Satellite telemetry of humpback whales off Madagascar reveals insights on breeding behavior and long-range movements within the southwest Indian Ocean

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ABSTRACT: Humpback whales breeding in the southwest Indian Ocean are thought to exhibit population substructure between Madagascar and east Africa. To investigate regional movements, breeding behavior and habitat utilization, 23 whales were satellite-tagged off Madagascar during peak breeding season off the northeast and southwest coasts. Mean tag duration was 24.2 d (3 to 58 d), during which time no individual remained near the immediate tagging sites and several displayed extensive long-range movements. We applied a switching state-space model to estimate behavioral modes of ‘transiting’ (b-mode approaching 1.0) vs. ‘localized’ (b-mode approaching 2.0) movement. A general linear mixed-effects model indicated females were more likely to display transiting behavior than males (mean b-mode females = 1.27, males = 1.65; p = 0.031). Whales tagged in the northeast displayed localized movements off the central east coast of Madagascar, whereas whales tagged in the southwest displayed localized movements on the southern coasts, with little overlap. Long-distance movements included north-westerly trajectories to eastern Africa and southerly transits to Walters Shoals and the Crozet Islands. Despite these long-range movements in short periods, no whale travelled to the northwest coast of Madagascar, nor to Mozambique or the Mascarene Islands. These results suggest there may be more interchange between Madagascar and central-east Africa than previously thought, and whales off east and west Madagascar may not use the same habitat within breeding seasons; important findings for defining sub-population structure and conservation management strategy. Furthermore, male mating strategy may involve more localized searching or displaying, whereas females travel more extensively during the breeding season, observations that are consistent with a large-scale lek mating system.

KEY WORDS: Humpback whales · Satellite telemetry · Madagascar · Breeding behavior · Movement patterns · Population structure

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INTRODUCTION

Humpback whales *Megaptera novaeangliae*, globally, are seasonal breeders, and with few exceptions, populations migrate yearly between high-latitude feeding regions in summer and low-latitude breeding regions in winter. Migrations are extensive, with latitudinal migratory movements of several thousand kilometers in different ocean basins (Mate et al. 1998, Zerbini et al. 2006, Rasmussen et al. 2007, Lagerquist et al. 2008, Kennedy et al. 2014, Rosenbaum et al. 2014). The mating system is generally agreed to be polygynous, with elements of lekking and male dominance polygyny (Herman & Tavolga 1980, Clapham 1996, 2000, Cerchio et al. 2005). Females reproduce on average once every 2 to 3 yr and likely have a short estrus period (Chittleborough 1958, 1965, Clapham & Mayo 1990), resulting in a severely skewed operational sex ratio (Clapham 2000). Gestation lasts 1 yr, with both conception and parturition generally occurring during a 5 mo period centered in winter. Thus, in a breeding region, females can be subdivided into 2 reproductive classes: adult females without calves and with relatively high fecundity (termed ‘non-parous females’ throughout) that have migrated solely to become fertilized; and post-partum females with a calf-of-the-year (termed ‘mothers’ throughout) that have relatively low fecundity since only a small fraction of females give birth in consecutive years (Clapham & Mayo 1990, Clapham 1996). Males exhibit alternative mating tactics while on the breeding grounds, including a male-limited acoustic display or song (Payne & McVay 1971), intense physical competition among groups of males for single estrous females in ‘competitive groups’ (Tyack & Whitehead 1983), and ‘escorting’ of mothers with a calf-of-the-year (despite their relatively low fecundity). Molecular paternity analysis has indicated polygyny with a slight but significant skew in reproductive success among males and that success is not strongly skewed towards a single tactic (Cerchio 2003, Cerchio et al. 2005).

The International Whaling Commission (IWC) designates 7 breeding stocks around the Southern Hemisphere, labeled A through G (IWC 1998, 2007). The breeding population wintering in the southwest Indian Ocean (SWIO) is labeled breeding stock C and is sub-divided into 4 designated sub-stocks (Fig. 1): in the eastern African coastal waters of South Africa to Kenya (C1), off the islands of the Mozambique Channel (C2), in the coastal waters of Madagascar (C3) and off the Mascarene Islands (C4) (IWC 2007). Modern whaling of humpback whales in the SWIO began in 1908 and expanded rapidly thereafter, depleting the population, with an estimated 19 000 whales taken up until 1963 (Findlay 2001). The first evidence of population structure in the region came from different whaling catch histories in C1 (South Africa/Mozambique, where there was early depletion prior to 1915) as compared to C3 (Madagascar, where catches remained relatively high after 1915), suggesting 2 distinct stocks that were differentially impacted by whaling pressure (Best 1994, Findlay 2001).

In the C1 sub-region, humpback whales migrate past eastern South Africa to a wintering ground off Mozambique, with apparent lower concentrations off Tanzania and Kenya (Findlay et al. 2011a,b, Amir et al. 2012). The relationship between whales in the southern range (Mozambique, C1S) and northern range (Tanzania, Kenya, C1N) is uncertain. In the C2 sub-region, humpback whales occur around the Comoros Islands and in lesser concentrations around the Seychelles Islands from Aldabra to the Mahe plateau (Kiszka et al. 2010, Ersts et al. 2011a). In the C3 sub-region, humpback whales are distributed around the entire island of Madagascar, with most information coming from the northeast (termed here C3E; Fig. 1) in Antongil Bay along with the nearby Ile Sainte (Ste.) Marie (Rosenbaum et al. 1997, Ersts & Rosenbaum 2003, Pomilla & Rosenbaum 2006, Cerchio et al. 2009, Vely et al. 2009). A low recapture rate between years
in Antongil Bay indicated a large population, but re-
captured individuals displayed highly regular migra-
tory timing sighted at the same time each year; in ad-
dition, recaptures within a season were few and of
short duration, suggesting short residency and con-
tinuous movement through the area (Cerchio et al.
2009). Concentrations of whales are also known in
the southeast and along the west coast, best docu-
mented in the southwest (termed here C3W; Fig. 1)
around Toliara, where encounter rates are similar to
Antongil Bay (S. Cerchio pers. obs.). The relationship
of animals utilizing the east coast (C3E) and west
coast (C3W) of Madagascar is unknown, so it is not
clear whether they represent the same or distinct sub-
stocks. In the C4 sub-region, humpback whales are
distributed around the islands of Réunion, Mauritius
and Rodrigues, but appear to be relatively recent ar-
rivals since approximately 2008, suggesting a range
expansion from other areas in the SWIO (Dulau-
Drouot et al. 2012). Within-year recapture rate off
Réunion is relatively high, with long apparent resi-
dency times for at least a subgroup of individuals;
however, whales sighted early in the season appear
more transient (Dulau-Drouot et al. 2011, 2012).

