Abstract: Adult male southern elephant seals instrumented in 2000 on King George Island \((n = 13)\), travelled both to the north \((n = 2)\) and to the east \((n = 6)\) of the Antarctic Peninsula. Five males remained within 500 km of the island focusing movements in the Bransfield Strait and around the Antarctic Peninsula. Sea surface temperatures encountered by these animals showed little variation. While animal trajectories appeared unaffected by sea ice cover, areas of shallow depths were frequented. Three males moved as far as 75°S to the east of the Peninsula with maximum distances of more than 1500 km from King George Island. They travelled into the Weddell Sea along the western continental shelf break until they reached the region of the Filchner Trough outflow. Here the sea floor consists of canyons and ridges that support intensive mixing between the warm saline waters of the Weddell Gyre, the very cold outflow waters and ice shelf water at the Antarctic Slope Front. The need for re-instrumentation of adult males from King George Island is highlighted to investigate whether males continue to travel to similar areas and to obtain higher resolution data.

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Key words: Antarctic Peninsula, area restricted movements, post-moult movements, satellite telemetry, sea ice

Introduction

In the last decade, studies of animal movements have provided insights into previously unexplored aspects of behaviour, ecology and biology (Schaffer et al. 2006). New technologies have provided data that facilitate the development of management actions and conservation of endangered species (Lombard et al. 2007). The use of animal movement data remains tentative for conservation applications in marine habitats, possibly due to the difficulty in establishing effective marine conservation areas (Sale et al. 2005), the quantification of marine habitat types (Gregr & Bodtker 2007), and the lack of multidisciplinary integration necessary for making wider management decisions. The study of marine top predators in relation to their physical environment may aid in marine conservation and provide a basis for further ecological studies. Southern elephant seals are top predators in Antarctic ecosystems and knowledge of their foraging ranges will aid in understanding their impacts within the Southern Ocean, as well as shedding light on the influences of oceanographic factors on higher order organisms.

Southern elephant seals have been tracked from South Georgia (McConnell et al. 1992, McConnell & Fedak 1996), Iles Kerguelen (Bailleul et al. 2007), Macquarie Island (Hindell et al. 1991, van den Hoff et al. 2002, Hindell et al. 2003) and Peninsula Valdés (Campagna et al. 2000, 2006, 2007). These large populations make up the major stocks of southern elephant seals worldwide and are augmented by populations from smaller surrounding islands. Southern elephant seals from King George Island form part of the South Georgia stock of elephant seals. Southern elephant seals from the South Georgia and Peninsula Valdés stocks have not declined in recent years (McMahon et al. 2005). The decline in stocks from other populations has been attributed to declines in juvenile and/or adult female survival, possibly due to limited food availability (Pistorius et al. 2004, McMahon et al. 2005). Southern elephant seal numbers on King George Island have remained stable for some time (SCAR 2000), although a decline has been noted in recent years (Carlini personal communication 2007). This population is characterized by heavier pups at weaning (Carlini et al. 1997) and larger females that show higher rates of mass gain whilst spending shorter periods at sea (Carlini et al. 2005) when compared to other populations. Ocean habitat usage of female and under-yearling elephant seals from South Georgia and King George islands are generally in the area west of the Antarctic Peninsula (McConnell et al. 2006, 2007).
Laws 1994). Future survival and reproductive success is
dependent upon blubber stores accumulated during
successful foraging. The large body size of adult male
southern elephant seals could possibly have two-fold
advantages: in addition to the ability of larger animals to
fast for longer durations, larger animals may also have the
ability to utilize a variety of oceanic habitats that are
inaccessible to smaller bodied female and juvenile animals.
The ability to adopt varied diving strategies and the need
to maximize foraging success may drive observed
variations in foraging strategies (Le Boeuf et al. 2000).

This study presents data showing the movements of adult
males from the King George Island population of southern
elephant seals. A preliminary method for quantifying
movement variation is also presented. Individual variation
in movements is shown. Evidence of the utilization of the
area to the south-east of the Peninsula, and particularly to
a highly specific area in the southern Weddell Sea, a
previously unobserved pattern (Biuw et al. 2007), is
presented.

