

Seabird associations with mesoscale eddies: the subtropical Indian Ocean

K. David Hyrenbach^{1,5,*}, Richard R. Veit², Henri Weimerskirch³, George L. Hunt Jr.^{4,5}

¹Duke University Marine Laboratory, 135 Duke Marine Lab. Rd., Beaufort, North Carolina 28516, USA

²Biology Department, CSI-CUNY, 2800 Victory Boulevard, Staten Island, New York 10314, USA

³Centre d'Etudes Biologiques de Chizé, Centre Nationale Research Scientifique, 79360 Villiers en Bois, France

⁴Department of Ecology and Evolutionary Biology, University of California Irvine, Irvine, California 92697, USA

⁵School of Aquatic and Fishery Sciences, University of Washington, Box 355020, Seattle, Washington 98195, USA

ABSTRACT: We investigated seabird–eddy associations in subtropical waters of the southern Indian Ocean during a summer (January) cruise from Amsterdam Island to Western Australia. To determine habitat associations and to enable predictions about foraging strategies, we related the distributions of satellite altimetry data (sea surface height anomalies) to the 3 most abundant taxa in the study area: terns (sooty *Sterna fuscata* and bridled *S. anaethetus*) and shearwaters (wedge-tailed *Puffinus pacificus*) were significantly more common and numerous within convergence zones. However, a step-wise logistic regression model revealed that these seabirds were more strongly associated with other habitat variables, including the distance to breeding colonies and warm sea surface temperature conditions. Great-winged petrels *Pterodroma macroptera* were widely scattered and did not show associations with convergences or divergences. These contrasting results highlight the need to consider smaller-scale hydrographic features, such as frontal systems, when quantifying seabird associations with mesoscale eddies. Large-scale patterns of ocean circulation and mesoscale variability also seem to play a critical role in determining the extent of seabird–eddy associations. In contrast with previous studies, which have focused on dynamic frontal regions and energetic current systems, we surveyed a region of weak mesoscale variability north of the Subtropical Convergence zone.

KEY WORDS: Altimetry · Indian Ocean · Mesoscale eddies · Remote sensing · Seabirds · Sea surface height anomalies

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INTRODUCTION

Eddies are large (~100 km diameter), long-lived (~months), and pervasive features in the world's oceans (Owen 1981, Robinson 1983, Oschlies & Garçon 1998). These conspicuous hydrographic features influence the horizontal and vertical distributions of the chemical (e.g. nutrients), physical (e.g. sea surface temperature [SST]) and biological (e.g. chlorophyll *a* [chl *a*] concentration) properties in pelagic systems (Yoder et al. 1981, McGillicuddy et al. 1998, Seki et al. 2001, Endo & Wiebe 2005). Moreover, areas of upwelling and downwelling driven by eddy vorticity stimulate primary productivity and aggregate buoyant and weakly swimming organ-

isms, including neuston, plankton, and fish (Olson & Backus 1985, Franks 1992, Strass 1992, Davis et al. 2002).

The ecological significance of these mesoscale features is underscored by evidence that regions where permanent (e.g. Costa Rica Dome; Fiedler et al. 1991) and transient (e.g. Oyashio-Kuroshio confluence; Uda & Ishino 1958) eddies predictably occur are characterized by elevated primary production, high zooplankton biomass, and the aggregation of upper-trophic predators (Yoder et al. 1981, Haney 1986a, Seki et al. 2001, Davis et al. 2002). The fertilizing effects of eddies may be especially important in tropical and subtropical regions, where winter mixing cannot replenish nutrients in the photic zone by eroding the deep (>50 m)

*Corresponding author. Email: khyrenba@u.washington.edu

thermocline (McGillicuddy et al. 1998, Oschlies & Garcon 1998, Seki et al. 2001). These relatively unproductive oceanic regions support far ranging upper-trophic predators that forage on widely dispersed resources and frequently exploit prey concentrated at the periphery of mesoscale eddies (Nel et al. 2001, Polovina et al. 2004, Weimerskirch et al. 2004).

Seabird distributions are clearly influenced by mesoscale hydrographic features (Hunt & Schneider 1987). In particular, many species forage and aggregate at hydrographic fronts and mesoscale eddies (Haney & McGilivray 1985, Abrams & Lutjeharms 1986, Haney 1986b, Nel et al. 2001, Weimerskirch et al. 2004). However, previous vessel-based studies have often focused on the importance of specific types of eddies (e.g. cyclonic vs. anti-cyclonic), without considering the spatial heterogeneity within and among these large habitat features. For example, the smaller-scale (~10s km) fronts at the periphery of mesoscale (~100 km) eddies may be important areas where seabirds aggregate to forage on concentrated prey. Because regions of surface convergence and divergence are characterized by gradients in SST and chl *a* concentration, it is imperative to consider seabird–eddy associations in the context of smaller-scale gradients in water-mass properties (Weller 1985, Haney 1986a,b, Yoder et al. 1994, Ribic et al. 1997).

