

Intrapopulation variation in habitat use by two abundant coastal fish species

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ABSTRACT: Decline of marine fisheries has become one of the most severe global environmental crises. In typical fishery management efforts, fish populations are often treated as homogeneous units, thereby tacitly ignoring potential intrapopulation variation within taxonomic groupings. We used acoustic telemetry and stable isotope analysis to examine movement patterns of 20 gray snapper *Lutjanus griseus* and 20 schoolmaster snapper *L. apodus* in a Bahamian tidal creek and wetland. In particular, we examined (1) if intrapopulation variation in fish habitat use and movement patterns existed, (2) whether that variation was a function of body size, and (3) if there was evidence of specialization in habitat use among individuals. We found that movement varied substantially among individuals, but was independent of body size. Some individuals exhibited frequent, repeated, movements to certain areas of the study site. $\delta^{13}\text{C}$ values of individual snapper were significantly related to movement metrics, suggesting that movement differences were related to specific patterns of foraging behavior. Our findings suggest the importance of incorporating intrapopulation niche variation—a source of variation that is often overlooked in traditional conservation and management strategies—into the study of coastal fish populations.

KEY WORDS: Behavioral ecology · Body size · Fisheries · Food web · Individual specialization · Movement patterns · Optimal foraging theory · Stable isotopes

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INTRODUCTION

Overexploitation of marine fishes is considered one of the most critical global environmental crises (Jackson et al. 2001, Worm et al. 2006). Stocks of economically and ecologically important species have been drastically reduced through commercial and recreational fisheries (Coleman et al. 2004, Lotze & Worm 2009). Although large pelagic predators are especially susceptible to declines (Pauly et al. 1998, Heithaus et al. 2008), populations of coastal mesopredators such as groupers (Serranidae) and snappers (Lutjanidae) are also threatened by overexploitation (Willis et al. 2001, Nieland et al. 2007, Graham et al. 2008). For example, populations of the Nassau grouper *Epinephelus striatus* have declined by up to 60% over the last 3 decades and the species has been added to the IUCN Red List (Baillie & Groombridge 1996, Sadovy 1997).

In order to provide a template for developing conservation and management strategies, scientists usually

seek to identify broad generalities that define the 'typical' individual of a particular population. Individuals are assumed to possess generally the same behavioral or feeding traits during certain life stages or size classes, and a population is thus treated as a homogeneous unit from a management perspective. One well-known example are juvenile grunts (Haemulidae) in Teague Bay, US Virgin Islands, which rest over coral heads during the day and move to seagrass beds as a group at night (Meyer et al. 1983). Similarly, gray snapper *Lutjanus griseus* in Biscayne Bay, Florida have been observed to move in groups from mangroves during the day to seagrass beds at night (Luo et al. 2009). With respect to ontogeny, individuals of many reef fishes are assumed to go through predictable sequences of habitat utilization (Nagelkerken et al. 2000). For instance, larval French grunts *Haemulon flavolineatum* in Piscadera Bay, Curaçao, settle in sand/rubble habitat at the mouth of bays, move further into the bay to utilize mangroves and seagrass beds as

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juveniles, and then shift to coral reefs as adults (Huijbers et al. 2008).

Despite the common perspective that individuals of a population are homogeneous units, substantial intrapopulation variation in behavior has been shown to occur across diverse taxonomic groups and may have important evolutionary, ecological and conservation implications (Van Valen 1965, Roughgarden 1972, 1974, Schindler et al. 1997, Bolnick et al. 2003). From a management perspective, focusing only on the core habitats of a population may ignore those individuals that utilize alternative habitats or food sources. This can be especially problematic when intrapopulation variation is a function of body size or age, since a demographically important subset of the population may be neglected (Durell 2000, Bolnick et al. 2003). Recognizing intrapopulation variation in fishery management plans may facilitate preserving species' ecological, phenotypic and genetic diversity (Moritz 1994, Coates 2000, Smith et al. 2001, Bolnick et al. 2003).

The aim of this study was to examine potential variation in habitat use and movement patterns among individuals of 2 ecologically and economically important nearshore snapper populations. Specifically, we investigated (1) if variation in habitat use and movement patterns occurred among individuals of gray snapper *Lutjanus griseus* and/or schoolmaster snapper *L. apodus*, (2) whether that variation was a function of body size (a well-acknowledged mechanism of intrapopulation variation), and (3) if there was evidence of specialization in habitat use for either focal population. We used acoustic telemetry to directly explore the movements of fishes at an individual level, as well as stable isotope ratios to provide indirect information as to potential feeding patterns. We endeavored to provide a framework for future research that acknowledges variation in behavioral attributes and foraging within marine populations.

