



Loss of suitable ocean habitat and phenological shifts among grouper and snapper spawning aggregations in the Greater Caribbean under climate change

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ABSTRACT: Phenological shifts have been observed among marine species due to climate change. Modeling changes in fish spawning aggregations (FSAs) under climate change can be useful for adaptive management, because it can allow managers to adjust conservation strategies in the context of specific life history and phenological responses. We modeled effects of climate change on the distribution and phenology of Caribbean FSAs, examining 4 snapper and 4 grouper species. An ecological niche model was used to link FSAs with environmental conditions from remote sensing and project FSA distribution and seasonality under RCP8.5. We found significant differences between groupers and snappers in response to warming. While there was variation among species, groupers experienced slight delays in spawning season, a greater loss of suitable ocean habitat (average loss: 72.75%), and poleward shifts in FSA distribution. Snappers had larger shifts towards earlier phenology, with a smaller loss of suitable ocean habitat (average loss: 24.25%), excluding gray snapper, which gained habitat. Snappers exhibited interspecific variability in latitudinal distribution shifts. Snapper FSAs appeared more resilient to climate change and occupy wider and warmer spawning temperature ranges, while groupers prefer cooler spawning seasons. Consequently, groupers may lose more suitable ocean spawning habitat sooner due to climate change. When comparing species, there were trade-offs among climate change responses in terms of distribution shifts, phenology changes, and declines in habitat suitability. Understanding such trade-offs can help managers prioritize marine protected area locations and determine the optimal timing of seasonal fishing restrictions to protect FSAs vulnerable to fishing pressure in a changing climate.

KEY WORDS: Fish spawning aggregation · Reef fish · Grouper · Snapper · Climate change · Species distribution · Phenology · Greater Caribbean

1. INTRODUCTION

Climate change has altered environmental conditions on a global scale and prompted changes in latitudinal distribution, depth range, and phenology of fish species in marine environments (Beaugrand et

al. 2003, Edwards & Richardson 2004, Poloczanska et al. 2013). In marine systems, climate change has affected the abundance, spatial distribution, and phenology of species from the base of the food web up to higher trophic level predators (Beaugrand et al. 2011). Impacts on tropical fish species are of particu-

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lar concern because warming temperatures may result in unprecedented conditions for fishes not experienced anywhere in the world over recent geological conditions (Asch et al. 2018, Reygondeau et al. 2020).

Fish spawning aggregations (FSAs) are temporary gatherings of large numbers of conspecific fish that form, often at predictable times and locations, for the sole purpose of reproduction (Erisman et al. 2017). Transient FSAs are a life history phenomenon in which individual fish migrate from within a large catchment area to congregate and spawn in high densities at very specific locations for relatively short periods (i.e. days to weeks; Heyman & Kjerfve 2008, Biggs et al. 2021). In the Greater Caribbean, at least 37 species from 10 fish families are known to form transient FSAs (Kobara et al. 2013). Fish populations that spawn in large aggregations are highly vulnerable to heavy fishing pressure due to the ability of fisheries to predict and intensively target seasonal spawning locations (Sadovy de Mitcheson & Erisman 2012, Pittman & Heyman 2020).

Climate change adds an additional challenge for transient spawning species, which have adapted spawning times to correspond with specific seasonal climatic patterns (Hare et al. 2016). Impacts of climate change are predicted to affect reproductive function of marine fish, with previous studies having identified spawners and embryos as the most temperature-sensitive stages in the life cycle of fishes (Pörtner & Peck 2010, Asch & Erisman 2018, Dahlke et al. 2020). Warmer than optimal temperatures can affect every stage of reproductive development, including spawning, potentially altering the physiology of spawning populations (Alix et al. 2020). Spawning habitat represents a subset of a species' distribution range (see Fig. 1). The timing of migrations and spawning events, or the thermal habitat suitability at the FSA site, may additionally be altered by warming conditions (Asch & Erisman 2018).

Many snapper and groupers (families Lutjanidae and Epinephelidae, respectively) are important species in the Greater Caribbean in terms of both their ecosystem role and as fishery resources (Polovina & Ralston 1987, Arreguin-Sanchez et al. 1996, Amorim et al. 2018). They are managed and harvested as a multi-species complex in the USA, because these families share similarities in life history and ecological characteristics (Farmer et al. 2016, Stevens et al. 2019). Large-bodied species in the snapper–grouper complex are typically long-lived, have late reproductive maturity, and spawn together in transient aggre-

gations (Coleman et al. 2000, de la Guardia et al. 2018). These common characteristics make many snapper and grouper species highly vulnerable to heavy fishing pressure and slow to recover if aggregation sites are overfished or extirpated (Heyman et al. 2013, Sadovy de Mitcheson et al. 2020). There have been substantial fishing impacts to snapper and grouper aggregations documented in the northern Caribbean, including Puerto Rico, the US Virgin Islands, Mexico, Belize, and the Florida Keys, with overall population declines (Claro & Lindeman 2003, Sadovy de Mitcheson & Erisman 2012). The size and structure of aggregations varies among snapper and grouper species, and the characteristics of these aggregations directly influences vulnerability to fishing pressures (Robinson et al. 2015, Biggs et al. 2021). For example, those species or populations that form a few large aggregations during brief periods tend to be more susceptible to rapid fishery declines than those that form many small aggregations over protracted periods (Erisman et al. 2011, Sadovy de Mitcheson & Erisman 2012). While empirical research focusing on climate change influence on fish reproduction is limited, studies on Caribbean fishes have suggested impacts to habitat availability, increased fishing vulnerability, and range shifts correlated with temperature sensitivity and climate change (Fodrie et al. 2010, Maharaj et al. 2018).

The goal of this study was to model potential shifts in spawning aggregation distribution as a result of climate change, examining 8 Caribbean reef fish species from the families Lutjanidae and Epinephelidae. Since temperature is a driving factor in species distribution and is directly affected by climate change, we explored how habitat preferences and spawning locations may change with species that spawn across varying temperature ranges. Temperature has been shown to be the predominant effect driving projected distribution changes in Nassau grouper (Asch & Erisman 2018), so we investigated whether this was the case for other reef fishes that form spawning aggregations in the Greater Caribbean. Species with lower thermal habitat preferences during spawning are hypothesized to be more sensitive to climate change and should exhibit larger changes in their distribution, ocean habitat suitability, and spawning phenology. Species with narrow thermal requirements for spawning may need to adjust their seasonality of spawning to a greater extent to stay within the preferred temperature range. With spring spawners, warming temperatures may cause species to spawn earlier in the year. Species spawning in the winter or fall may experience phenological shifts to spawning

later in the year as a result of waiting for seasonal temperatures to cool (Pankhurst & Porter 2003).

In the Caribbean, groupers tend to spawn during cooler, winter months, while snappers typically spawn during the warmer spring and summer (Heyman & Kjerfve 2008, Kobara et al. 2013, SCRFA 2014). We hypothesized that the differences in spawning seasonality and their thermal spawning preferences may result in grouper species experiencing greater changes to their distribution and timing of spawning as a result of climate change, compared to snappers. A multivariate approach was also taken to determine the influences of additional environmental factors on FSAs beyond temperature. Over a centennial scale, we applied an ecological niche modeling approach to compare FSAs of multiple species to assess distributional and phenological shifts, as well as changes in overall ocean habitat suitability, under climate change.

2. MATERIALS AND METHODS

2.1. Study area and species

To obtain records of spawning aggregation sites for all of our study species, we used a database developed through collaboration with specialists and con-

taining a comprehensive list of known FSA sites (Kobara et al. 2013, Asch & Erisman 2018). The database included observations and records of FSAs that were verified by direct observation or pubcount (our Fig. 1; see also Heyman et al. 2013, Kobara et al. 2013). The data aggregate multiple studies done throughout time periods ranging from 1992 to 2011 (Table 1). This database, along with a literature review and consultation from experts on the target species, was used as a reference for this research to identify spawning aggregation sites for each of the study species in the Greater Caribbean within 11.7° to 32.2°N and 61.6° to 92.4°W (Table 1). Geographic descriptions of spawning locations from the database were used to make minor adjustments to coordinates of FSA sites, since data on spawning aggregation locations were frequently rounded for security purposes to prevent exploitation of spawning aggregations by fishers. These adjustments included moving coordinates off land to the nearest stretch of the coastline and inspecting location descriptions in both the database and the original literature referenced. Latitudinal and longitudinal coordinates were fine-tuned to match these descriptions from the literature. Any gaps in the data and missing information on spawning months from the database were filled and cross-checked with the scientific literature to confirm

Table 1. Study species with fish spawning aggregation (FSA) characteristics and conservation status

Scientific name	Common name	Spawning months	Spawning duration (d)	Spawning depth (m)	FSA type	IUCN status
<i>Lutjanus cyanopterus</i>	Cubera snapper	March–September ^{a–f}	15	9–30	Transient	Vulnerable
<i>L. analis</i>	Mutton snapper	March–September ^{a–c,e–g}	8–10	0–40	Transient	Near Threatened
<i>L. synagris</i>	Lane snapper	May–October ^{b,c}	8–10	0–50	Resident	Near Threatened
<i>L. griseus</i>	Gray snapper	June–September ^{b,c}	Unknown	10–85	Transient	Least Concern
<i>Epinephelus striatus</i>	Nassau grouper	December–April, June–July (Bermuda) ^{c–e,h–j}	1–12	0–120	Transient	Critically Endangered
<i>Mycteroperca venenosa</i>	Yellowfin grouper	January–May ^{c–e,g–j}	3–10	0–120	Transient	Near Threatened
<i>M. bonaci</i>	Black grouper	December–March, May–August (Bermuda) ^{c,e,g,k–n}	8–10	0–90	Transient	Near Threatened
<i>E. guttatus</i>	Red hind	December–February, May–August (Bermuda) ^{a,i,j,o–t}	5–7	0–45	Transient	Least Concern

Sources: ^aDomeier & Colin (1997); ^bLindeman et al. (2000); ^cClaro & Lindeman (2003); ^dNemeth et al. (2004); ^eHeyman & Kjerfve (2008); ^fBoomhower et al. (2010); ^gD. M. Rielinger (Reefkeeper International, pers. comm.); ^hColin (1992); ⁱSala et al. (2001); ^jCushion et al. (2008); ^kLuckhurst (2010); ^lCarter & Perrine (1994); ^mEklund et al. (2000); ⁿPaz & Sedberry (2008); ^oLuckhurst (2011); ^pKadison et al. (2010); ^qEristhee et al. (2006); ^rNemeth (2005); ^sNemeth et al. (2006); ^tNemeth et al. (2007)

