



*Journal of Fish Biology* (2017) **90**, 265–282 doi:10.1111/jfb.13180, available online at wileyonlinelibrary.com

# Environmental drivers of diurnal visits by transient predatory fishes to Caribbean patch reefs

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(Received 3 May 2016, Accepted 7 September 2016)

Video cameras recorded the diurnal visitation rates of transient (large home range) piscivorous fishes to coral patch reefs in The Bahamas and identified 11 species. Visits by bar jack Caranx ruber, mutton snapper Lutjanus analis, yellowtail snapper Ocyurus chrysurus, barracuda Sphyraena barracuda and cero Scomberomorus regalis were sufficiently frequent to correlate with a range of biophysical factors. Patch-reef visitation rates and fish abundances varied with distance from shore and all species except S. regalis were seen more frequently inshore. This pattern is likely to be caused by factors including close proximity to additional foraging areas in mangroves and on fore-reefs and higher abundances close to inshore nursery habitats. Visitation rates and abundances of C. ruber, L. analis, O. chrysurus and S. regalis also varied seasonally (spring v. winter), possibly as fishes responded to temperature changes or undertook spawning migrations. The abundance of each transient predator species on the patch reefs generally exhibited limited diurnal variability, but L. analis was seen more frequently towards dusk. This study demonstrates that the distribution of transient predators is correlated spatially and temporally with a range of factors, even within a single lagoon, and these drivers are species specific. Transient predators are considered an important source of mortality shaping reef-fish assemblages and their abundance, in combination with the biomass of resident predators, was negatively correlated with the density of prey fishes. Furthermore, transient predators are often targeted by fishers and understanding how they utilize seascapes is critical for protecting them within reserves.

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Key words: barracuda; mangroves; marine reserves; snapper; The Bahamas; video analysis.

# INTRODUCTION

Predation is a key influence on the structure of reef-fish assemblages (Hixon, 1991) and has led to an extensive literature on the importance of predator refuges for prey (Hixon

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& Beets, 1993), the consequences of predator removal on trophic cascades (Stallings, 2008) and population regulation through density-dependent mortality (White et al., 2010). While the effects of predation have been examined through a series of correlative and manipulative studies, there are fewer data on the behaviour of predatory fishes. The advent of acoustic tracking technology has led to a growing literature on shark movements (Schlaff et al., 2014), but the behaviour of large, teleost piscivores and smaller meso-predators is less well known (Lédée et al., 2015). The few studies that have been conducted on fishes such as Carangidae, large Lutjanidae, Epinephelinae and Sphyraenidae, however, have highlighted how their movement varies seasonally, diurnally, across tidal cycles and among habitats (Meyer et al., 2007a, b; O'Toole et al., 2011; Matley et al., 2015). Further elucidating the movement of predatory fishes is critical to better understand their functional role in reef ecosystems. In addition, predators are typically the most heavily targeted fishes in tropical fisheries and are important to maintain income from recreational fishers and divers (Stallings, 2009; Hammerschlag et al., 2010) and are a key target of conservation initiatives such as marine reserves (Russ, 2002). Consequently, information on how and why predators move around seascapes is necessary to allow researchers to ensure that spatially explicit management efforts are as effective as possible (Meyer et al., 2007a; Pittman et al., 2014; Lédée et al., 2015).

The need to study the movement of piscivores is particularly true for transient predators [alternatively 'pelagic' predators; Ford & Swearer (2013a)], namely those species that chase prey and forage widely across multiple habitat patches and at spatial scales much larger than their prey home ranges (Carr & Hixon, 1995; Overholtzer-McLeod, 2004). Although there is increasing evidence that even large, mobile species such as carangids may not move as extensively among individual reefs as was presumed (Meyer et al., 2007a; Lédée et al., 2015), their daily movements are typically more expansive and varied than resident predators that remain within predictable and relatively limited home ranges [e.g. epinephelids, Carr & Hixon (1995)]. Consequently, resident and transient species are often quantified separately because of their different home ranges, especially in manipulative studies where resident fishes can be removed from treatment reefs, but transients represent either an uncontrolled predatory threat (Overholtzer-McLeod, 2004) or are excluded through the use of cages (Hixon & Carr, 1997; Ford & Swearer, 2013a). This work has demonstrated that transient predators are an important cause of mortality to fishes on Caribbean (Carr & Hixon, 1995), Pacific (Hoey & McCormick, 2004), sub-tropical (Holmes et al., 2012) and temperate reefs (Ford & Swearer, 2013a). Furthermore, a combination of both transient and resident predators may be necessary for density-dependent mortality of prey fishes and potentially population regulation (Hixon & Carr, 1997).

Like all species, the movement of transient predators will be influenced by interactions with other species, such as prey availability and avoidance of their own predators. Furthermore, the location of a habitat within the seascape can have important influences on the abundance of predatory species, caused by factors such as nursery habitat availability (Mumby *et al.*, 2004). The present study focuses on abiotic factors that receive less attention than biological and benthic structural variables, but may be the most important influences on piscivorous fish abundance patterns (Karnauskas *et al.*, 2012). For example, as ectothermic organisms, fishes have activities intrinsically linked to water temperatures, which will affect spatial resource use, daily activity patterns and seasonal changes in fish behaviours (Lédée *et al.*, 2015). Furthermore, time

of day influences the feeding rates of small predators, with greater activity of moon wrasse *Thalassoma lunare* (L. 1758) during the mid-afternoon (Holmes *et al.*, 2012) and small epinephelids being more active during crepuscular periods (Randall, 1967). Water movement also has important influences on the species seen across gradients from sheltered to exposed habitats, because only some species with particular fin morphologies are able to cope with high wave energy environments (Fulton *et al.*, 2005).

This study considers the effects of abiotic and some biotic variables on one aspect of the movement of transient predators within a Caribbean lagoon complex, particularly their visits to patch reefs. Although transient predators visit both natural and artificial patch reefs (Carr & Hixon, 1995; Hixon & Carr, 1997; Overholtzer-McLeod, 2004; Karnauskas *et al.*, 2012), the factors determining species-specific visitation rates are rarely studied. Predation rates on patch reefs are important because this habitat functions as a site of direct settlement for juvenile fishes (Carr & Hixon, 1995) and as an intermediate habitat during ontogenetic shifts by nursery using species from seagrass and mangrove areas to adult habitats (Mumby *et al.*, 2004). Consequently, predator-driven fish mortality rates on patch reefs may have important demographic consequences, especially as lagoons may be the preferred habitat of some transient species (O'Toole *et al.*, 2011). Furthermore, the spatial separation of prey-rich patch reefs within an environment dominated by fish-depauperate, soft-sediment habitats provides an opportunity to investigate variability in the abundance of transient predators within a complex foraging seascape.

Cameras were used to record diurnal visitation rates and abundances of transient predators to patch reefs across a gradient of increasing distance from shore, which encompassed a range of seascape-scale variables. In addition, by deploying the cameras at different times of day and tidal states, visitation rates could be linked to current speed and direction and hours after sunrise. Finally, by filming the patch reefs in both winter and spring, the study aimed to detect differences in transient predator abundances over seasonal time scales. Linking the diurnal visitation rates and abundances of transient predators to actual mortality rates of prey fishes is problematic because of the difficulty of detecting rare predation events. Therefore, this study focuses on detecting which variables are most important for influencing visitation rates by transient predators, which is assumed to be a proxy of their diurnal predation pressure. It was hypothesized that this estimate of predation pressure would be consistent across the patch reefs within a section of a single lagoon because of the extensive home ranges of transient predators.