There is significant genetic differentiation between
C1 and C3 animals at both mitochondrial and nuclear
microsatellite loci; however, differentiation is less pro-
nounced than between the SWIO and adjacent stocks
in the southeast Atlantic and southeast Indian Oceans
(Rosenbaum et al. 2009, Kershaw et al. 2016). Con-
verse to the C1–C3 relationship, there is no significant
differentiation between C2 and C3 (Rosenbaum et al.
2009, Kershaw et al. 2016). Similarly, capture-recap-
ture studies indicate low probability of exchange of
individuals between C1 and C3 (Cerchio et al. 2008)
and apparently greater exchange between C2 and C3
(Ersts et al. 2011b) and between C3 and C4 (Dulau-
Drouot et al. 2011). Despite significant population
structure and low probability of interchange based on
capture-recapture analyses, estimated long-term gene
flow rates and effective migrants per generation be-
tween sub-stocks C1 and C3 were relatively high in
comparison to different adjacent stocks (Rosenbaum
et al. 2009, Kershaw et al. 2016).

Thus, current information suggests that humpback
whales are heterogeneously distributed throughout
the SWIO, and there is distinct population structure
between at least sub-populations in the C1 and C3
sub-regions. The sub-population identity of C2 and
C4 is less clear; however, it is possible that each may
have greater interaction with C3 and perhaps repre-
sent range expansions from C3. Alternatively, due to
its proximity in the Mozambique Channel, C2 may
have more interaction with C1 than currently docu-
mented and may act as a conduit for exchange of ani-
ments (and genes) between C1 and C3. Migratory cor-
rridors, specific routes of travel, and the degree of
route fidelity among animals from the same or differ-
ent sub-region, are completely unknown. These
unknowns carry consequence for our understanding
of population status (i.e. abundance and growth rate),
delineation of sensitive habitat, and management
and protection of the population.

The purpose of this study was to use satellite
telemetry to describe movements of individuals dur-
ing the breeding season and thereby make inferen-
ces on the breeding behavior, preferred breeding
habitat and sub-population connectivity of hump-
back whales in the southwestern Indian Ocean. These
factors were assessed by tracking the movements of
individual, satellite-tagged whales on both the east
and west coast of Madagascar which represent the
C3 designated breeding sub-stock. Implantable sat-
ellite tags have been used to remotely monitor move-
ments of various large whale species, in cluding
humpback whales (Zerbini et al. 2006, 2011, Dalla
Rosa et al. 2008, Garrigue et al. 2010, Hauser et al.
2010). Satellite telemetry is ideal for providing infor-
mation on the behavior of individuals and improving
the understanding of a population substructure
which has been previously inferred through genetic
and recapture studies, as well as for addressing
previously unexplored questions such as those
regarding behavioral differences between sexes and
wide-scale habitat utilization during the breeding
season. By tagging whales off both the east and west
coasts of Madagascar during the peak breeding sea-
son, we assessed the relationship between animals in
the sub-regions C3E and C3W and their interactions
with the other sub-stocks, i.e. C1, C2 and C4.

MATERIALS AND METHODS

Tag technology and deployment

The tags used were Wildlife Computers (Redmond,
WA, USA) SPOT 5 transmitters within implantable
cylinder housings (Wildlife Computers Mold 193) and
anchoring systems were a modified version of the
Gales et al. (2009) design. The tags are designed to
penetrate to a maximum of 290 mm into the dorsal
surface of the whale, generally just forward and to the
side of the dorsal fin, and to anchor within the variable
muscle and connective tissue matrix (the fascia) that
underlies the blubber. Tag retention is maintained by
passively deployed sets of barbs. All external components of the tag are built from surgical-quality stainless steel. Tag deployment was carried out from the bow of a small boat at distances of 3 to 5 m using a tagging pole (Heide-Jørgensen et al. 2003, Zerbini et al. 2006) or a modified pneumatic tagging device (Heide-Jørgensen et al. 2001, Gales et al. 2009).

**Sampling design**

Of priority interest were movements of whales during the breeding season and resultant implications for stock structure and breeding behavior, with lower priority given to migration routes to summer feeding destinations. For this reason, we tagged during the period of peak whale density in the breeding season, as opposed to late season which would maximize potential for migratory movements and increased tag duration but miss within-region movements. Tagging was conducted off the northeast coast of Madagascar from Ile Sainte Marie in 2012 and off the southwest coast from the village of Anakao in 2013 (see Fig. 1). These regions were known to have high densities of humpback whales during the breeding season.

An attempt was made to tag equal proportions of adult females and males in order to evaluate differences between the sexes. Both mothers and non-parous females were targeted in equal proportions since reproductive status may affect intra-regional movements and timing of migration. Males and females are impossible to morphologically distinguish from a boat, presenting a challenge for this targeted sample design. Mothers are readily distinguished from males because they are accompanied by a small calf in close association. A non-parous female can be distinguished based upon behavioral cues when the female is the focus of competition between males in a ‘competitive group’ (Tyack & Whitehead 1983, Clapham et al. 1992). In a typical competitive group, a female, termed the ‘nuclear animal’ (NA), is closely guarded by an adult male, termed the ‘principal escort’ (PE), while a number of additional males, termed ‘secondary escorts’ (SE), follow the pair; some SEs attempt to supplant the PE and are designated ‘challengers’ (CH). With adequate observation of position and behaviors, the different roles can be diagnosed allowing inference about sex for both non-parous females (NA) and adult males (PE, SE and CH). In addition to these behavioral categories, whales were tagged as escorts accompanying mothers (esc), as members of a pair of whales (pair) and when found as a solitary individual (solo). Because competition among males can be intense and very physical, it was recognized that tags placed on competitive males could be damaged, shortening the tag duration. We further attempted to tag males and females in associated pairs, either as the ‘NA’ and ‘PE’ in a competitive group, or as a mother and her escort in a mother/calf/escort trio. Tagged whales were biopsied using standard biopsy equipment (crossbow with biopsy dart) and photographed for individual identification using tail flukes (when possible) and dorsal fins. Whole genomic DNA was extracted from biopsy skin tissue samples using the DNeasy Blood and Tissue Kit (Qiagen) and the sex of whales was confirmed using simultaneous PCR amplification of SRY and ZFX/ZFY gene fragments (Jayasankar et al. 2008).