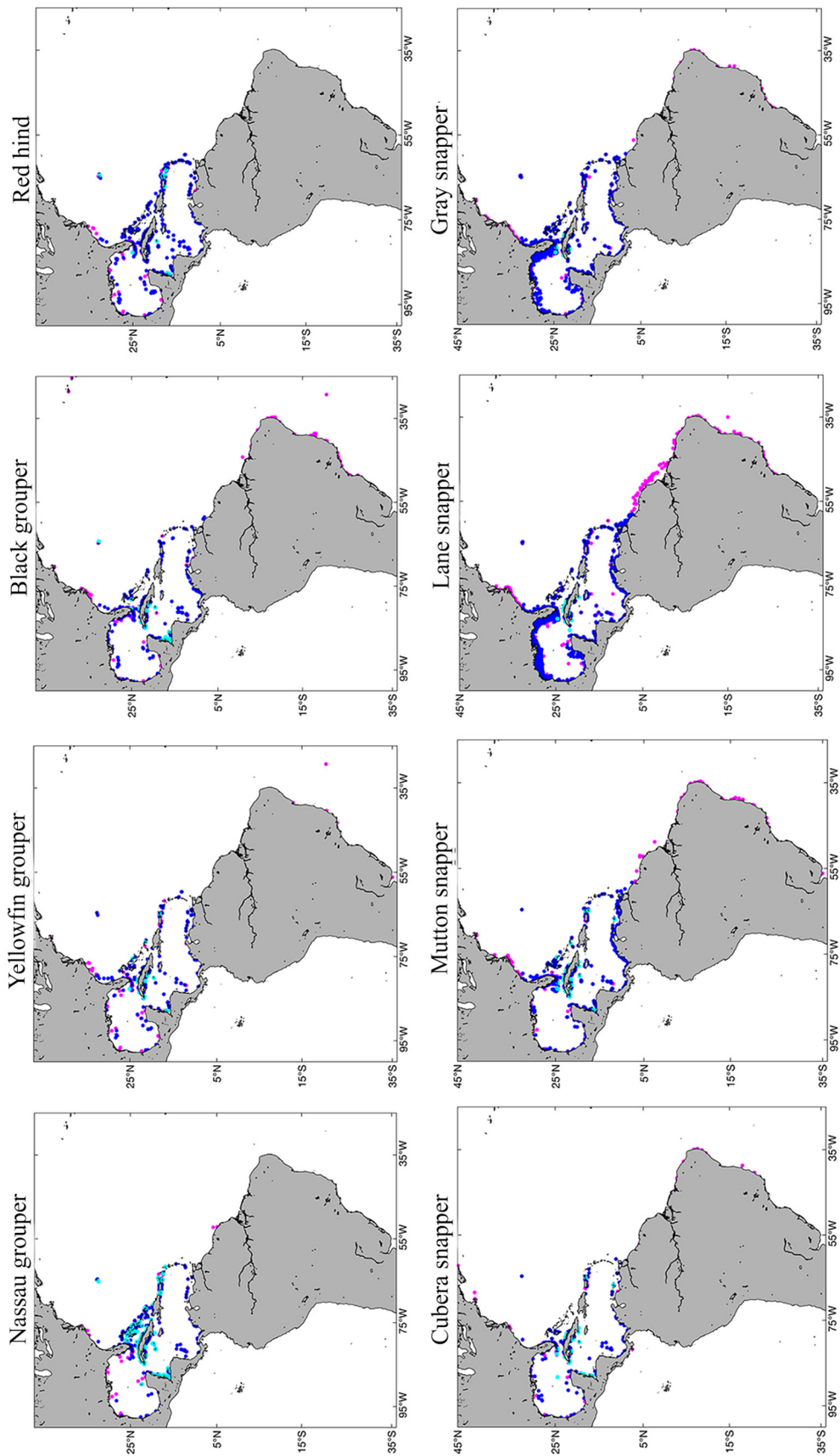


Fig. 1. Spawning sites from Kobara et al. (2013) and other references included in Table 1. To show that spawning aggregation sites occur across a more limited area of a species range, we also show data in these maps on the range-wide distribution of each species outside of the spawning season. These data are from 2 sources: the Ocean Biodiversity Information System (OBIS; <https://obis.org>) and Robertson & Van Tassel (2019). The latter reference has had all observations of species validated based on museum collections. OBIS is a more voluminous database, but taxonomic identification in this database is not validated with the same degree of expertise. OBIS also includes observations of larval and juvenile fishes that may have been advected into areas by currents, such as the Gulfstream. Therefore, these 2 datasets are complementary, given these distinctions. Cyan dots: spawning aggregation sites; blue dots: Robertson & Van Tassel (2019) species range; magenta dots: OBIS species range

the final list of spawning sites. The families Lutjanidae and Epinephelidae were among the most commonly listed species in existing FSA databases and our literature review, making these 2 groups ideal study species for this project. The study species in Lutjanidae included cubera snapper *Lutjanus cyanopterus*, mutton snapper *L. analis*, lane snapper *L. synagris*, and gray snapper *L. griseus*, while Epinephelidae included yellowfin grouper *Mycteroperca venenosa*, black grouper *M. bonaci*, Nassau grouper *Epinephelus striatus*, and red hind *E. guttatus*. The ecology of lane snapper differed from the other target species in that lane snapper is a resident spawner that does not travel long distances to reach FSA sites. Their spawning migration is to the closest reef edge adjacent to their primary habitat (Donahue et al. 2015). Species were selected based on data availability and number of observations within the Kobara et al. (2013) database. Sample size varied between species and among grouper and snapper FSAs. Sample size was defined based on the combined number of FSA sites and months when spawning occurred. Grouper sample size overall had an average of 113.5 (± 117.4 SD) observations, while snappers had 58.75 (± 31.6 SD) (see Tables 2 & 3). Lane and gray snapper had fewer recorded FSA observations compared to other species, but they were included in the list of study species so there could be a balanced comparison between the number of species in both families. Nassau grouper FSAs were previously examined by Asch & Erisman (2018) to develop a prototype species distribution model to examine climate change impacts. Data on this species were included herein for comparative purposes to assess more completely how thermal spawning preferences compared across these families of fishes.

2.2. Environmental variables

Satellite data on 7 environmental variables were used to assess their effect on the probability of occurrence of spawning aggregations. These variables included sea surface temperature (SST), seasonal SST gradients, geostrophic currents in the east–west and north–south directions, eddy kinetic energy (EKE), chlorophyll *a* concentration, and vertical velocity of seawater (i.e. upwelling and downwelling). SST was examined because it influences the distribution of many fish species (Pörtner & Peck 2010, Pinsky et al. 2013). Previous research showed SST to be a driving factor for the timing and distribution of spawning aggregations of Nassau grouper (Asch & Erisman

2018). Seasonal SST gradients were examined to determine temperature differences between subsequent months since some species may be triggered to spawn by directional increases or decreases in temperature rather than by an absolute temperature (Wootton & Smith 2014, Asch & Erisman 2018). Chlorophyll *a* concentration was used as a proxy for biological productivity at spawning aggregation sites, while EKE, vertical velocity, and geostrophic currents are connected to currents that can influence larval fish retention and dispersal, which can in turn influence the suitability of a potential spawning site (Karnauskas et al. 2011, Donahue et al. 2015).

The monthly climatology of SST sensed remotely throughout the Caribbean was obtained through the Advanced Very High Resolution Radiometer (AVHRR) Pathfinder, version 5.0 (NODC 2021). Infrared radiometers, such as AVHRR, only sense the upper 10–20 μm of the ocean (Nardelli et al. 2005). However, we used both day and nighttime measurements to minimize the effect of thermal stratification that may occur at the ocean surface during the day. Chlorophyll data were taken from the Hermes GlobColour dataset version 3.2 (ACRI-ST 2021), which was used to produce a monthly chlorophyll climatology. Vertical velocities were calculated based on data from the QuikScat SeaWinds scatterometer. This dataset was downloaded from the National Oceanic and Atmospheric Administration's Environmental Research Division's Data Access Program (NOAA 2021). Geostrophic current anomalies and EKE data were calculated from mean sea-level anomalies (MSLA) from the AVISO satellite altimetry data repository (AVISO 2021). We used a physical oceanographic naming convention where geostrophic flows in the north–south direction are labeled v , while currents flowing in the east–west direction are labeled with u . Chlorophyll concentration and EKE were \log_{10} -transformed prior to inclusion in modeling due to their log normal distribution. The Reefs at Risk database was used to obtain information on coral reef distribution to constrain projections of spawning occurrence to areas containing reefs (Burke et al. 2011). While the species distribution analysis did not include potential changes in spawning depth, shallow-water coral reef ecosystems are typically limited to a depth of 30 m or less.

2.3. Modeling and data analysis

The modeling and data analysis were conducted using Matlab software version R2018a. The Non-

Parametric Probabilistic Ecological Niche (NPPEN) model was used to model relationship data on FSA location and timing and information on environmental conditions from satellite data. The NPPEN model was designed to work with presence-only data and is based on a modified version of the Multiple Response Permutation Procedure (Beaugrand et al. 2011). Using a model that handles presence-only data was necessary for this study because there are no confirmed absences of spawning aggregations. A previous study comparing modeling methods of species distribution with presence-only data showed that a technique based on the Mahalanobis distance had one of the best performances when predicting species distribution based on an independent dataset (Tsoar et al. 2007). Since NPPEN is also based on Mahalanobis distance, NPPEN was expected to produce high model skill compared to alternative methods.

NPPEN was used to evaluate ocean habitat suitability for each species and assess what conditions were preferred for reproduction. All possible combinations of environmental variables were used in the NPPEN model initially, and the model with the set of environmental variables that minimized the corrected Akaike information criterion (AIC_c) was selected to use for developing future projections under climate change (Hurvich & Tsai 1989, Asch & Erisman 2018). Running all combinations of the model with the 7 environmental variables produced 128 possible models, including the null model. For species that had an AIC_c with the top models separated by values <2 , NPPEN results from each model were used to make future projections, and then the multi-model mean was used in subsequent statistical analysis (Burnham & Anderson 2002). Akaike weights were calculated for each variable for all species to determine the weighted influence of environmental covariates on distribution of FSAs (Burnham & Anderson 2002). Residual deviance explained (D) was used to assess model skill for each species. Deviance was calculated based on the following formula:

$$\text{Deviance} = AIC_c - 2 \cdot df \cdot \phi \quad (1)$$

where df is degrees of freedom, and ϕ is the dispersion parameter (Quinn & Keough 2002). A ϕ of 1 was used, since counts of FSAs should follow a Poisson distribution. D was then calculated as the difference between the null model deviance and the deviance of the selected model.

Future climate projections were based on the Intergovernmental Panel on Climate Change (IPCC) scenarios using Representative Concentration Pathways (RCPs). We made projections with RCP8.5, which is

considered a high-emissions climate scenario with an 8.5 W m^{-2} change in radiative forcing by 2100 resulting from anthropogenic impacts on climate (IPCC 2013). This RCP was used for analysis since it is the emissions scenario that recent greenhouse gas emissions have most closely tracked (Peters et al. 2013), although there has been a divergence of current emissions from this pathway in the last few years (Hausfather & Peters 2020). An earth system model (ESM) was used to examine how spawning aggregation sites for each species shift under the RCP8.5 climate scenario from 2081 to 2100 and compared against a historical scenario from 1981 to 2000. This analysis considered 20 yr climatologies and conditions were analyzed under both the RCP8.5 and the historical simulation. The principal model used for this study was developed by the NOAA Geophysical Fluid Dynamics Laboratory Earth System Model (GFDL ESM2M) (Dunne et al. 2012, 2013). The GFDL ESM2M model was selected due to the moderate equilibrium climate sensitivity compared to the other atmosphere–ocean circulation models included in the 5th Coupled Model Intercomparison Project (CMIP5) ensemble (Cheung et al. 2016).

The GFDL ESM2M model had a resolution of 1° at high latitudes, gradually becoming finer scale with a $1/3^\circ$ latitudinal resolution near the equator (Dunne et al. 2013, Cheung et al. 2016). The physical oceanography component of the GFDL ESM2M model utilized the Modular Ocean Model version 4.1 (Dunne et al. 2012, 2013). The marine biogeochemical model used in GFDL ESM2M was Tracers of Ocean Phytoplankton with Allometric Zooplankton, version 2.0, which included a nutrient–phytoplankton–zooplankton–detritus model with 3 phytoplankton functional groups (diazotrophs, small phytoplankton [pico- and nanoplankton], and large phytoplankton [e.g. diatoms]) (Dunne et al. 2013).