### MATERIALS AND METHODS

### STUDY SITE

Data were collected in November to December 2011 (winter) and April to May 2012 (spring) from patch reefs close to Cape Eleuthera, The Bahamas (Fig. 1). The lagoon area east of Cape Eleuthera is c. 2-3 m deep and predominantly comprises soft-sediment habitats, but also contains hundreds of patch reefs of various sizes. This study focused on six small ( $\leq 30 \,\mathrm{m}^2$ ) reefs (Fig. S1, Supporting Information). Small reefs were chosen as they are common in the area (Fig. S2, Supporting Information) and allowed a camera to film the whole reef and immediate surrounding area from a distance that was close enough to allow for species-level identification. The reefs were positioned along a gradient of increasing distance from the shoreline (offshore gradient). The reefs were an average of 1.07 km (minimum 0.31, maximum 1.56 km) apart and

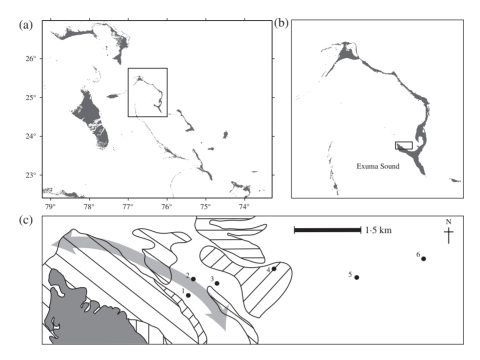


Fig. 1. Location of (a) Eleuthera within The Bahamas, (b) the study area and (c) the six patch reefs used in the study (1–6) and the surrounding habitats. Deep water and fore-reef habitats are found just to the west of Eleuthera, bordering Exuma Sound. ↑, approximate directions of strong, inshore tidal currents; □, land; □, sparse seagrass and algae; □, sand with *Batophora*; □, sand, *Sargassum*, seagrass, sponges and other algae; □, sand and sparse seagrass; □, sand, algae, sparse seagrass; sponges, gorgonians and corals.

ranged from 1·45 to 6·78 km from the shoreline. By focusing on patch reefs of varying distance from shore, this study examined whether transient predator visitation rates were affected by a range of biophysical variables. Eleuthera has a semi-diurnal tidal regime with a maximum range of only c. 80 cm (Murchie et al., 2010), but the geomorphology of Cape Eleuthera leads to strong ebbing and flooding currents close to the shoreline (Fig. 1). Maximum current speed then decreases with increasing distance from shore. The variation in maximum current speeds is a significant influence on the distribution of lagoon habitats, meaning that the habitats surrounding each patch reef vary with increasing distance from shore (Fig. 1). Furthermore, the shoreline of Cape Eleuthera supports a series of mangrove creeks that provide important nursery areas and foraging grounds for juvenile and adult transient predators respectively (Mumby et al., 2004; Harborne et al., 2016). As these variables co-vary across the offshore gradient any patterns cannot be definitively attributed to a single factor. If visitation rates varied significantly across this gradient, however, the study provides a list of potential drivers and hypotheses of transient predator distributions that will inform further research in seascapes where their effects may be evaluated independently.

Each reef was surveyed in detail prior to filming to quantify any systematic differences in the characteristics of the patch reefs across the offshore gradient. These surveys quantified the biomass of other meso-predators, including serranids [mainly sub-adult Nassau grouper *Epinephelus striatus* (Bloch 1792) and graysby *Cephalopholis cruentata* (Lacépède 1802)] and the invasive lionfish *Pterois volitans* (L. 1758) and *Pterois miles* (Bennett 1828) that are abundant on the patch reefs of Eleuthera. Predator biomasses were calculated from *in situ* visual estimates of length using allometric relationships (Froese & Pauly, 2010). Visual surveys also determined the densities of potential prey items (Pomacentridae, small Labridae and juvenile Scaridae and Haemulidae). Coral and algal cover was measured using replicate (n = 10-15)

 $0.25 \text{ m}^2$  video quadrats placed randomly on each reef. Furthermore, the length, width and maximum height of each reef were measured, along with replicate (n = 8 - 21) 1 m chain transects to measure habitat complexity (Luckhurst & Luckhurst, 1978). Variables of patch-reef characteristics were logit [for proportional data; Warton & Hui (2011)] or ln transformed where necessary to fulfil linear modelling assumptions prior to linear regression against the distance from shore.

Tidal-flow rates and temperatures at each patch reef could not be monitored continuously during the video deployments and therefore models of current flow were generated using data collected in March 2015. A TCM-1 Tilt Current Meter (Lowell Instruments, LLC; http://www.lowellinstruments.com/) was deployed at each reef for a mean of 9.4 tidal cycles, (with the exception of reef 2, which was modelled using an average of the data from reefs 1 and 3 because of their proximity). A regression model was fitted to current-speed data from each reef using the explanatory variables of time since previous slack, a quadratic term of time since previous slack and height of the previous slack tide as predicted by tide tables. These models were then used to predict the current speed at each reef every 15 min during the camera deployment period. The current meter also recorded the temperature at each reef at midday. Although temperature was not monitored continuously during the study, a temperature logger (HOBO Pendant Temperature/Light; http://www.onsetcomp.com/) recorded seasonal changes in temperature every 30 min from March 2014 to March 2015 at patch reef 6. Data were summarized as the mean temperature each day, along with the maximum daily variation.

### VIDEO DATA COLLECTION

Video cameras were used in this study because they have numerous advantages over underwater visual censuses for transient (and often rare) predators, including the ability to monitor multiple locations for relatively long periods simultaneously, creating a permanent record of each fish seen and reducing *in situ* disturbance. A GoPro camera (www.gopro.com) was placed 3 m from each reef (total of six cameras day<sup>-1</sup> of filming) and typically ran for c. 4 h (mean  $\pm$  s.D. = 239·8  $\pm$  56·3 min). Filming was undertaken using a crossed experimental design to record transient fishes at both different times of day (from soon after sunrise to sunset) and different tidal states. All transient predators passing over or close to a patch reef (field of view c. 5·6 m at 3 m from the camera representing a filmed area of 8·4 m<sup>2</sup>) were identified and counted.

The majority of fishes were only seen briefly  $(<30 \,\mathrm{s})$  while swimming past the patch reefs, but some remained in the field of view for longer periods. While care was taken to try and not record the same fish multiple times, tracking highly mobile individuals was problematic across the entire duration of filming, particularly for transient species that form large shoals [e.g. bar jack Caranx ruber (Bloch 1793)]. Therefore, counts of each species of transient predator were conducted for consecutive 5 min time periods. Because each replicate count represented the abundance of transient predators on each patch reef per unit of time (5 min) individual fish only needed to be tracked for a short time. Consequently, a transient predator remaining around a patch reef for an extended (>5 min) time period was only recorded once within the 5 min count when it first arrived at the reef, but could also be recorded in subsequent time periods. Recording a fish in multiple 5 min periods was consistent with the aims of the study, which were to identify variability in the potential predation pressure caused by transient predators. Thus a fish spending an extended period of time at a reef was a greater threat to prey species than an individual moving quickly past the reef and this residence time was reflected in the data set in a way that would not have occurred if only arrival time had been recorded. While tractable for analysing the video footage, the temporal resolution of this technique is limited to 5 min (i.e. fish present for <1 min are not distinguished from fish present for 4-5 min), but it was assumed that visits <5 min were functionally equivalent in terms of predation risk. Although the focus of this study was on assessing visitation rates of transient predators to the reefs and the videos were not sufficiently detailed to record predation events, any apparent hunting or feeding behaviour by the predators was documented.