In the Indian Ocean, seabird–eddy associations have been documented in tropical (e.g. Mozambique Current, 35 to 44° E) and subtropical waters (e.g. Agulhas Return Current, 20 to 80° E) (Abrams & Lutjeharms 1986, Nel et al. 2001, Weimerskirch et al. 2004). The present study addresses seabird–eddy associations within a region of weak mesoscale variability in the eastern Indian Ocean (80 to 115° E) (Park & Gambéroni 1995). First, we set out to determine whether marine birds are associated with mesoscale hydrographic features, defined in terms of sea surface height anomalies (SSHA). Second, we quantify which habitat features (color and temperature fronts, convergence/divergence zones) best explain seabird distributions. To determine whether different seabird groups with specific morphological and behavioural specialisations differ in their oceanographic habitat associations, we relate the sightings of 3 focal taxa (the most abundant species) to the concurrent patterns of SSHA, SST, and chl *a* derived from remote sensing imagery.

MATERIALS AND METHODS

Study area. We analyzed a dataset collected during a 5 d cruise (January 17 to 21, 2003) from Amsterdam Island (37° 52' S, 77° 32' E) to Perth, Western Australia (WA) (32° 01' S, 115° 47' E). The cruise track, which spanned from 410 km northeast of Amsterdam Island to 233 km southwest of Western Australia, surveyed deep (3 to 5 km depth) waters north of the Subtropical Convergence zone (Fig. 1).

Seabird surveys. Two of us (K.D.H. & R.R.V.) surveyed marine birds during daylight hours from the flying bridge of the RV 'Marion Dufresne', eye height of 26 m above the sea surface, while the vessel cruised at speeds between 10 and 15 knots (18.5 to 27.7 km h⁻¹). Following standardized strip transect methods (Tasker et al. 1984), we logged all birds sighted within 300 m of the side of the vessel with best visibility (i.e. least sun glare or wind) into a field computer with a temporal resolution of 0.1 min. Individuals that followed the ship were recorded when first encountered and ignored thereafter. We aggregated the continuous seabird observations into 9 km bins (198), and quantified the occurrence (presence/absence) and the density of each species by dividing the total number of individuals sighted (all behaviors combined) by the area surveyed (km²). Our analyses focus on the 3 numerically dominant taxa (Table 1).

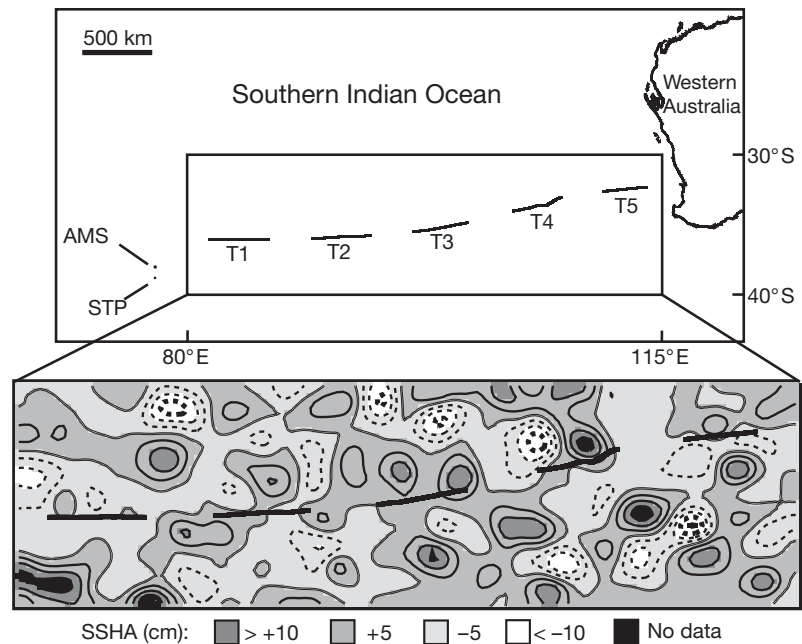


Fig. 1. Map of the study area showing the location of Amsterdam Island (AMS) and St. Paul Island (STP) and the 5 daily survey transects (T1 to T5). The insert depicts the contours of sea surface height anomalies (SSHA) from satellite altimetry, indicative of the distribution of mesoscale eddies, at the midpoint of the survey (January 19, 2003)

Table 1. Summary of seabird taxa recorded across the southern Indian Ocean (January 17 to 21, 2003), showing the total number of birds and the proportion (%) within areas of convergence and divergence, defined by daily sea surface height anomalies (SSHA)