To assess projection uncertainty due to the use of different ESMs, we also examined how NPPEN projections differed among 3 different ESMs. These additional analyses were performed only on cubera snapper, which was used as a demonstration species to assess how inter-model differences might affect spawning habitat projections. Cubera snapper was selected as the representative snapper species due to its high number of observations of spawning aggregation sites.

In addition to GFDL ESM2M, we examined the Max Planck Institute MPI-ESM-MR model (Ilyina et al. 2013) and the Institut Pierre Simon Laplace IPSL-CM5A-MR model (Dufresne et al. 2013). These models were selected since, together with GFDL

ESM2M, they span the full range of equilibrium climate sensitivity from across the full CMIP5 ensemble. This combination of models has also been used frequently when developing climate projections for other living marine resources and, therefore, lows comparability of our results with those studies (Cheung et al. 2016, Muhling et al. 2017, Asch et al. 2018, Smith et al. 2021). While all 3 models had a 1° latitude/longitude resolution, each used a separate climate model grid. Environmental data from each model were re-gridded to a common 0.5° grid to allow for greater comparability (Cheung et al. 2016). Average annual NPPEN output from each model was then averaged across grid cells for model comparison. We also assessed the direction of changes in spawning habitat suitability in each grid cell to visualize the degree of agreement among the 3 ESMs. This analysis was completed for both the RCP8.5 and RCP4.5 scenarios to provide an initial assessment of how results might vary depending on the extent of climate mitigation. By the year 2100, RCP4.5 entails a 47% reduction in changes in radiative forcing compared to the high-emissions RCP8.5 scenario.

Model bias correction was used to optimize the comparability between the climate models and satellite observations. This bias correction was based on the monthly mean value of each individual environmental variable throughout the study area. Bias corrections have been shown to be useful in studying climate change impacts on several tropical marine systems (Logan et al. 2014, Matear et al. 2015). This allows for comparisons with observational data and increased model accuracy as bias corrections create a statistical relationship between the original data and modeled values for each covariate and then apply the resulting correction function to the modeled data (McHenry et al. 2019). Monthly mean correction factors were developed for the historical baseline period by comparing climatological values from ESMs and satellite observations, with these correction factors then applied to future periods when making climate change projections.

2.4. Statistics

To further analyze the results from future projections of FSA ocean habitat suitability, statistical metrics were calculated for each species and compared across species. Mean latitudinal shift was calculated as the change in weighted average distribution of FSA projections in kilometers per decade for each species. Weights were based on the area of each

ESM grid cell. Phenological shifts in month of spawning from historical and future projections indicated the extent to which spawning seasonality could be affected under the impacts of climate change. Central tendency (CT) of the spawning season was calculated as an indicator of phenological change that corresponded to the near center of a species' seasonal distribution. CT can be used to compare skewed seasonal distributions of spawning habitat (Edwards & Richardson 2004). CT was defined based on the following formula:

$$CT = \frac{\sum_{i=1}^{12} (S_i \cdot i)}{\sum_{i=1}^{12} S_i} \quad (2)$$

where i is the month, and S_i is the probability of suitable ocean spawning habitat in month i . These results were expressed as a phenological shift with units of days per decade. An integrated habitat suitability (IHS) score was calculated as a metric of future habitat loss, defined by the loss or gain of habitat based on an array of environmental factors included in the future projections under the RCP8.5 climate scenario. This integral was calculated as a sum across months and locations for probability of spawning habitat to obtain the total ocean habitat suitability across a species range. IHS was defined by the following formula:

$$IHS = \sum_{j=1}^{i_{\max}} \sum_{i=1}^{i_{\max}} h(i, j) \quad (3)$$

where $h(i, j)$ is the habitat suitability score in model grid cell i and month j , and i_{\max} is the total number of grid cells where coral reef habitat occurs.

This was compared between the historical and future scenarios. Change in IHS was measured as a percent change between the 2 scenarios and then converted to an odds ratio to perform a statistical test to assess the hypothesis that there were differences in IHS change across taxonomic groups. For the other 2 metrics measuring climate change impacts (i.e. CT and mean latitudinal shift), an independent, 2-sample t -test was used to examine if differences in sensitivity to climate change occurred between groupers and snappers. The 95% confidence intervals for changes in central tendency, weighted mean latitude, and IHS were calculated using a bootstrap method (Efron & Tibshirani 1998). All sites with NPPEN projections of spawning habitat suitability were resampled with replacement using a uniform distribution. Sample size for each bootstrap iteration was kept the same as in the original analysis. Central tendency, mean latitude, and IHS were calculated for the future and historical periods from this resampled dataset. Spawning habitat suitability probabilities from all months were used for calculating central

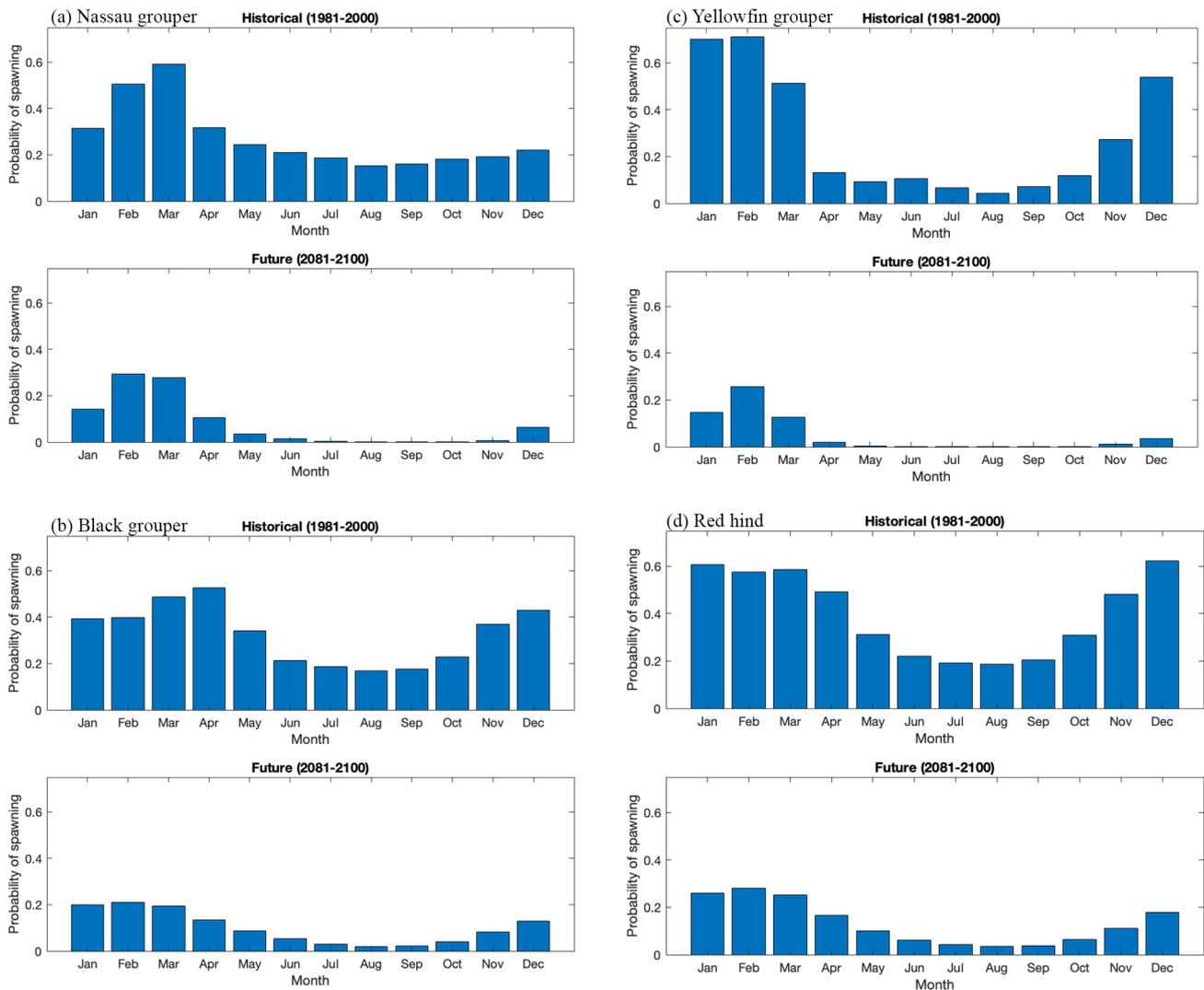


Fig. 2. Changes in spawning phenology and the probability of spawning across seasons for grouper species from the Geophysical Fluid Dynamics Laboratory Earth System Model (GFDL ESM2M) between historical and future periods. (a) Nassau grouper, (b) black grouper, (c) yellowfin grouper, (d) red hind

tendency and IHS, whereas weighted mean latitude was calculated based only on months with documented spawning during the historical period. This process was repeated 500 times with different bootstrap samples. Based on this distribution of samples, we identified the 2.5% and 97.5% quantiles to calculate 95% confidence intervals for each metric.

3. RESULTS

3.1. Comparison between grouper and snapper FSAs

Data presented herein showed that groupers tended to prefer cooler winter months and historically spawned between December and April throughout most of

their range, with the exception of Bermuda (Fig. 2). The selected snappers typically spawned later in the spring and summer between March and September, preferring warmer temperatures (Fig. 3, Fig. S1 in the Supplement at www.int-res.com/articles/suppl/m699p091_supp.pdf). Modeled spawning probability was maximized for groupers between 24 and 28°C (Fig. S1). For seasonal SST gradients, the maxima modeled probability of encountering FSAs fell between -1.5 and 1°C for groupers, with the exception of red hind, for which the model of best fit did not include seasonal SST gradients (Fig. S2). The modeled probability of spawning habitat for the snapper group was maximized at SSTs of 26.5–31°C and between -1 and 2°C for seasonal SST gradients (Figs. S1 & S2).

