

## Research



**Cite this article:** Williams JJ, Papastamatiou YP, Caselle JE, Bradley D, Jacoby DMP. 2018 Mobile marine predators: an understudied source of nutrients to coral reefs in an unfished atoll. *Proc. R. Soc. B* **285**: 20172456. <http://dx.doi.org/10.1098/rspb.2017.2456>

Received: 2 November 2017

Accepted: 9 February 2018

**Subject Category:**

Ecology

**Subject Areas:**

ecology, environmental science, behaviour

**Keywords:**

acoustic telemetry, grey reef sharks, *Carcharhinus amblyrhynchos*, network theory, nitrogen cycle, Palmyra Atoll

**Authors for correspondence:**

Jessica J. Williams

e-mail: [jessica.williams.16@ucl.ac.uk](mailto:jessica.williams.16@ucl.ac.uk)

David M. P. Jacoby

e-mail: [david.jacoby@ioz.ac.uk](mailto:david.jacoby@ioz.ac.uk)

Electronic supplementary material is available online at <https://dx.doi.org/10.6084/m9.figshare.c.4007623>.

# Mobile marine predators: an understudied source of nutrients to coral reefs in an unfished atoll

Jessica J. Williams<sup>1</sup>, Yannis P. Papastamatiou<sup>2</sup>, Jennifer E. Caselle<sup>3</sup>, Darcy Bradley<sup>4</sup> and David M. P. Jacoby<sup>5</sup>

<sup>1</sup>Department of Life Sciences, Imperial College London, Silwood Park Campus, Buckhurst Road, Ascot, Berkshire SL5 7PY, UK

<sup>2</sup>Department of Biological Sciences, Florida International University, North Miami, FL 33181, USA

<sup>3</sup>Marine Science Institute, and <sup>4</sup>Bren School of Environmental Science and Management, University of California Santa Barbara, Santa Barbara, CA 93106, USA

<sup>5</sup>Zoological Society of London, Institute of Zoology, Regent's Park, London NW1 4RY, UK

**id** JJW, 0000-0002-8275-7597; YPP, 0000-0002-6091-6841; DMPJ, 0000-0003-2729-3811

Animal movements can facilitate important ecological processes, and wide-ranging marine predators, such as sharks, potentially contribute significantly towards nutrient transfer between habitats. We applied network theory to 4 years of acoustic telemetry data for grey reef sharks (*Carcharhinus amblyrhynchos*) at Palmyra, an unfished atoll, to assess their potential role in nutrient dynamics throughout this remote ecosystem. We evaluated the dynamics of habitat connectivity and used network metrics to quantify shark-mediated nutrient distribution. Predator movements were consistent within year, but differed between years and by sex. Females used higher numbers of routes throughout the system, distributing nutrients over a larger proportion of the atoll. Extrapolations of tagged sharks to the population level suggest that prey consumption and subsequent egestion leads to the heterogeneous deposition of 94.5 kg d<sup>-1</sup> of nitrogen around the atoll, with approximately 86% of this probably derived from pelagic resources. These results suggest that sharks may contribute substantially to nutrient transfer from offshore waters to near-shore reefs, subsidies that are important for coral reef health.

## 1. Introduction

The movement patterns of animals can be highly complex, being influenced by both their social and physical environments. Consequently, understanding and accurately measuring population dynamics can be challenging [1–3]. Understanding movement patterns, however, is vital for identifying habitats critical for population connectivity or migration [4], for predicting how stochastic or future environmental conditions will affect populations [5] and for evaluating the effectiveness of protected areas [6]. It cannot simply be assumed that the habitat in which a population is most commonly observed, is that which provides a critical function (a source of food or location for reproduction; e.g. [7]). Therefore, to develop effective conservation approaches, it is essential to understand how a species's behaviour and movement varies across and between habitats [8,9].

In both terrestrial and marine environments, movements of predators can directly and indirectly influence ecological processes such as nutrient cycling and trophic interactions [10,11]. Indirectly, predators can affect nutrient cycling through interactions with prey species; for example, the foraging activities of grazing amphipods and isopods were shifted in response to the presence of predatory blue crabs (*Callinectes sapidus*), contributing to increases in labile organic matter within sea grass ecosystems [12]. More directly, animals can act as nutrient and organic matter vectors, by egesting material within the same habitats in

which the food was consumed, or across habitat boundaries (translocation [10,13,14]). In Alaska, freshwater and/or marine-derived nutrients released by brown bears facilitate growth in white spruce up to 1 km from riparian zones [15]. By foraging at depth and then excreting faeces within the euphotic zone, marine mammals such as humpback and fin whales have been found to replenish nitrogen concentrations at the ocean's surface, thereby enhancing primary productivity (termed the upwards 'whale pump' [16]). Wide-ranging predators such as some whales and sharks also have the potential to contribute significantly to the horizontal transport of nutrients between habitats within marine ecosystems [9,17].

Measuring nutrient transfer between areas and assessing the stability of such flow, however, is non-trivial. There is substantial evidence that the disruption of animal movements can negatively impact productivity through the loss of certain species, posing considerable threat to an ecosystem's long-term resilience [11,18,19]. For example, in the Aleutian archipelago, seabirds act as vectors, transporting nutrients from the ocean to land [20]. However, since the introduction of arctic foxes (*Alopex lagopus*), which have preyed upon seabirds and thus reduced this important nutrient supply, plant communities have been transformed and productivity has decreased [20]. Understanding how predators link habitats and transport nutrients through their environment is, therefore, crucial for ecosystem management.