MATERIALS AND METHODS

Study system and species. Broad Creek (26° 29' 35" N, 77° 02' 34" W) is an estuarine tidal creek located on the east side of Abaco Island, The Bahamas (Fig. 1).

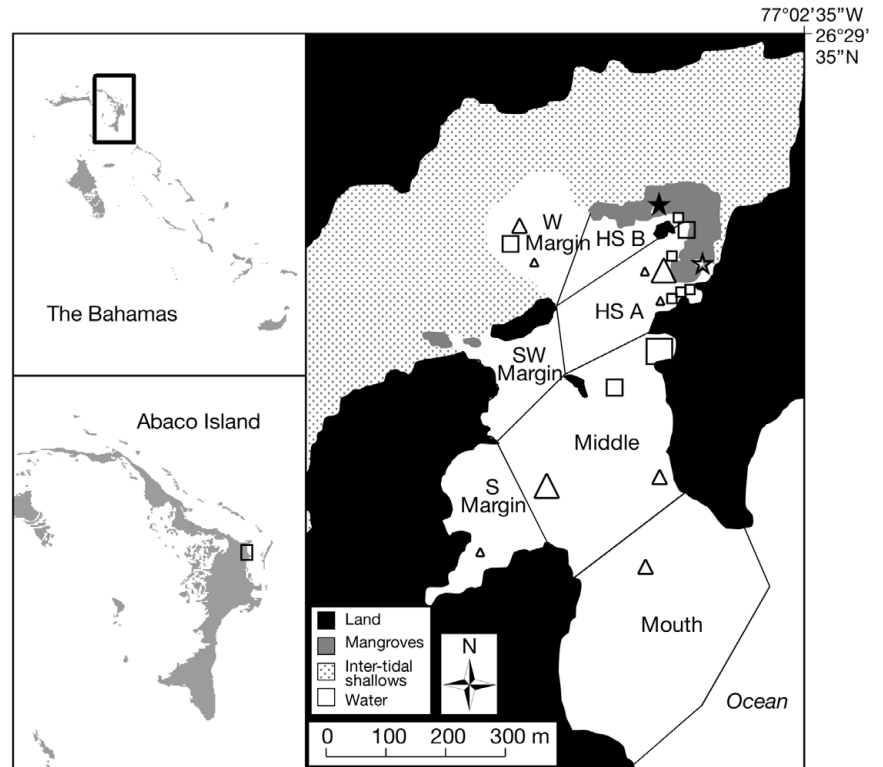


Fig. 1. Broad Creek system coded by land/seascape type. (Δ, □) Furthest distance away from home sites travelled by individual gray snapper *Lutjanus griseus* and schoolmaster snapper *L. apodus*, respectively, during the study period. Size of triangles and squares reflects the size of the fish (small: 110–150 mm standard length, SL; medium: 151–200 mm SL; big: >200 mm SL. (☆, ★) Home sites A and B (HS A and HS B). Panels on the left are maps of The Bahamas (top left) and Abaco Island (bottom left), with rectangle representing the location of Broad Creek

Tidal creeks in The Bahamas range in size from several hectares with maximum low tide depths of 1 m, to thousands of hectares with maximum depths >10 m (Layman et al. 2007, Rypel et al. 2007, Valentine-Rose et al. 2007b); Broad Creek falls at the lower end of this range. The system has a semi-diurnal tidal regime and a mean daily tidal amplitude of ~0.8 m. There is little freshwater input to this system; thus, it is marine-dominated throughout its extent (Valentine-Rose et al. 2007a).

Broad Creek consists of extensive, shallow, intertidal flats. These flats are primarily composed of a silt substrate with interspersed mangroves (mainly *Rhizophora mangle*). The most prevalent benthic habitat types are seagrass beds (primarily *Thalassia testudinum*), submerged mangrove prop roots (mostly *R. mangle*), hard bottom with soft corals (mostly *Gorgonia* spp.) and sand. At low tide, water depths ~>0.3 m are only found in 2 pools (~10 m in diameter) in the northwest corner of Broad Creek (Fig. 1). The tidal pools are depressions in the karst substrate that stay inundated

even at the lowest tides, thereby providing habitat for fishes at all tidal phases (Rypel et al. 2007, Valentine-Rose et al. 2007b). The majority of the creek area (>99%) is <0.3 m at low tide, and is generally too shallow for adult and subadult snapper to traverse (Rypel et al. 2007).