Common name	Species	Total	Convergence (%)	Divergence (%)
Wedge-tailed shearwater	<i>Puffinus pacificus</i>	823	89.2	10.8
Great-winged petrel	<i>Pterodroma macroptera</i>	351	41.1	58.9
Dark tern	<i>Sterna fuscata / anaethetus</i>	118	100.0	0.0
Little shearwater	<i>Puffinus assimilis</i>	19	52.6	47.4
Flesh-footed shearwater	<i>Puffinus carneipes</i>	11	45.5	54.5
Black-bellied storm-petrel	<i>Fregetta tropica</i>	8	100.0	0.0
Unidentified petrel	<i>Procellariidae</i> spp.	6	33.3	66.7
Wilson's storm-petrel	<i>Oceanites oceanicus</i>	5	60.0	40.0
Indian yellow-nosed albatross	<i>Thalassarche carteri</i>	5	40.0	60.0
Long-tailed jaeger	<i>Stercorarius longicaudus</i>	5	80.0	20.0
White-chinned petrel	<i>Procellaria aequinoctialis</i>	4	50.0	50.0
Light-mantled sooty albatross	<i>Phoebastria palpebrata</i>	3	100.0	0.0
White-faced storm-petrel	<i>Pelagodroma marina</i>	2	100.0	0.0
Unidentified giant petrel	<i>Macronectes</i> spp.	1	100.0	0.0
White-necked petrel	<i>Pterodroma externa cervicalis</i>	1	0.0	100.0
9 km survey bins		198	40.9	59.1

Oceanographic data. We used altimetry data from the merged TOPEX/Poseidon and ERS product, with a spatial resolution of 0.25° latitude/longitude and a 10 d repeat cycle, available from the Colorado Centre for AstroDynamics Research (CCAR) (www-ccar.colorado.edu). These altimetry data, which quantify the temporal variability in the height of the sea surface driven by changes in currents (location and speed) and eddies, are indicative of *in situ* surface and subsurface oceanographic conditions (Park & Gambéroni 1995, Davis et al. 2002). We superimposed the survey transects over concurrent daily altimetry maps, and extracted the SSHA values for each 9 km survey bin. This approach assumes that the SSHA values at the centroid of each survey bin are representative of the conditions within an overlying 9 km segment of survey track.

We also relied on satellite imagery to characterize the 2D distributions and gradients in SST and chl *a* along the cruise track. To match the 10 d repeat cycle of the altimetry data, we created 10 d imagery composites for each survey day (January 17 to 21, 2003), by calculating the average SST and chl *a* pixel value during the same 10 d time period sampled by the satellite altimeter. We then matched the seabird survey bins to the imagery composites centered on the same day. That is, the composite spanned from 5 d before to 4 d after the seabird survey.

We used 9 km night Pathfinder SST imagery from the Advanced Very High Resolution Radiometer (AVHRR) (http://podaac.jpl.nasa.gov/pub/sea_surface_temperature/avhrr/pathfinder/data_v4.1/), which on average underestimates *in situ* water temperatures by 0.09°C (Casey & Cornillon 1999). We derived chl *a* along the cruise track from 9 km Sea-viewing Wide

Field-of-view Sensor (SeaWiFS) imagery (<http://seawifs.gsfc.nasa.gov/SEAWIFS.html>), which are within 35 % of *in situ* measurements in the range of 0.05 to 50 mg m⁻³ (Hooker & McClain 2000). The 9 km pixel size allowed us to co-register the SST and chl *a* imagery from the 2 different satellites. This resolution was comparable to the spatial scale (~10 km hydrographic station spacing) successfully used to characterize the temperature and color fronts at the periphery of mesoscale eddies (Ribic et al. 1997, Davis et al. 2002).

In addition to measuring the mean SST and chl *a* for each 9 km survey bin, we quantified the proportional change (PC) in these properties within a surrounding 27 × 27 km (3 × 3 pixel) grid. This gradient metric, which expresses the relative magnitude of change in the habitat variable scaled to the maximum value, ranges from 0 to 100% and is calculated as follows: $PC = [(maximum\ value/minimum\ value) \times 100]/(maximum\ value)$.

Scales of analysis. Because the cruise track was dissected by night-time periods of no survey effort, we combined adjacent 9 km survey bins into 5 disjunct transects (mean transect distance = 356.4 ± 35.8 SD) (Table 2). We assessed the spatial scales (autocorrelation) of environmental variability along each daily transect using Pearson correlation coefficient for lags of 1 to 11 bins (9 to 99 km) (Sokal & Oden 1978). We defined the characteristic patch scale of each environmental variable (SST in units of °C, chl *a* in units of mg m⁻³, SSHA in units of cm) as the point along the space domain where the correlogram first crossed the x-axis (correlation = 0) (Reid et al. 2004).