**Data collection and analysis**

Tags were duty-cycled in 2012 to be on 6 h during day (07:00 h to 13:00 h local) and 6 h at night (19:00 h to 01:00 h local) and then increased in 2013 to be on 9 h during day (05:00 h to 14:00 h local) and 9 h at night (17:00 h to 02:00 h local). Location data were obtained from Service ARGOS by instruments on satellites from the NOAA and the European Organization for the Exploration of Meteorological Satellites (ARGOS User Manual, © 2007–2016 CLS, www.argos-system.org/manual/). Raw data included a location class defined by increasing number of messages and decreasing uncertainty from Z, B, A, 0, 1, 2 to 3 and an estimate of spatial elliptical error. Prior to analysis, we removed all locations with class of Z (the least accurate) and applied a speed filter that removed all locations that resulted in a leg speed >12 km h$^{-1}$ (Garrigue et al. 2010), in order to remove the most likely unrealistic positions.

We fitted a behavioral switching state-space model (SSSM; Jonsen et al. 2003, 2005, 2007, Breed et al. 2009) that estimates model parameters by Markov chain Monte Carlo (MCMC) to the locations of each tagged whale using software R v.2.11.1 (R Core Team 2015) and WinBUGS v.1.4 (Bayesian inference Using Gibbs Sampling Project). SSSMs simultaneously solve a model of observation error and a mechanistic model of animal movement (Jonsen et al. 2005) and yield better estimates of the locations and the uncertainty in those locations than raw tracking data. This is because the SSSM draws on all of the data and the animal’s behavior (e.g. speed and turning angles) to predict the probability of an animal being found at a certain location (Jonsen et al. 2003, 2005). To make data comparable between the 2 years and to combine
years for a comparison of sexes, 2013 data were sub-
sampled to match the 6 h on:6 h off duty cycle of 2012
data prior to running the SSSM. We estimated loca-
tions and associated credibility intervals twice daily,
one during the day and once at night, equivalent to a
12 h time step. Behavioral state was classified on the
basis of 2 parameters: mean turning angle (θ) and
autocorrelation in speed and direction (γ). We ran 2
MCMC chains in parallel, each for 50,000 iterations,
including a burn-in of 20,000 iterations which were
discarded. The remaining samples were further
thinned by retaining every 30th sample to reduce au-
tocorrelation; the retained iterations were used to es-
timate the mean and variance for each location and
its respective behavioral state. Thus, the posterior
distribution for each parameter was based on 1000
samples from each chain, giving a total of 2000 inde-
pendent samples. The 2 behavioral states were differ-
entiated by a separation of the values of the 2 param-
eters (θ and γ), resulting in ‘b-mode’ state values of 1
or 2 for each iteration and a mean b-mode calculated
for each estimated position across all iterations. We
nominally referred to the 2 states as: ‘transiting’,
which were highly directional (turning angles near 0°)
and consistent long-distance movements, likely rep-
 resenting transits through breeding habitat or be-
tween distinct breeding habitats (b-mode state of 1);
and ‘localized’, which were more variable movements
with a higher rate of acute turning angles (near 180°),
likely representing searching behavior or meandering
within breeding habitats (b-mode state of 2). Spatial
distribution of behavioral states were evaluated by
plotting the standardized mean of b-modes on a 0.3°
grid, so that each grid cell contained a mean of indi-
vidual means for all positions in that cell.

Linear mixed-effects models (Pinheiro & Bates
2000, Zuur et al. 2009) were used to evaluate the
influence of diel period (day and night), Julian day,
moon phase, tagging day, sex and tagging site on the
SSSM-estimated behavioral mode using the package
nlme (Pinheiro et al. 2015) with R v.3.2.0 (R Core
Team 2015). The response variable (behavioral mode)
was logit transformed prior to analysis, ‘tagged indi-
viduals’ were used as a random effect and an
ARMA(2,0) autocorrelation structure was used to
account for lack of temporal independence within
telemetry data for each whale. Choice of the best cor-
relation structure and model selection followed the
procedures outlined by Zuur et al. (2009): nested
models were fit using maximum likelihood, candi-
date models were ranked using the Akaiake informa-
tion criterion (AIC), and the most supported model
was then fitted using restricted maximum likelihood.

**RESULTS**

**Tag results summary**

Twelve humpback whales were satellite-tagged off
Ile Ste. Marie, Madagascar, between 24 July and 3
August 2012 and 11 whales were tagged off Anakao,
Madagascar, between 16 and 28 July 2013 (Table 1).
We succeeded in tagging a cross-section of sexes,
reproductive classes and behavioral subclasses, with
relatively equal representation of sexes despite the
relatively small sample size (see Table 1 for details on
sex and subclass composition). Tagged whales con-
sisted of 7 males and 5 females off Ile Ste. Marie and
3 males and 7 females off Anakao, confirmed by
molecular sexing, and 1 of unknown sex off Anakao
for which a biopsy was not obtained. Among tagged
females, 5 were mothers and 7 were non-parous
females.

Mean tag transmission duration was (mean ± SD)
24.2 ± 16.1 d (range 2 to 58) for all whales, sexes com-
bined, and 30.1 ± 18.1 d (range 2 to 58) for females and
17.9 ± 11.6 d (range 3 to 34) for males. In 2 cases, the
tag temporarily stopped transmitting for an extended
period, resulting in a gap of 16 d during the 32 d
duration for Tag 1 (a solo male) and a gap of 12 d dur-
ing the 25 d duration for Tag 4 (a mother). To esti-
mate total track distance for these tags, the shortest
Navigable distance was measured between the loca-
tions on either side of the transmission gap (and
added to distances of track segments for total dis-
tances, Table 1); for state-space modeling of these
individuals, only the temporally longer segment for
each tag was modeled. The total number of raw loca-
tions per tag ranged from 8 to 807, and speed filtering
reduced raw locations by 5 to 31% of the original
total (Table 1). Filtered track lengths ranged from
73 km for the shortest duration tag (Tag 16, female,
in pair, 2 d) to 5631 km for the longest duration tag
(Tag 9, female, NA, 58 d).