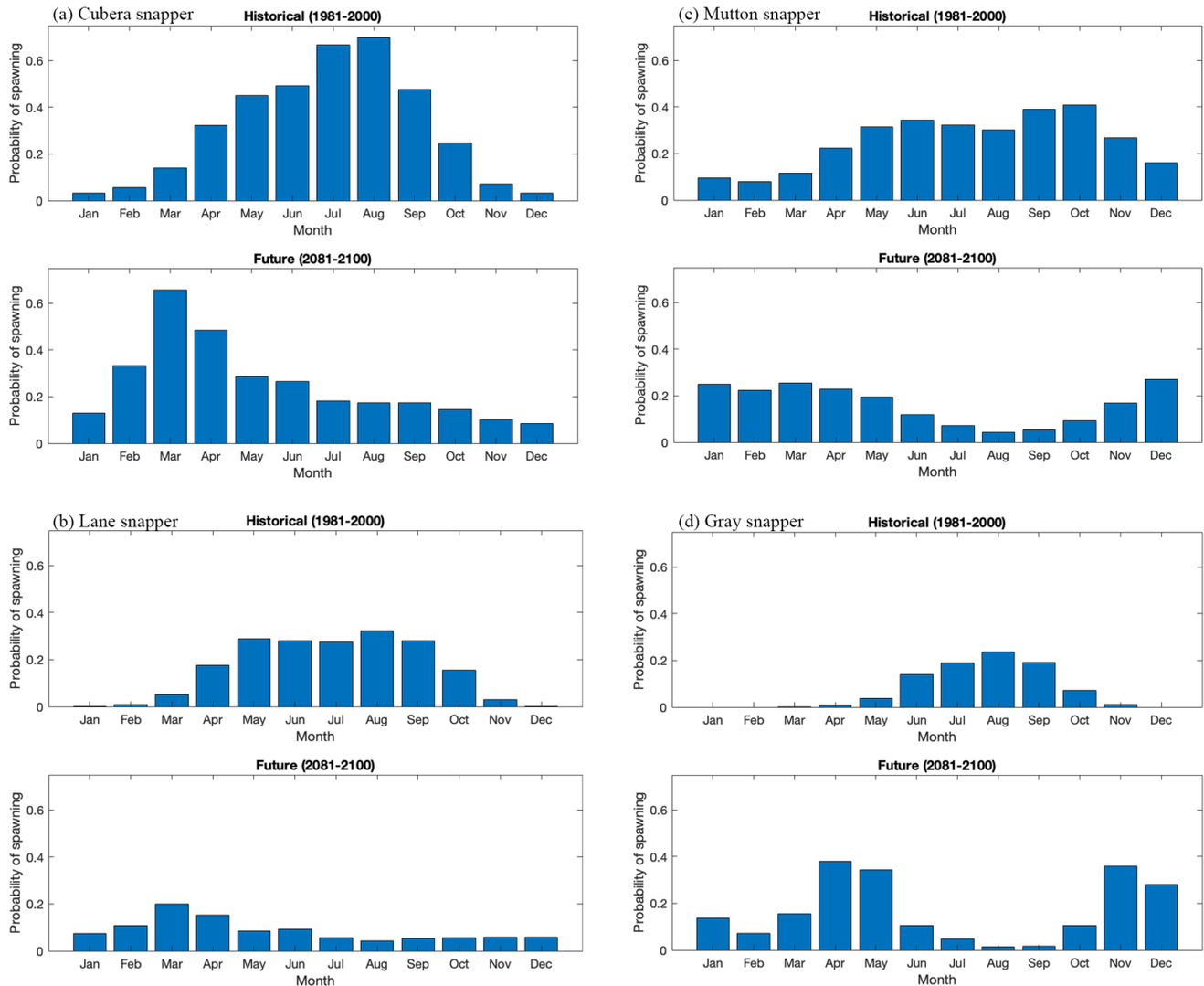


Fig. 3. Changes in spawning phenology and the probability of spawning across seasons for snapper species from the GFDL ESM2M between historical and future periods. (a) Cubera snapper, (b) lane snapper, (c) mutton snapper, (d) gray snapper

Significant differences between taxonomic groups were seen in CT of phenological shifts and IHS, while the differences between taxonomic groups for mean latitudinal shift were not significant (Table 2). Groupers had a mean spawning phenology shift of $2.8 \text{ d decade}^{-1}$ between historical and future scenarios, while snappers had a shift of $-4.7 \text{ d decade}^{-1}$ (Table 2). Changes in CT were significantly different across taxonomic groups at $p < 0.05$ ($t = 4.6$, $df = 6$). The positive mean phenology change for groupers suggests that these fishes will shift to spawn later in the season, while the negative value for snappers signifies that the spawning season for snappers will move to earlier in the year. The results of the statistical analysis also indicate a difference between snappers and groupers in terms of changes in the proba-

bility of suitable ocean spawning habitat ($p < 0.05$, $t = 3.8$, $df = 6$). Groupers appeared more affected by a greater loss of suitable ocean habitat in comparison to snappers. On average, groupers had a poleward shift in distribution of $15.6 \text{ km decade}^{-1}$ compared to a $-3.7 \text{ km decade}^{-1}$ equatorward shift among the snappers (Table 3), but the difference between families was not statistically significant ($p > 0.05$, $t = 1.9$, $df = 6$).

3.2. Groupers (Family Epinephelidae)

3.2.1. Yellowfin grouper *Mycteroperca venenosa*

NPPEN selected a model of best fit for yellowfin grouper with 3 environmental variables including

Table 2. Independent 2-sample *t*-tests and odds-ratio tests comparing grouper and snapper metrics from historical to future periods under the RCP8.5 climate scenario. Central tendency results show shifts in spawning seasonality by month. Integrated habitat suitability (IHS) was measured using an odds ratio comparing change in suitable ocean spawning habitat. Analysis of mean latitudinal shift of fish spawning aggregation sites was compared in days per decade

	Groupers	Snappers
Central tendency shift		
Mean	2.77	-4.66
df	6	
Test statistic (<i>t</i>)	7.27	
p	0.0003	
IHS odds ratio		
Mean	1.32	0.12
df	6	
Test statistic (<i>t</i>)	3.76	
p	0.0197	
Mean latitudinal shift		
Mean	15.61	-3.72
df	6	
Test statistic (<i>t</i>)	1.89	
p	0.1322	

SST, seasonal SST gradients, and *u*-geostrophic current anomalies ($D = 63.5$, $df = 36$, Figs. S1–S3, Table 4). Those 3 variables had high Akaike weights, between 0.99 and 1.00, suggesting high influence on FSA distribution and a high AIC weight of the selected model (Table S1). Spawning occurred primarily between January and April each year based on empirical observations, which was consistent with model predictions. Future projections indicated an overall decrease in spawning probability, with a peak during January through April (Figs. 4 & 2c). CT displayed changes of 4.9 d

Table 3. Mean latitudinal shift (MLS) for each species for historical and future scenarios in degrees of latitude; differences between periods measured in km decade⁻¹ (95% CIs in parentheses). Positive values: poleward shift; negative values: equatorward shift

	Historical MLS 1981–2000	Future MLS 2081–2100	Latitudinal change (km decade ⁻¹)
Nassau grouper	17.42	21.02	39.82 (31.56 to 44.87)
Yellowfin grouper	17.09	19.13	22.49 (15.08 to 27.48)
Black grouper	20.26	20.31	0.15 (-10.33 to 3.77)
Red hind	20.37	20.22	-0.01 (-7.17 to 8.01)
Cubera snapper	16.62	16.74	1.33 (-6.57 to 4.57)
Mutton snapper	18.84	19.11	3.00 (-13.62 to 5.32)
Lane snapper	18.32	17.36	-10.65 (-27.22 to -2.56)
Gray snapper	18.66	17.89	-8.54 (-20.34 to -3.22)

decade⁻¹ between the historical and future time periods, with the positive change in CT indicating a trend towards later spawning (Table 5, Fig. 5). This was the largest shift in phenology among the grouper species. There was a 71.1% projected loss in suitable ocean habitat for spawning based on the IHS score (Table 6, Fig. 6). Mean latitude of FSA sites was projected to shift by 22.5 km decade⁻¹ in the poleward direction (Table 3, Fig. 7).

3.2.2. Black grouper *M. bonaci*

Four variables had a substantial influence on black grouper FSA projections (Akaike weights between 0.98 and 1.0) and were selected by NPPEN in the best fit model: SST, seasonal SST gradients, and *u*- and *v*-geostrophic current anomalies ($D = 139.7$, $df = 99$) (Table S1). Both empirical observations and modeled results illustrate that black grouper spawned between December through March at all sites, except Bermuda, where spawning occurred between May and November. Future projections indicate that spawning seasonality will continue to peak in December through March but will be less seasonally variable (Fig. 2b). Black grouper was projected to shift spawning to later in the season at a rate of 1.8 d decade⁻¹ as a result of its flattened distribution curve for seasonal spawning (Table 5, Fig. 5). Based on the IHS scores from the model, black grouper is projected to lose 69.2% of its potential spawning habitat and have lower probability of suitable ocean spawning habitat overall (Table 6, Figs. 4c & 6). FSA sites were projected to shift slightly northward by 0.2 km decade⁻¹ (Table 3, Figs. 4 & 7).

3.2.3. Red hind *Epinephelus guttatus*

The environmental variables selected in the model of best fit for red hind were SST, *v*- and *u*-geostrophic current anomalies, and log₁₀ EKE ($D = 50.0$, $df = 26$). Red hind was the only species in NPPEN to have EKE selected as a variable influencing FSA distribution (Fig. S4). Akaike weights were above 0.99 for SST and *v*- and *u*-geostrophic current anomalies, with the Akaike weight for log₁₀ EKE at 0.79 (Table S1). Spawning occurred in Bermuda during May through August,

Table 4. Top 3 models for each grouper species with the environmental variables selected from the Non-Parametric Probabilistic Ecological Niche (NPPEN) model. Results based on minimization of corrected Akaike information criterion (AIC_c). SST: sea surface temperature; SSTG: seasonal SST gradients; v : east–west geostrophic current anomalies; u : north–south geostrophic current anomalies; w : vertical velocity; EKE: eddy kinetic energy; Chl: chlorophyll. For environmental variables, 1 = inclusion of the variable in a NPPEN model; 0 = exclusion from the model

	Sample size	Null model AIC_c	Selected model AIC_c	ΔAIC_c	Akaike weights	SST	SSTG	v	u	w	EKE	Chl
Nassau grouper	283	4368.80	4038.51	0.00	1.00	1	1	1	0	0	0	0
			4053.08	14.57	0.00	1	1	1	0	0	1	0
			4053.48	14.97	0.00	1	0	1	0	0	0	0
Yellowfin grouper	39	593.10	529.65	0.00	0.98	1	1	0	1	0	0	0
			538.93	9.27	0.01	1	1	0	1	0	0	1
			540.49	10.84	0.00	1	0	0	1	0	0	1
Black grouper	103	1557.50	1417.80	0.00	0.93	1	1	1	1	0	0	0
			1423.76	5.96	0.05	1	1	0	1	0	1	1
			1426.59	8.79	0.01	1	0	0	0	0	1	1
Red hind	30	445.90	395.27	0.00	0.29	1	0	1	1	0	1	0
			397.85	2.58	0.08	1	1	1	0	0	1	1
			398.18	2.91	0.07	1	0	1	1	0	1	1

while all other FSA sites were characterized by spawning between December and February. The historical model varied slightly from these observations for red hind, with a low probability of spawning habitat (<0.2) during September, October, March, and April. Future projections indicated seasonality may remain relatively consistent, but with a reduced probability of spawning habitat each month (Figs. 2d & 4d). CT indicated a phenological change of 1.3 d decade⁻¹ later in the season (Table 5, Fig. 5). Shifts to spawn later in the season were consistent with results from other grouper species. IHS scores from

the model projected a 66.8% loss in suitable ocean habitat between historical and future periods (Table 6, Fig. 7). Red hind had the smallest mean latitudinal change of all species, with a shift of -0.01 km decade⁻¹ equatorward (Table 3, Fig. 7).