Seabird response to mesoscale features. To quantify seabird associations with eddies, we contrasted the

Table 2. Summary of environmental conditions and seabird sightings along the 5 daily transects across the southern Indian Ocean (January 17 to 21, 2003). DKTE: dark tern; GWPT: great-winged petrel; WTSW: wedge-tailed shearwater

Transect	Survey length (cm)	SSHA range (km)	Hydrographic feature	Mean (CV)			—Patch scale (km)—			Mean abund. (CV) (birds km ⁻²)	Proportional abundance (%)		
				SSH (cm)	SST (°C)	Chl <i>a</i> (mg m ⁻³)	SSH (cm)	SST (°C)	Chl <i>a</i> (mg m ⁻³)		DKTE	GWPT	WTSW
1	387	-0.7/3.5	Weak convergence	+1.08 (129.9)	18.03 (2.1)	0.18 (11.5)	72	90	63	0.69 (96.4)	0.0	88.9	0.0
2	378	-3.7/2.7	Weak dipole	-0.55 (352.0)	18.34 (1.0)	0.13 (10.8)	72	36	72	0.76 (120.5)	0.0	86.2	0.0
3	369	-9.2/-0.1	Strong divergence	-4.17 (51.0)	18.84 (1.7)	0.25 (29.9)	135	54	90	0.73 (120.9)	0.0	90.0	0.0
4	351	-5.7/1.3	Cyclone edge	-2.26 (103.8)	19.85 (2.9)	0.07 (17.0)	81	135	135	0.84 (114.2)	0.0	79.8	1.1
5	297	-1.1/6.7	Anticyclone edge	+2.67 (96.8)	20.49 (2.8)	0.12 (12.6)	63	108	108	11.48 (305.4)	11.50	5.8	80.4

incidence (e.g. presence/absence) and the abundance of the 3 focal taxa within 4 types of mesoscale features, defined on the basis of SSHA (Ribic et al. 1997, Davis et al. 2002): anticyclones ($> +10$ cm), cyclones (< -10 cm), convergence zones (0 to $+10$ cm), and divergence zones (0 to -10 cm).

Multivariate seabird habitats. To determine which oceanographic conditions most strongly influenced the distribution of the 3 focal taxa, we used logistic regression to relate bird occurrence (binary dependent variable, presence/absence) to 6 continuous predictor variables: SSHA (cm), chl *a* (mg m⁻³), chl *a* gradients (%), SST (°C), SST gradients (%), and the distance to Western Australia (km), where an estimated 2 million wedge-tailed shearwaters and 500 000 sooty terns breed (Surnam & Wooller 2003).

Before we analyzed the seabird–eddy associations, we accounted for the potential autocorrelation in the distributions of the 3 focal taxa and for co-variation in the explanatory environmental variables (Tabachnik & Fidell 1996). We quantified the co-variation of the 6 environmental variables using pair-wise Spearman rank correlations. We also tested whether the 9 km survey bins represented independent sampling units. We quantified whether the presence/absence of each focal taxa within contiguous survey bins was correlated using the joint-count statistic (Sokal & Oden 1978). We used the Rookcase Excel Add-In software, with the rook adjacency rule, to compute the observed and expected number of presence/presence, absence/absence, and presence/absence events, given the total number of bird occurrences along the track (Sawada 1999). Under the null hypothesis of no autocorrelation (independent seabird sightings along the survey track), the observed and expected counts should not differ. Conversely, if the birds aggregate into patches, subsequent observations will be dependent on previous sightings, and the observed and expected frequen-

cies should differ. We contrasted these proportions for each transect separately using *G*-tests.

RESULTS

Oceanographic observations

Our cruise track spanned a heterogeneous subtropical (SST $> 17^{\circ}\text{C}$) seascape, with each daily transect surveying distinct hydrographic conditions (Table 2). Transect 1 spanned a region of weak convergence, with a small SSHA range (4.2 cm) and intermediate chl *a* (mean = 0.18 mg m^{-3}). Transect 2 crossed a weak dipole, a warm-core and a cold-core eddy pair, as evidenced by the variable SSHA conditions (range = 6.4 cm, CV = 352%). Transect 3 surveyed a region of strong divergence, characterized by a broad range of SSHA values (9.1 cm), a large magnitude negative mean SSHA (-4.2 cm), and the highest chl *a* observed during the cruise (mean = 0.25 mg m^{-3}). Transects 4 and 5 sampled the edges of a cyclone (divergence) and an anticyclone (convergence), respectively. Overall, we sampled regions of surface convergence and divergence, with SSHA values ranging from -9.2 to $+6.7$ cm. Because we did not survey the core of any anticyclones ($\geq +10$ cm) or cyclones (≤ -10 cm), our analysis focuses on convergence (SSHA > 0) and divergence (SSHA < 0) zones (Davis et al. 2002).

The spatial patterns of SSHA varied across the 5 daily transects. The region of strong divergence (Transect 3) was characterized by the largest patch scale (135 km), and the edge of the anticyclone (Transect 5) was characterized by the smallest (63 km). The scales of SSHA variability along the other 3 transects were intermediate, with the weak convergence (Transect 1), the weak dipole (Transect 2), and the edge of the cyclone (Transect 4) having patch scales of 81, 81 and 72 km, respectively (Table 2).