Palmyra Atoll is a remote, relatively undisturbed coral reef ecosystem, and is part of a US National Wildlife Refuge within the central Pacific Ocean [21,22]. Owing to its protected status, Palmyra has a healthy predator population, with grey reef sharks (*Carcharhinus amblyrhynchos*) being the most abundant predator on the fore-reefs [23,24]. The grey reef shark population at Palmyra is probably at carrying capacity [24], and may play a significant role in the transportation and flow of nutrients onto the reef and throughout the atoll. Grey reef sharks are often detected on (and suggested to favour) outer-reef slopes and drop-off habitats, but on occasion are detected within lagoons [25–27]. Previous research has also highlighted sexual segregation in some grey reef shark populations, which suggests that males and females may connect habitats differently and thus transfer nutrients in differing quantities [25]. Stable isotope analyses at Palmyra have demonstrated that grey reef sharks acquire resources from different habitats, including from pelagic and near-shore environments [7]. However, the use of these habitats for foraging is uneven, with around 86% of grey reef shark biomass being derived from pelagic resources [7]. Mobile species that transport nutrients between habitats have the potential to impact new primary productivity and contribute to the modification of the physical environment [28]. How mobile marine predators such as sharks facilitate this nutrient transport, how much they contribute and how this is subsequently distributed across shallow, productive reef habitats remain unexplored [25,29]. Thus, by transporting materials onto reef habitats that were produced elsewhere, grey reef sharks may generate important linkages between ecosystems and possibly play an ecologically important role in nutrient connectivity. Grey reef sharks show quite strong residency to core areas of the reef, and low rates of movement between reef habitats [27]. However, they are probably transporting pelagic nutrients to fore-reef and potentially back-reef habitats.

Using acoustic telemetry and network analyses, we measure the connectivity generated by the intra- and inter-habitat

movements of predatory grey reef sharks. We then quantify estimates of potential nitrogen transport onto the reef by this species at Palmyra Atoll to understand how nitrogen is probably distributed along different routes of the movement network. We use recent population estimates [24] to extrapolate to the population level in order to assess how significant this nutrient subsidy is likely to be to reef productivity. Owing to previous evidence of sexual segregation in this species in Palmyra Atoll (Y. Papastamatiou and D. Bradley 2012–2014, personal observation), we hypothesize that male and female sharks will have a different influence on nutrient dynamics.

## 2. Method

### (a) Study site and species

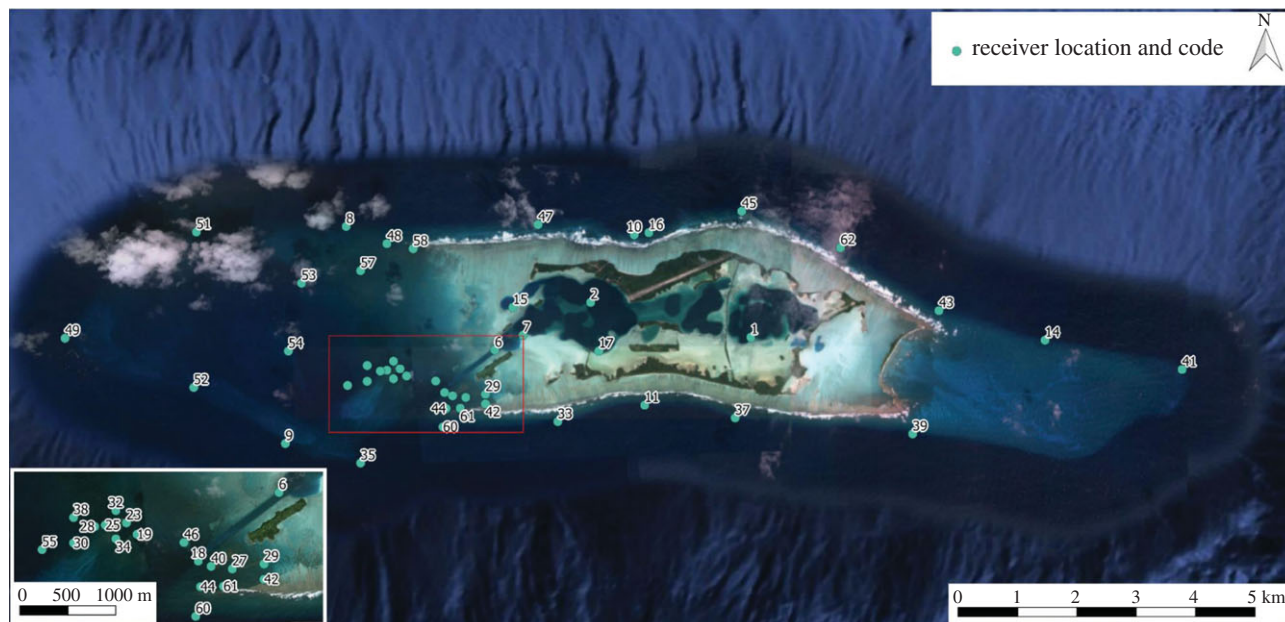
Palmyra Atoll (5°53' N, 162°05' W) is situated in the northern Line Islands in the central Pacific Ocean (figure 1). In 2001, the atoll became a US National Wildlife Refuge, prohibiting take of marine organisms. Since 2001, the only inhabitants have been small (less than 20) groups of researchers and refuge staff [21]. Within the wildlife refuge there is a spatial array of 65 VR2W acoustic receivers (Vemco, Halifax, Nova Scotia, Canada; figure 1). Receivers detect animal-borne, acoustic transmitters at an approximate range of 250–300 m; each time transmitters are detected, the identification number, date and time are recorded by the receiver. Receiver habitat was broadly classified by its geographical zone (lagoon, back-reef or fore-reef).

Detections were recorded from 41 grey reef sharks, comprising approximately 0.49% of the grey reef shark population at Palmyra Atoll [24]. These sharks were tagged with acoustic transmitters (Vemco V16 and V13 coded transmitters) that had been surgically inserted into their body cavities (for details on the method of shark capture and tagging, see [21]). Shark tagging took place on 10 days between 2010 and 2013 at various locations around the atoll. For each individual tagged, we recorded sex and size. Weekly sea surface temperature readings from Palmyra Atoll were obtained from the Coral Reef Ecosystem Program, National Oceanic and Atmospheric Administration (NOAA; electronic supplementary material, S1).

### (b) Network analyses

Applying network theory to acoustic telemetry data allows the movement of sharks to be viewed as a system of connections, in which acoustic receivers are linked by shark movements (for further details on network theory and how it is applied to telemetry data, see [30]). This technique offers insight into how species move between and thus connect habitats [6].