Gray snapper *Lutjanus griseus* (Linnaeus 1758) and schoolmaster snapper *Lutjanus apodus* (Walbaum 1972) are abundant generalist fish predators that coexist in many nearshore ecosystems in the tropics and subtropics of the Atlantic (Starck & Schroeder 1971, Allen 1985). In Broad Creek, as is common in shallow creeks of The Bahamas, these fishes typically inhabit deeper pools or deep mangrove-lined shorelines, herein referred to as 'home sites' (e.g. HS A and HS B in Fig. 1). Individual fish may leave these home sites with rising water during flood tides, likely to feed, and typically return to the home sites as water levels fall with ebbing tide (Rypel et al. 2007). These repeated daily movements to and from the home sites in Broad Creek provide a unique opportunity to quantitatively evaluate individual level variation in local scale movement patterns.

Tagging and tracking. Twenty gray and 20 schoolmaster snapper were caught at home sites A and B at low tide between 20–28 April 2009 using baited hooks and fish traps. Sizes were representative of those fishes that are found in these systems (Layman et al. 2007, Luo et al. 2009). Small acoustic transmitters (V7, Vemco; 19 × 7 mm, 1.6 g in air, 77 d battery life), which pinged every 15 to 45 s, were surgically implanted into each fish. Each transmitter had a unique code to identify individual fish. Individual fish were anesthetized in a mixture of fresh seawater and clove oil (active ingredient: eugenol), which is a commonly used fish anesthetic (Sladky et al. 2001, Hiscock et al. 2002, Parsons et al. 2003, Cotter & Rodruck 2006), and measured for standard length (SL). The transmitter was inserted into the body cavity through a 10 mm incision that was made between the pelvic and anal fins following a procedure similar to Nowak & Quinn (2002). The incision was closed with 2 stitches using a C-curved needle with attached suture. A small sample of the dorsal fin tissue (~1 cm²) was removed from each fish for stable isotope analysis (see below) before the fish was transferred to a cooler with fresh seawater for recovery. After ~1 h, the fish were released at the capture location. Fin clips were transported on ice to a field station, stored in a freezer and later processed at Florida International University.

To quantify presence/absence patterns at the 2 home sites, 1 stationary omnidirectional hydrophone (VR2, Vemco) was placed at each location. The receivers were secured to cinder blocks and placed on the substrate in the middle of both home sites at ~1.5 m low

tide depth. Since both home sites were surrounded by dense mangroves, detection ranges did not extend beyond the edges of the pools (C. Hammerschlag-Peyer unpubl. data).

To estimate movement outside the home sites, tagged fish were located from an inflatable Zodiac boat using a hand-held receiver and hydrophone (VR100, Vemco). The hydrophone was situated in the water column about halfway between the substrate and the water surface to minimize acoustic disturbances. Once a fish was detected, the coordinates of its position and the time of detection were recorded by the manual receiver. Mobile tracking was conducted every day at high tide for an entire lunar cycle, from 26 April to 24 May 2009. Because most parts of the creek are largely intertidal (and thus very shallow at low tide, with no fish movement), manual tracking was conducted 2 h before until 2 h after high tide. Tracking paths were chosen haphazardly during each session.

Data analysis. Continuous data from stationary receivers was categorized into 'time in' and 'time out' of the home site for each individual fish. If an individual was not detected by a stationary receiver for >60 min, the fish was considered to have left the tidal pool. For each fish, the home site, date, time in and time out were identified. Tracking with the manual receiver was used to identify the location of fish outside the home sites during each daily high tide. If >1 location fix (outside of the home sites) was obtained for an individual per tidal cycle, we used the fix with the highest detection strength (dB) as an estimate of its most precise location during that given detection period. This procedure eliminated the problem of autocorrelation of successive detections (Van den Avyle & Evans 1990, Sackett et al. 2007).

To determine mean distance moved, maximum distance moved and furthest distance away from the home site for each individual, the stationary and mobile tracking data were combined. Euclidean distance between 2 successive locations (typically between a home site and the strongest detection from the manual receiver) was estimated using ArcMap GIS version 9.3.1 (ESRI 2008). The mean distance moved by an individual was calculated by dividing its total distance moved during the study period by the total number of tidal cycles in which a movement of that individual occurred (e.g. Roth & Greene 2006, Carfagno & Weatherhead 2008). The maximum distance moved by an individual was the longest distance it moved during a single tidal cycle. This measure usually included a movement from one of the tidal pools to some location in the tidal creek and then back to a tidal pool during the same tidal cycle. The furthest distance away from the home site was the greatest distance that a fish was ever detected away from the home site during the study period.

Activity space was defined as the area that a fish utilized during daytime tracking for the duration of the entire study period. It was estimated using the minimum convex polygon (MCP), i.e. the area of the smallest convex polygon that contains all observed positions of an individual fish (Anderson 1982). MCP estimations were obtained using Animal Movements extension (Hooge & Eichenlaub 1997) for ArcView GIS version 3.2 (ESRI 1999). Areas of MCP that extended onto land were clipped using ArcMap Geoprocessing tools. The total activity space of each population was calculated by combining movement data for all 20 individuals of each species.