Methods

Fifteen adult male (≥ 6 years old) southern elephant seals
were instrumented between March and April 2000 at
Stranger Point, King George Island (62°14’S, 58°40’W)
(Fig. 1a & b). This work follows on an earlier project that
entailed the instrumentation of 13 adult females and seven
under-yearling seals (Bornemann et al. 2000).

All the animals were instrumented with satellite-linked
depth recorders (SDR T-6 or SDR T-10 Wildlife
Computers, Redmond, WA, USA). These two transmitter
types were designed to provide processed data on the
locations and diving behaviour (maximum depth, dive
duration, dive depth frequency and time-at-depth) in the
form of 6-hourly histograms. The transmitters varied in
the resolution of dive measurements that were taken. The
present paper focuses specifically on location data, and
the dive data is not discussed. The transmission protocol of
location data was exactly the same for both transmitter
types, transmitting data at 90 sec intervals between 00h00
(GMT) and 23h00 (GMT) when animals were at the surface.

The immobilization procedure used in this study has been
described elsewhere (Ramdohr et al. 2001). Animals were
first remotely darted using automatically evacuating
syringes (Telinject, Römerberg, Germany) filled with Large
Animal (LA) Immobilon to achieve initial immobilization.
Ketamine hydrochloride was subsequently administered
intramuscularly by hand to maintain immobilization. Both
the dosage, and the number of injections of ketamine was
determined by the stage of narcosis (Ramdohr et al. 2001).
The transmitters were attached using a star-shaped gauze
mat that was glued onto the head of the animal using quick-
setting epoxy resin (Bornemann et al. 2000).

Incoming ARGOS data was decoded using SatPack
software (Wildlife Computers). Location data of all quality
classes were filtered according to a 10 km.h⁻¹ velocity
algorithm (McConnell et al. 1992) which removes locations requiring unrealistic swimming speeds of greater than 10 km.h\(^{-1}\). All the locations were averaged to a daily location to provide regular trajectories over time. All locations, regardless of location quality, were used because long range movements (>10 km) were analysed (White & Sjöberg 2002). All primary data derived from ARGOS locations are archived and available in open access through the data library PANGAEA—Publishing Network for Geoscientific and Environmental data (http://doi:pangaea.de/10.1594/PANGAEA.692856). Of the 14 instrumented animals, only eight animals were tracked for more than 40 days (mean transmission time = 112 days; range = 8–338 days). These tracks were retained in the analysis to give an overall impression of habitat usage by these animals due to the fact that they included at least one transit phase and one area of restricted movement.

Individual tracks were analysed separately to identify area restricted movement (ARM) as opposed to transit movements. By choosing the animal as a sampling unit, issues of pseudo-replication that arise from the use of locations as sampling units were avoided (Otis & White 1999). A simple analysis of the cumulative distance from the origin of the track was performed to identify areas of restricted movement. As time-constrained central place foragers, southern elephant seal tracks are characterized by a spatially discrete area of altered activity flanked by direct transit phases (e.g. McConnell et al. 1992, McConnell & Fedak 1996, Jonker & Bester 1998, Campagna et al. 2007). Area restricted movement was defined as all the consecutive locations that displayed a minimum variability in terms of distance from the origin of the track that were flanked by transit movements, which had high variability in terms of distance from the origin of the track (Fig. 2). Although relatively crude and subjective to directional bias, this method is adequate for this set of location data which present very clear distinctions between periods of travel to and from such areas. In order to rank the degree of restriction displayed in movements, an index of movement variability (MV) was created for each segment of each track:

\[
MV\{n:1\} = SD\{n:1\}/\text{no. days}\{n:1\}
\]

where \(SD\) is the standard deviation in distance from the point of origin, in this case King George Island and \(\text{no. days}\) is the number of days spent in a specific sector of movement \((n:1)\). Quantifying movement variation in this way was deemed suitable for tracks where segments of area restricted movement were clearly delineated.