Seabird observations

We surveyed 1782 km of cruise track over 5 d, sighted 1361 seabirds belonging to 17 taxa, and identified 91.8% to species level (Table 1). To avoid misidentifications, we combined the counts of sooty *Sterna fuscata*, bridled *S. anaethetus*, and unidentified dark *S. fuscata* or *S. anaethetus* terns, which accounted for 8.5, 2.5, and 89.0%, respectively, of dark-colored tern sightings, into what we termed ‘dark terns’. However, those flocks that we approached closely enough to obtain definite species-specific identification, contained both sooty and bridled terns. Wedge-tailed shearwater *Puffinus pacificus*, great-winged petrels *Pterodroma macroptera*, and dark terns were the 3 most numerous taxa, accounting for approximately 95% of all the sighted birds. The extent to which these species followed the ship was negligible, with the 3 focal taxa showing vessel attraction indices (AI) of -1 , indicating that they were not recorded following the vessel (K. D. Hyrenbach et al. unpubl.).

The behaviors of the 3 focal species differed significantly. The shearwaters were sighted feeding (45.08%) and sitting on the water (41.56%) in similar proportions, and were less frequently seen in flight (13.36%). About 3 quarters of all the petrels sighted were in flight (74.07%), with the rest sitting on the water (21.37%), and feeding (4.56%). Almost all of the terns we sighted were feeding (98.31%), with the rest (1.69%) in flight. The incidence of feeding (G -test: $G = 154.75$, $df = 2$, $p < 0.001$) and flying ($G = 223.93$, $df = 2$, $p < 0.001$) differed significantly across the 3 focal taxa.

Overall, seabird abundance (all species combined) increased from west to east, with the highest densities off Western Australia (Table 2). In particular, Transect 5 was characterized by bird densities that were an order of magnitude (10 times) higher. Additionally, the wedge-tailed shearwater replaced the great-winged petrel as the numerically dominant species off Western Australia (Transect 5).

Ocean habitats

The environmental variables considered in this analysis were cross-correlated (Table 3). Of the 15 possible pair-wise comparisons, 8 yielded significant results ($p < 0.05$). SSHA was correlated positively with SST and negatively with chl *a*, suggesting that areas of surface divergence were characterized by ‘greener’ and cooler waters. SSHA was also negatively correlated with gradients in SST and chl *a*, indicating that regions with strong surface divergence

(e.g. Transect 3) were more homogeneous. Furthermore, chl *a* and SST were negatively correlated, suggesting that regions of localized upwelling supported enhanced primary productivity. We did not detect significant spatial trends in SSHA or in the gradients of SST and chl *a* along the cruise track, as revealed by the lack of significant correlations between these variables and the distance from Western Australia. We did, however, document ‘greener’ (higher chl *a*) and cooler (lower SST) waters to the west, farther from Western Australia. Finally, SST gradients were marginally ($0.10 > p > 0.05$) correlated with SSHA, indicating that warmer regions were characterized by lower temperature gradients.

Seabird–SSHA associations

Overall, 81 (40.9%) and 117 (59.1%) of the 9 km survey bins occurred in convergence and divergence zones, respectively. When we contrasted the relative occurrence (presence/absence) and abundance of seabirds within convergences (SSHA > 0) and divergences (SSHA < 0), 2 focal taxa showed significant associations. For the wedge-tailed shearwater, the number of occurrences (73.7%) and birds sighted (89.2%) were disproportionately more numerous within convergence zones (G -test, $df = 1$, $p < 0.001$ for both comparisons). This pattern was even more extreme for the tern, which occurred exclusively in convergence zones (G -test, $df = 1$, $p < 0.001$ for both comparisons). For the great-winged petrel, the number of occurrences (42.0%) and individuals (41.1%) within convergence zones were proportional to the survey effort (40.9%), suggesting that there were no significant differences in their relative sighting rates or abundances across hydrographic features (G -test, $df = 1$, $p > 0.50$ for both comparisons).

Table 3. Summary of cross-correlations between the environmental variables used to characterize seabird distributions. Chlorophyll *a* (chl *a*) concentration, sea surface temperature (SST), sea surface height anomaly (SSHA), distance to Western Australia (DIST), temperature gradient (TGR), and chlorophyll gradient (CGR). For each pair-wise combination, the matrix shows the sign and the magnitude of the Spearman rank correlation coefficient, and the associated significance level. The bold font denotes significant results (r_s critical = 0.138, $n = 198$). Upper diagonal: p ; lower diagonal: r_s -coefficients