3.2.4. Nassau grouper *E. striatus*

Primary oceanographic factors affecting Nassau grouper spawning distribution across large spatial scales included SST, seasonal SST gradients, and v -geostrophic current anomalies in the north–south direction ($D = 330.3$, $df = 280$). Akaike weights indicated these top 3 variables exerted high influence on FSA distribution (Table S1). Spawning for Nassau grouper typically occurred between December and April, apart from Bermuda, where they spawned during June and July. CT of spawning phenology shifted at a rate of 3.1 d decade⁻¹ later in the season between historical and future scenarios (Table 5, Fig. 5). The 2081–2100 projections showed spawning will primarily occur between January to March (Fig. 2a). The IHS score indicated Nassau grouper may experience an 82% loss in suitable ocean spawning habitat (Table 6, Fig. 6). Mean latitude of FSA

Table 5. Central tendency (CT) of seasonal spawning time for each species from the model for historical and future scenarios. CT corresponds to the weighted average month of spawning (1 = spawning in January; 2 = spawning in February; etc.). Interpolation is used to translate fractional CT values into calendar dates, with values <1 indicating spawning between December 1st and January 1st. Differences between climate scenarios expressed as changes in d decade⁻¹ (95% CIs in parentheses). Positive values: shift to later in the season; negative values: shift to earlier in the spawning season

	Historical CT 1981–2000	Future CT 2081–2100	Phenology change (d decade ⁻¹)
Nassau grouper	0.82 (Dec 25)	1.84 (Jan 25)	3.10 (2.54 to 3.77)
Yellowfin grouper	0.53 (Dec 16)	2.15 (Feb 5)	4.92 (4.75 to 5.32)
Black grouper	0.57 (Dec 17)	1.16 (Jan 5)	1.80 (1.73 to 2.76)
Red hind	0.46 (Dec 14)	0.87 (Dec 26)	1.25 (0.48 to 2.43)
Cubera snapper	6.85 (Jun 26)	5.15 (May 5)	-5.16 (-5.42 to -4.92)
Mutton snapper	7.37 (Jul 11)	5.80 (May 24)	-4.77 (-5.56 to -3.95)
Lane snapper	7.16 (Jul 5)	5.26 (May 8)	-5.78 (-6.98 to -4.61)
Gray snapper	7.76 (Jul 23)	6.80 (Jun 24)	-2.92 (-3.77 to 1.96)

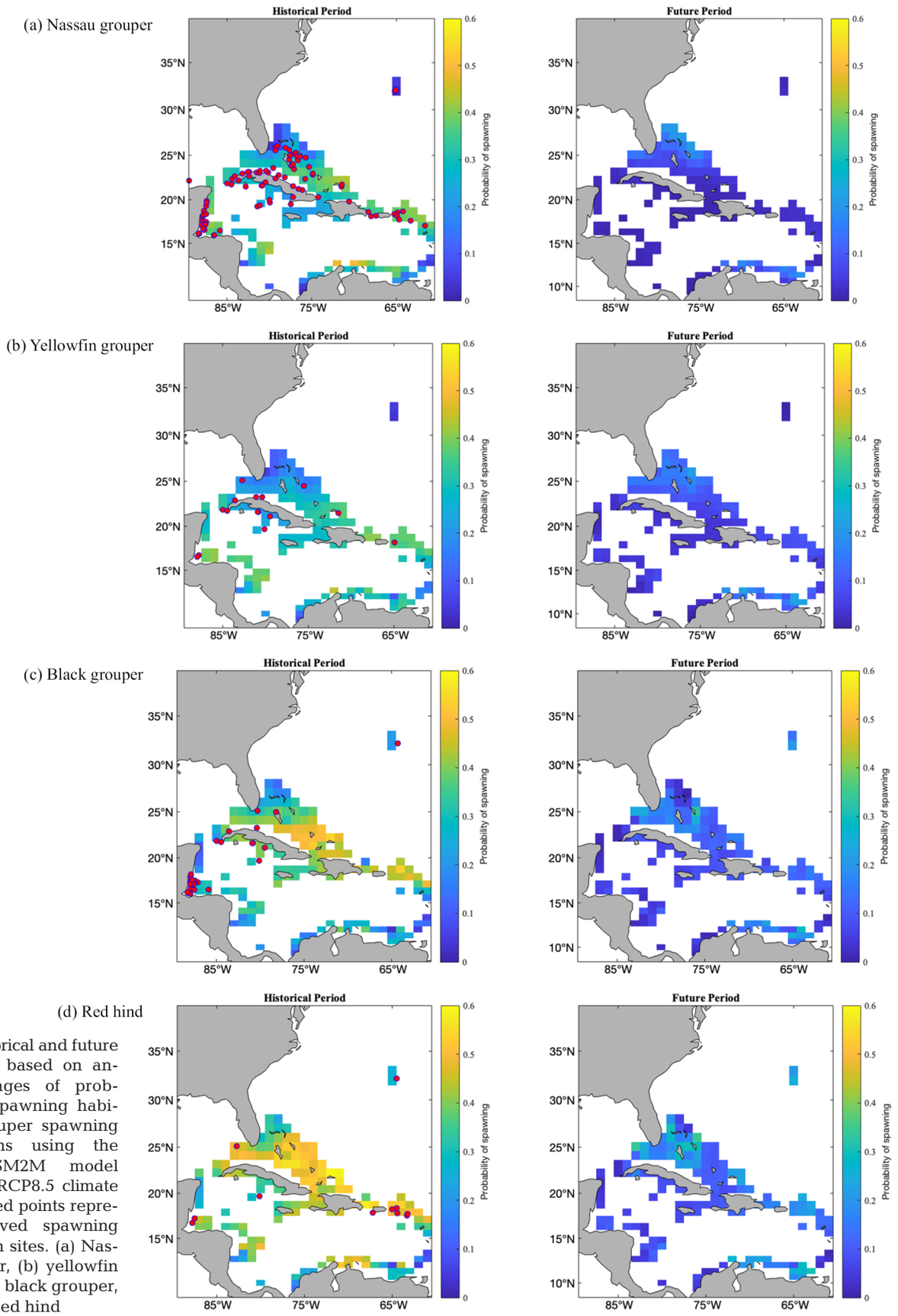


Fig. 4. Historical and future projections based on annual averages of probability of spawning habitat for grouper spawning aggregations using the GFDL ESM2M model under the RCP8.5 climate scenario. Red points represent observed spawning aggregation sites. (a) Nassau grouper, (b) yellowfin grouper, (c) black grouper, (d) red hind

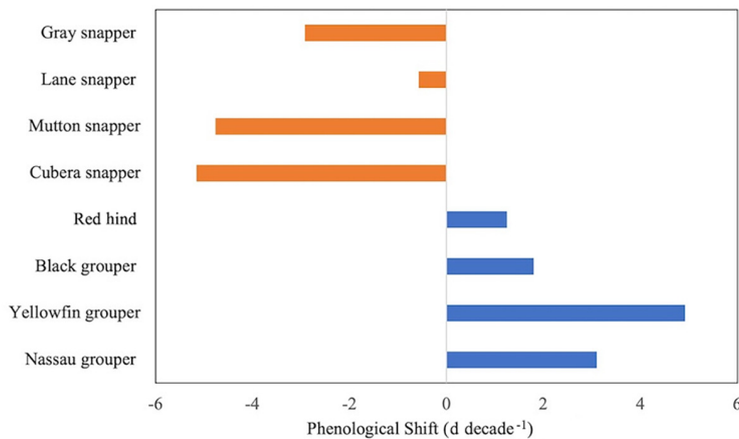


Fig. 5. Phenological shift among species from historical to future periods measured as central tendency. Negative numbers: shift towards earlier spawning; positive numbers: delay in spawning; orange: snappers; blue: groupers

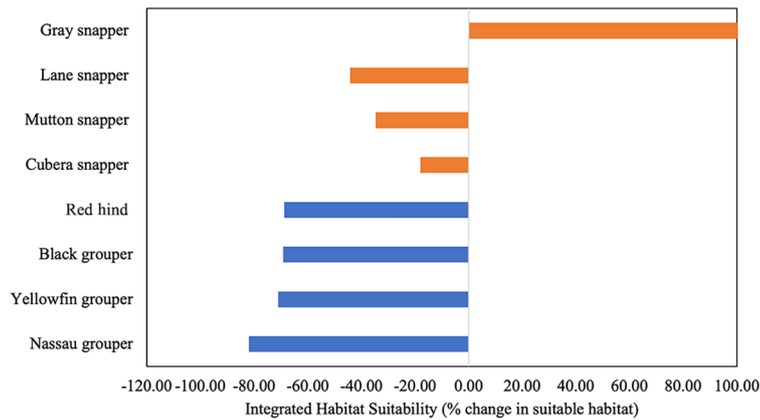


Fig. 6. Integrated habitat suitability (IHS) for species measured as loss of suitable ocean habitat and expressed in percentage of suitable habitat change from historical to future models. Positive numbers: gain in IHS; negative numbers: loss in IHS; orange: snappers; blue: groupers

occurrence projected a poleward shift at a rate of $39.8 \text{ km decade}^{-1}$ (Table 3, Fig. 7). Of the 8 study species examined, Nassau grouper experienced the greatest loss in suitable ocean spawning habitat and the largest mean latitudinal shift (Fig. 4a).

3.3. Snappers (Family Lutjanidae)

3.3.1. Cubera snapper *Lutjanus cyanopterus*

The primary oceanographic factors affecting spawning for cubera snapper

across a distribution-wide spatial scale were SST, seasonal SST gradients, and geostrophic current anomalies in the north–south direction ($D = 34.4$, $df = 64$, Table 7). These 3 environmental variables had Akaike weights above 0.9 (Table S2). Cubera snapper tended to use warm temperatures to spawn, with an average of 29°C (Fig. S1e). With respect to geostrophic velocity, cubera snapper spawn in areas with little-to-no current velocity present (Fig. S5).

Historical models and observations are consistent in indicating that spawning occurred primarily during April through October. Under RCP8.5, future projections indicate spawning will shift earlier in the year to peak between February and June by 2081–2100 (Fig. 3a). CT displayed an earlier shift of spawning season by $-5.2 \text{ d decade}^{-1}$ (Table 5, Fig. 5). This reflects approximately a 6 wk change between the historical and future time periods. Cubera snapper experienced the greatest shift in spawning seasonality compared to other species that were modeled. Its IHS score showed an 18.1% decline in spawning habitat, which was the smallest absolute change compared to other species (Table 6, Fig. 6). However, projections still showed a declining probability of spawning habitat occurrence between historical and future periods (Fig. 8). Of all the snapper species, cubera snapper also had the lowest mean latitudinal shift, $1.3 \text{ km decade}^{-1}$ in the poleward direction (Table 3, Fig. 7).