Elephant seal locations were overlaid onto meso-scale environmental data. Bathymetric data, including sea floor depth and bathymetric feature names, were obtained from the GEBCO Digital Atlas (IOC 2003). The data were interpolated to a 25 km \(\times\) 25 km grid, the same as the sea ice data obtained by the Special Sensor Microwave Imager (SSM/I) satellite, as retrieved by the NASA-Team-Algorithm (Cavaliere et al. 1996). The average rate of travel for all the animals in this study was 20 km day\(^{-1}\), making comparisons at this resolution reasonable because the resolution of the auxiliary data is the same as the resolution of the track data. All elephant seal locations were interpolated to monthly sea ice values in order to highlight seasonal variability in sea ice concentrations for the locations considered in this study. Weekly optimally

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 Ala Fig. 1. The tracks of adult male southern elephant seals (Mirounga leonina) from King George Island (1) instrumented in 2000. **a.** The tracks of AM3 (–\(\circ\)–), AM6 (–\(\Delta\)–), AM8 (–\(\bullet\)–), AM11 (–\(\blacksquare\)–) and AM13 (–\(\square\)–). These animals spent extended periods of time in close proximity to King George Island. **b.** Tracks of AM9 (–\(\bullet\)–), AM10 (–\(\blacktriangle\)–), and AM14 (–\(\bigodot\)–) locations were recorded from April 2000 to February 2001. King George Island (1), the Bransfield Strait (2) and Elephant Island (3) as well as major bathymetric features in the Weddell Sea are indicated on the map.
integrated sea surface temperatures (OISST) were also interpolated to the elephant seal locations. These data are supplied in the corrected format (ftp://oceans.gsfc.nasa.gov/OISST) and are derived from \textit{in situ}, satellite and sea ice simulated sea surface temperatures (Reynolds et al. 2002).

Results

Eight animals provided tracks of more than 40 days; these tracks were retained in the analysis in order to represent at least one transit phase and one area of restricted movement for each animal. The duration of the selected tracks varied between 48 days and 338 days, with only two animals providing tracks that were shorter than 100 days. All of these animals displayed area restricted movements (ARM) at some point during their tracks. Basic movement parameters, environmental values encountered during tracking, and average movement variations for areas of restricted movements and transit locations are presented in Table I.

Area restricted movement indices (MV) varied substantially between animals, with AM10 displaying the largest $\text{MV} = 1.48 \text{ km day}^{-1}$ and AM3 displaying the smallest $\text{MV} = 0.05 \text{ km day}^{-1}$ during its only period of ARM. MV values during transit varied between 25.93 km day$^{-1}$ (AM8) and 7.14 km day$^{-1}$ (AM3) (Table I). Two animals had more than one area of restricted movement. AM11 and AM14 both returned to King George Island for the breeding season and retained their transmitters during the post-breeding trip. MV values, for both animals, were greater for the areas of restricted movement after the breeding period than after the moult (Fig. 2).

Individual movements varied considerably, with maximum distances from King George Island varying between 156 km (AM3) and 1800 km (AM14) and total trip distances varying between 850 km (AM3) and 10782 km (AM14).

One animal (AM3) spent 4.8 months in the Bransfield Strait, no further than 156 km from King George Island.

![Distance (km) from King George Island over time for a. AM11, and b. AM14, indicating the movement variation (MV) during the post-moul t area of restricted movement (ARM1) and the post-breeding area of restricted movement (ARM2).](image)

Table I. Standard length, basic location statistics, average values and standard deviations (SD) of environmental variables encountered and movement variation indices (Areas of Restricted Movement = ARM) for the adult male southern elephant seals ($n = 8$) instrumented during March and April 2000 at King George Island.