Fin tissues were analyzed for their $\delta^{13}\text{C}$ values, as $\delta^{13}\text{C}$ reflects long-term diet and may be used to indicate foraging areas (Layman 2007). In Bahamas tidal creek systems, prey $\delta^{13}\text{C}$ values vary predictably from upper reaches of creek systems to creek mouths (authors' unpubl. data). Snapper prey at the creek mouth are relatively enriched in $\delta^{13}\text{C}$ relative to prey that are typically found in the upper reaches of creeks (see Fig. 4); thus, $\delta^{13}\text{C}$ values of snapper fin tissue may reflect long-term feeding in particular parts of the creek system. Analysis was based on Post et al. (2007) and was conducted at the Yale Earth System Center for Stable Isotopic Studies (ESCSIS). All stable isotope values are reported in the δ notation, where

$$\delta^{13}\text{C} = [(R_{\text{sample}}/R_{\text{standard}}) - 1] \times 1000,$$

and R is the $^{13}\text{C}/^{12}\text{C}$ ratio.

Statistical analysis. For some individuals, we had a small number of observations likely because a fish might have died or left the study area. Linear regression revealed that all of our response measures were not significantly related to sample size when individuals with <9 observation points were excluded from each analysis ($p > 0.05$), so each of these individuals was excluded. Additionally, it is also important to note that the number of individuals for a given analysis sometimes differed because of the way the individual observation points were used. For example, while 2 data points of each fish were used in the analysis of activity space and individual specialization, the same 2 points resulted in only 1 distance estimate for that fish. In sum, 14 gray and 15 schoolmaster snapper were used for the activity space and individual specialization analysis (see below), and 9 gray and 9 schoolmaster snapper were used for distance analysis.

Movement distances and activity space were examined as a function of body size using linear regression in SigmaPlot 10.0. When necessary to meet model assumptions, data were $\ln(x+0.5)$ transformed. Frequency histograms of movement distances and activity space were performed in R version 2.9.2. We examined

a potential relationship between activity space and $\delta^{13}\text{C}$ using linear regression.

Individual specialization. Recent studies have shown that intrapopulation variation in resource use can exist at a single ontogenetic life history stage (Bolnick et al. 2003). Individual specialization, which is defined as a significantly smaller (dietary) niche of an individual than the niche of its population for reasons not attributable to its sex, body size, or discrete morphology, has been identified in a number of diverse taxa (Ringler 1983, Magurran 1986, Schindler et al. 1997, Bolnick et al. 2003). One way to infer the occurrence of individual specialists in a population is by examining components of niche variation following Roughgarden (1972, 1974, 1979). In this approach, the population's total niche width (TNW) in terms of resource use is subdivided into a within-individual component (WIC) and a between-individual component (BIC). Individual specialists occur in a population when the TNW consists mostly of the BIC, such that WIC/TNW is small (Roughgarden 1972, 1974, 1979). Most studies on individual specialization have focused on differences in diet and morphology (Bolnick et al. 2003, Araujo et al. 2007, Svanbäck & Bolnick 2007, Svanbäck et al. 2008, Quevedo et al. 2009). Herein, we extend this framework to examine movement data.

In this analysis, the tidal creek was divided into 7 zones (the equivalent of individual diet items in traditional specialization studies) according to their habitat composition (Fig. S1 in the supplement available at www.int-res.com/articles/suppl/m415p211_supp.pdf), water depth, and distance from home sites A and B (Fig. 1). To calculate WIC/TNW, we used the number of detections of an individual fish in each zone during the entire study period. Individual specialization is likely to be present if the WIC/TNW value differs significantly from a null expectation. To this end, the movement-based WIC/TNW value for each population was tested against a null-model using a nonparametric Bootstrap Monte Carlo simulation (Text S1 in the supplement). Calculations were performed in IndSpec1 (Bolnick et al. 2002).