To construct the movement networks, data were initially filtered to include only detections relating to movements of individuals between receivers (i.e. departures and arrivals). However, residency patterns of sharks at each receiver location were calculated from the full dataset (see below for details on residency). To limit transmitters being detected by more than one receiver at the same time (due to some overlap in detection range for a few receivers), the receivers with the greatest overlap were removed from analyses, ensuring that, within the same habitat classification, no two receivers were closer than 150 m (detection distance determined during range testing). Following data filtering, detections from 47 acoustic receivers between January 2011 and December 2014 (1461 days) were included in the analyses. Network theory was employed to analyse these detections, where movement networks measure the relationship between nodes (the acoustic receivers), which are linked by edges (shark movements) [30]. A key temporal measure associated with an edge is its duration: the time between an individual's last detection at one receiver and its first detection at a different



**Figure 1.** Spatial array of acoustic receivers at Palmyra Atoll; only receivers included within the analyses performed in this study are shown. Satellite image from Google Earth. (Online version in colour.)

receiver (time taken to make the movement [3]). As we were interested in movements that potentially led to the transfer of nutrients around Palmyra Atoll, we filtered the data to only include movements within  $\leq 110$  h time windows. This duration represents the length of time, post-feeding, that lemon sharks (*Negaprion brevirostris*) have been observed to continue producing faeces following prey consumption [31]. Lemon sharks are the only species, to our knowledge, for which faecal production time has been measured, and this quantity has been used in previous studies to filter edge duration when focusing on nutrient transfer by marine predators (e.g. [9]). From January 2011 to December 2014, tagged grey reef sharks were detected 848 100 times by the 47 acoustic receivers; this included 99 752 movements between receivers of which 99 342 were  $\leq 110$  h apart (table 1). To explore temporal dynamics, the data were divided into four 'seasons' by examining sea surface temperature data to determine thermally similar three-month periods (December–February, March–May, June–August and September–November).

### (i) Connectivity within the network

To assess the connectivity within the reef ecosystem generated by grey reef shark movements, monthly network edge densities were extracted for each sex for every month of the study. Edge density is the proportion of edges (movements connecting receivers) existing in the network, out of the total number of edges possible for that network (if all receivers were linked by movements to every other receiver [30]). Movement networks with higher edge densities are more densely connected and, thus, individuals have a greater number of routes they can choose from to move through the system [2]. Analyses of variance were used to explore whether network edge densities (the dependent variable) differed between sexes, seasons and years (the independent variables).

### (ii) Estimating nitrogen transfer throughout the atoll

To estimate shark-mediated nutrient flow and highlight areas important to nutrient connectivity, we calculated the potential quantity of nitrogen (N) that these predators may distribute within Palmyra Atoll during the  $\leq 110$  h filtered movement network. Length–weight relationships from Wetherbee *et al.* [32] were used to estimate the weight of each tagged individual

(for all but one female for which total length was not recorded). Egestion rates of N for each individual within Palmyra Atoll per day were then calculated by using the upper limit of 2% (for carcharhinid sharks) of body weight ingested per day [33]. Absorption efficiency was set at 76% for organic matter based on estimates from lemon sharks [31]. We estimated N transfer using the method described by Nelson *et al.* [34], where the total egested kg N per day is the product of the biomass ingested by an individual shark, the biomass egested, the absorption efficiency and the per cent N found in grey reef shark tissue at Palmyra Atoll ( $14.84 \pm 0.065\%$  N mean  $\pm$  s.e. [7]). For every tagged individual, we multiplied their estimated daily egested N (kg) by their residency time within the array, to give a cumulative estimate over the entire study period. To extrapolate to the population level, we took the average length of a male (138.7 cm) and female (146 cm) grey reef shark from [35], and the species abundance and sex ratio estimates from [24]. For all equations used, see electronic supplementary material S2.

Finally, the estimates of daily egested N by tagged sharks were then mapped spatially to explore which areas around Palmyra Atoll were likely to experience the largest influx of shark-derived N. To incorporate the movement of the grey reef sharks with the time they spent in different areas of the atoll, we calculated a sex-dependent dynamic residency score for each receiver. This score incorporated a residency index (the proportion of days the receiver detected a male/female shark over the study period), and the receiver's node strength (which combines the number of connections a node had (i.e. weighted degree) and the relative frequency with which those connections were used). Estimates of N distribution by the tagged male and female sharks were then spatially mapped according to the relative dynamic residency score of each receiver; see electronic supplementary material, S2 for further information. Statistical analyses were completed in R [36] and mapping in QGIS v. 2.14.0 [37].

## 3. Results

### (a) Connectivity within the network

Across the study period, combining movements of both sexes, just under half of all edges possible in the network were



**Table 1.** Summary of the tagged grey reef sharks and their movements detected by the acoustic receivers in Palmyra Atoll; mean values displayed in the table are presented with one standard deviation.

	all sharks	female	male
number of individuals detected			
2011	27	20	7
2012	38	25	13
2013	33	22	11
2014	30	20	10
total	41	28	13
number of movements $\leq 110$ h			
2011	16 665	13 559	3106
2012	21 750	17 177	4573
2013	29 639	24 648	4991
2014	31 288	24 933	6355
total	99 342	80 317	19 025
days at liberty between 2011 and 2014 <sup>a</sup>			
min	21	21	40
max	1439	1437	1439
mean (s.d.)	985 (396)	977 (410)	1001 (380)
fork length (m)			
min	0.86 <sup>b</sup>	0.86 <sup>b</sup>	1.03
max	1.47 <sup>b</sup>	1.47 <sup>b</sup>	1.31
mean (s.d.)	1.18 (0.15) <sup>b</sup>	1.21 (0.17) <sup>b</sup>	1.13 (0.10)

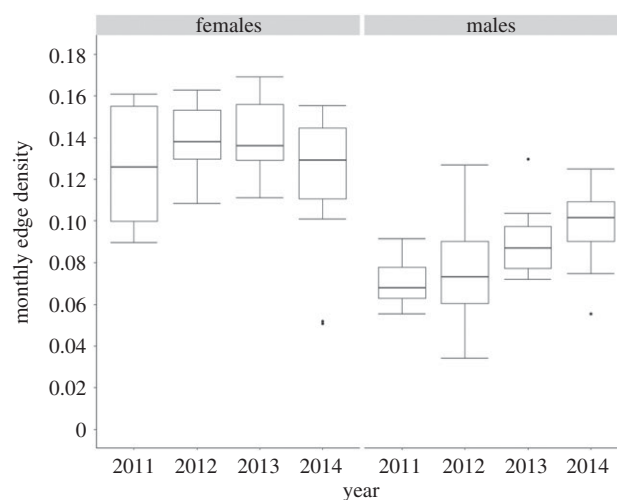
<sup>a</sup>Days at liberty are the number of days between an individual's first and last detection in the dataset.