**Description of movements**

Locations and tracks of tagged individuals are
 grouped by region and sex in Fig. 2 and shown
separately for each individual in Fig. S1 in the
Supplement at www.int-res.com/articles/suppl/m562
p193_suppl.pdf, using the speed-filtered location
data. In some cases, a location occurred on land
when an animal was traveling close to the coast due
to associated error, and we chose not to delete these
locations because they still provide information on
the complete track of the individual. No individuals remained in the immediate vicinity (within 50 km) of the tagging sites for >3 d and most moved out of the area within 1 d. Observed movements were categorized into 4 subclasses: (1) movement along the central east coast of Madagascar, primarily from Ile Ste. Marie, (2) movement along the south and southeast coast of Madagascar, primarily from Anakao, (3) northbound movements and departure from Madagascar, only from Ile Ste. Marie, and (4) southbound movements and departure from Madagascar, from both Ile Ste. Marie and Anakao. Descriptions of movements in each category are as follows (with tag numbers corresponding to those listed in Table 1 and depicted in Fig. S1).

Central east coast movements

Five males (Tags 2, 3, 6, 8 and 11) tagged off Ile Ste. Marie in 2012 spent the entire duration (ranging from 3 to 29 d between 30 July and 29 August 2012) along an approximately 550 km stretch of the Madagascar central east coast south of Ile Ste. Marie and north of latitude 22°S, a region not previously recognized as active breeding habitat (Fig. 2B). Three females (Tags 9, 10 and 12) also spent time in the same general stretch of coast south of Ile Ste. Marie, as part of more extensive travel during relatively similar amounts of time as the 5 males (Fig. 2A,B), including departure from Madagascar to the south and north (Tags 9 and 12, respectively), described below. One mother (Tag 10) traveled steadily north from 1 to 8 August 2012 approximately 640 km to the north tip of Madagascar and then traveled back south approximately 1000 km from 9 to 30 August, with some meandering off the central east coast region. Only 2 individuals tagged off Anakao in 2013 traveled up the east coast of Madagascar into this region, and notably they were the only mothers tagged off Anakao (Tags 20 and 23), such that no male or non-parous female displayed these movements (Fig. 2C).

One mother (Tag 23) reversed direction about 140 km south of Ile Ste. Marie and traveled back to the southeast corner of Madagascar before the tag ceased transmitting.

Table 1. Summary data from 23 humpback whales satellite tagged off Madagascar, at sites in the northeast, Ile St. Marie (SM), and southwest, Anakao (AO). Subclass: Single individual (solo), 1 of 2 individuals in a pair (pair), competitive group (CG), principal escort in a CG (PE), secondary escort in a CG (SE), challenger in a CG (CH), nuclear animal in a CG (NA), mother with calf-of-the-year (mother), escort to a mother (esc). SSSM: switching state-space model, sth migr: southward migration, n/a: not available
Fig. 2. Tracks of all tagged whales (A) from Ile Ste. Marie in the northeast during 2012, with (B) detail of movements off the central east coast, and (C) from Anakao in the southwest during 2013, with (D) detail of movements off the southwest to southeast coasts. Each track is represented by dots for speed-filtered locations (see 'Data collection and analysis') and a line connecting temporally consecutive locations (legs). Females are represented in yellow, males in red, and a single whale of unknown sex in green. Separate figures for each individual are presented in Fig. S1 in the Supplement.
South and southeast coast movements

Only 1 whale tagged off Ile Ste. Marie in 2012 visited the south or southeast of Madagascar, a female tagged as an NA (Tag 9) that moved directly to the south and lingered for 6 d on the Madagascar Plateau before more extensive southerly movements described below (Fig. 2A). Conversely, all but 1 individual tagged off Anakao in 2013 that transmitted for more than a few days travelled to the south and southeast coast of Madagascar (Fig. 2C,D; Tags 14, 15 and 17–23). A non-parous female tagged in a pair (Tag 15) traveled approximately 1000 km around the south of Madagascar and up the east coast to latitude ca. 21°S between 17 July and 1 August 2013, reversed the route back to Anakao and then repeated the same general path before the tag stopped transmitting while on the Madagascar Plateau on 28 August 2013. A male tagged in a pair (Tag 17) followed a similar pattern, travelling to approximately the same point up the southeast coast by 5 August 2013, before turning back south and eventually departing Madagascar on 16 August 2013. Two males (Tags 18 and 19) and 2 non-parous females (Tags 21 and 22) also traveled to the south coast and meandered on the Madagascar Plateau to varying degrees between 21 July and 15 August 2013.

Northbound movements

Five animals, 2 males and 3 females, traveled north from Ile Ste. Marie, 4 of which departed Madagascar on similar northwesterly trajectories (Fig. 2A). Two females (a mother, Tag 7, and a non-parous NA, Tag 12) travelled north ultimately to a location beyond Aldabra Atoll when the tags stopped transmitting, traveling approximately 450 km in 4 d and 500 km in 3 d after departing Madagascar on 8 August 2013 and 21 August 2012, respectively. The NA (Tag 12) appeared to linger around Aldabra for 1 d; however, both females passed through the Aldabra region and were still moving to the northwest when the tags stop transmitting. This represents over 1100 km covered in 13 d for the Tag 7 mother and over 2300 km in 23 d for the Tag 12 female.