<table>
<thead>
<tr>
<th>Seal</th>
<th>Standard length (m)</th>
<th>Date of deployment</th>
<th>Track duration (d)</th>
<th>No. locations (n)</th>
<th>Haulout</th>
<th>Maximum distance from King George Island (km)</th>
<th>Total distance travelled (km)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AM3</td>
<td>3.9</td>
<td>09/03/2000</td>
<td>146</td>
<td>131</td>
<td></td>
<td>155.80</td>
<td>849.27</td>
</tr>
<tr>
<td>AM6</td>
<td>3.9</td>
<td>18/03/2000</td>
<td>100</td>
<td>89</td>
<td></td>
<td>264.42</td>
<td>1797.27</td>
</tr>
<tr>
<td>AM8</td>
<td>4.32</td>
<td>31/03/2000</td>
<td>136</td>
<td>126</td>
<td></td>
<td>378.98</td>
<td>1279.45</td>
</tr>
<tr>
<td>AM9</td>
<td>4.6</td>
<td>07/04/2000</td>
<td>197</td>
<td>165</td>
<td>South Georgia$^B$</td>
<td>1582.27</td>
<td>5129.91</td>
</tr>
<tr>
<td>AM10</td>
<td>4.3</td>
<td>12/04/2000</td>
<td>48</td>
<td>30</td>
<td></td>
<td>1715.49</td>
<td>2000.25</td>
</tr>
<tr>
<td>AM11</td>
<td>4.2</td>
<td>12/04/2000</td>
<td>294</td>
<td>225</td>
<td>King George Island$^B$</td>
<td>505.98</td>
<td>3420.63</td>
</tr>
<tr>
<td>AM13</td>
<td>4.05</td>
<td>21/04/2000</td>
<td>92</td>
<td>89</td>
<td></td>
<td>332.48</td>
<td>1463.96</td>
</tr>
<tr>
<td>AM14</td>
<td>4.38</td>
<td>21/04/2000</td>
<td>338</td>
<td>203</td>
<td>King George Island$^B&amp;M$</td>
<td>1799.59</td>
<td>10781.53</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Sea Ice [%]</th>
<th>Bathymetry (m)</th>
<th>Sea surface temperature ($^\circ$C)</th>
<th>Movement variation$\text{ARM}$</th>
<th>Movement variation$\text{Transit}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mean ± SD</td>
<td>Mean ± SD</td>
<td>Mean ± SD</td>
<td></td>
<td></td>
</tr>
<tr>
<td>AM3</td>
<td>0.47 ± 3.82</td>
<td>-315.15 ± 235.09</td>
<td>-1.66 ± 0.5</td>
<td>0.05</td>
</tr>
<tr>
<td>AM6</td>
<td>21.07 ± 21.27</td>
<td>-288.98 ± 275.93</td>
<td>-1.61 ± 0.36</td>
<td>0.23</td>
</tr>
<tr>
<td>AM8</td>
<td>19.86 ± 19.74</td>
<td>-500.06 ± 280.98</td>
<td>2.58 ± 1.87</td>
<td>0.07</td>
</tr>
<tr>
<td>AM9</td>
<td>82.13 ± 34.21</td>
<td>-1284.33 ± 1212.57</td>
<td>-1.47 ± 0.91</td>
<td>0.25</td>
</tr>
<tr>
<td>AM10</td>
<td>70.67 ± 40.87</td>
<td>-1735.47 ± 1534.94</td>
<td>0.01 ± 1.63</td>
<td>1.48</td>
</tr>
<tr>
<td>AM11</td>
<td>60.61 ± 37.61</td>
<td>-385.60 ± 269.78</td>
<td>-1.39 ± 0.56</td>
<td>0.44</td>
</tr>
<tr>
<td>AM13</td>
<td>38.03 ± 20.21</td>
<td>-496.24 ± 325.15</td>
<td>0.36 ± 0.74</td>
<td>0.33</td>
</tr>
<tr>
<td>AM14</td>
<td>71.62 ± 35.87</td>
<td>-824.98 ± 712.95</td>
<td>-0.86 ± 1.46</td>
<td>0.43</td>
</tr>
</tbody>
</table>

Notes: $^B$Breeding, $^M$Moulting.
AM3 encountered shallow depths at all locations, sea ice concentrations were low and constant, and sea surface temperatures also showed little variability (Table I). Two animals (AM8, AM13) moved to the north of King George Island, spending considerable periods of time to the east of Elephant Island (Fig. 1a). AM8 travelled further north than AM13 and encountered warmer waters and less sea ice than AM13 (Table I). Both animals encountered shallow waters (Table I).

Two animals (AM6, AM11) displayed a tendency to forage within the topographically heterogeneous environment of the Antarctic Peninsula (Fig. 1a). AM8 travelled further north than AM13 and encountered warmer waters and less sea ice than AM13 (Table I). Both animals encountered shallow waters (Table I).