<sup>b</sup>The fork length for one female was missing, thus for the measurements of all sharks,  $n = 40$ , and for females,  $n = 27$ .

present each year (edge density per year, mean (s.d.) = 0.477 (0.014); electronic supplementary material, S3). Females used a larger proportion of possible routes within the atoll than males, indicated by the female's significantly higher monthly edge density ( $F_{1,94} = 98.8$ ,  $p < 0.01$ ; figure 2). Unlike females, the monthly edge density of the male network differed significantly between years (females,  $F_{1,46} = 0.358$ ,  $p = 0.55$ ; males,  $F_{1,46} = 17.3$ ,  $p < 0.01$ ; figure 2), suggesting that, over the study period, males were less consistent in the linkages generated across the atoll. Between seasons, there was no significant difference in monthly edge density for either sex (females,  $F_{3,44} = 0.920$ ,  $p = 0.44$ ; males,  $F_{3,44} = 0.960$ ,  $p = 0.42$ ).

### (b) Estimating nitrogen transfer throughout the atoll

In total, over the 4-year duration of the study, tagged male ( $n = 13$ ) and female ( $n = 27$ ) grey reef sharks were estimated to have egested  $42.11 \pm 0.19$  and  $119.05 \pm 0.52$  kg, respectively, of N across the atoll and near-shore ecosystem. Given that 86% of biomass consumed by grey reef sharks at Palmyra is thought to be derived from pelagic resources [7], we predict that over the study period (1461 days) there was a maximum potential biomass subsidy of  $138.60 \pm 0.61$  kg N transported onto Palmyra Atoll reefs by the tagged individuals. Based on the frequency with which the tagged sharks moved between



**Figure 2.** The monthly edge densities of movement networks for female and male grey reef sharks (*Carcharhinus amblyrhynchos*) over each year of the study period; these differed significantly between the sexes ( $p < 0.01$ ); the boxplots present the median and quartile values, the circles denote outliers.

and spent time at specific receiver locations, derived from our sex-specific movement networks, we then predict where the nitrogen is likely to be deposited (table 2; figure 3). We also visualize the relative frequency of shark movements between each geographical zone (back-reef, fore-reef and lagoon), to further explore the relative fluxes of N between different habitats (figure 3).

Using the mean length of male and female grey reef sharks sampled by Bradley *et al.* [35], an average male and female individual was estimated to egest as much as  $0.008 \pm 0.00004$  and  $0.011 \pm 0.00005$  kg  $d^{-1}$  of pelagic nitrogen in Palmyra Atoll, respectively. Taking recent population estimates and sex ratios (8344 individuals, 44% male, 56% female) from Bradley *et al.* [24], we extrapolate to the population, estimating a total biomass transfer of  $94.52 \pm 0.42$  kg N  $d^{-1}$ , of which as much as  $81.28 \pm 0.36$  kg N  $d^{-1}$  is a subsidy from pelagic resources brought to the reef by grey reef sharks.

## 4. Discussion

Reef sharks transfer a significant amount of nitrogen to and within an isolated atoll, but the degree of connectivity differs between the sexes, with females using a higher number of routes throughout the near-shore ecosystem. Thus, in doing so, females, which are also typically larger than males, transfer nutrients more broadly across the atoll. Enhancing the understanding of these habitat linkages within reef ecosystems is critical to assist management and conservation strategies, protect movement corridors and respond to potential changes in nutrient dynamics [7].

Higher coverage of N distribution across the atoll by female sharks is probably due in part to female grey reef sharks being larger than males [35], as well as having higher movement rates within near-shore waters. By calculating kernel utilization distributions of acoustically tagged grey reef sharks at Palmyra Atoll, Bradley *et al.* [24] found that, compared with females, the activity space of male sharks was slightly larger, which is supported here by the distribution of dynamic residency scores (figure 3). Along with our results, this suggests that males may disperse

**Table 2.** The five nodes around which the greatest quantity of nitrogen (N) is estimated to have been distributed by the tagged female or male grey reef sharks per day, based on the dynamic residency score of each node; see figure 1 for location of nodes.

acoustic receiver (geographical zone <sup>a</sup> )	node strength	residency index (%)	dynamic residency score	quantity of nitrogen potentially distributed by the tagged grey reef sharks ( $\text{g d}^{-1}$ )
females				
18 (FR)	11 674	89.73	10 474.62	15.66
40 (BR)	9023	81.23	7329.64	10.96
16 (FR)	7360	84.11	6190.47	9.26
10 (FR)	7122	79.66	5673.21	8.48
60 (FR)	4094	92.19	3774.33	5.64
males				
16 (FR)	1704	82.26	1401.72	5.24
60 (FR)	1702	75.55	1285.83	4.80
10 (FR)	1567	55.34	867.22	3.24
18 (FR)	1727	39.11	675.43	2.52
40 (BR)	1413	32.26	455.84	1.70