Table 6. Integrated habitat suitability (IHS) scores expressed in percent change for each species (95% CIs in parentheses). Positive values: gain of suitable ocean spawning habitat; negative values: loss of habitat. Note that IHS scores are unitless and dependent on area of integration, but they are comparable across species and periods since the same model grid was used for integration

	Historical IHS 1981–2000	Future IHS 2081–2100	Change (%)
Nassau grouper	516.08	92.82	-82.01 (-85.16 to -78.35)
Yellowfin grouper	501.20	145.07	-71.06 (-74.59 to -67.93)
Black grouper	599.25	184.80	-69.16 (-71.91 to -66.20)
Red hind	676.82	211.32	-68.78 (-64.22 to 69.34)
Cubera snapper	564.22	461.98	-18.12 (-23.40 to -11.88)
Mutton snapper	462.44	301.98	-34.70 (-42.60 to -25.15)
Lane snapper	264.02	147.30	-44.21 (-52.09 to -36.60)
Gray snapper	147.27	301.50	104.73 (85.41 to 184.68)

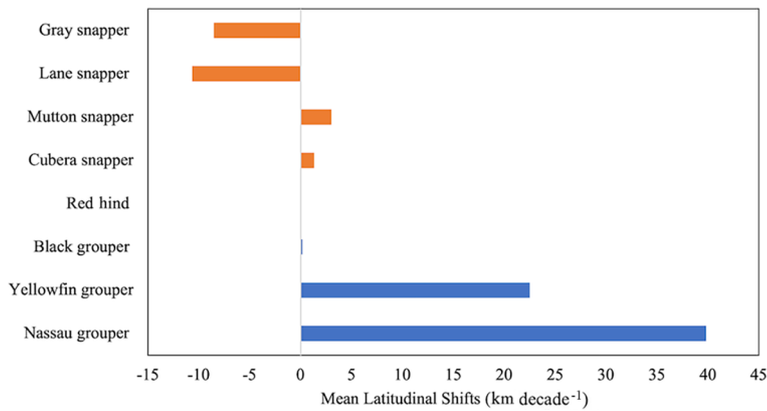


Fig. 7. Distributional shifts of study species from historical to future periods measured as mean latitudinal shifts. Positive numbers: poleward shifts; negative numbers: equatorward shifts; orange: snappers; blue: groupers

3.3.2. Mutton snapper *L. analis*

The 4 environmental variables selected in the model for mutton snapper spawning habitat included SST, seasonal SST gradients, and *v*- and *u*-geostrophic current anomalies ($D = 190.9$, $df = 88$). Akaike weights for all selected variables exerted a high influence on FSA distribution, with values around 1.0 (Table S2). Based on the NPPEN model, historical spawning patterns during the baseline period showed increases in suitable ocean spawning habitat during May through October, although FSA observations did not include October for mutton snapper spawning aggregation formation. Future projections indi-

cate a decrease in spawning particularly during June through October, with the maximal amount of spawning projected to occur during December through March by the end of the 21st century (Fig. 3c). This is supported by the CT results, which showed a shift to earlier spawning by $-4.8 \text{ d decade}^{-1}$ (Table 5, Fig. 5). Overall seasonality for mutton snapper spawning still encompassed a wide number of months (Fig. 3c). IHS scores from the model indicate a 34.7% decline in ocean habitat suitability under RCP8.5 (Table 6, Figs. 6 & 8). A poleward change in FSA distribution was projected to occur at a rate of $3.0 \text{ km decade}^{-1}$ (Table 3, Fig. 7).

3.3.3. Lane snapper *L. synagris*

The model of best fit for this species had 3 variables, including SST, seasonal SST gradients, and *u*-geostrophic current anomalies ($D = 184.7$, $df = 57$, Table S1). SST, *u*-, and seasonal SST gradients had high Akaike weights of around 1.0 (Table S2). Comparison of spawning phenology between historical and future periods showed a shift in historical peak spawning from May through September to January through March, as well as a lowered probability of occurrence of suitable ocean spawning habitat (Fig. 3b). Observations of spawning from April to

Table 7. Top models for each snapper species with the environmental variables selected from the NPPEN model. Results based on minimization of AIC_c. Top 3 models listed for all species, except for gray snapper, where 6 models were all within 2 units of the AIC_c from each other. See Table 4 for abbreviations and inclusion of environmental variables in models

	Sample size	Null model AIC _c	Selected model AIC _c	ΔAIC _c	Akaike weights	SST	SSTG	<i>v</i>	<i>u</i>	<i>w</i>	EKE	Chl
Cubera snapper	67	1020.80	986.42	0.00	0.84	1	1	1	0	0	0	0
			992.08	5.67	0.05	1	1	0	1	0	0	0
			992.21	5.79	0.05	1	1	1	0	0	0	1
Mutton snapper	92	1403.90	1212.97	0.00	1.00	1	1	1	1	0	0	0
			1236.46	23.49	0.00	1	1	1	1	0	1	0
			1236.94	23.97	0.00	1	1	1	1	1	0	0
Lane snapper	60	923.92	739.25	0.00	0.87	1	1	0	1	0	0	0
			743.45	4.20	0.11	1	1	0	1	0	1	0
			747.25	8.01	0.02	1	1	0	1	0	0	1
Gray snapper	16	242.90	203.42	0.00	0.14	1	0	0	0	1	0	0
			203.92	0.50	0.11	1	1	1	0	1	0	0
			203.97	0.55	0.10	1	0	1	0	1	0	0
			204.23	0.82	0.10	1	1	1	1	0	0	0
			204.31	0.82	0.10	1	0	1	0	0	0	0
			204.39	0.89	0.10	1	0	0	1	0	0	0

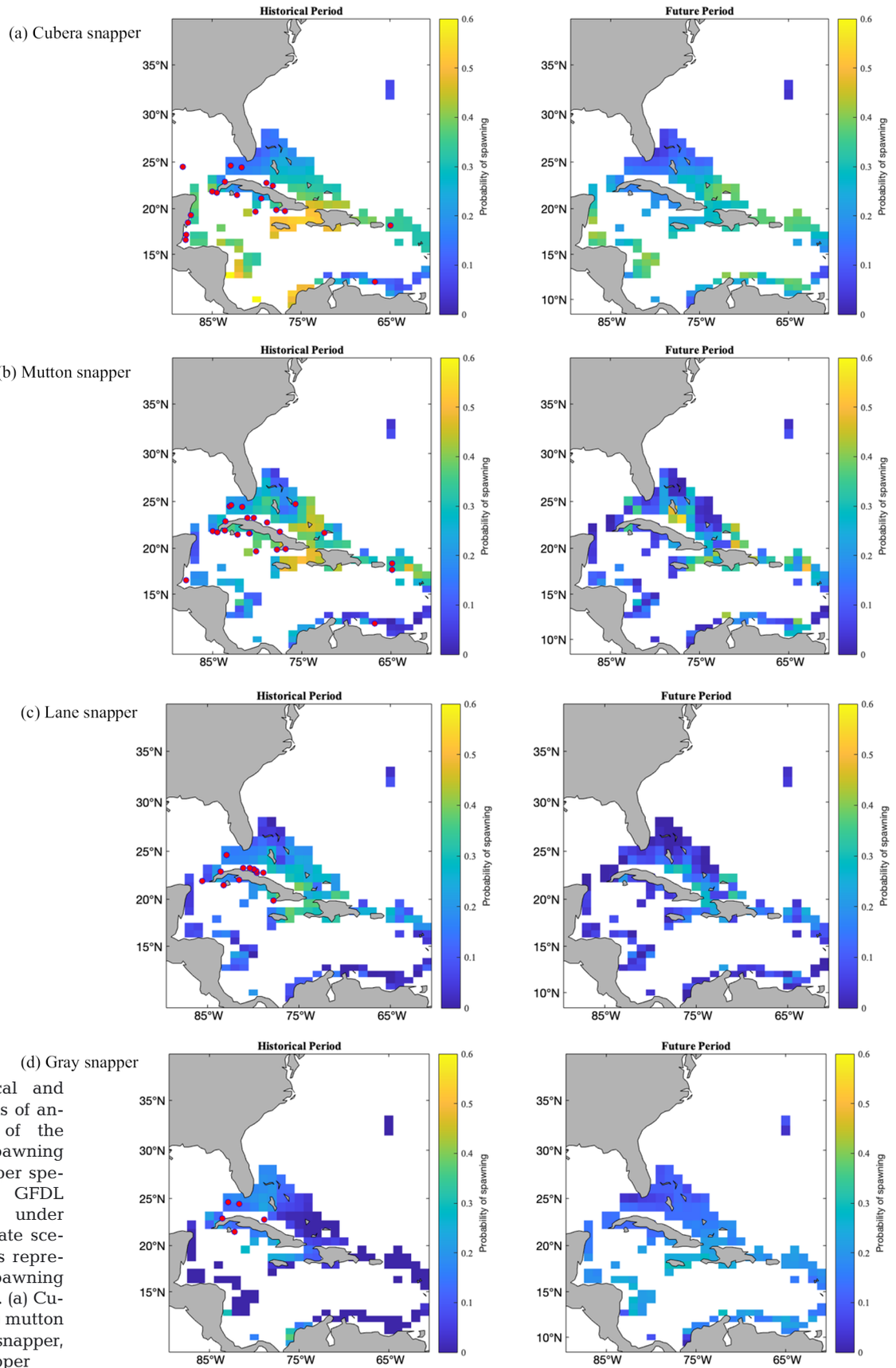


Fig. 8. Historical and future projections of annual averages of the probability of spawning habitat for snapper species using the GFDL ESM2M model under the RCP8.5 climate scenario. Red points represent observed spawning aggregation sites. (a) Cubera snapper, (b) mutton snapper, (c) lane snapper, (d) gray snapper

October were consistent with the historical spawning period from the model. CT showed a phenological shift to earlier in the season at -5.8 d decade⁻¹ (Table 5, Fig. 5). The NPPEN model calculated a 44.2% loss in suitable ocean spawning habitat from the IHS scores between the historical and future periods (Fig. 6). Lane snapper had the greatest equatorward change in mean latitude across all species at a rate of 10.7 km decade⁻¹ (Table 3, Fig. 7).

3.3.4. Gray snapper *L. griseus*

Gray snapper was the only study species to have a Δ AIC of <2 across the first 6 models of best fit, so steps were taken to average the NPPEN results of those top models. The first model included SST and vertical velocity (w) as the 2 variables influencing distribution ($D = 39.5$, $df = 14$). Results from vertical velocity data indicate gray snapper occurred in areas with slight downwelling (Fig. S6). The remaining models with a Δ AIC < 2 included a combination of 5 variables. The variables included were not only SST and w , but also seasonal SST gradients and u - and v -geostrophic current anomalies (Table 3). Akaike weights for environmental covariates were more variable for gray snapper compared to other species. SST was the primary influence, with an Akaike weight of 1.0, while the remaining selected variables had weights that ranged from approximately 0.4 to 0.6 (Table S2).

Consistent with the historical model, gray snapper was observed to spawn primarily between June through September. The future scenario from the model suggests spawning will become more variable, decreasing during summer and spiking during November–December and April–May (Fig. 3d). This species is projected to develop a bimodal spawning season under the RCP8.5 scenario. Spawning seasonality is projected to shift by -2.9 d decade⁻¹ earlier based on CT scores (Table 5, Fig. 5). IHS for gray snapper changed differently compared to other species. While other species experienced 18–82% loss of spawning habitat, gray snapper showed a 104.7% increase of suitable ocean habitat (Table 6, Fig. 6). An equatorward shift in spawning habitat distribution is projected at a rate of 8.5 km decade⁻¹ (Table 3, Fig. 7).

3.4. Comparison of models for cubera snapper

Three ESMs were used to assess how choice of model and climate scenario might impact our results. This analysis was performed solely on cubera snap-

per to demonstrate the degree of inter-model and scenario variability. Models agreed in the direction of change in spawning probability by the end of the century for a majority of grid cells under both RCP4.5 (71.2% of grid cells) and RCP8.5 (54.6%) (Fig. 9). Additionally, most grid cells with disagreement amongst models in terms of the direction of change were for smaller changes close to zero. For the RCP4.5 scenario, 53.6% of grid cells with disagreements in projected changes had changes less than 0.2 probability. For RCP8.5, 86.3% of disagreements in projected changes occurred in areas with a probability of spawning habitat of 0.2 or less. Overall, the IPSL-CM5A-MR model projected the largest changes in integrated habitat suitability across the study area, while the MPI-ESM-MR model projected the least amount of change in spawning habitat (Fig. S7).