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Two animals (AM8, AM13) moved to the north of King George Island, spending considerable periods of time to the east of Elephant Island (Fig. 1a). AM8 travelled further north than AM13 and encountered warmer waters and less sea ice than AM13 (Table I). Both animals encountered shallow waters (Table I). The three remaining animals (AM9, AM10, AM14) travelled into the Weddell Sea moving along the shelf margin east of the Antarctic Peninsula, until they reached their southernmost positions at approximately 75°S, a bathymetrically heterogeneous region (Berkner Bank, Akademik Fedorov Canyon and the Filchner Trough) (Fig. 1b). Movements through this region were characterized by high sea ice concentrations, varied bathymetry, as well as cold sea-surface temperatures, resulting in high variability for sea surface temperatures encountered throughout the recorded tracks (Table I). The regions in which ARM was recorded were adjacent to the dense pack ice that exists in the middle of the Weddell Sea (Fig. 3). An animation of sea ice cover and seal locations visualizes seal movements relative to overall sea ice distribution in a daily time series from March 2000 to March 2001 (Martin et al. 2006). This indicates substantial differences between winter and summer ice conditions in the southern Weddell Sea resulting in very high and low to intermediate sea ice concentrations during the respective ARMs (Fig. 4). AM9 remained in a localized 100 km wide shelf-slope area between the Berkner Bank and the Akademik Fedorov Canyon for 102 days. On 25 August the seal headed north through heavy pack ice to South Georgia to breed; last transmissions were received from the southernmost tip of the island on 23 October 2000. AM14 travelled on a path remarkably similar to that of AM9,
arriving 22 days later (26 May) at exactly the same localized area between the Berkner Bank and the Akademik Fedorov Canyon and remained there for 100 days. On 3 September AM14 moved back to King George Island for the breeding season and embarked on a post-breeding journey to approximately the same area as his post-moult trip. It remained in this area for 63 days (24 December–25 February) in a large coastal polynya and then returned to King George Island to moult, before transmissions ended on 20 March 2001 (Fig. 1b).

Discussion

Adult male elephant seals instrumented on King George Island in 2000 displayed different movements compared to female southern elephant seals instrumented on the same island in 1997 (Bornemann et al. 2000). Female elephant seals travelled to the area west of the Antarctic Peninsula in 1997 (Bornemann et al. 2000). It is difficult to explain the possible reasons for their different travel patterns, especially since individuals of both sexes were not instrumented during the same time period. This makes it difficult to uncover whether observed behaviours are in response to environmental conditions or if large amounts of individual variation within a relatively small sample of animals are being observed. Continued instrumentation of animals is necessary to see if these differential migratory patterns are persistent over time and if sexual segregation of foraging areas actually does exist.

Quantifying movement variation allows for the comparison of movements between different areas of restricted movement catalogued during a track (Tremblay et al. 2007). For example, AMs 11 and 14 both showed higher movement variation indices during the post-breeding areas of restricted movement than post-moult areas of restricted movement (Fig. 2). Seasonal variation in foraging behaviour, based on diving characteristics, has been described for animals from South Georgia (Bennet et al. 2001). This variation could arise from: 1) seasonality in the distribution of resources, 2) metabolic constraints (Bennet et al. 2001), or in this case, 3) from restricted access to breathing holes/cracks due to higher sea ice concentrations during the post-moult trip, which generally occurred in the late autumn and early winter. The collection of good quality tracks over time and the quantification of mass gain for animals travelling to different areas would help to evaluate the potential benefits of utilizing different foraging areas. Quantifying movement variation allows for comparison between populations and would prove valuable in evaluating the effects of dynamic oceanographic variables on the movements of these animals.

The environmental variables that were investigated in this study fall into two categories, namely static variables, such as bathymetry, and dynamic variables that vary continuously over space and time. These dynamic variables appear to be the biggest constraint in modelling marine habitat suitability (Skov et al. 2008). It is impossible to generalize to the whole population from a few animals instrumented in a particular year that display a large amount of individual variation. A few animals avoided sea ice altogether and others seemed to exploit the increased primary productivity that is associated with fragmented sea ice and ice edges (Bargagli 2005). The tendency to feed along the continental shelf does not seem to be unique to elephant seals from King George Island. Animals from South Georgia (McConnell & Fedak 1996) and Patagonia (Campagna et al. 1999) also displayed the tendency to feed close to their respective haulout areas and made use of shallow, localized feeding areas along the respective shelf margins.