For each daily camera deployment, 5 min fish counts (ranging in number from one to 12 individual counts) were averaged to calculate the mean number of fish per species 5 min<sup>-1</sup> present at each patch reef during each hour surveyed after sunrise (time segment; first time segment from sunrise to 1 h after sunrise, second time segment from 1 to 2 h after sunrise and so on). Each hour-long segment was associated with a time after sunrise by calculating the time from sunrise

to the mid-point of each time segment (*e.g.* 30 min after sunrise for the first time segment). The current speed and direction (ebb or flood) for each hour-long segment was then estimated from the model prediction at the mid-point of the segment. A schematic overview of the derivation of the data from the videos is provided (Fig. S3, Supporting Information).

### DATA ANALYSIS

For even the most common transient species, fishes were recorded during <70% of the 1 h time segments, leading to zero-inflated and left-skewed data distributions. Therefore, the data for each species were modelled using two-part (hurdle) models to account for zero inflation (Fletcher et al., 2005; Zuur et al., 2009). Firstly, the data were transformed from number of fish seen 5 min<sup>-1</sup> to presence – absence 1 h<sup>-1</sup> segment and analysed using generalized linear models with binomial error structures and the logit-link function. Explanatory variables were season (spring or winter), distance from shore (including a quadratic term to examine curvilinear relationships), hours after sunrise, current speed and current direction (ebbing or flooding), plus the interaction between the two water-flow variables. Because the number of 5 min intervals recorded in each 1 h segment varied and could affect the probability of recording a visit by each species, the number of intervals (ranging from one to 12) was also included as a covariate in the model. Since data from individual 1 h segments were nested within daily camera deployments (i.e. each day's camera deployment generated data for multiple time segments), a random variable representing camera deployment was included within the analysis. Therefore, generalized linear mixed-effects models were performed using the lme4 package in R (www.r-project.org; Bates et al., 2014).

The second part of the hurdle model analysed the mean number of fish recorded  $5 \, \mathrm{min}^{-1}$  within each 1 h time segment, but only when that species did visit a patch reef (*i.e.* modelling non-zero values only). The explanatory and random variables were as for the binomial model, with the exception of the variable representing the number of  $5 \, \mathrm{min}$  intervals surveyed that was omitted because the response variable of number of fish  $5 \, \mathrm{min}^{-1}$  was independent of survey effort. These were fitted using linear-mixed effects models, with Gaussian error structures and the identity link function. Fish abundances were only modelled if the species occurred in > 10% of 1 h segments. Response variables were ln or reciprocal-root transformed when necessary to improve normality of residuals. Where required, the product of the predicted probability of a visit by each species (from the binomial models) and the predicted number of fish during a visit (from the Gaussian models) were used to predict the number of fishes at each patch reef at any time.

Finally, to investigate any predator-prey relationships, the density of prey species was regressed against both the modelled abundance of transient predators and the surveyed biomass of resident predators at each patch reef and an interaction term between predator abundances.

Both binomial and Gaussian models were fitted using the procedure outlined by Crawley (2007). Briefly, a maximal model was fitted including all factors and the interaction. Least significant terms were then removed in turn, starting with the interaction. After each term was removed, models were compared to ensure that term removal did not lead to an increase of >2 of the Akaike information criterion (AIC) (Burnham & Anderson, 2004) or a significant increase in deviance. Terms were removed until the model contained only significant terms or removal of any non-significant terms caused a significant increase in deviance or an increase of >2 of AIC (minimal adequate model). Minimal adequate models were checked for violations of assumptions.

### RESULTS

# ABIOTIC AND BIOTIC CHARACTERISTICS OF THE PATCH REEFS

The physical characteristics of the six patch reefs were relatively homogeneous, with each having a similar size, height above the bottom and complexity (Table I).

TABLE I. Regression coefficients for abiotic and biotic characteristics of the six patch reefs filmed for transient predator visitation rates, and the results of regression analysis of each factor against distance from shore (regression coefficient and significance shown). Prey category comprise Pomacentridae, small Labridae and juvenile Scaridae and Haemulidae

off Reef (	Distance offshore (km)	Height (m)	Area (m <sup>2</sup> )	Rugosity	Coral cover (%)	Algal cover (%)	Pterois spp. biomass (g m <sup>-2</sup> )	Epinephelid biomass (g m <sup>-2</sup> )	Prey density (m <sup>-2</sup> )
	1.45	0.85		1.32	4.7	35.4	0.0	0.0	2.2
2	1.76	1.20	22.7	1.37	9.1	59.4		14.2	8.0
8	2.19	0.64		1.25	0.6	8.7		12.6	1.3
4	3.50	0.75		4.	11.6	45.5		25.3	0.3
ν.	5.22	0.77		1.28	14.6	24.5		0.0	4.6
9	8.49	0.83		1.34	34.6	22.6		29.3	0.7
Mean ± S.D. 3.48	$3.48 \pm 2.13$	$0.84 \pm 0.19$	CI	$1.33 \pm 0.07$	$13.9 \pm 10.6$	$32.7 \pm 18.1$	6.1	$13.6 \pm 12.3$	$1.7 \pm 1.6$
Regression $\nu$ . distance	1	$-0.024^{a}$		$<0.001^{a}$	0.347**	$-0.106^{a}$		$58.460^{a}$	$-0.003^{a}$

a, Not significant (P > 0.05). \*\*P < 0.01.

Furthermore, these characteristics did not vary systematically with increasing distance from shore. The biological characteristics of the reefs were more heterogeneous, with some reefs having much higher biomasses of *Pterois* spp. and native epinephelids and densities of prey species, but the abundance of resident predators and prey fishes were not significantly correlated with distance from shore (Table I). Coral cover was lower than macroalgal cover (13.9 and 32.7% respectively) and the coral assemblage was dominated by *Porites astreoides*, *Porites porites* and *Siderastrea siderea*. Coral cover was the only variable that differed systematically with location and was significantly positively correlated with distance from shore (adjusted  $r^2 = 0.847$ ; Table I).

Multiple regression of current speeds against time since the most recent low or high tide and tidal height, provided good model fits ( $r^2 > 0.60$ ). Current speeds generally decreased with distance offshore and at peak times varied from c. 30 to  $40\,\mathrm{cm\,s^{-1}}$  at patch reefs 1-3 compared with c.  $12\,\mathrm{cm\,s^{-1}}$  at patch reef 6. The models allowed predictions of current speeds at each reef throughout the period of the study (Fig. S4, Supporting Information). The temperature logger at patch reef 6 recorded a variation in daily mean temperatures of  $12.6^{\circ}$  C over the 1 year time span, with a mean daily variation of  $1.4^{\circ}$  C (Fig. S4, Supporting Information). Temperatures were generally higher and increasing during May (spring), compared with the lower and decreasing temperatures in November (winter), which is likely to have also occurred during filming in 2011-2012. Mean midday temperature was not correlated with distance offshore (P > 0.05).