Two individuals, a male (solo) and a female (mother), traveled to the central east African coast, but in both cases, the tag did not transmit for the entire transit, so the path can only be inferred (Fig. 2A). The mother (Tag 4) travelled approximately 600 km north between 31 July and 7 August 2012 to the north tip of Madagascar before the tag temporarily stopped transmitting. The tag began to report again on 20 August 2012, when the mother was approximately 15 km off Pate Island, off the north coast of Kenya. The shortest navigable distance between the 2 end locations is 1497 km, so it is likely that she moved steadily on a direct course during the 12 d that would have followed the path of the females Tag 7 and 12 past Aldabra. She then remained within 50 km of Pate Island for 5 d before transmissions ceased; the total transit covered over 2100 km in 25 d. The male tagged as a solo (Tag 1) travelled north 110 km to the mouth of Antongil Bay before the tag temporarily stopped transmitting on 26 July 2012. The tag began to report again on 12 August 2012, when the male was approximately 390 km off the coast of Kenya at latitude 5° 18.9' S. The shortest navigable distance between the 2 end locations is 1540 km, so it is likely that he moved steadily on a direct course during the 17 d, again following the same path of the 3 females past Aldabra. This male then covered 450 km to join the African coast in south Somalia at 1° 25.9' S on 19 August 2012. Thereafter, he moved steadily north up the Somalia coast for 750 km in 5 d, crossing the equator before the tag stop transmitting at 2° 59.9' N. During 32 d, he covered over 2800 km (over 3120 km of trackline), ending within 1200 km of the Gulf of Aden.

Southbound movement

A non-parous female tagged off Ile Ste. Marie as an NA (Tag 9) traveled south immediately after tagging, traveling a somewhat meandering course along the coast approximately 1200 km between 1 and 21 August 2012 to the south tip of Madagascar. This female lingered on the Madagascar Plateau for 6 d before departing Madagascar on a meandering southwesterly course west of the Madagascar ridge between 28 August and 9 September to the west edge of Walters Shoals approximately 900 km south of Madagascar (Fig. 2A). She then turned north and returned to Madagascar between 10 and 18 September, following the west edge of the Madagascar ridge, lingered for 3 d off the southern tip of Madagascar and finally moved south again 700 km before the tag stopped transmitting on 27 September. In total, she covered over 5600 km in 58 d.

A non-parous female tagged off Anakao on 27 July, 2013 in a non-competitive group of 3 whales (Tag 22) moved immediately to the south coast of Madagascar where she lingered on the Madagascar Plateau for 19 d before departing Madagascar to the south on August 15. This female followed a somewhat mean-
dering path for 24 d before arriving on the Crozet Island Plateau on 8 September 2013, approximately 2200 km south of Madagascar (Fig. 2C). The female lingered on the western edge of the plateau approximately 50 km west of the Crozet Islands before departing in a north-northwesterly direction on 18 September; the tag stopped transmitting 2 d later.

Aside from these 4 general movement categories, 1 whale of unknown sex tagged in Anakao (Tag 13) traveled on a somewhat meandering path 470 km north of Anakao between 16 and 25 July 2013, before turning back south and departing the coast of Madagascar on a southwesterly course on 27 July (Fig. 2C). The tag stopped transmitting after another 5 d and 380 km, when the whale was off Europa Island. This was the only case both of a whale moving north up the west coast and departing Madagascar across the Mozambique Channel.

**Behavioral observations of tagged male–female pairs**

On 3 occasions, an associated male and female were tagged in the same group, all off Ile Ste. Marie: once as a mother and escort (Tags 4 and 5, respectively), and twice as a PE to an NA in 2 different competitive groups (Tags 8 & 9 and Tags 11 & 12). In the case of the mother and escort, the pair appeared to remain together for the entire period of the time that both tags were transmitting (5 d, the period of the shorter duration Tag 5 on the male). In both cases of the PE and NA paired tag events, the 2 whales appeared to remain together for the first 24 h after tagging, but then diverged on distinctively separate paths thereafter. In addition to these 3 pairs that were tagged within the same group, there were 2 whales tagged separately off Anakao that appeared to join and travel together many days after the tagging event. A male tagged in a pair on 17 July 2013 (Tag 17) and a non-parous female tagged in a non-competitive group on 27 July 2013 (Tag 22) followed very different paths during the first 30 and 20 d of the tags’ duration, respectively. Then on 16 August 2013, they appear to converge ~40 km off the edge of the Madagascar Plateau and tightly follow the same positions, course and timing for 3 d and 285 km until Tag 17 stopped transmitting on August 19 (Fig. 2C,D; see the 2 tracks departing Madagascar in southeasterly direction). Although it is impossible to be certain if they were associated due to error in tag positions, it appears likely that they were traveling together, given the timing and close proximity of the locations.

**SSSM results**

For the behavioral SSSM analysis, tags that provided <8 d of data or <50 locations (Tags 5, 8, 14 and 16) were not considered, and only the longer segment of those tags that had a large gap in transmission (Tags 1 and 4) was considered. Females displayed a lower mean b-mode value, 1.27 (1.30 when removing southward migration tracks), compared to males, with a mean b-mode of 1.65, suggesting that females engaged primarily in directional transiting movement, whereas males displayed more localized and meandering movement. Average b-modes were similar for animals tagged at either site: 1.45 for Ile Ste. Marie and 1.42 for Anakao (1.46 and 1.46, respectively, when removing southward migration tracks). Spatial plots of the standardized mean b-modes clearly show an overall distinction between sexes when comparing all females (Fig. 3A) with all males (Fig. 3B), with male behavior strongly skewed toward localized movements and female movements strongly skewed toward transiting. Females tended to display areas of more localized movement along the central east coast region and south/southeast coast region (Fig. 3C) where males were also displaying the highest degree of localized movements (Fig. 3D). Thus, although females were transiting through these regions, they tended to linger and meander more than when they moved outside of these regions. The movements of whales that left the coastal shelf of Madagascar mostly conformed to transiting behavior, as might be expected. Exceptions to this include male Tag 1, which displayed a period of localized movement approximately 300 km off the Tanzanian coast (Fig. 3B), female Tag 9, which displayed localized movement at the most southern extent in the vicinity of Walters Shoals and female Tag 22, which displayed localized movement at the most southern extent on the Crozet Plateau (Fig. 3A).