RESULTS

The 20 gray and 20 schoolmaster snapper had mean SLs of 168.5 ± 43.4 mm (mean \pm SD; range 111–276 mm; Table S1 in the supplement) and 148.3 ± 33.7 mm (range: 110–272 mm), respectively. Based on otolith readings, the body size range in this study included fish of ages 1 to 7 for gray snapper (Rypel & Layman 2008) and 2 to 8 for schoolmaster snapper (A. Rypel, University of Alabama unpubl. data), and included sexually mature individuals (>185–195 mm SL for gray

Table 1. *Lutjanus griseus* and *L. apodus*. Influence of body size on habitat use measures for gray snapper (GS) and schoolmaster snapper (SM), based on linear regressions. HS: home site

Measure	Species	R ²	Slope	p-value
Mean distance moved	GS	0.03	-0.25	0.64
	SM	0.4	1.03	0.07
Max distance moved	GS	0.03	-0.67	0.68
	SM	0.36	2.17	0.09
Furthest distance from HS	GS	0.06	-0.84	0.55
	SM	0.07	0.48	0.49
Minimum convex polygon	GS	0.04	-0.01	0.50
	SM	0.15	0.02	0.15

snapper, Starck & Schroeder 1971, and >250 mm FL for schoolmaster snapper, Munro 1983). Based on length-weight regressions from tidal creeks on Abaco Island (C. Hammerschlag-Peyer unpubl. data), the SL range corresponded to body weights between 39 and 533 g for gray snapper and between 37 and 571 g for schoolmaster snapper. A total of 295 621 individual detections were recorded by stationary and manual receivers for these 40 fishes. All individuals were detected at least once at the home sites by the stationary receivers, and 17 gray and 18 schoolmaster snapper were detected outside the home sites using the manual receiver. Body size had no significant effect on any habitat use measure in both gray and schoolmaster snapper (Table 1). Most schoolmaster snapper stayed near the home sites throughout the study period. Only 3 of the 20 schoolmaster snapper and 7 of the 20 gray snapper were detected outside the zones immediately adjacent to the home sites (Fig. 1).

A few individuals in both populations utilized large areas, whereas most individuals were characterized by small activity spaces (Fig. 2G,H). The MCP estimates of the total activity space were 145 837 m² for 14 gray snapper and 46 565 m² for 15 schoolmaster snapper. Eleven gray and 12 schoolmaster snapper used <10% of the total activity space for each population, while 3 gray and 3 schoolmaster snapper used between 40 to 60% and 20 to 35% of the total activity space, respectively. Importantly, activity spaces differed also among individuals by their spatial orientation and location in the creek (e.g. Fig. 3). These data suggest that movements and habitat use among individuals varied substantially.

The WIC/TNW of gray and schoolmaster snapper equaled 0.34 and 0.47, respectively, with the BIC being larger than the WIC in both cases. The WIC/TNW index for both populations was significantly different from the null model ($p < 0.001$), which provides evidence for individual specialization in habitat use (Bolnick et al. 2002). Specialization is evident when look-

ing more closely at the differential habitat use of individual fish. For instance, 4 gray and 5 schoolmaster snapper only utilized 1 home site area during the entire study period and were never observed outside the home site areas. The 7 gray and 3 schoolmaster snapper that utilized other zones of the creek tended to move to the same areas repeatedly, with areas differing among individuals (Table 2). Four gray and 2 schoolmaster snapper moved to only 1 zone in addition to the home sites.

The $\delta^{13}\text{C}$ values of schoolmaster snapper were significantly related to the size of the activity space ($R^2 = 0.31$, $p = 0.03$), with individuals with small activity spaces having more depleted $\delta^{13}\text{C}$ values (Fig. 4B). The $\delta^{13}\text{C}$ values of gray snapper became more enriched with increased activity space utilization, but this positive relationship was not statistically significant ($R^2 = 0.15$, $p = 0.19$; Fig. 4A). These patterns are consistent with the $\delta^{13}\text{C}$ values of prey in upper and lower portions of the creek. Individual snapper that moved further (i.e. had large activity spaces extending toward the ocean) tended to have enriched $\delta^{13}\text{C}$ values similar to prey collected at the mouths of creeks (i.e. marine influences). This likely reflects a food web module originating from seagrass or seagrass epiphytes (that tend to be relatively enriched in $\delta^{13}\text{C}$). Snapper individuals that moved less had more depleted $\delta^{13}\text{C}$ values. These values were similar to those of prey collected in portions of creeks adjacent to the home sites, likely reflecting feeding pathways originating from micro- and macroalgae that are abundant in these areas (Kieckbusch et al. 2004).

DISCUSSION

Populations are usually considered to be homogeneous entities, with variation among individuals often not being incorporated into ecological studies. Our data suggest that there may be substantial small-scale intrapopulation variability in movement patterns in gray and schoolmaster snapper. In both populations, some individuals moved repeatedly outside of the home sites, while others remained in or near the deep pools for the entire study period. The activity space of individual fish also differed based on spatial orientation and location. Such intrapopulation variation has been shown to be important in several freshwater fish taxa (Bourke et al. 1997, Morbey et al. 2006, Kobler et al. 2009), and we provide one of the first extensions of this research framework to marine fishes (cf. Egli & Babcock 2004).