### VISITS BY TRANSIENT PREDATORS

A total of 15 camera deployments (six cameras deployed on each of 15 days) were conducted, resulting in 347.7 h of video footage (details of deployments in Table SI, Supporting Information). These deployments were distributed across 420 1 h segments after sunrise (not every segment included a full hour of footage) and fish counts from 3752 5 min intervals. Video footage from these deployments recorded the presence of 11 species that were categorized as transient predators based on previous categorisations (Beets, 1997), home range sizes (Farmer & Ault, 2011) and the inclusion of fishes in their diets (Randall, 1967; Froese & Pauly, 2010), although they are not all obligate piscivores. There were a total of 10763 fishes within the 5 min counts. The transient predators were yellow jack Carangoides bartholomaei Cuvier 1833, C. ruber, blue runner Caranx crysos (Mitchell 1815), reef shark Carcharhinus perezii (Poey 1876), nurse shark Ginglymostoma cirratum (Bonnaterre 1788), mutton snapper Lutianus analis (Cuvier 1828), lemon shark Negaprion brevirostris (Poey 1868), yellowtail snapper Ocyurus chrysurus (Bloch 1791), cero Scomberomorus regalis (Bloch 1793), barracuda Sphyraena barracuda (Edwards 1771) and houndfish Tylosurus crocodilus (Péron & LeSueur 1821). Six of these species were too rare for detailed analyses (≤16 individuals, <0.005 fish  $5 \,\mathrm{min^{-1}}$  seen in <0.05 of 1 h segments): C. bartholomaei, C. crysos, C. perezii, G. cirratum, N. brevirostris and T. crocodilus. Therefore, the focus of this study was on patch-reef visitation rates by the remaining species: C. ruber (5991 fish, 1.55 fish  $5 \,\mathrm{min}^{-1}$ , seen in 0.43 of 1 h segments), L. analis (2827.00, 0.81, 0.67), O. chrysurus (1793·00, 0·51, 0·32), S. barracuda (59·00, 0·02, 0·08) and S. regalis (58.00, 0.01, 0.05). Only C. ruber, L. analis and O. chrysurus were sufficiently abundant to allow hurdle models of both the probability of each species visiting the patch reefs and the number of fishes seen when they were recorded on the reefs. Obvious

Table II. Model coefficient values for minimal adequate generalized linear mixed-effects models (fixed effects only) for the presence—absence h<sup>-1</sup> of each transient predator species on patch reefs along an offshore gradient. Coefficients for categorical variables are for winter in comparison with spring (season) and for flooding in comparison to ebbing tide (current direction). ×: interaction term

Variable	Caranx ruber	Lutjanus analis	Ocyurus chrysurus	Sphyraena barracuda	Scomberomorus regalis	All five species
Intercept	0.915NS	-1.997*	15.575***	-3.974***	-10.953***	3.386***
Season	0.753**	_	-6.755***	_	2.325**	_
Distance from shore	-1.416***	0.803*	-6.315***	1.543NSa	2.736**	-0.710***
Distance from shore <sup>2</sup>	0.135**	-0.153**	_	-0.276*	-0.240*	_
Hours since sunrise	_	0.186**	_	_	_	0.260***
Current speed (CSPD)	_	_	_	_	-	=
Current direction (CDIR)	_	_	_	_	-1.225*	=
Number of 5 min intervals	0.126**	0.147***	0-182NSa	_	-	=
CSPD×CDIR	-	-	-	-		-

<sup>-,</sup> Not significant, variable not contained in minimal adequate model.

feeding or hunting behaviour was only detected in <0.01% of visits by these transient predators.

Generalized linear mixed-effects models demonstrated that the probability of each transient species, or any of the five focal species, visiting a reef was significantly correlated with one or more of the physical drivers and survey effort (Table II and Figs 2-4). The strongest trend was that the probability of recording each species varied along the offshore gradient, with visitation rates generally decreasing with distance from shore except for S. regalis that was more frequently seen at the furthest reefs [Fig. 4(b)]. For example, not a single O. chrysurus was seen on patch reefs 4-6 [Fig. 3(a)]. Visitation rates also varied seasonally, with C. ruber and S. regalis seen more frequently during the winter and O. chrysurus more frequent in the spring. Time of day only significantly affected sightings of L. analis, which was recorded more often later in the day [Fig. 2(d)]. The metrics of tidal flow were generally not correlated with visits by transient species, but S. regalis was seen more often on ebbing tides. These species-level patterns combined to lead to a decrease in the probability of a visit by any of the five focal species with increasing distance offshore and an increase in the probability of a visit with increasing time since sunrise [Fig. 3(d)].

For the three species that were sufficiently common to also investigate the number of fishes seen during patch-reef visits, linear mixed-effect models suggested that not only were patch reefs visits by *C. ruber* more frequent during the winter and closer to shore, but these variables were significantly positively correlated with the number of fish recorded [Table III and Fig. 2(b)]. In addition, the number of *C. ruber* was positively correlated with current speed. Unlike the probability of a visit, the number of *L. analis* recorded increased during the winter and were higher midway along the offshore gradient [Table III and Fig. 2(e)]. The model for the number of *O. chrysurus* was qualitatively the same as the model for the probability of a visit, with fewer fish seen

NS, Not significant.

<sup>\*</sup>P < 0.05.

<sup>\*\*</sup>P < 0.01.

<sup>\*\*\*</sup>P < 0.001.

NSa, Removal of non-significant terms led to a significant increase in model deviance and Akaike information criterion.

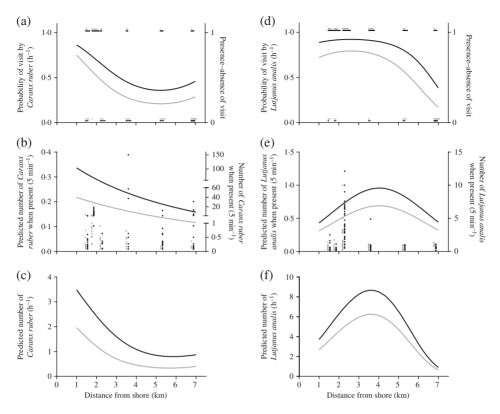


Fig. 2. Scatter plots between the distance of patch reefs from shore and the patch reef visitation rates of (a), (b), (c) *Caranx ruber* and (d), (e), (f) *Lutjanus analis* separated by (a), (d) probability of a visit, (b), (e) number of fish seen during a visit and (c), (f) predicted number of fish seen (combining the probability of a visit and the number of fish seen during the visit). Predictions are (b), (c) at slack tide (current speed = 0 cm s<sup>-1</sup>) and (f) in the morning (time after sunrise = 3 h). Predicted values from statistical models (left-hand axis) are \_\_\_\_ (winter) and \_\_\_ (spring), except for (d) where \_\_\_ (morning) and \_\_\_\_ (afternoon). Observations (RH axis); (a), (b), (e) ● (winter) and ● (spring); (d) ● (P.M.) and ● (A.M.).

further offshore and during the winter [Table III and Fig. 3(b)]. These species-level patterns combined to lead to a decreased abundance (when seen) of the five focal species with increasing distance offshore [Fig. 3(e)]. Furthermore, predictions of the number of transient predators visiting the patch reefs decreases with distance offshore [Fig. 3(f)] with a minimal influence of season or time of day, reflecting the species-level patterns of generally fewer visits further offshore and a mixture of responses to the change of season [Figs 2(c), (f) and 3(c)].