4. DISCUSSION

4.1. Variables influencing FSAs

For all species studied, spawning habitat was modeled by a combination of SST and a hydrographic variable, including u - or v -geostrophic current anomalies, EKE, and vertical velocity. A previous study analyzing spawning aggregation distribution shifts for Nassau grouper under climate change was replicated for comparison with other species (Asch & Erisman 2018). Our results were consistent with Asch & Erisman (2018), with SST as the most frequent metric selected in the models as influencing species distribution, indicating the importance of temperature in ocean habitat selection and its influence on FSA distribution changes. Our findings were also consistent with previous work showing that seasonal deviations in monthly SST were among the most important influences on the distribution on large-bodied fish species (Mellin et al. 2016). We found that grouper species spawned at similar temperatures (24–28°C). However, Nassau grouper were the most selective in their range with regards to spawning temperature, while red hind was the most variable (Fig. S1). Snappers spawned within similar geographic ranges, but at warmer temperatures (26.5–31°C) compared to groupers (Fig. 2).

Seasonal SST gradients were included in best-fit models for all species, except for red hind, indicating that the change in temperature across seasons was a factor influencing spawning habitat in addition to the effect of absolute temperature. Certain reef fish species spawn during either warming or cooling periods

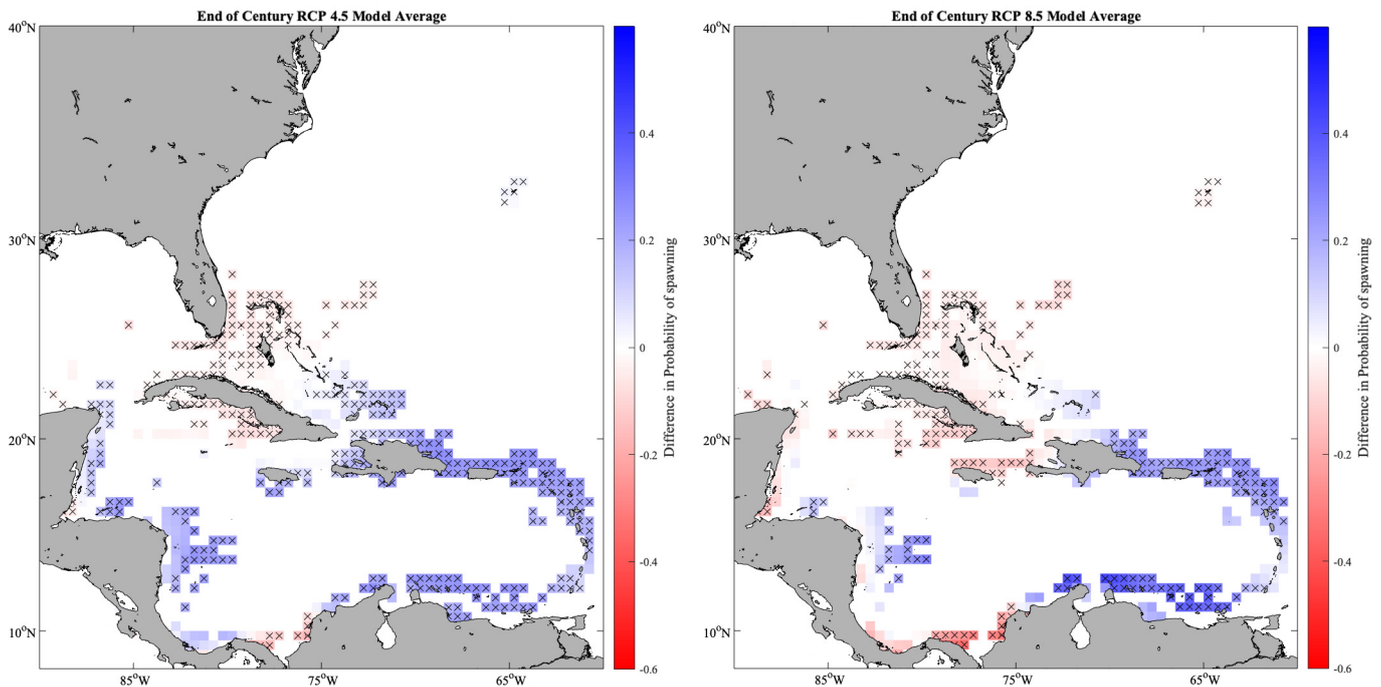


Fig. 9. Multi-model mean projected changes in spawning habitat suitability for cubera snapper *Lutjanus cyanopterus* between the future and historical periods for the RCP4.5 and 8.5 scenarios. Colors in the heat map indicate average changes across the GFDL, IPSL, and MPI models. Cross-hatching: locations where all 3 earth system models projected changes in the same direction

rather than at a specific temperature (Wootton & Smith 2014). Snappers tended to favor warming temperatures with seasonal SST gradients of -1 to 2°C compared to groupers, which most frequently utilized gradients of -1.5 to 1°C , but with high variability between species (Fig. 3).

Four snapper and 3 grouper species had v -geostrophic current anomalies as a metric selected in their best fit model, while 3 grouper and 2 snapper species had u -geostrophic current anomalies selected. For both variables, the probability of suitable ocean spawning habitat was maximized at current anomalies centered around zero, suggesting spawning occurs in areas with slow currents that may result in a greater probability of self-recruitment. Based on genetic evidence and biophysical models, self-recruitment has been documented to be common among grouper and snapper populations (Paris et al. 2005, Almany et al. 2013, Kough et al. 2016). Alternatively, this may be due to the fact that this study looked at climatologies of geostrophic currents anomalies averaged over a 20 yr period, which resulted in the majority of values for geostrophic current anomalies being close to zero. Due to the use of climatologies, another interpretation of the results is that deviations from typical conditions in currents were not conducive to spawning.

EKE was selected as a metric in the model of best fit for both red hind and gray snapper (Fig. S1), while vertical velocity was only detected as an important metric for gray snapper (Fig. S2). EKE and vertical velocity may affect FSA distribution through creation of conditions that influence larval fish feeding, as well as the advection and dispersal of eggs and larvae. In a previous study, drifters released during fish spawning periods were retained in eddies, indicating eddies may act to retain larvae closer to suitable coastal habitat (Heppell et al. 2008). In another study, drifters released at FSA sites moved more quickly offshore than those released at adjacent sites, which suggested that rapid offshore movement lowered predation risk on fish eggs and larvae (Gladstone 2007). However, red hind and gray snapper, the only species to include EKE and vertical velocity in their selected models, had low Akaike weights of 0.29 and 0.14 in their top models, respectively (Tables 4 & 7). This indicates lower confidence for model selection. Compared to other species, gray snapper had a low number of spawning observations, with a sample size of 16, while the sample size for red hind was 30. This may have impacted results. Alternatively, the reduced influence of EKE and vertical velocity on FSA habitat may reflect the spatial scale of our analysis, which focused on changes at the scale of species

distribution. Since EKE and vertical velocity exhibit heightened variability at the scale of kilometers to tens of kilometers, finer-scale spatial data may be needed to detect their influence on FSAs.

We initially hypothesized that results may be characterized by greater uncertainty for lane snapper due to the fact that lane snapper is classified as a resident spawner rather than a transient spawner. This could cause its distribution to be less tightly influenced by environmental conditions in the model. Rather than migrating to specific sites with certain environmental conditions, this species stays within its home range for spawning. Resident spawning sites may serve to minimize costs of migration and time exposed to increased predation risk (Donahue et al. 2015). Also, smaller-bodied reef fish may not have the physical and physiological capacity to migrate over long distances, so they may have evolved to adapt to spawn in more variable conditions close to their home range rather than seeking out specific conditions. Despite the expectation that lane snapper FSAs would be less tied to environmental variables due to resident spawning, the percentage of residual deviance explained by the environmentally driven NPPEN model was higher for lane snapper than for any other species. This surprising observation is a subject worthy of future investigation to better understand the underlying mechanisms.

Chlorophyll was not selected in models of best fit for any species, indicating that spawning habitat may not be influenced by biological productivity. This measure alone may not be enough to capture the influence of productivity in the system, or variability in chlorophyll may be too low to impact model results. Biological productivity is potentially important to FSA locations, but it may not be adequately captured by chlorophyll at the spatial scale of this analysis. Different measures of productivity, such as primary production, zooplankton productivity, or export of energy to the benthos, may be more appropriate to include as a variable in future modeling efforts (Stock et al. 2017). Friedland et al. (2012) found that metrics such as particle-export ratio and the ratio of secondary-to-primary production provided greater explanatory power as factors controlling fisheries production than chlorophyll concentration. Additional research could include examining the influence of the chlorophyll maxima at the thermocline on spawning aggregations, in addition to surface chlorophyll.

Overall, spawning habitat suitability for every snapper and grouper species examined herein could be modeled as a function of oceanographic variables. Deviance explained by environmental variables

ranged between 39.5 and 330.3 compared to a null model where equal spawning habitat suitability was assigned to all areas with coral reefs. However, the percentage deviance explained by these variables relative to the null model was low to moderate (3.4–20.0%). Also, in Figs. 4 & 8, modeled peak habitat use during the historical period was not always equal to areas with observed aggregations, which implies a level of uncertainty in the projected distribution among certain species. Looking at the magnitude of mismatch of the FSA observations, 1 observation was not predicted well by the models for snapper, while approximately 3 observations were not predicted well for grouper (Figs. 4 & 8).

These factors suggest that additional variables may need to be considered to fully understand spawning dynamics. For example, geomorphology, coral reef condition, lunar phase, and tidal dynamics have been previously shown to affect FSA occurrence among these species (Kobara et al. 2013, SCRFA 2014). Moreover, since our research focused on large-scale, climatological conditions, it is possible that additional variation in FSA occurrence could be explained if we were able to incorporate *in situ* data collected synchronously with spawning (Mannocei et al. 2017). However, due to the patchy nature of FSA monitoring programs across Caribbean countries, such environmental data were not uniformly available across the full distribution range of these species. Also, a number of biotic factors likely influence spawning habitat suitability. Population size will determine what percentage of suitable ocean spawning habitat is used (Planque et al. 2007). Since many FSA forming species are currently depleted compared to their carrying capacity, this may have influenced our ability to model the fullest extent of suitable ocean spawning habitat. Interspecific interactions, such as predation risk and availability of suitable prey, may also influence FSA selection (Fernandes et al. 2013, de Araújo et al. 2014). Lastly, some reef fishes exhibit behavioral dynamics where the return to FSA sites might be influenced by the presence of older fish who guide first-time spawners on their migration (NOAA 2013, MacCall et al. 2019).

4.2. Phenological shifts in FSA occurrence

When modeling phenology (Figs. 2 & 3), the maximum probability of spawning habitat generally corresponded well with observed spawning phenology (Gokturk 2021). However, low, but non-zero, probabilities of spawning habitat occurred in model output

during months when no spawning has historically been observed. This is indicative that there is a non-zero probability of occurrence of the physical oceanographic conditions associated with spawning, but that spawning may be constrained during these months by other physiological, ecological, or evolutionary factors, such as the time needed for oocyte development and vitellogenesis.