Intrapopulation variations in movement patterns and habitat use were generally not explained by body size even though the body size range in this study included

juvenile and adult individuals. Body size is widely accepted as a driver of intrapopulation variation in fish habitat utilization (e.g. Minns 1995). However, our results support recent research findings that indicated no effect of body size on intraspecific patterns of home range/activity space (Lowry & Suthers 1998, Weller & Winter 2001, Sakaris et al. 2003, Morbey et al. 2006)

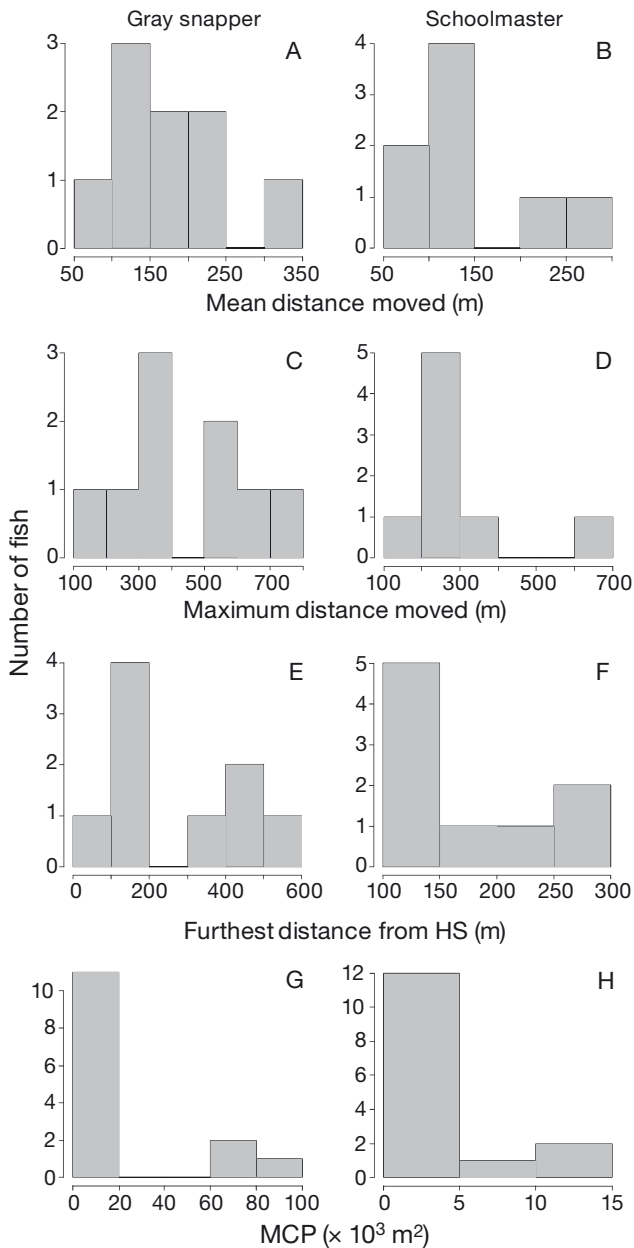


Fig. 2. *Lutjanus griseus* and *L. apodus*. Frequency histograms of (1) mean distance moved (average of daily movement distances, 26 April – 24 May 2009) (A–B); (2) maximum distance moved (C–D); (3) furthest distance away from home sites (HS; E–F); and (4) activity space based on minimum convex polygon (MCP) estimates (G–H). Left panels: results for gray snapper, right panels: results for schoolmaster snapper

and movement patterns (Bourke et al. 1997, Egli & Babcock 2004, Ng et al. 2007, Childs et al. 2008, Koster & Crook 2008). Only the mean and maximum distances moved by schoolmaster snapper were marginally significant ($0.1 > p > 0.05$; Table 1), likely because 3 of the larger individuals moved outside the home site zones (Fig. 1). These findings could be affected by the inherently small sample size that is common in telemetry studies due to costs associated with tagging technology (Luo et al. 2009). Despite this limitation, we demonstrated that intrapopulation differences in movement patterns of marine fishes may be greater than has previously been recognized and that factors other than body size may drive much of this variation.