There was a negative relationship between the abundance of transient and resident predators and surveyed prey density on the six patch reefs (intercept coefficient 4.958, P < 0.01; resident coefficient -0.119, P < 0.01; transient coefficient -0.106, P < 0.05; interaction P > 0.05; adjusted  $r^2 = 0.89$ ). The abundance of transient predators was the predicted abundances of the five focal transient species at each patch reef [Fig. 3(f)]. Resident predators were included in the regression using their biomass recorded at each reef during visual surveys. Models including only one of the predator types had no significant correlation with prey density (P > 0.05).

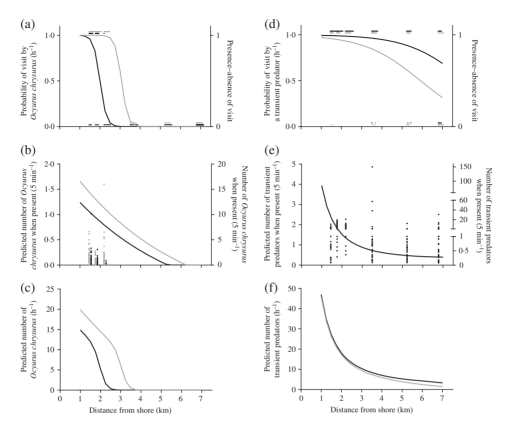


Fig. 3. Scatter plots between the distance of patch reefs from shore and the patch-reef visitation rates of (a)–(c) Ocyurus chrysurus and (d)–(f) any of the five focal transient predators separated by (a), (d) probability of a visit, (b), (e) number of fish seen during a visit and (c), (f) predicted number of fish seen (combining the probability of a visit and the number of fish seen during the visit). Predicted values from statistical models (left-hand axis) are (a), (b), (c) \_\_\_ (winter) and \_\_\_ (spring), and (d), (f) \_\_\_ PM and \_\_\_ AM. Observations (RH axis); (a), (b) ● (winter) and ● (spring); (d) ● (P.M.) and ● (A.M.).

### DISCUSSION

Increasingly accessible technologies, such as acoustic tagging, are demonstrating that wide-ranging carnivorous fish species do not represent a uniform predatory threat across seascapes, but are responding to a range of biophysical drivers (Meyer *et al.*, 2007*a*, *b*; Lédée *et al.*, 2015). The present study contributes to this growing literature by highlighting that the abundances of transient predators visiting patch reefs varies significantly even within a single lagoon, despite the species considered having previously documented home ranges of >4·2 km², migrating among sites >42 km apart and covering distances >12 km in a single day (Farmer & Ault, 2011; O'Toole *et al.*, 2011; Pittman *et al.*, 2014). This study also demonstrates that in addition to significant intra-habitat variability in the abundance of transient predators, there were interspecific differences in the drivers influencing visits to patch reefs. For example, although most species were more abundant close to shore, *S. regalis* was seen more frequently further offshore. Finally, although it should be interpreted with care given the limited number of patch reefs and difficulties of inferring prey mortality rates and influxes without

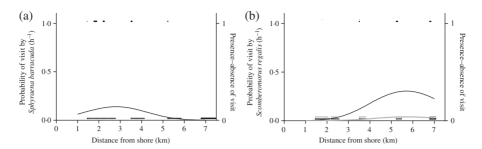


Fig. 4. Scatter plots between the distance of patch reefs from shore and the probability of a visit by (a) *Sphyraena barracuda* and (b) *Scomberomorus regalis*; predictions for *S. regalis* are on an ebbing tide. Predicted values from statistical models (left-hand axis); (b) \_\_\_\_ (winter) and \_\_\_\_ (spring). Observations on RH axis.

any temporal replication, the negative correlation between the abundance of transient predators and prey densities suggests that the spatial variability of transient species influences the demographics of small patch-reef fishes. Furthermore, the effect on prey species of varying abundances of transient species appeared only to be significant when combined with the abundance of resident predators, supporting the hypothesis that there is synergistic predation between these two functional groups (Hixon & Carr, 1997).

The change in visitation rates and abundances on patch reefs along the offshore gradient was apparent in all common transient predators, but this pattern cannot easily be attributed to a single factor because multiple variables co-vary with increasing distance from shore. Similarly, the relative importance of this gradient is likely to vary among

Table III. Coefficients from minimal adequate linear mixed-effects models (fixed effects only) for the number 5 min<sup>-1</sup> (when present) of the three most abundant transient predator species, and all five focal species combined (*i.e.* including *Sphyraena barracuda* and *Scomberomorus regalis*), on patch reefs along an offshore gradient. Coefficients for categorical variables are for winter in comparison with spring (season) and for flooding in comparison to ebbing tide (current direction). ×: interaction term

Variable	Caranx ruber	Lutjanus analis	Ocyurus chrysurus	All five species
Intercept	-0.011***	-1.773***	1.164***	-0·155NS
Season	0.418*	0.328*	-0·171NSa	_
Distance from shore	-0.132**	0.702**	-0.186**	-0.380***
Distance from shore <sup>2</sup>	_	-0.087**	_	0.025*
Hours since sunrise	_	_	_	_
Current speed (CSPD)	0.027**	_	_	_
Current direction (CDIR)	_	_	_	_
CSPD×CDIR	_	_	_	_

<sup>-,</sup> Not significant, variable not contained in minimal adequate model.

NSa, Removal of non-significant term led to a significant increase in model deviance and Akaike information criterion.

NS, Not significant.

<sup>\*</sup>P < 0.05.

<sup>\*\*</sup>*P* < 0.01.

<sup>\*\*\*</sup>P < 0.001

lagoons with different biophysical characteristics. It seems likely, however, that proximity to mangrove creeks is an important cause of this pattern. There are three major mangrove creeks close to the studied patch reefs and at least three of the common transient predators (C. ruber, O. chrysurus and S. barracuda) use these creeks for foraging and refuge (Harborne et al., 2016). Therefore, the complex of creeks and inshore patch reefs may represent a centre of activity for many transient predators. Use of the creeks by these species also varies across tidal cycles (Harborne et al., 2016) and fishes may forage or refuge on the closest patch reefs when the creeks are inaccessible at low tide. Furthermore, four of the species (C. ruber, L. analis, O. chrysurus and S. barracuda) use mangroves and seagrass beds as nursery habitats (Nagelkerken et al., 2000) and their abundances are typically higher on patch reefs close to their settlement habitats (Mumby et al., 2004). Finally, patch reefs closer to shore may receive higher influxes of small prey fishes that also used mangroves nurseries. The increase in visitation rates by S. regalis to patch reefs further from the shore is consistent with the proximity to mangroves being an important driver of the distribution of other transient predators, as this species is not recorded as using mangrove creeks to forage (Harborne et al., 2016) and does not appear to be found in surveys of mangroves (Serafy et al., 2003), as might be expected for a species using this habitat as a nursery.