The movements of AM8 and AM13 may not be unique, as similar results have been found for animals instrumented at Elephant Island in 1999 (Muelbert et al. 2004). Both findings coincide with the area just to the north of the Antarctic Peninsula, where the Antarctic Circumpolar Current is directed through the Drake Passage into the Scotia Sea (Holm-Hansen et al. 2004). The results of the present study show that at least three of the eight animals preferred the steep continental slope areas far south in the Weddell basin during the study period. Here, the southern branch of the Weddell Gyre meets the Filchner Trough outflow, which together with the rough bathymetry causes intensive mixing of the contributing water masses.

The particular region utilized by AM9 and AM14 corresponds closely with the region identified by Foldvik et al. (2004) as being important for the production of Weddell Sea Bottom Water (WSBW). The production of WSBW is important for the export of Antarctic Bottom Water (AABW), the water body along which cold, dense and oxygenated water is exported from the Antarctic to the surrounding oceans. This area is also characterized by elevated trace element concentrations dissolved from the shelf ice into the Ice Shelf Water. This water mass is transported to the north through the Filchner Trough into the Weddell Gyre circulation system where it is heavily mixed within the Antarctic Slope Front. The Filchner outflow is one of, if not the only, steady annual source of shelf water that is responsible for the deep water and bottom water production in the Weddell Sea. The restricted area movements displayed in this region illustrate a connection between important physical oceanographic features and biological processes of a top-level predator.

The tendency of southern elephant seals to forage on the Antarctic Continental Shelf, within the pack ice, has been illustrated for seals from Iles Kerguen (Bailleul et al. 2007) and Macquarie Island (Bradshaw et al. 2003). Weddell seal foraging behaviour has been linked to the presence of the Antarctic silverfish (Pleuragramma antarcticum Boulenger) in the Weddell Sea (Plötz et al. 2001). The presence of this fish, a dominant pelagic
species (Hubold 1985), has also been suggested to influence the movement patterns of female elephant seals (Bornemann et al. 2000) - and forms an important part of the pelagic fish diet of southern elephant seals from King George Island (Daneri & Carlini 2002). The presence of the Antarctic silverfish in the ice-bound Weddell Sea has been linked to the high densities of krill, Euphausia superba Dana, that congregate under ice sheets and take advantage of melting ice releasing algae into underlying water bodies (Stretch et al. 1988). Given that male southern elephant seals perform both benthic and pelagic foraging dives (Hindell et al. 1991), a pattern also observed for juvenile and adult male northern elephant seals (Le Boeuf et al. 1996), the presence of *P. antarcticum* may explain the tendency of male southern elephant seals from King George Island to travel through dense pack ice into high Antarctic latitudes in order to find an abundant food source.

**Outlook**

The Intergovernmental Panel on Climate Change (IPCC) predicts that climate change will influence changes in temperature, which in turn will influence ocean circulation, ice coverage and sea levels (McCarthy et al. 2001). All these factors, often associated with broad-scale climate change, can have immediate and direct impacts on marine mammals by influencing food availability. At present, the link between seal population numbers and environmental conditions remains tentative (Sun et al. 2004), especially in terms of directionality. Climate cooling may cause sea ice cover to increase, protecting biotas that are able to survive under the ice from top-level predators such as whales and seals. Conversely, climate warming may cause a reduction in sea ice cover, increasing primary and secondary production and providing access for top-level predators through the creation of breathing holes. This research shows the dependence of male southern elephant seals from King George Island on ocean circulation systems, such as the Weddell Gyre, as well as an intimate association of these animals with sea ice. Populations of these animals can easily be monitored owing to their terrestrial breeding habits, fidelity to natal sites and large body sizes and are thus ideal climate change monitoring species (Simmonds & Isaac 2007).

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**References**