In back reef systems, seascape attributes have been shown to be a primary determinant of habitat use of fishes (Pittman et al. 2007). Many coastal fishes, including snapper, generally prefer mangroves surrounded by dense seagrass (Pittman et al. 2007) since mangrove prop roots provide shelter (Hammerschlag et al. 2010) and seagrass beds contain high densities of prey items for coastal fishes (Orth et al. 1984). In the present study system, potential resource pools, such as seagrass, were patchily distributed across the creek system. Such a heterogeneous matrix of resources provides a large scope for individuals to develop specialized movement patterns. This contrasts with systems

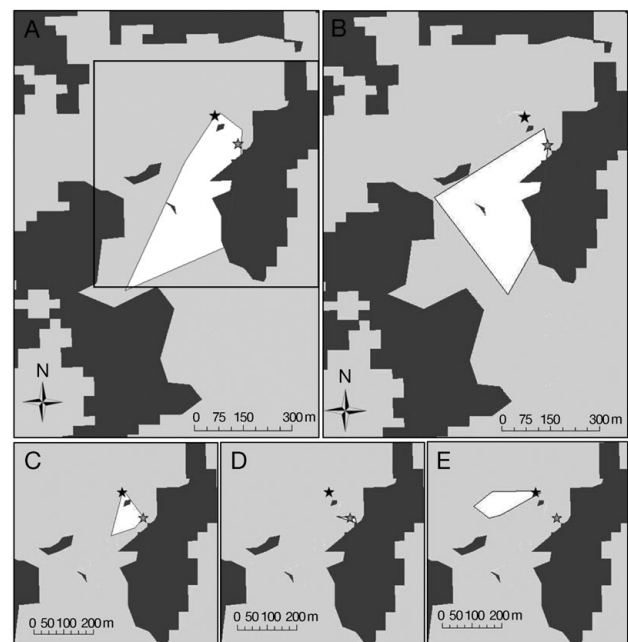


Fig. 3. *Lutjanus griseus* and *L. apodus*. Examples of the activity space for 5 ind. (gray snappers: A–C; schoolmaster snappers: D–E) in the study area (dark gray: land, light gray: aquatic habitat) from 26 April to 24 May 2009. (Black frame) in (A) corresponds to the area represented in the smaller maps (C–E)

Table 2. *Lutjanus griseus* and *L. apodus*. Total number of days on which gray snapper (GS) and schoolmaster snapper (SM) moved outside the home sites and the percentage of trips when each individual visited the different creek zones (see 'Results. Individual specialization' and Fig. 1 for more details). Total number of days is smaller than the actual number of observation points per individual because observations in home site areas are excluded here (see Table S1 in the supplement available at www.int-res.com/articles/suppl/m415p211_supp.pdf for more details)

ID	Species	Total no. of days	% to south west margin	% to west margin	% to south margin	% to middle	% to mouth
57223	GS	9	0	100	0	0	0
57224	GS	1	0	100	0	0	0
57229	GS	10	0	100	0	0	0
57230	GS	3	0	0	33	67	0
57245	GS	3	0	0	0	100	0
58468	GS	7	29	0	0	57	14
58474	GS	17	88	0	0	12	0
57221	SM	9	0	100	0	0	0
57222	SM	2	0	50	0	50	0
58469	SM	1	0	0	0	100	0

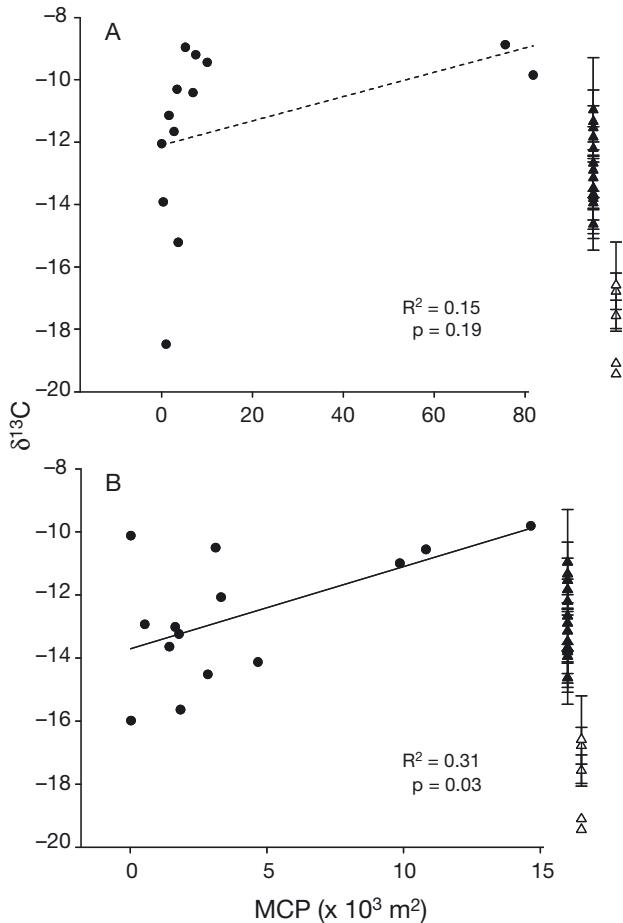


Fig. 4. Linear regression between $\delta^{13}\text{C}$ and activity space (minimum convex polygon, MCP) for (A) gray snapper and (B) schoolmaster snapper. (●) Data for an individual snapper. Triangles represent mean $\delta^{13}\text{C}$ values for individual taxa that are known to be snapper prey (error bars: SDs). The $\delta^{13}\text{C}$ values of prey items vary between (▲) creek mouth and (△) upstream areas. The $\delta^{13}\text{C}$ values of prey items were corrected for trophic discrimination (1‰, according to Post 2002)