In addition to being close to the mangrove creeks and other inshore habitats that may be important nurseries, the patch reefs closer to shore are closer to the deeper fore-reefs just west of Eleuthera. Transient species that move widely across seascapes are likely to use both patch reefs and fore-reefs for foraging and may also use the lagoon as a refuge from their own predators. When foraging or refuging on lagoon patch reefs, these fishes may prefer to visit patch reefs closer to the deeper reefs in order to conserve energy. For example, pelagic predators in a temperate bay were four times more abundant on reefs close to an oceanic entrance than on reefs further inside the bay (Ford & Swearer, 2013b). Similarly, the distance to a channel connecting a lagoon and adjacent fore-reefs explained nearly half of the variation in fish assemblages on Belizean patch reefs, with species such as *S. barracuda* being more abundant close to channels (Karnauskas et al., 2012). Furthermore, fishes moving between the deeper reefs and patch reefs closer to shore can utilize the higher speed tidal currents to swim efficiently between these habitats, as observed in other large transient predators (Meyer et al., 2007b) and many shark species (Schlaff et al., 2014).

Changes in current speed across tidal cycles were not a significant variable in the models of fish visitation rates, although groups of *C. ruber* were larger when visiting reefs at higher current speeds. High abundances of *C. crysos* were also strongly associated with areas of high water flow in Brazil (Floeter *et al.*, 2007). Other transient fishes, however, may preferentially associate with patch reefs that have higher maximum current speeds, which could contribute to the negative relationship between fish visitation rates and distance offshore since maximum current speed decreases with increasing distance from shore. In addition to facilitating movement to deeper reefs, the higher current speeds may provide fusiform predators with an advantage over prey that are less well adapted to swimming in rapidly moving water and are thus less able to escape. Such asymmetries in predator–prey interactions are rarely studied, but there is some evidence that smaller fishes with less swimming ability are more susceptible to predators where water currents are strongest (Holmes & McCormick, 2006). In contrast, a preference for more moderate maximum current speeds may lead to the higher abundance of *L. analis* towards the middle of the offshore gradient.

In addition to close proximity to mangroves and deep water and higher maximum current speeds, the mosaic of habitats surrounding each patch reef varied and the habitats surrounding reefs close to shore may offer better foraging than those further offshore. While prey fishes are concentrated on the patch reefs, some juveniles settle and remain in soft-bottom habitats, particularly seagrass (Mumby *et al.*, 2004) and may offer important food resources to transient predators. The distribution of hard-bottom habitats surrounding patch reefs has also been demonstrated to affect the abundance of lutjanids (Karnauskas *et al.*, 2012). Finally, reefs closer to the shore in Eleuthera may be visited more frequently because of factors such as their size distribution and proximity to neighbouring patches. For example, *O. chrysurus* does visit aggregated artificial reefs (5 m apart) more frequently than isolated reefs 50 m apart (Overholtzer-McLeod, 2006) and the abundance of *O. chrysurus* and *C. ruber* was significantly influenced by the proximity to other patch reefs in a Belizean lagoon (Karnauskas *et al.*, 2012).

Most other characteristics of the patch reefs did not change systematically with increasing distance from shore and seem unlikely to be significant influences on transient predator visitation rates. The only variable that did change systematically along the offshore gradient was coral cover. Coral cover increased, however, with increasing distance offshore, while transient predator visitation rates decreased. Fishes typically become less abundant on reefs with lower coral cover (Pratchett *et al.*, 2008), but transient predators may hunt more frequently on low coral-cover reefs where prey have fewer refuges. Because the reefs in this study are typically characterized by encrusting corals, however, increasing coral cover did not cause systematic differences in rugosity, which is typically a more important control of the abundance of small prey fishes (Gratwicke & Speight, 2005).

In addition to changing their visitation rates to patch reefs along the offshore gradient, the probability of a visit by *C. ruber*, *O. chrysurus* and *S. regalis* and the group size of *L. analis*, varied significantly among seasons. All species except *O. chrysurus* were more abundant around the patch reefs during the winter and this may be in response to fishes seeking optimal temperatures. Temperatures on deeper reefs are typically more stable, in contrast to shallower water where there is a greater variability and more extreme highs and lows (Potts & Swart, 1984). As this study considers diurnal activity, fishes may be seeking energetically beneficial warmer water in the shallow lagoon during the winter days. Seasonal variations in transient predator visitation rates may also be caused by fishes migrating to deeper reefs to spawn, or preparing to undertake spawning migrations. Despite the significant decrease in spring visits to patch reefs by transient species, further work is necessary to determine how predator abundance and prey mortality rates are coupled. Although predators may be more abundant during the winter, species such as *S. barracuda* may feed less intensely at this time of year because of lower water temperatures (Hammerschlag *et al.*, 2010).

Lutjanus analis was the only species that displayed a significant diurnal pattern and visited the patch reefs more frequently later during the day. Sharks feed optimally at dusk because they have a sensory advantage in low-light conditions and their body temperatures are higher than that of their prey (Papastamatiou et al., 2015) and L. analis may have a similar metabolic benefit. Lutjanus analis may also be returning to the patch reefs later in the day in order to seek nocturnal shelter. Relatively limited video observations of visits to patch reefs elsewhere in The Bahamas have previously suggested that C. ruber and S. barracuda may be seen more frequently towards the middle of the day compared with at dawn and dusk (Carr & Hixon, 1995) and acoustic

data from Eleuthera indicated that *S. barracuda* moves from lagoon habitats to deeper fore-reefs from mid-morning to mid-afternoon before returning in the late afternoon (O'Toole *et al.*, 2010). These patterns were not apparent in the present study and may suggest significant variability within species, across seasons and among locations. In addition to changing their patch-reef visitation rates, transient predators may also alter their feeding rates throughout the day. For example, *L. analis* appears to feed less frequently at midday compared with during mornings or evenings (Mueller *et al.*, 1994). Combining the spatial and temporal movements of transient predators with their foraging behaviour and feeding rates represents an important next step in elucidating the functional role of these species within Caribbean seascapes.

This study has provided new insights into the abundance of five transient predators, but the use of video cameras limited observations at night when their distributions may be different. For example, there is some evidence from artificial-reef studies that L. analis may disperse away from habitat structures at night (Eggleston et al., 1990). Indeed many transient predators may make diel migrations across the seascape, as documented for Pacific carangids and lutjanids (Meyer et al., 2007a, b). There are potential solutions to the problem of filming at night (Holmes et al., 2012), but acoustic tracking may be better for quantifying nocturnal movement. Data are also required from different life stages of each species, as some transient predators change their foraging behaviour ontogenetically (Mueller et al., 1994), which may affect their behavioural patterns as it does in sharks (Schlaff et al., 2014). Furthermore, the importance of the offshore gradient for influencing transient predator distributions suggests there is a need for additional research at locations where the importance of individual factors, such as mangrove proximity and distance to deeper reefs, can be examined independently to test the hypotheses suggested by this study. Finally, studies are required to elucidate where the transient predators swim to when not in lagoons in order to fully understand their movements and elucidate their role in seascape-scale connectivity among habitats (McCauley et al., 2012).