Before linear mixed-effects models were applied to assess variables influencing behavioral state (diel period, tag day, moon phase, tagging day, sex and

| Model        | df | AIC     | ΔAIC |
|--------------|----|---------|------|
| Sex          | 6  | 1981.83 | 0.00 |
| Sex + TagDay | 7  | 1983.47 | 1.64 |
| Sex + DielPer| 7  | 1983.75 | 1.92 |
Fig. 3. Spatial distribution of switching state-space model b-mode values, represented as standardized mean values in a 0.3° grid. Shown are (A) all females and (B) all males for the entire range of tracks, as well as details off the east and south coasts of Madagascar for (C) all females and (D) all males. B-mode is represented on a color gradient from red (1.0, transiting movement) to yellow (2.0, localized movement)
region), portions of tracks for those animals that departed Madagascar from the south were trimmed because these portions were interpreted as southward migratory movements and we sought to assess movements while in the breeding area. The most supported linear mixed-effects model (Table 2) indicated that sex was the only variable to significantly influence the behavioral states of the tagged whales in their breeding grounds off Madagascar. Model coefficients indicated that male b-modes are significantly higher than females (Table 3, Fig. 4), which is consistent with an interpretation that males moved slower and in a more meandering fashion than females during the breeding season.

### DISCUSSION

In this study, we used satellite telemetry in the southwest Indian Ocean to describe movements of humpback whales during the height of the breeding season, providing some novel inferences on exchange between sub-populations within the region, as well as on breeding behavior and definition of breeding habitat. Despite caveats related to sample size constraints, our individuals displayed a remarkable variety of movement patterns, with consistency among multiple individuals for several different patterns and some unexpected long-range movements in relatively short periods.

#### Implications for population movements and structure

Several long-distance movements within the SWIO were documented over relatively short periods of time, all tagged off Ile Ste. Marie. Most notably, 40% of the 10 whales tagged in Ile Ste. Marie that transmitted >10 d displayed long-range movements on the same northwest trajectory towards east Africa. These included a male and a mother that made it to the central east African coast and a non-parous female and a mother that stopped transmitting near Aldabra. These movements to east Africa suggest greater interchange of C1 and C3 sub-populations than previously inferred. Studies of genetic differentiation and photographic mark-recapture suggest a population subdivision between whales from C1 and C3, although notably with relatively high rates of gene flow compared to that between populations in different ocean basins (Cerchio et al. 2008, Rosenbaum et al. 2009, Kershaw et al. 2016). It is important to note that, in both the genetic and photographic comparisons with C3, the C1 samples were collected in the southern portion of the range, in South Africa (a migratory corridor) and southern Mozambique (C1S). The satellite-tagged whales moved directly from Madagascar to the extreme northern portion of the C1 range, Kenya and Somalia (C1N). There are currently no genetic or photo-identification data elucidating the relationship between whales that aggregate in the northern and the southern extremes of the C1 range. Therefore, it is possible that there may be 2 different or partially overlapping migratory streams to the east coast of Africa, one that follows the coastline past South Africa to Mozambique and beyond (as is traditionally described as C1) and a second that passes through the central west Indian Ocean, passing through Madagascar waters en route (as suggested by the individuals tagged in this study). Based on these tagging results, along with population genetic and recapture analyses taken collectively, population substructure and interchange appears to be complex. Recently, the

| Parameter | Estimate | SE   | df | t    | p     |
|-----------|----------|------|----|------|-------|
| Intercept | -0.425   | 0.166| 840| -2.568| 0.0104|
| Sex       | 0.612    | 0.260| 16 | 2.359| 0.0314|

Table 3. Estimates of the fixed-effect coefficient in the most supported mixed-effects model for humpback whale behavioral state in the breeding grounds off Madagascar.

![Fig. 4. Boxplot showing distribution of behavioral states of female and male humpback whales in the breeding grounds off Madagascar. Values are the median of the data (central bar), the lower quartiles (box), upper quartiles (whiskers) and outliers (open circles).](image-url)
IWC completed a comprehensive assessment of this population that estimated post-whaling recovery using models with relatively simple assumptions about population substructure and exchange (Jackson et al. 2015). New inferences implied by these data on exchange between C3 and C1 would likely affect the conclusions of the assessment and further photo-identification, genetic and telemetry studies should be done to elucidate the affiliation of whales off the central East African coast when developing future models.

In addition to the new perspectives that these results imply for sub-stock mixing, novel information was obtained on the specific travel routes through the region. Virtually nothing is known about the direct paths that whales take throughout the west Indian Ocean, beyond the coastal movement along South Africa (Findlay et al. 2011a,b) and scant information from the Mozambique channel based on a few acoustic recordings of song (Best et al. 1998). Here, we have documented a route of travel from Madagascar to Kenya/Somalia through the Outer Seychelles Islands. The large proportion of whales tagged off Ile Ste. Marie that followed this route suggests this may be a prominent movement pattern. Defining important habitat and specific travel routes has particular conservation relevance in relation to threats and habitat degradation associated with the recent discovery of oil and gas reserves and expansion of petroleum industry activities along the east African Coast and in the Mozambique Channel (Rusk & Bertagne 2014, Tyrrell et al. 2015).

Regarding destinations of these whales, of particular consequence is the movement of a male up the Somali coast, across the equator and into Northern Hemisphere waters to 3°N latitude, the first trans-equatorial movement documented for the Indian Ocean. Aggregations of breeding humpback whales are known from Tanzania and Kenya (Findlay et al. 2011b, Amir et al. 2012), but less is known about regions further north. There are historical and modern observations of whales off the north Somali coast, in the Gulf of Aden and the Red Sea during the Austral winter, and suggestions that they may represent southern hemisphere seasonal migrants or vagrants (Brown 1957, Notarbartolo di Sciara et al. 2014). Trans-equatorial migration of whales from the Southern Hemisphere into breeding areas in low latitudes of the Northern Hemisphere has been documented in the eastern South Atlantic (Rosenbaum et al. 2014) and the eastern South Pacific (Rasmussen et al. 2007). This tagged male had traveled over 2800 km in 32 d, and when the tag stopped transmitting, the male was within 1200 km (and 8 d at last travel rate) of the Gulf of Aden, so it is possible if not likely that it traveled into the Arabian Sea region.