	Chl <i>a</i>	SST	SSHA	DIST	TGR	CGR
Chl <i>a</i>	–	<0.001	0.05–0.02	<0.001	0.02–0.01	0.50–0.20
SST	–0.408	–	0.05–0.02	<0.001	0.10–0.05	0.50–0.20
SSHA	–0.147	+0.148	–	>0.50	<0.001	0.05–0.02
DIST	+0.373	–0.911	+0.022	–	>0.50	0.50–0.20
TGR	+0.172	–0.122	–0.289	+0.042	–	>0.50
CGR	+0.079	–0.059	–0.145	+0.063	+0.012	–

Multivariate seabird habitats

The autocorrelation analyses for the 3 focal taxa revealed no significant patterns for the shearwater (Transect 5, $G = 2.633$, $0.50 > p > 0.25$), for the petrel (Transects 1 to 5, $G = 0.179 - 4.673$, $p > 0.05$ for all 5 comparisons), or for the tern (Transect 5, $G = 0.281$, $0.90 > p > 0.75$) (G -critical = 5.991, $df = 2$ for all comparisons). Thus, we considered the 9 km survey bins as independent samples for the logistic regression analysis. To account for the cross-correlations between the environmental variables (Table 3), we used a step-wise approach to assemble the best-fit model iteratively, by rejecting/retaining the variables with the lowest/highest explanatory power one at a time (Davis et al. 2002).

The best-fit forwards and backwards step-wise models selected the same explanatory variables. A different environmental variable was selected for each focal taxon, with SSHA not being incorporated into any of the best-fit models. The shearwater and the tern were more numerous in warm SST waters closer to Western Australia (Table 4). For these 2 taxa, the best-fit models were highly significant and explained over 90 % of their occurrences. In contrast, the petrel was only marginally correlated ($p = 0.087$) with SST gradients, with the best-fit model only correctly assigning 55.9 % of its occurrences along the cruise track.

DISCUSSION

There is mounting evidence that mesoscale eddies are characterized by enhanced ocean productivity, the retention of zooplankton, and the aggregation of sea turtles, predatory fishes, cetaceans, and seabirds (Nel et al. 2001, Seki et al. 2001, 2002, Davis et al. 2002, Polovina et al. 2004). Previous studies, however, have often focused on the significance of specific hydrographic features (e.g. cyclonic eddies) defined using a single habitat variable (e.g. SST, SSHA). Nevertheless, because mesoscale eddies are characterized by strong gradients in water mass properties (e.g. temperature

and color fronts), it is inherently difficult to determine which hydrographic features most strongly influence seabird distributions (Haney & McGillivray 1985, Haney 1986b, Strass 1992, Seki et al. 2001). This is particularly the case along the periphery of eddies, where many pelagic predators forage (Nel et al. 2001, Polovina et al. 2004, Weimerskirch et al. 2004).

In the present study, we used 2 approaches to assess the association of subtropical seabirds with mesoscale eddies. We first compared the relative occurrence (presence/absence) and abundance (proportion of individuals) of 3 focal taxa within convergence and divergence zones. Next, we used a step-wise procedure to determine which environmental variables, including SSHA and SST/chl *a* gradients, most accurately characterized seabird distributions across our study area.

These comparisons revealed statistically significant associations for 2 taxa. The wedge-tailed shearwater, which was restricted to the 2 eastern most transects (4 and 5), occurred disproportionately more frequently and in higher abundances within convergence regions (SSHA > 0). The logistic regression model selected the distance to Western Australia as the best predictor of wedge-tailed shearwater occurrence within the study area, highlighting the importance of this major breeding colony location as a determinant of the oceanic distribution of this species in the eastern Indian Ocean (Table 4). The wedge-tailed shearwater ranges throughout the tropical and subtropical Indian Ocean, and breeds to the east and west of the study area, though not on Amsterdam or St. Paul islands (Harrison 1987). An estimated 2 million of these birds breed off Western Australia (115–116°E, 31–36°S), where they are the most numerous shearwater species during summer (Surnam & Wooller 2000, 2003). Previous studies suggest that this species breeds during the summer, and feeds predominantly on fish and squid, often in association with subsurface predators (dolphins, tunas) (Harrison et al. 1983, Jaquemet et al. 2004). During the cruise of the present study, these shearwaters most commonly occurred as solitary birds, though they also aggregated into large

Table 4. Results of the step-wise logistic regression, describing the occurrence (presence/absence) for the wedge-tailed shearwater (WTSH), the great-winged petrel (GWPT), and the dark tern (DKTE). For each focal taxa, the significant variables, the associated test statistic and significance levels, and the qualitative description of the model results are shown. Additionally, the log-likelihood results and the proportion of the observations correctly assigned by the best-fit model are reported. WA: Western Australia; SST: sea surface temperature

Focal taxon	Variable	Coefficient (±SE)	t-ratio	p	Result	Log likelihood	df	p	Correct (%)
WTSH	Distance WA	-0.007 ± 0.003	-2.614	0.009	Closer to WA	74.070	1	<0.001	91.4
GWPT	Temp. gradient	-0.165 ± 0.096	-1.714	0.087	Homogeneous SST	2.949	1	0.086	55.9
DKTE	SST	+2.602 ± 1.211	+2.148	0.032	Warm SST	14.815	1	0.022	97.5

feeding flocks (median number of birds per sighting = 1, range = 1 to 200, $n = 67$ sightings).