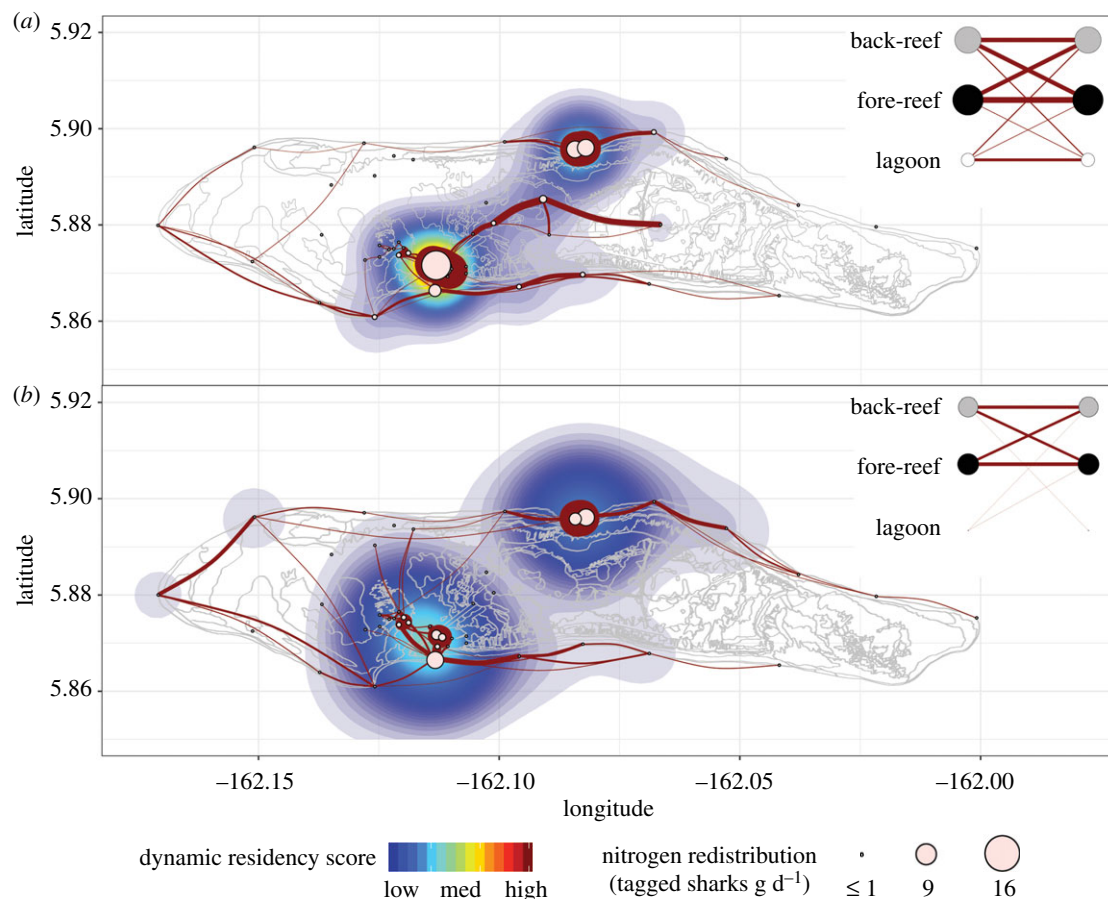
<sup>a</sup>Geographical zones include the fore-reef (FR), back-reef (BR) and lagoon.

more than females, potentially spending more time offshore, beyond the receiver array. Sexual segregation in the near-shore and offshore environment has been recorded in other populations of grey reef sharks, as well as other carcharhinids, and is suggested to be due to differential reproductive or foraging strategies [8]. The difference in routes used between sexes and their differing role in nutrient transfer needs to be incorporated into conservation plans, so areas important to or preferred by males and females are managed appropriately, ensuring each sex maintains their role in fostering connections throughout the ecosystem. Although male sharks used fewer routes, they also increased the proportion of routes they used over the years, even though the number of males detected decreased after 2012. This suggests the movement corridors used are not consistent over time for subsets of the population. This also demonstrates that measuring the efficacy of management strategies such as marine protected areas will require ongoing monitoring, because as animal movement patterns change, spatial strategies may need to be modified to ensure movement corridors remain protected. This may become even more important as marine ecosystems experience rapid effects of climate change.

There were no differences between seasons in the proportion of routes used. Grey reef shark movements on the Great Barrier Reef were not driven by environmental factors such as water temperature, rainfall or wind speed, and more probably related to biotic factors such as reproduction [39]. These results either reflect a resilience to change in environmental conditions within the movement network, or that environmental conditions experienced in these tropical systems were not variable enough to have an impact (average temperature for each season ranged from 27.2°C in March–May 2012 to 29.5°C in September–November 2014). Owing to reef sharks potentially being isolated from alternative suitable habitat, they may have higher tolerances to the range in local environmental conditions to avoid changing movement patterns [39].

Owing to an extensive 8-year tag–recapture programme at Palmyra that has led to accurate estimates of population size [24], we were in a unique position to be able to quantify population-level estimates of N distribution. The within-geographical-zone movements are potentially assisting nutrient recycling, as sharks may be egesting nutrients in the same habitat in which they were consumed [28]. For instance, from our analyses it can be seen that, in some cases, there was a high level of connectivity between nearby receivers (such as between acoustic receivers 10 and 16); this is also reflected in the high proportion of within-geographical-zone movements (figure 3). Moreover, it has recently been shown that grey reef sharks demonstrate strong residency within specific areas at the sub-habitat level [27]. In addition, grey reef sharks demonstrate vertical movement [27,40]; thus, the predators may be transporting nutrients vertically within habitats also [9]. Some routes and receiver locations along which the largest inputs of N were estimated to have occurred by the tagged sharks crossed between reef zones, demonstrating the potential for grey reef sharks to contribute to nutrient translocation. For example, just over 35% of movements by the tagged grey reef sharks that were recorded by the receiver array occurred between the fore-reef and back-reef (figure 3).