The movement of a SWIO whale into this region has potential implications for our understanding of the north Indian Ocean population of humpback whales in the Arabian Sea. To the best of our knowledge, the Arabian Sea population is isolated from the populations in the Southern Hemisphere, as documented through strong genetic differentiation, lack of any photographic recaptures, small population size, reduced genetic diversity, lack of seasonal migration and a Northern Hemisphere breeding cycle (Rosenbaum et al. 2009, Minton et al. 2010, 2011, Pomilla et al. 2014). Although the Arabian Sea population likely originated from the southern Indian Ocean, it has been isolated for approximately 70 000 yr (Pomilla et al. 2014). Given this reported isolation of the Arabian Sea population, and the asynchrony of breeding cycles between Southern and Northern Hemisphere populations, we find it unlikely that this animal’s track signifies substantial mixing between SWIO and Arabian Sea animals. However, it does raise questions over potential range overlap, in part due to range expansion by an increasing SWIO population as witnessed in other areas (Dulau-Drouot et al. 2012).

Despite the documented long-range movements within the SWIO in relatively short periods, no whale traveled to the northwestern coast of Madagascar, Mozambique, the Comoros Archipelago or the Mascarene Islands, where breeding aggregations are well-documented. This may simply be a matter of sample size or timing of tagging, and larger samples along with tagging at different times of season may reveal a greater variety of movements and destinations. At least one whale tagged off Anakao started to cross the Mozambique Channel before the tag stopped transmitting, so there is clearly the potential for movement to Mozambique. Fossette et al. (2014) tagged late in the breeding season in the Comoros Archipelago and documented several whales moving to the west and east coast of Madagascar; therefore, there is exchange between these regions at least during southward migration. In a satellite telemetry study off Reunion Island, a majority of whales tagged during peak season moved west to the east coast of Madagascar (V. Dulau pers. comm.). Therefore, there appears extensive exchange between some whales visiting the Mascarene Islands and Madagascar; however, movement may be only or primarily in a westerly direction.
Southern migration movements

Although our peak-season timing of tagging deliberately deprioritized detection of late-season migration, we expected to increase the probability of documenting a southward migration in targeting non-parous females. It is logically expected that a female, once impregnated, would return to high-latitude feeding grounds as soon as possible, and non-parous females have been documented with shorter residency times on and earlier departure dates from the breeding grounds (Craig et al. 2003). Two non-parous tagged females displayed what can be considered southward migratory movements. The movement of a female to the Crozet Islands Plateau likely represents a southward migration to feeding habitat, and the localized movement behavior near the Plateau indicated by the SSSM suggests a stop for feeding behavior (see also Trudelle et al. 2016). It is not clear whether this represented a short stop on the way to other feeding grounds, however, because the animal did leave the Crozet Plateau before the tag stopped transmitting. Fossette et al. (2014) also documented a migratory transit to the Crozet Plateau from the Comoros Archipelago, so it appears that this may be a relatively common feeding destination or stopover on the way to other feeding grounds.

The movements of the non-parous female Tag 9 (an NA in a competitive group), which left Madagascar for a long-distance transit to Walter Shoals and then returned 21 d later, was particularly unexpected. This type of departure and return has not been previously observed in any other humpback whale breeding area from which there are telemetry data, representing a substantial total sample size of tagged whales (Mate et al. 1998, Zerbini et al. 2006, Lagerquist et al. 2008, Horton et al. 2011, Kennedy et al. 2014), Chittleborough (1958) documented through examination of ovaries that females can have multiple ovulations within a season due to failure of fertilization or early failure of pregnancy. This behavior pattern would be an expected prediction of Chittleborough’s observation, i.e. if the tagged female was in fact fertilized, but the pregnancy failed, and thus she reentered estrus and returned to again search for mates.

Implications for breeding habitat definition

The tracks of tagged whales and results of the state-spaced models suggest that the central east coast of Madagascar represents important breeding habitat for humpback whales tagged off Ile Ste. Marie, whereas the south and southeast coasts represent important breeding habitat for whales tagged off Anakao. Given the distribution of whales around Madagascar, with aggregations in the northeast, southeast and southwest, it was expected that whales would at least transit through the central east coast region; however, our results indicate that whales are utilizing it as preferred breeding habitat. This was not previously recognized, largely due to a lack of research effort in the region and absence of data. The recent development of mining industry activities in Tamatave, located within the stretch of coast utilized by these tagged whales, is of particular conservation consequence due to the potential for coastal pollution from effluents and expanded development.

Existing information for the east coast of Madagascar comes primarily from efforts in Antongil Bay. Concentrations of whales, some with repeated interannual recapture histories, and active breeding habitat have been clearly documented in Antongil Bay (Rosenbaum et al. 1997, Ersts & Rosenbaum 2003, Pomilla & Rosenbaum 2006); however, Cerchio et al. (2009) suggested that animals had short residency and were regularly moving through the bay, based on photographic mark-recapture data from 2000 to 2006. This is congruent with our satellite telemetry results indicating that whales move extensively and are utilizing a larger area. Notably, no individuals tagged in Madagascar were documented entering Antongil Bay, despite its close proximity to the tagging location Ile Ste. Marie; similarly, of 7 whales tagged in 2013 off Réunion Island that moved to Madagascar, 5 of them visited the northeast region off Ile Ste. Marie, but none entered Antongil Bay (V. Dulau pers. comm.). Prior to tagging on Ile Ste. Marie, 3 d (21 to 23 July 2012) were spent in Antongil Bay with no sightings of humpback whales, and anecdotal information from whale-watch tourist groups in the bay indicated that 2012 was 1 of 3 yr in the past 10 yr when very few whales were sighted (A. Saloma pers. comm.). On face value, this lack of use of Antongil Bay is unusual given extensive documentation of high densities in the bay during peak season in previous years, 1996 to 2006 (Rosenbaum et al. 1997, Ersts & Rosenbaum 2003, Cerchio et al. 2008, 2009). Therefore, some variation may exist between years or there may have been a recent shift in distribution patterns, which will require more data and renewed effort in the bay to elucidate.