While dark terns were only sighted within convergence zones, the multivariate analysis revealed that their distribution was most closely related to the higher SST (Table 4). This result is consistent with evidence from the mid-Atlantic United States, where bridled terns aggregated within warm filaments associated with the Gulf Stream (Haney 1986a). Both sooty and bridled terns range and breed throughout the tropical Indian Ocean, though only rarely on Amsterdam or St. Paul islands (Harrison 1987). An estimated 250 000 pairs of sooty terns breed on islands off Western Australia, where their diet consists of cephalopods, euphausiids, and fish (Surnam & Wooller 2003). Some of these prey species concentrate at convergence zones along the periphery of anti-cyclonic eddies (e.g. myctophids, Olson & Backus 1985) and fronts (e.g. by-the-wind-sailor *Velella velella*, Polovina et al. 2004). Additionally, sooty terns often forage in association with subsurface predators (dolphins, tunas) (Jaquemet et al. 2004). During our cruise, these terns sometimes aggregated into large feeding flocks (median number of birds per sighting = 3.5, range = 1 to 100, $n = 8$ sightings).

The third focal species, the great-winged petrel, was widely distributed across the study area (Table 2), with neither its occurrence nor its abundance being significantly related to SSHA. Moreover, the multivariate analysis revealed that the distribution of this petrel was most strongly related to homogeneous areas of low SST gradients (Table 4). These results are consistent with the natural history and the foraging ecology of this species. Great-winged petrels breed in winter to the east (Western Australia) and to the west (Kerguelen) of the study area, and are broadly distributed throughout the subtropical Indian Ocean (Harrison 1987, Surnam & Wooller 2000). This species feeds primarily on cephalopods, and secondarily on fish and crustaceans, captured at the sea surface (Harper et al. 1985). However, because many of its prey are vertically migrating meso-pelagic species (myctophid fishes, ommastrephid squids, euphausiid crustaceans), Imber (1973) hypothesized that this petrel scavenges buoyant squid during the day and takes bioluminescent prey at night. Thus, we would expect this species to forage along convergence zones, where buoyant carcasses should become concentrated. However, our analysis suggests that this species does not associate with specific oceanographic habitats, at least in the daytime. During the cruise of the present study, we largely sighted solitary petrels (median number of birds per sighting = 1, range = 1 to 5, $n = 276$ sightings) engaged in flight (74.07%) or sitting on the water (21.37%). The nocturnal feeding habits of this species may be responsible for the very low incidence of feed-

ing behaviour (4.56%), its dispersed distribution, and the lack of an association with mesoscale eddies or finer-scale hydrographic features.

Our exploratory analysis offers 3 conclusions concerning the study of seabird–eddy associations. First, it highlights the need to clearly define and characterize mesoscale hydrographic features on the basis of quantitative remotely sensed or *in situ* oceanographic measurements (e.g. SSHA, thermocline depth), and to consider smaller-scale convergence and divergence regions in addition to large sized eddies (Ribic et al. 1997, Davis et al. 2002, Weimerskirch et al. 2004). For instance, in this study we considered 4 types of mesoscale features, defined in terms of SSHA: anticyclones ($\geq +10$ cm), cyclones (≤ -10 cm), convergence zones (from 0 to +10 cm), and divergence zones (from 0 to -10 cm). Additionally, we characterized the heterogeneous nature of these mesoscale features by quantifying the variability in SSHA characteristics (e.g. patch scale, magnitude).

Second, our results stress the importance of considering the influence of finer-scale water mass properties, such as temperature and color fronts. In particular, frontal meanders and jets give rise to smaller-scale convergences and divergences, which in turn concentrate upper-trophic predators and their prey (Haney 1986a, Fiedler & Bernard 1987, Polovina et al. 2004). In spite of their short-lived nature, these ephemeral fronts may enhance foraging opportunities for upper-trophic predators (Pitman & Ballance 1990, Podesta et al. 1993), particularly in pelagic systems where prey patches are over-dispersed. Because eddies are characterized by gradients in SST and chl *a*, it may be difficult to ascribe upper-trophic predator occurrence and aggregation solely to the presence of mesoscale eddies (Haney & McGillivray 1985, Davis et al. 2002, Polovina et al. 2004). To deal with these confounding factors, it is essential to quantify the co-variation of the environmental variables, and to use multivariate statistical approaches capable of separating the influence of each individual factor. In this study, univariate analyses suggested that 2 focal taxa aggregated at convergence zones (SSHA > 0). However, the multivariate analyses revealed that these seabirds were more strongly associated with other environmental variables.