With approximately 86% of grey reef shark biomass derived from pelagic resources [7], these sharks may be distributing large quantities of nutrients onto the reef that could not have been produced within the atoll itself. Coral reefs are located in nutrient-limited oceanic waters, yet often support very high biodiversity and productivity [41]. While previous focus has been on tight nutrient cycling, research has shown that within coral reefs, fish are an important nutrient reservoir; both coral growth and primary production are enhanced by fish storing nutrients (in biomass) and egesting them [42–44]. New research indicates that, within reef systems, these fish-derived nutrients may play an important role in the maintenance of ecosystem dynamics [45]. Just how important might these shark-derived nutrient subsidies



**Figure 3.** The 4-year movement networks of (a) female ( $n = 28$ ) and (b) male ( $n = 13$ ) grey reef sharks overlaid on kernel densities that represent dynamic residency at each receiver. Networks include only movements that took  $\leq 110$  h and have been filtered to show the 75 most frequently used routes by each sex. Edge thickness represents the frequency of movements (male range = 36–2711; female range = 129–13 131). The dynamic residency score was calculated as the node strength ( $S_i$ ) of each receiver divided by 100 and multiplied by a standard residency index,  $R$  (male range = 1–1401; female range = 2–10 474). The size of each node represents the potential N redistribution by the tagged grey reef sharks (table 2). The inset habitat networks illustrate the relative frequency of shark movements within and between geographical zones, with the size of the nodes representing the relative detection frequencies in each habitat; the left-hand nodes represent the zone the sharks moved into after last being detected in the habitat depicted by the right-hand node. The base map of Palmyra Atoll was acquired from NOAA National Ocean Service, National Centers for Coastal Ocean Science [38]. (Online version in colour.)

be in Palmyra? Palmyra Atoll has been recorded to have an average of  $1.75 \mu\text{M}$  of dissolved inorganic nitrogen (ammonium, nitrates and nitrites) [46], which corresponds to between 32 and  $109 \mu\text{g}$  of nitrogen-containing compounds per litre of water. In addition, during *in situ* nutrient sampling at locations around Palmyra Atoll (from 2006 to 2012) of nitrate and nitrite, a combined maximum of  $15.21 \mu\text{M}$  was recorded ( $n = 125$ , range  $0.08$ – $15.21 \mu\text{M}$  [47]), corresponding to  $942 \mu\text{g}$  of nitrate and nitrite per litre of water. Therefore, our estimations of an average-sized individual male and female grey reef shark subsidizing the reef with as much as  $0.008 \pm 0.00004$  and  $0.011 \pm 0.00005 \text{ kg d}^{-1}$ , respectively, of pelagic-derived N into the atoll potentially provides a substantial contribution to reef primary productivity. While the precise effects of this nutrient subsidy on Palmyra's benthic communities remain to be explored, changes in grey reef shark population size will probably lead to disruptions in nutrient transport dynamics on this typical, nutrient-limited coral reef. Interspecific interactions between grey reef sharks and blacktip reef sharks may also alter nutrient dynamics, due to strong spatial partitioning between the two species [27]. Removal or reduction of one species may change the degree of among-habitat movements by the other, potentially altering nutrient deposition. For example, a loss of blacktip

reef sharks may cause increased deposition of pelagic N into the lagoons by the grey reef sharks [27].

We recognize that these results should be interpreted with caution due to the fact that we do not know exactly where sharks go once they leave one receiver and arrive at another, and that not all egested material will be deposited within Palmyra Atoll; hence, we stress that these are estimates of *potential* nutrient flow. However, this is the first study to explicitly attempt to measure shark-derived nutrient transfer using a model that incorporates both the movement dynamics and residency patterns of free-ranging sharks. With the current available data and limited knowledge on shark daily rations, absorption and faecal production rates, this study's method enhances our understanding of the role grey reef sharks may play in nutrient connectivity.

Acoustic telemetry data and network theory are emerging as particularly useful tools for exploring habitat use and animal movements [6]. However, acoustic telemetry does have limitations. For example, here, as in many telemetry studies, the number of individuals with active tags was not consistent over the entire study period. This was partly due to some individuals being tagged after the beginning of 2011. In addition, by focusing on movements between different receivers, if sharks left Palmyra Atoll's fore-reef to feed



within pelagic waters and then were next detected on the fore-reef by the same receiver (i.e. self-loops), the movement would not have been included within the analyses. Therefore, the number of movements made by the tagged individuals is likely to be on the conservative side. In addition, acoustic tagging of sharks was spatially non-uniform due to weather-dependent access to sampling sites. This will not affect the quantitative estimates of total N transferred, but it needs to be stressed that the visual representation of N redistribution (figure 3) is a spatial estimation for our tagged sharks only, not the population. Despite the limitations, acoustic telemetry can serve as a powerful instrument to quantify the movements of marine predators, particularly in remote or challenging environments, as well as over large areas [3,6,48].

In light of the fundamental influence that marine predators have on the functioning of ecosystems, understanding how these animals foster within- and cross-system connections is crucial to produce effective conservation and management strategies [7,49]. Palmyra Atoll, one of a limited number of near-pristine atolls, offered a valuable opportunity to assess unrestricted within-system connectivity fostered by grey reef shark movements and their potential role in nutrient transport [7,23]. This study offers a useful comparison for assessments of predator-initiated connections within exploited reefs, to predict the effects of exploitation on undisturbed reefs [50]. Further, it extends our understanding of grey reef shark movements across various reef systems, which is crucial for developing effective conservation approaches and species vulnerability assessments. Finally, it provides the first quantitative estimate, to our knowledge, of population-level nutrient transport in marine predators with implications for the long-term resilience of coral reef ecosystems.