where resources are concentrated in a single location. For example, schoolmaster snapper in Spanish Water Bay, Curaçao, typically move <5 m during the day (Verweij et al. 2007), perhaps because dense seagrass beds are found immediately adjacent to fringing mangroves. In this context, distinct movement patterns among individuals may be less likely to develop because of the concentration of food resources. Alternatively, the relatively short movement distances of schoolmaster snapper in Spanish Water Bay could also be because they feed and seek shelter in mangroves during the day, and feed in adjacent seagrass beds at night (Nagelkerken & van der Velde 2004, Verweij et al. 2006), probably to minimize predation risk (e.g. Laegdsgaard & Johnson 1995). Predation pressure might be higher in Spanish Water Bay than in our study system (we did not quantify this variable), and is another factor that must be considered when analyzing movement distances of individual fish.

Several contexts could be used to elucidate the intrapopulation variation in habitat use. The simplest explanation is that individuals moved randomly, thereby creating habitat use variation among individuals. However, repeated use of certain zones by the same individuals does not seem to support this explanation. Alternatively, movement patterns of individuals may relate to optimal foraging (MacArthur & Pianka 1966) in conjunction with differential learning among individuals (Brown & Laland 2003). In this context, it could be more beneficial for certain individuals to forage in proximity to the home sites, while others maximize their intake by foraging in other areas of the creek. Differential habitat use patterns among individuals are also consistent with the 'boldness versus shyness' dichotomy of behavioral traits (Gosling & John 1999, Gosling 2001), which is a pattern that is found in

many fish species (Sih et al. 2004a, Sih et al. 2004b). Bold individuals would be those that frequently explore other habitats or zones, while shy ones remain in proximity to their home sites (Russell 1983, Wilson et al. 1993).

An underlying driver for the above-mentioned contexts of intrapopulation habitat variation could be differential responses of individuals to competition. Specialization in movement patterns and habitat use is most likely when intraspecific competition is high (Svanbäck et al. 2008), and our empirical data may provide such an example in a marine system. Because all individuals are forced into the home sites at low tide, competition for food resources is likely high (Whitham 1978). Fishes that move further may be able to access underutilized food resources in patchily distributed seagrass beds outside the home site zones, with a trade-off of increased predation risk and energy expenditure (MacArthur & Pianka 1966).

Snapper $\delta^{13}\text{C}$ values tended to be more enriched in individuals that had large activity spaces. For both species, individuals that used large activity spaces (i.e. moved further toward the ocean) had similar $\delta^{13}\text{C}$ values to prey collected from creek mouths, while individuals with small activity spaces tended to have more depleted $\delta^{13}\text{C}$ values. These findings suggest that individuals with large activity spaces were feeding from a more marine-based food web module, likely originating in seagrass beds ($\delta^{13}\text{C} = -10.5 \pm 2.5$; Kieckbusch et al. 2004). In contrast, individuals with small activity spaces seem to feed on prey in the upstream portion of the creek (at and around the home sites), likely based on diverse macro- and microalgal assemblages ($\delta^{13}\text{C} = -18.0 \pm 5.4$, Kieckbusch et al. 2004). Patterns in tissue $\delta^{13}\text{C}$ provide strong evidence that the movement patterns that were observed over the course of the 4 wk study period were reflective of longer-term feeding trends (Hesslein et al. 1993, Suring & Wing 2009).

Fisheries management strategies have often ignored intrapopulation variation in marine systems. Simply concentrating on the 'typical' habitats of a fish population may inherently ignore those individuals that use alternative habitats or food sources (Durell 2000, Bolnick et al. 2003). For instance, in a New Zealand marine protected area (MPA), ~50% of New Zealand snapper *Pagrus auratus* remained within the area during the research period. However, the remaining individuals spent most of their time outside the reserve (Egli & Babcock 2004). This MPA would only provide protection for those individuals that moved less through time. Such cases and the present study suggest that a shift in perspective—from a population to an individual level—may be warranted in the management of some marine fisheries. Such an approach may help preserve species' ecological, phenotypic and

genetic diversity, and thus their ability to adapt to environmental change and to human impacts in marine ecosystems (Moritz 1994, Coates 2000, Smith et al. 2001, Bolnick et al. 2003).

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