Third, it is worth noting that remote sensing imagery can only provide a 2D perspective of 3D mesoscale eddies, and the associated smaller-scale fronts and convergence/divergence zones (Weller 1985, Franks 1992). However, much of the biological enhancement resulting from the presence of mesoscale eddies and frontal features takes place below the surface, where they cannot be monitored using satellite sensors (Strass 1992, Seki et al. 2002). Therefore, mechanistic studies are required to better understand the physical–biological coupling and the links between lower and upper trophic eco-

system constituents that take place at eddies and other mesoscale features.

In summary, this study underscores the need to consider seabird–eddy associations in a broader oceanographic context. It is critical to consider the large-scale (1000s km) patterns of mesoscale variability across ocean basins, which are influenced by ocean currents (Park & Gambéroni 1995). In contrast with previous studies which have focused on dynamic frontal regions (e.g. Gulf Stream fronts off the northeastern United States; Haney 1986) and energetic current systems (e.g. Agulhas Return Current off East Africa, Abrams & Lutjeharms 1986), we surveyed a region of weak mesoscale variability north of the Subtropical Convergence zone (Fig. 2). Our study found that 2 of the 3 focal species we considered (wedge-tailed shearwater and dark terns) concentrated along convergence zones, even though these associations were not significant when we included additional habitat variables in the analysis. This result raises a question about the extent to which oceanic birds exploit mesoscale hydrographic features in subtropical regions. Comparative studies of seabird–eddy associations within regions of enhanced and diminished mesoscale activity are needed to gain a broader understanding of the global significance of these habitat features for oceanic birds (Haney 1986a, Hunt & Schneider 1987).

Because eddy formation is only one of a number of physical processes that arise in dynamic water mass boundaries, it is critical to incorporate an understanding of the smaller-scale (10s km) interactions of eddies with adjacent features (e.g. ocean currents, fronts). In particular, recent modelling efforts suggest that the entrainment of cool nutrient-rich waters from a cold-core cyclone into an adjacent warm-core anticyclone stimulates high primary and secondary productivity, and the aggregation of biomass along the outer edges and trailing filaments of eddies (Logerwell et al. 2001, Lima et al. 2002). Interestingly, satellite tracking studies have revealed that surface feeding albatrosses and frigatebirds forage for squid and fish along the periphery of warm-core eddies, while avoiding the highest SSHA areas at the center (Nel et al. 2001, Weimerskirch et al. 2004).

Future studies should consider the spatial variability in mesoscale variability across ocean basins (e.g. boundary currents vs. subtropical gyres), the spatial heterogeneity within eddies (e.g. periphery vs. core) and the variability across features (e.g. age, size, vorticity). These attributes

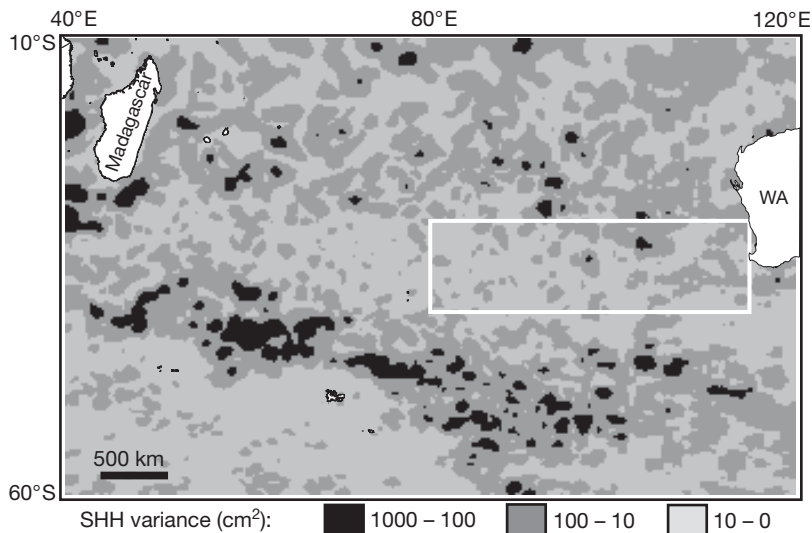


Fig. 2. Map of meso-scale activity in the Southern Indian Ocean during January 2003. Regions of high (black), mid (dark grey), and low (light grey) variability are identified using the variance in absolute sea surface height (SSH) from merged AVISO satellite altimetry data (<http://www.jason.oceanobs.com/>), with a spatial resolution of 0.25° latitude/longitude and a weekly temporal resolution.

The box depicts the study area as shown in Fig. 1. WA: Western Australia

may account for the high variability observed in the extent of seabird aggregation at eddies (Haney 1986a,b). In particular, the 'history' of an eddy (e.g. water mass of origin, interactions with other mesoscale features) likely influences its productivity, and biotic communities (e.g. for discussion of Gulf Stream anticyclonic eddies see Endo & Wiebe 2005).

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