## References

- Patterson TA, Thomas L, Wilcox C, Ovaskainen O, Matthiopoulos J. 2008 State-space models of individual animal movement. *Trends Ecol. Evol.* **23**, 87–94. (doi:10.1016/j.tree.2007.10.009)
- Lédée EJ, Heupel MR, Tobin AJ, Knip DM, Simpfendorfer CA. 2015 A comparison between traditional kernel-based methods and network analysis: an example from two nearshore shark species. *Anim. Behav.* **103**, 17–28. (doi:10.1016/j.anbehav.2015.01.039)
- Jacoby DMP, Freeman R. 2016 Emerging network-based tools in movement ecology. *Trends Ecol. Evol.* **31**, 301–314. (doi:10.1016/j.tree.2016.01.011)
- Heupel MR, Simpfendorfer CA, Espinoza M, Smoothey AF, Tobin A, Peddemors V. 2015 Conservation challenges of sharks with continental scale migrations. *Front. Mar. Sci.* **2**, 1–7. (doi:10.3389/fmars.2015.00012)
- Fortuna MA, Gomez-Rodriguez C, Bascompte J. 2006 Spatial network structure and amphibian persistence in stochastic environments. *Proc. R. Soc. B* **273**, 1429–1434. (doi:10.1098/rspb.2005.3448)
- Espinoza M, Lédée EJ, Simpfendorfer CA, Tobin AJ, Heupel MR. 2015 Contrasting movements and connectivity of reef-associated sharks using acoustic telemetry: implications for management. *Ecol. Appl.* **25**, 2101–2118. (doi:10.1890/14-2293.1)
- McCauley DJ, Young HS, Dunbar RB, Estes JA, Semmens BX, Micheli F. 2012 Assessing the effects of large mobile predators on ecosystem connectivity. *Ecol. Appl.* **22**, 1711–1717. (doi:10.1890/11-1653.1)
- Espinoza M, Heupel MR, Tobin AJ, Simpfendorfer CA. 2015 Residency patterns and movements of grey reef sharks (*Carcharhinus amblyrhynchos*) in semi-isolated coral reef habitats. *Mar. Biol.* **162**, 343–358. (doi:10.1007/s00227-014-2572-x)
- Papastamatiou YP, Meyer CG, Kosaki RK, Wallsgrrove NJ, Popp BN. 2015 Movements and foraging of predators associated with mesophotic coral reefs and their potential for linking ecological habitats. *Mar. Ecol. Prog. Ser.* **521**, 155–170. (doi:10.3354/meps11110)
- Schmitz OJ, Hawlena D, Trussell GC. 2010 Predator control of ecosystem nutrient dynamics. *Ecol. Lett.* **13**, 1199–1209. (doi:10.1111/j.1461-0248.2010.01511.x)
- Saunders M, Brown C, Foley MM, Febria CM, Albright R, Mehling MG, Kavanaugh MT, Burfeind DD. 2015 Human impacts on connectivity in marine and freshwater ecosystems assessed using graph theory: a review. *Mar. Freshw. Res.* **67**, 277–290. (doi:10.1071/MF14358)
- Canuel EA, Spivak AC, Waterson EJ, Duffy JE. 2007 Biodiversity and food web structure influence short-term accumulation of sediment organic matter in an experimental seagrass system. *Limnol. Oceanogr.* **52**, 590–602. (doi:10.4319/lo.2007.52.2.0590)
- Jardine TD, Pusey BJ, Hamilton SK, Pettit NE, Davies PM, Douglas MM, Sinnamon V, Halliday IA, Bunn SE. 2012 Fish mediate high food web connectivity in the lower reaches of a tropical floodplain river. *Oecologia* **168**, 829–838. (doi:10.1007/s00442-011-2148-0)
- Hübner L, Pennings SC, Zimmer M. 2015 Sex- and habitat-specific movement of an omnivorous semi-terrestrial crab controls habitat connectivity and subsidies: a multi-parameter approach. *Oecologia* **178**, 999–1015. (doi:10.1007/s00442-015-3271-0)
- Hilderbrand GV, Hanley TA, Robbins CT, Schwartz CC. 1999 Role of brown bears (*Ursus arctos*) in the flow of marine nitrogen into a terrestrial ecosystem. *Oecologia* **121**, 546–550. (doi:10.1007/s004420050961)

