in waters with a temperature range of 11–17°C revealed a myctophid community dominated by *Symbolophorus barnardi*, *Lampanyctus australis*, *Electrona paucirastra* and *Lampichthys procerus* (G. Duhamel, P. Prouvost, J. Boutain unpubl. data). These fish species were more abundant in the northern vicinity of the STF. Therefore, the fact that SES dive to deeper depths to reach water temperatures lower than 8°C suggests that they target a different assemblage of myctophids. This community is likely to be dominated by *Electrona calisbergi*, *E. subaspera*, *Gymnoscopelus piabilis*, *G. fraseri* and *G. bolini*, and this myctophid community distribution deepens with increasing temperature north of the SAF (Hulley 1981).

As mentioned earlier, a reduced diurnal difference in the diving depths of SES was also detected in Antarctic waters. This behaviour was also detected on very few occasions between the SACCF and the PF. In Antarctic waters, *Electrona carlsbergi*, a schooling species (Zasel'sliy et al. 1985), is commonly encountered close to the surface during the day (Hulley 1986, Fielding et al. 2012). *Electrona antarctica*, a non-schooling species, which is slightly larger than *E. carlsbergi*, is also found in SES stomach contents and may also represent an important prey item, along with *E. carlsbergi*.

The difference in schooling behaviour of *Electrona* species might influence the foraging strategy of SES. Indeed, 3-dimensional reconstruction of dive tracks reveals that the sinuosity of the path at the bottom of the dive increases with the number of PEEs. Individuals exhibit an area-restricted search behaviour at the bottom of their dive and for several consecutive dives when large numbers of PEEs occur (C. G., J. V.-G., Y. L., J. Jouma'a unpubl. data), suggesting that SES are targeting schooling resources in these dives. However, on a few dives, large numbers of PEEs were observed despite the fact that the seals were travelling in a nearly constant direction, suggesting a more dispersed prey resource, possibly *E. antarctica*. In Antarctic waters, krill is also observed within the first 100 m of the water column, and could be targeted by some seals feeding close to the surface during the day, as krill is suspected to be preyed upon by some seals (Y. Chérel pers. comm.).

The lack of relationship between the mean T_{200} encountered by the SES during their foraging trip and either the absolute mass gain or the daily mass gain is supportive of a change in prey quality and/or size according to the water masses visited. This suggests that the lower catch rate observed in warmer waters, as well as the greater proportion of the swimming effort allocated to a greater diving depth to access those prey, might be compensated by the consumption of larger and/or higher quality prey in those water masses. For example, larger myctophids such as *Gymnoscopelus piabilis*, *G. nicholsi* and possibly *G. bolini* may represent a major component of SES female diet in these warmer northern waters. Indeed, in a recent study, G. Richard, J. V.-G. J. Jouma'a, C. G. (unpubl. data) found that the decrease in SES density (i.e. increase of lipid content) was positively related to the number of PEEs and was found to vary according to the foraging habitat. SES foraging north of the SAF improved their body condition at a similar rate compared with those foraging further south, despite a lower PEE rate and a greater swimming effort due to greater diving depth. This result is supportive of the hypothesis that SES females foraging north of the SAF were likely to feed on larger prey items, such as *Gymnoscopelus* spp., compared with those foraging south of the SAF.

Another interesting finding of this study regarding foraging success was that PEE rate both during the dive and at the bottom of the dive varied according to diurnal rhythm, with SES being more efficient during twilight compared with at night and during the day. Individuals were also found to have a significantly higher PEE rate at night compared with during the day. While we were expecting a lower foraging efficiency during daytime dives (complete dive) due to the greater diving depths recorded, we were not expecting such differences when only taking into account the PEE rate calculated at the bottom of the dive. In fact, we were expecting the opposite relationship, with a greater success rate during the bottom phases of daytime dives. Indeed, while some myctophids species are known to be actively feeding during both day and night, several studies have shown that during the day myctophids tend to aggregate in denser schools, behaving more lethargically, possibly making them more vulnerable to predation (Barham 1966, Kinzer et al. 1993, Kaartvedt et al. 2009). The present study suggests the opposite, indicating that elephant seals might be more efficient when their prey is supposedly actively foraging at night closer to the surface or performing nycthemeral migrations. Therefore, we hypothesise that actively,

vertically migrating (at dusk and dawn) and/or foraging myctophids (at night) are easier to detect and/or locate, either because they reach aggregation levels allowing the maximization of encounter rates by predators or because their active behaviour makes them more vulnerable to predation. Actively moving prey might be easier to locate by the foraging seals. Indeed, as prey move they create hydrodynamic trails likely to be detected by SES through their vibrissae (Dehnhardt et al. 2001).

Alternatively, a recent modelling study has shown that predators reach their maximum PEE efficiency at intermediate aggregation levels of prey, with highly aggregated patches being too difficult to locate, while dispersed prey are diluted in the environment and do not allow the predator to be efficient (Massardier 2013).

Prey may also produce sounds detected by elephant seals, which are known to have excellent underwater hearing ability (Kastak & Schusterman 1998). Indeed, over a 12-yr study, McCauley & Cato (pers. comm.) found recently that fish choruses, most likely to be myctophids, always occurred shortly after dusk and pre-dawn, with most energy in the 1–3 kHz range. Furthermore, while active, myctophids might be more likely to signal their presence by producing bioluminescence that seals could detect visually (Vacquié-Garcia et al. 2012). Such hypotheses should be further investigated in future studies and should provide new and valuable insights into the ecology and distribution of SES prey.

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