The focus of this study was on providing new insights into the variability of visits by transient predators to patch reefs, but these data also provide indications of the potential effects of environmental change. Firstly, temperature is likely to be at least partly causing the seasonal variation seen in some of the species and warmer sea-surface temperatures under global climate change scenarios may affect the movement of transient predators (Currey et al., 2015). The consequences on fish abundances of clearing mangrove nursery and foraging habitats have frequently been stated (Valiela et al., 2001) and the present study also suggests that changes to the distribution of transient predators may occur if inshore habitat connectivity is affected by coastal development. The functional role of transient predators will also be affected by fishing [e.g. O. chrysurus is increasingly being targeted in Belize, Mumby et al. (2012)]. In addition, the removal of apex predators might increase the abundance of smaller transient predators and also affect their behaviour if the threat of predation decreases (Preisser et al., 2005). Predicting how all these changes may cascade spatially and temporally through tropical food webs is challenging, but it is clear that a better understanding of the behaviour of transient predators is important to conserve and manage the ecosystem services provided by coral reefs.

This paper was made possible by funding to A.R.H. from the U.K. Natural Environment Research Council (fellowship NE/F015704/1), the Australian Research Council (fellowship

DE120102459) and the Earthwatch Institute. We are grateful to the staff of the Cape Eleuthera Institute for their help during fieldwork and L. Pointon for help with video analysis. This is contribution 16 of the Marine Education and Research Center in the Institute for Water and Environment at Florida International University.

# **Supporting Information**

Supporting Information may be found in the online version of this paper:

- Fig. S1. Video stills of each of the patch reefs used in the study.
- Fig. S2. Size distribution of 195 patch reefs measured within the study area.
- Fig. S3. Schematic overview of the process of deriving mean abundance and presence or absence of each species for a single 1 h time segment at a study patch reef.
- Fig. S4. Example of predicted current speeds at each patch reef and annual changes in mean daily temperature from March 2014 to March 2015.
  - TABLE SI. Details of the camera deployments on each day of the study.

# References

- Beets, J. (1997). Effects of a predatory fish on the recruitment and abundance of Caribbean coral reef fishes. *Marine Ecology Progress Series* **148**, 11–21.
- Burnham, K. P. & Anderson, D. R. (2004). Multimodel inference understanding AIC and BIC in model selection. *Sociological Methods & Research* **33**, 261–304.
- Carr, M. H. & Hixon, M. A. (1995). Predation effects on early post-settlement survivorship of coral-reef fishes. *Marine Ecology Progress Series* **124**, 31–42.
- Crawley, M. J. (2007). The R Book. Chichester: John Wiley & Sons Ltd..
- Currey, L. M., Heupel, M. R., Simpfendorfer, C. A. & Williams, A. J. (2015). Assessing environmental correlates of fish movement on a coral reef. *Coral Reefs* **34**, 1267–1277.
- Eggleston, D. B., Lipcius, R. N., Miller, D. L. & Cobacetina, L. (1990). Shelter scaling regulates survival of juvenile Caribbean spiny lobster *Panulirus argus*. *Marine Ecology Progress Series* **62**, 79–88.
- Farmer, N. A. & Ault, J. S. (2011). Grouper and snapper movements and habitat use in Dry Tortugas, Florida. *Marine Ecology Progress Series* **433**, 169–184.
- Fletcher, D., MacKenzie, D. & Villouta, E. (2005). Modelling skewed data with many zeros: a simple approach combining ordinary and logistic regression. *Environmental and Ecological Statistics* **12**, 45–54.
- Floeter, S. R., Krohling, W., Gasparini, J. L., Ferreira, C. E. L. & Zalmon, I. R. (2007). Reef fish community structure on coastal islands of the southeastern Brazil: the influence of exposure and benthic cover. *Environmental Biology of Fishes* **78**, 147–160.
- Ford, J. R. & Swearer, S. E. (2013a). Shoaling behaviour enhances risk of predation from multiple predator guilds in a marine fish. *Oecologia* **172**, 387–397.
- Ford, J. R. & Swearer, S. E. (2013b). Two's company, three's a crowd: food and shelter limitation outweigh the benefits of group living in a shoaling fish. *Ecology* **94**, 1069–1077.
- Fulton, C. J., Bellwood, D. R. & Wainwright, P. C. (2005). Wave energy and swimming performance shape coral reef fish assemblages. *Proceedings of the Royal Society B* 272, 827–832.
- Gratwicke, B. & Speight, M. R. (2005). The relationship between fish species richness, abundance and habitat complexity in a range of shallow tropical marine habitats. *Journal of Fish Biology* **66**, 650–667.
- Hammerschlag, N., Ovando, D. & Serafy, J. E. (2010). Seasonal diet and feeding habits of juvenile fishes foraging along a subtropical marine ecotone. *Aquatic Biology* **9**, 279–290.
- Harborne, A. R., Talwar, B. & Brooks, E. J. (2016). The conservation implications of spatial and temporal variability in the diurnal use of Bahamian tidal mangrove creeks by transient predatory fishes. *Aquatic Conservation: Marine and Freshwater Ecosystems* **26**, 202–211.