16. Roman J, McCarthy JJ. 2010 The whale pump: Marine mammals enhance primary productivity in a coastal basin. *PLoS ONE* **5**, e13255. (doi:10.1371/journal.pone.0013255)
17. Roman J *et al.* 2014 Whales as marine ecosystem engineers. *Front. Ecol. Environ.* **12**, 377–385. (doi:10.1890/130220)
18. Doughty CE, Wolf A, Malhi Y. 2013 The legacy of the Pleistocene megafauna extinctions on nutrient availability in Amazonia. *Nat. Geosci.* **6**, 761–764. (doi:10.1038/ngeo1895)
19. Momigliano P, Harcourt R, Robbins WD, Stow A. 2015 Connectivity in grey reef sharks (*Carcharhinus amblyrhynchos*) determined using empirical and simulated genetic data. *Sci. Rep.* **5**, 13229. (doi:10.1038/srep13229)
20. Croll D, Maron J, Estes JA, Danner E, Byrd G. 2005 Introduced predators transform subarctic islands from grassland to tundra. *Science* **307**, 1959–1961. (doi:10.1126/science.1108485)
21. Papastamatiou YP, Friedlander AM, Caselle JE, Lowe CG. 2010 Long-term movement patterns and trophic ecology of blacktip reef sharks (*Carcharhinus melanopterus*) at Palmyra Atoll. *J. Exp. Mar. Bio. Ecol.* **386**, 94–102. (doi:10.1016/j.jembe.2010.02.009)
22. Davis K, Carlson PM, Bradley D, Warner RR, Caselle JE. 2017 Predation risk influences feeding rates but competition structures space use for a common Pacific parrotfish. *Oecologia* **184**, 139–149. (doi:10.1007/s00442-017-3857-9)
23. Papastamatiou YP, Lowe CG, Caselle JE, Friedlander AM. 2009 Scale-dependent effects of habitat on movements and path structure of reef sharks at a predator-dominated atoll. *Ecology* **90**, 996–1008. (doi:10.1890/08-0491.1)
24. Bradley D, Conklin E, Papastamatiou YP, McCauley DJ, Pollock K, Pollock A, Kendall BE, Gaines SD, Caselle JE. 2017 Resetting predator baselines in coral reef ecosystems. *Sci. Rep.* **7**, 43131. (doi:10.1038/srep43131)
25. Field IC, Meekan MG, Speed CW, White W, Bradshaw CJA. 2011 Quantifying movement patterns for shark conservation at remote coral atolls in the Indian Ocean. *Coral Reefs* **30**, 61–71. (doi:10.1007/s00338-010-0699-x)
26. Lea JSE, Humphries NE, von Brandis RG, Clarke CR, Sims DW. 2016 Acoustic telemetry and network analysis reveal the space use of multiple reef predators and enhance marine protected area design. *Proc. R. Soc. B* **283**, 20160717. (doi:10.1098/rspb.2016.0717)
27. Papastamatiou YP, Bodey TW, Friedlander AM, Lowe CG, Bradley D, Weng K, Priestly V, Caselle JE. In press. Spatial separation without territoriality in shark communities. *Oikos*. (doi:10.1111/oik.04289)
28. Vanni MJ. 2002 Nutrient cycling by animals in freshwater ecosystems. *Annu. Rev. Ecol. Syst.* **33**, 341–370. (doi:10.1146/annurev.ecolsys.33.010802.150519)
29. McCauley DJ, DeSalles PA, Young HS, Gardner JPA, Micheli F. 2016 Use of high-resolution acoustic cameras to study reef shark behavioral ecology. *J. Exp. Mar. Bio. Ecol.* **482**, 128–133. (doi:10.1016/j.jembe.2016.04.012)
30. Jacoby DMP, Brooks EJ, Croft DP, Sims DW. 2012 Developing a deeper understanding of animal movements and spatial dynamics through novel application of network analyses. *Methods Ecol. Evol.* **3**, 574–583. (doi:10.1111/j.2041-210X.2012.00187.x)
31. Wetherbee BM, Gruber SH. 1993 Absorption efficiency of the lemon shark *Negaprion brevirostris* at varying rates of energy intake. *Copeia* **1993**, 416–425. (doi:10.2307/1447140)
32. Wetherbee BM, Crow GL, Lowe CG. 1997 Distribution, reproduction and diet of the gray reef shark *Carcharhinus amblyrhynchos* in Hawaii. *Mar. Ecol. Prog. Ser.* **151**, 181–189. (doi:10.3354/meps151181)
33. Wetherbee B, Cortés E, Bizzarro J. 2012 Food consumption and feeding habits. In *Biology of sharks and their relatives* (eds JC Carrier, JA Musick, MR Heithaus), pp. 239–264. Boca Raton, FL: CRC Press.
34. Nelson JA, Stallings CD, Landing WM, Chanton J. 2013 Biomass transfer subsidizes nitrogen to offshore food webs. *Ecosystems* **16**, 1130–1138. (doi:10.1007/s10021-013-9672-1)
35. Bradley D, Conklin E, Papastamatiou YP, McCauley DJ, Pollock K, Kendall BE, Gaines SD, Caselle JE. 2017 Growth and life history variability of the grey reef shark (*Carcharhinus amblyrhynchos*) across its range. *PLoS ONE* **12**, e0172370. (doi:10.1371/journal.pone.0172370)
36. R Core Team. 2016 *R: a language and environment for statistical computing*. Vienna, Austria: R Foundation for Statistical Computing. See <https://www.R-project.org/>.
37. QGIS Development Team. 2016 QGIS geographic information system. Open Source Geospatial Foundation Project. See <http://www.qgis.org/> (accessed April 2016).
38. NOAA National Ocean Service, National Centers for Coastal Ocean Science. 2016 Project details. Benthic Habitat Mapping of Palmyra Atoll. See <https://coastalscience.noaa.gov/projects/detail?key=70> (accessed 1 May 2016).
39. Heupel MR, Simpfendorfer CA. 2014 Importance of environmental and biological drivers in the presence and space use of a reef-associated shark. *Mar. Ecol. Prog. Ser.* **496**, 47–57. (doi:10.3354/meps10529)
40. Vianna GMS, Meekan MG, Meeuwig JJ, Speed CW. 2013 Environmental influences on patterns of vertical movement and site fidelity of grey reef sharks (*Carcharhinus amblyrhynchos*) at aggregation sites. *PLoS ONE* **8**, e60331. (doi:10.1371/journal.pone.0060331)
41. Hearn CJ, Atkinson MJ, Falter JL. 2001 A physical derivation of nutrient-uptake rates in coral reefs: effects of roughness and waves. *Coral Reefs* **20**, 347–356. (doi:10.1007/s00338-001-0185-6)
42. Meyer JL, Schultz ET, Helfman GS. 1983 Fish schools: an asset to corals. *Science* **220**, 1047–1049. (doi:10.1126/science.220.4601.1047)
43. Allgeier JE, Yeager LA, Layman CA. 2013 Consumers regulate nutrient limitation regimes and primary production in seagrass ecosystems. *Ecology* **94**, 521–529. (doi:10.1890/12-1122.1)
44. Allgeier JE, Valdivia A, Cox C, Layman CA. 2016 Fishing down nutrients on coral reefs. *Nat. Commun.* **7**, 12461. (doi:10.1038/ncomms12461)
45. Allgeier JE, Burkipple DE, Layman CA. 2017 Animal pee in the sea: consumer-mediated nutrient dynamics in the world's changing oceans. *Glob. Chang. Biol.* **23**, 2166–2178. (doi:10.1111/gcb.13625)
46. Sandin SA *et al.* 2008 Baselines and degradation of coral reefs in the Northern Line Islands. *PLoS ONE* **3**, e1548. (doi:10.1371/journal.pone.0001548)
47. Gove JM *et al.* 2016 Near-island biological hotspots in barren ocean basins. *Nat. Commun.* **7**, 1–8. (doi:10.1038/ncomms10581)
48. Papastamatiou YP, Meyer CG, Carvalho F, Dale JJ, Hutchinson MR, Holland KN. 2013 Telemetry and random-walk models reveal complex patterns of partial migration in a large marine predator. *Ecology* **94**, 2595–2606. (doi:10.1890/12-2014.1)
49. Heithaus MR, Frid A, Wirsing AJ, Worm B. 2008 Predicting ecological consequences of marine top predator declines. *Trends Ecol. Evol.* **23**, 202–210. (doi:10.1016/j.tree.2008.01.003)
50. Barnett A, Abrantes K, Seymour J, Fitzpatrick R. 2012 Residency and spatial use by reef sharks of an isolated seamount and its implications for conservation. *PLoS ONE* **7**, e36574. (doi:10.1371/journal.pone.0036574)