There was minimal overlap between tracks of individuals tagged off Ile Ste. Marie and Anakao, and a lack of movement to the mid-west and northwest coast of Madagascar, despite such movements representing much shorter transits than the documented
long-range movements. Whales tagged off Ile Ste. Marie largely favored the central east coast of Madagascar, whereas those tagged off Anakao tended to favor the south and southeast coasts. Those whales that did cross into the other tagging area were females (particularly mothers) that tended to show more transiting movement in general. It appears that whales coming to either side of Madagascar may not use the same breeding habitat within a season or possibly movement patterns may vary substantially at different points in the season and whales may move between east and west Madagascar at other times (i.e. earlier or later in the breeding season, similarly to the movement documented from the Comoros Archipelago to Madagascar at the end of the breeding season; Fossette et al. 2014).

**Implications for breeding behavior**

The defined breeding habitat regions were differentially utilized by males and females, with key differences in movement patterns between the sexes in general. Males displayed more variable and localized movement in the defined habitat, whereas females tended to transit through the breeding habitat, covering more distance in shorter periods of time and traveling greater distances overall. However, despite this transiting tendency, females tended toward more localized movement when moving through areas where males were displaying strongly localized movement (the central east coast and the south/ southeast coast regions). This difference between males and females suggests varying mating strategies between the sexes, with males focusing more effort on prospecting a local area for females. Conversely, females may be covering more distance in an effort to expose themselves to a higher number of males, but lingering in areas with higher concentrations of prospecting males. A lek mating system has been suggested for humpback whales (Herman & Tavolga 1980, Clapham 1996, 2000, Connor et al. 2000), which would entail aggregations of males displaying through song and females moving through aggregations assessing male displays and ultimately exerting female mate choice (Höglund & Alatalo 1995) where the lekking arena is relatively small, localized and fixed, the lek for humpback whales might be very large, as well as mobile, as suggested by Clapham’s (1996) ‘floating lek’ proposal. It is noteworthy that no clear distinction was apparent between mothers and non-parous females, with examples of both traveling extensive distances in transiting mode. The sample is too small to draw conclusions and further data are required before relevant comparisons can be made between reproductive classes.

Associations among individual humpback whales on the breeding grounds, including between potentially courting males and females, are thought to be transient in nature (Mobley & Herman 1985, Clapham 1996, 2000). Among 3 male–female pairs tagged in this study, 2 associations appeared to last no more than 1 d; this was not surprising because in both cases the association was between an NA and PE within competitive groups (Tags 8 & 9 and 11 & 12) and thus subjected to disruption by competing males. In the third case, an escort to a mother (Tags 4 and 5), the association appeared to be prolonged for at least 5 d before the tagged male stopped transmitting. In addition, a male and female not tagged together appeared to join post-tagging and remain together for several days (Tags 17 and 22). Andriolo et al. (2014) found similar periods of association (5 and 4 d) from 2 of 4 pairs tagged off Brazil and also documented pairs apparently coming back together post-tagging. Cerchio (2003), in a molecular paternity assessment off the Revillagigedos Archipelago, Mexico, found 2 cases in which mated male–female pairs were sighted together during the year in which the female was impregnated by that male; each mated pair was seen in association twice over the course of 3 and 9 d, respectively, in 2 CG sightings and 2 pair sightings. Although associations on breeding grounds are clearly fluid, and long-term associations may be rare or absent, it is likely that medium-term associations over at least several days may be important in courtship behavior and reproductive success and may be more common than currently realized.

**Future work and perspectives**

This first effort of tracking whales from Madagascar using satellite telemetry has provided new information and some unexpected results. It is clear that there is much more to be learned about this population and region. Tagging during the height of the breeding season provided indications of exchange between breeding areas and definition of preferred
breeding habitat that would not have been obtained otherwise. The composition of our sample of individuals suggests that it is possible to deliberately target a representative sample across the different sexes and reproductive classes; this is a valuable lesson for future work given the tendency for small sample size in telemetry studies. Future satellite tagging effort should focus on increasing the sample size in each subclass, allowing comparisons across subclasses. Given the documented movement patterns, future tagging off Madagascar should also expand spatial effort, both repeating tagging in the northeast and southwest regions and adding, in particular, the northwest coast of Madagascar, where we documented no movements despite the presence of whales in the late breeding season (fossette et al. 2014, S. Cerchio pers. obs.). Moreover, these results demonstrate the importance of broad-scale regional collaboration throughout the SWIO and the north Indian Ocean, given the documented extent of whale movements and potential for population mixing and complex substructure. Increasing the available information on movement patterns of humpback whales throughout the Indian Ocean will provide valuable guidance for future research and conservation efforts. The identification of population substructure and connectivity, as well as the preferred habitat throughout the region, are critical for the development of effective conservation management strategies as threats, such as extractive industry and coastal development, increase.

Acknowledgements. Many individuals made this work possible. This study represents a collaborative effort between the Wildlife Conservation Society (WCS, USA), Institute of Neurosciences Paris Saclay (NeuroPSI, France), Association Cetamada (Madagascar) and NOAA National Marine Mammal Lab (NMML, USA), Maria Faria, Henry Bellon, Boris Andriolo anantenaina and Andrew Wilson provided critical field logistic support and participated in field work; further logistic and administrative support was provided by Sophia Rakotohari malala, Anjara Salomavola, Sylviane Raharivelio, Luccianie Raonison, Cesaire Ramlison, Rina Ralison, Devon Litherland, Victoria Cordi, Ambroise Brenier and Christopher Holmes. For logistical support in the Anaakao region, we thank the staff of IHSM in Tolara, in particular Daniel Ramampihirika and Thierry Laviira, Thierry Bouronville and the staff of Landaka Lodge, Michel Agou and Madame Diamontra from Le Prince Hotel, and the Ministry of Fisheries PACP project. Amy Kennedy provided valuable guidance in analysis. Loraine Mendez, Eric Alfonso, Éléonore Méheust, Stephen Gaughran and François-Gilles Carpenter provided assistance with molecular analysis. Field work was approved under permit from the Madagascar Ministry of the Environment issued to Cetamada for work around Ile Sainte Marie and to WCS for work around Anaakao. Funding for satellite tagging was provided by generous support from the Total Foundation to NeuroPSI and by individuals and foundations to the WCS Ocean Giants Program.

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Editorial responsibility: Peter Corkeron, Woods Hole, Massachusetts, USA

Submitted: April 11, 2016; Accepted: October 25, 2016
Proofs received from author(s): December 21, 2016