- Hixon, M. A. (1991). Predation as a process structuring coral reef fish communities. In *The Ecology of Fishes on Coral Reefs* (Sale, P. F., ed), pp. 475–508. London: Academic Press.
- Hixon, M. A. & Beets, J. P. (1993). Predation, prey refuges and the structure of coral-reef fish assemblages. *Ecological Monographs* **63**, 77–101.
- Hixon, M. A. & Carr, M. H. (1997). Synergistic predation, density dependence and population regulation in marine fish. *Science* **277**, 946–949.
- Hoey, A. S. & McCormick, M. I. (2004). Selective predation for low body condition at the larval-juvenile transition of a coral reef fish. *Oecologia* **139**, 23–29.
- Holmes, T. H. & McCormick, M. I. (2006). Location influences size-selective predation on newly settled reef fish. *Marine Ecology Progress Series* **317**, 203–209.
- Holmes, T. H., Wilson, S. K., Vanderklift, M., Babcock, R. & Fraser, M. (2012). The role of *Thalassoma lunare* as a predator of juvenile fish on a sub-tropical coral reef. *Coral Reefs* **31**, 1113–1123.
- Karnauskas, M., Chérubin, L. M., Huntington, B. E., Babcock, E. A. & Thoney, D. A. (2012). Physical forces influence the trophic structure of reef fish communities on a remote atoll. *Limnology and Oceanography* 57, 1403–1414.
- Lédée, E. J., Heupel, M. R., Tobin, A. J. & Simpfendorfer, C. A. (2015). Movements and space use of giant trevally in coral reef habitats and the importance of environmental drivers. *Animal Biotelemetry* 3, 1–14.
- Luckhurst, B. E. & Luckhurst, K. (1978). Analysis of the influence of substrate variables on coral reef fish communities. *Marine Biology* **49**, 317–323.
- Matley, J. K., Heupel, M. R. & Simpfendorfer, C. A. (2015). Depth and space use of leopard coralgrouper *Plectropomus leopardus* using passive acoustic tracking. *Marine Ecology Progress Series* **521**, 201–216.
- McCauley, D. J., Young, H. S., Dunbar, R. B., Estes, J. A., Semmens, B. X. & Micheli, F. (2012). Assessing the effects of large mobile predators on ecosystem connectivity. *Ecological Applications* **22**, 1711–1717.
- Meyer, C. G., Holland, K. N. & Papastamatiou, Y. P. (2007a). Seasonal and diel movements of giant trevally *Caranx ignobilis* at remote Hawaiian atolls: implications for the design of marine protected areas. *Marine Ecology Progress Series* 333, 13–25.
- Meyer, C. G., Papastamatiou, Y. P. & Holland, K. N. (2007b). Seasonal, diel and tidal movements of green jobfish (*Aprion virescens*, Lutjanidae) at remote Hawaiian atolls: implications for marine protected area design. *Marine Biology* **151**, 2133–2143.
- Mueller, K. W., Dennis, G. D., Eggleston, D. B. & Wicklund, R. I. (1994). Size-specific social interactions and foraging styles in a shallow water population of mutton snapper, *Lutjanus analis* (Pisces, Lutjanidae), in the central Bahamas. *Environmental Biology of Fishes* **40**, 175–188.
- Mumby, P. J., Edwards, A. J., Arias-González, J. E., Lindeman, K. C., Blackwell, P. G., Gall, A., Gorczynska, M. I., Harborne, A. R., Pescod, C. L., Renken, H., Wabnitz, C. C. C. & Llewellyn, G. (2004). Mangroves enhance the biomass of coral reef fish communities in the Caribbean. *Nature* 427, 533–536.
- Mumby, P. J., Steneck, R. S., Edwards, A. J., Ferrari, R., Coleman, R., Harborne, A. R. & Gibson, J. P. (2012). Fishing down a Caribbean food web relaxes trophic cascades. *Marine Ecology Progress Series* **445**, 13–24.
- Murchie, K. J., Schwager, E., Cooke, S. J., Danylchuk, A. J., Danylchuk, S. E., Goldberg, T. L., Suski, C. D. & Philipp, D. P. (2010). Spatial ecology of juvenile lemon sharks (*Negaprion brevirostris*) in tidal creeks and coastal waters of Eleuthera, The Bahamas. *Environmental Biology of Fishes* **89**, 95–104.
- Nagelkerken, I., Dorenbosch, M., Verberk, W. C. E. P., Cocheret de la Morinière, E. & van der Velde, G. (2000). Importance of shallow-water biotopes of a Caribbean bay for juvenile coral reef fishes: patterns in biotope association, community structure and spatial distribution. *Marine Ecology Progress Series* 202, 175–192.
- O'Toole, A. C., Murchie, K. J., Pullen, C., Hanson, K. C., Suski, C. D., Danylchuk, A. J. & Cooke, S. J. (2010). Locomotory activity and depth distribution of adult great barracuda (*Sphyraena barracuda*) in Bahamian coastal habitats determined using acceleration and pressure biotelemetry transmitters. *Marine and Freshwater Research* **61**, 1446–1456.
- O'Toole, A. C., Danylchuk, A. J., Goldberg, T. L., Suski, C. D., Philipp, D. P., Brooks, E. & Cooke, S. J. (2011). Spatial ecology and residency patterns of adult great barracuda

- (Sphyraena barracuda) in coastal waters of The Bahamas. Marine Biology 158, 2227-2237.
- Overholtzer-McLeod, K. L. (2004). Variance in reef spatial structure masks density dependence in coral-reef fish populations on natural versus artificial reefs. *Marine Ecology Progress Series* **276**, 269–280.
- Overholtzer-McLeod, K. L. (2006). Consequences of patch reef spacing for density-dependent mortality of coral-reef fishes. *Ecology* **87**, 1017–1026.
- Papastamatiou, Y. P., Watanabe, Y. Y., Bradley, D., Dee, L. E., Weng, K., Lowe, C. G. & Caselle, J. E. (2015). Drivers of daily routines in an ectothermic marine predator: hunt warm, rest warmer? *PLoS ONE* **10**, e0127807. doi: 10.1371/journal.pone.0127807
- Pittman, S. J., Monaco, M. E., Friedlander, A. M., Legare, B., Nemeth, R. S., Kendall, M. S., Poti, M., Clark, R. D., Wedding, L. M. & Caldow, C. (2014). Fish with chips: tracking reef fish movements to evaluate size and connectivity of Caribbean marine protected areas. *PLoS ONE* **9**, e96028. doi: 10.1371/journal.pone.0096028
- Potts, D. C. & Swart, P. K. (1984). Water temperature as an indicator of environmental variability on a coral reef. *Limnology and Oceanography* **29**, 504–516.
- Pratchett, M. S., Munday, P. L., Wilson, S. K., Graham, N. A. J., Cinner, J. E., Bellwood, D. R., Jones, G. P., Polunin, N. V. C. & McClanahan, T. R. (2008). Effects of climate-induced coral bleaching on coral-reef fishes ecological and economic consequences. *Oceanography and Marine Biology: An Annual Review* **46**, 251–296.
- Preisser, E. L., Bolnick, D. I. & Benard, M. F. (2005). Scared to death? The effects of intimidation and consumption in predator–prey interactions. *Ecology* **86**, 501–509.
- Randall, J. E. (1967). Food habitats of reef fishes of the West Indies. *Studies in Tropical Oceanography* **5**, 665–847.
- Russ, G. R. (2002). Yet another review of marine reserves as reef fishery management tools. In *Coral Reef Fishes: Dynamics and Diversity in a Complex Ecosystem* (Sale, P. F., ed), pp. 421–443. San Diego, CA: Academic Press.
- Schlaff, A. M., Heupel, M. R. & Simpfendorfer, C. A. (2014). Influence of environmental factors on shark and ray movement, behaviour and habitat use: a review. *Reviews in Fish Biology and Fisheries* **24**, 1089–1103.
- Serafy, J. E., Faunce, C. H. & Lorenz, J. J. (2003). Mangrove shoreline fishes of Biscayne Bay, Florida. *Bulletin of Marine Science* **72**, 161–180.
- Stallings, C. D. (2008). Indirect effects of an exploited predator on recruitment of coral-reef fishes. *Ecology* **89**, 2090–2095.
- Stallings, C. D. (2009). Fishery-independent data reveal negative effect of human population density on Caribbean predatory fish communities. *PLoS ONE* **4**, e5333. doi: 10.1371/journal.pone.0005333
- Valiela, I., Bowen, J. L. & York, J. K. (2001). Mangrove forests: one of the world's threatened major tropical environments. *Bioscience* **51**, 807–815.
- Warton, D. I. & Hui, F. K. C. (2011). The arc sine is asinine: the analysis of proportions in ecology. *Ecology* **92**, 3–10.
- White, J. W., Samhouri, J. F., Stier, A. C., Wormald, C. L., Hamilton, S. L. & Sandin, S. A. (2010). Synthesizing mechanisms of density dependence in reef fishes: behavior, habitat configuration and observational scale. *Ecology* **91**, 1949–1961.
- Zuur, A. F., Ieno, E. N., Walker, N. J., Saveliev, A. A. & Smith, G. M. (2009). *Mixed Effects Models and Extensions in Ecology with R*. New York, NY: Springer.

### **Electronic References**

- Bates, D., Maechler, M., Bolker, B. & Walker, S. (2014). lme4: Linear mixed-effects models using Eigen and S4. R package version 1.1-7. Available at http://CRAN.R-project.org/package=lme4/ (accessed September 2015).
- Froese, R. & Pauly, D. (2010). FishBase. Available at http://www.fishbase.org/ (accessed September 2015).