There was a clear pattern in phenological shifts between groupers and snappers that fell in line with the original hypotheses that there would be differences in phenological change between taxonomic groups. Groupers shifted phenology slightly later in the season. Groupers spawn during colder months and consequently cannot shift the timing of spawning by very much to track seasonal climate velocity since they are spawning already in the coldest months of the year. This is consistent with previous research suggesting that climate change will cause marine organisms to track the velocity of climate change and shifts in seasonal timing of temperatures (Burrows et al. 2011, Poloczanska et al. 2013). Since marine biodiversity is greater in the tropics, this is of particular conservation concern, as these areas tend to have greater phenological velocities of climate change (Burrows et al. 2011). In contrast to the groupers, snappers were projected to shift spawning earlier in the year, with peak spawning months moving from summer to spring and even late winter, with some variability between species. Gray snapper was the only species to exhibit a bimodal response in phenology, with climate velocities pointing in both directions, meaning shifts in spawning occurred both earlier and later in the year. This difference between how groupers and snappers react to climatic forcing has been observed in other settings. For example, in the Gulf of California, yellow snapper *Lutjanus argentiventris* and leopard grouper *Mycteroperca rosacea* exhibited different reactions to El Niño events, with recruitment and landings of grouper augmented during cool La Niña years and snapper recruitment and landings peaking during warm and wet El Niño years (Aburto-Oropeza et al. 2010).

Several snapper species were projected to undergo extremely large changes in reproductive phenology. This includes cubera snapper and lane snapper, which both experienced a 5 mo advancement in their projected peak spawning (Fig. 3). Similarly, projected seasonality of mutton snapper spawning was nearly reversed between historical and future periods (i.e. months with the lowest probability of spawning became months with the highest probability).

However, when viewing these patterns in terms of CT rather peak spawning month, rates of change were more modest (i.e. -4.8 to -5.8 d decade⁻¹ for these 3 species; Table 5), which is just slightly higher than the mean rate of observed change among marine species (Poloczanska et al. 2013). In contrast, the 5 mo change in peak spawning over a centennial scale corresponded to a rate of -15 d decade⁻¹. These differences reflect the fact that CT is a conservative phenological metric since it accounts for data from all months of the year (Ji et al. 2010). Generally, changes in seasonal extremes (i.e. peak, first, or last occurrence of a species) are subject to quicker phenological change than measures of mean occurrence (Langan et al. 2021).

A key question is whether the rapid shifts in month of peak spawning among snappers is biologically realistic or simply a function of NPPEN assuming that seasonal distributions will track a species' thermal niche. While the gonadosomatic index (i.e. an indicator of preparation for spawning) has been shown to closely track cumulative temperature exposure in many fishes (Ware & Tanasichuk 1989, Lange & Greve 1997, Gillet & Quélin 2006, Neuheimer & MacKenzie 2014), there are likely thermal limits beyond which this relationship would break down. The realism of rapid changes in spawning phenology might also depend upon the source of energy for spawning. Among fish that use recent food intake to provide energy for spawning (i.e. income spawners), there might be a greater capacity and need to rapidly adapt reproductive phenology to environmental changes compared to fishes that utilize stored energy reserves (i.e. capital spawners; Varpe et al. 2009). However, capital spawning may also allow fish with greater flexibility to use stored energy to react to unexpected conditions. Resident spawners, such as lane snapper (Table 1), who do not need to travel long distances to spawning sites, are more likely to be income spawners, whereas species that form transient spawning aggregations are more likely to be capital spawners.

Nonetheless, phenological changes as large as those projected for cubera, lane, and mutton snapper seem feasible, since observed changes of a similar magnitude have been seen among marine species as diverse as copepods, ctenophores, nudibranchs, shrimp, fish, and seabirds (Mackas et al. 1998, Bertram et al. 2001, Philippart et al. 2003, Juanes et al. 2004, Schlüter et al. 2010, Lambert 2013, Langan et al. 2021). Examples in these papers show observed rates of phenological change exceeding a magnitude of 15 d decade⁻¹ (i.e. the modeled rate of change in

peak snapper spawning). These examples of extreme phenological change were frequently linked to increases in temperature, with several of case studies coming from climate change hot spots (Lambert 2013, Langan et al. 2021). However, in several cases, additional factors influenced these extreme rates of phenological change (e.g. introduction of invasive species, changes in fisheries management, alteration of predator–prey dynamics; Juanes et al. 2004, Schlüter et al. 2010, Lambert 2013). Among these examples of large phenological changes, several of them negatively affected the survival, abundance, or recruitment of protected or commercially valuable species that interacted with the organism undergoing the sizable phenological change (Bertram et al. 2001, Philippart et al. 2003, Mackas et al. 2007).

4.3. Latitudinal shifts in FSA sites

Compared to phenological changes where there were distinct differences among fish families, the results of this study are less clear-cut when looking at projected latitudinal shifts in groupers and snappers. Results for latitudinal shifts differed among species and ranged from -11 to 40 km decade⁻¹ (Table 3). Groupers tended to have larger poleward shifts in distribution, but this also varied from species to species. Similar to the pattern seen with phenological shifts, Nassau and yellowfin grouper had the greatest latitudinal distribution shifts within the groupers. Equatorward shifts were observed with lane and gray snapper, whereas red hind was projected to have an extremely small shift in distribution (0.01 km decade⁻¹) (Table 3). Models from 5 of the 8 study species that all had high confidence based on Akaike weights showed a poleward shift in distribution. This is consistent with the global trend towards poleward distribution shifts observed for many fish species (Cheung et al. 2009, Morley et al. 2018).

Local extinctions and invasions can be caused by shifts in latitudinal ranges (Parmesan 2006, Cheung et al. 2009). These shifts in distribution have generally extended in the poleward direction with increasing temperatures, resulting in greater potential for climate-induced invasions and higher invasion intensity at higher latitudes (Hiddink & Hofstede 2008, Cheung et al. 2009, Fodrie et al. 2010). Conditions that exceed the temperature range of fish species may make reproduction at preferred FSA sites and seasons no longer possible, forcing them to adapt or shift spawning to cooler regions and seasons to avoid extinction (Dahlke et al. 2020).

While this was the first study to examine climate change impacts on multi-species spawning aggregation locations of Greater Caribbean groupers and snappers, several previous studies have investigated distribution shifts among these species outside of their spawning season. Morley et al. (2018) showed less agreement in distribution shift direction of species with low certainty in some models, which was the case for gray snapper, lane snapper, and red hind. The southward shift for gray snapper identified in this study contrasts with previous research indicating a poleward distribution shift (Hare et al. 2012). However, the magnitude of distribution shifts for FSAs in general and gray snapper in particular are dependent on the climate change scenarios examined in each study. Also, the smaller sample size for gray snapper FSA sites and low certainty in model selection may have influenced results. Additionally, this study limited spawning observations to sites with presence of current-day coral reefs and only examined the spawning life history stage, which differed from Hare et al. (2012). Since our study modeled gray snapper and other species as obligate reef spawners and did not make projections of future changes in coral reef distribution, this likely limited the extent of poleward distribution shifts. However, gray snapper can use coral reefs facultatively during spawning and may alternatively use other structured, hard-bottom substrate (SEDAR 2018). Future work should explore how this affects projected FSA distribution by parameterizing NPPEN to make projections of spawning in areas with coral reefs, other natural hard substrates (e.g. rock reefs), and artificial substrates, such as oil rigs and pipelines, that can serve as artificial reefs (Paxton et al. 2020).

4.4. Loss in suitable FSA ocean habitat

Our integrated habitat suitability metric explained gains and losses in marine spawning habitat independent of shifts in fish distribution and changes in phenology. Overall loss of suitable FSA habitat between historical and future periods under the RCP8.5 scenario ranged between 68 and 82% for groupers (Table 6). This contrasts with the smaller 18–44% loss of habitat among lane snapper, mutton snapper, and cubera snapper, producing significant differences between these 2 groups of fishes (Table 2). Gray snapper differed from all other study species and showed a gain of suitable ocean spawning habitat of nearly 105% by 2100 (Table 6). Previous work on gray snapper has also found that this species is projected to ex-

pand its range under the impacts of climate change as temperatures increase (Hare et al. 2012, Morley et al. 2018). For example, a study conducted on gray snapper in the Gulf of Mexico found a 71% increase in thermal habitat (Morley et al. 2018). Range expansions may increase local biodiversity on short-term timescales as poleward-retreating species are outpaced by poleward-advancing species. However, in the long term, these climate range expansions may act similarly to nonnative species invasions, modifying local dynamics related to competition, predation, herbivory, and parasitism (Fodrie et al. 2010). Populations shifting to favorable environmental conditions may still experience novel selection pressures from altered biotic interactions and unprecedented combinations of photoperiod cues and climatic effects (Moran & Alexander 2014).

Some research has suggested that rather than range shifts, it may be possible for fishes to respond to climate change by adjusting thermal range through individual acclimatization or evolutionary adaptation across generations (Angilletta 2009, Dahlke et al. 2020). This could be applicable to the study species in their ability to adapt to changing temperatures and spawn in warmer conditions rather than shifting their spawning habitat and range. The changes in suitable ocean habitat for FSAs may also be tied to other ecological needs for spawning habitat in addition to temperature requirements. Spawning sites may provide necessary substrates for egg deposition or hydrographic features that assist with egg and larval dispersal, including current speeds and flow direction (Heyman & Kjerfve 2008, Wootton & Smith 2014, Dahlke et al. 2020). As a result, even if a range shift would provide more suitable temperature conditions for species forming FSAs, additional requirements needed for reproduction may be missing, including factors related to hydrographic conditions, reef geomorphology, and biotic conditions like predator absence and food availability. Habitat fragmentation as a result of range expansion could result in decreased larval connectivity (Moran & Alexander 2014). Studies using coupled physical–biological models suggest larval transport could influence the range of marine species irrespective of local environmental conditions like temperature (Gaylord & Gaines 2000, Thompson et al. 2018). Changes and shifts in spawning habitat have further ecological implications, since predation, competition, and prey availability can change in response to co-occurrence of temperate, subtropical, and tropical species (Sax et al. 2007, Fodrie et al. 2010). These ecological interactions may influence whether it is possible to establish new FSAs in locations with suitable climatic conditions.

4.5. Future research

Further research on transient FSAs is needed, as there may be sites that are unmanaged or undiscovered currently (Kobara et al. 2013). Knowledge gaps on FSAs limit the effectiveness of management strategies meant to protect FSA habitat and productive fisheries, consequently limiting the performance or design of marine protected areas (MPAs) (Sale et al. 2005, Crowder & Norse 2008, Kobara et al. 2013). Fish populations can be sustained effectively through no-take reserves if spawning occurs within the boundaries or there is connectivity between FSA sites and marine reserves (Sale et al. 2010).

As establishment of MPAs and implementation of fishing restrictions can be depth-dependent, change in depth of spawning should be considered for future research. Similar research conducted has projected species to become restricted to deeper habitats based on historic observations (Pinsky et al. 2013, Kleisner et al. 2016, Morley et al. 2018). This could be an important metric to consider for species that may adjust depth in response to warming temperatures rather than adjusting their phenology or latitudinal distribution as was the focus of this study.

4.6. Conclusions

Researchers should continue to identify and monitor transient FSA sites following standardized protocols and sharing results via a cooperative FSA monitoring network (Heyman et al. 2019, Pittman & Heyman 2020). Future work exploring the life history characteristics of different species may provide key insights into responses under the impacts of climate change, and consequently managers may need to adjust current management strategies. Different life stages of fish species are not equally sensitive to temperature and the effects of climate change. These differences make it critical to develop modeling approaches that consider the interactions between multiple life history stages under the same framework.

Overall, there was evidence from this study supporting that climate change will impact the phenology, distribution, and ocean habitat suitability of reef-fish transient FSAs. Among these effects, strong differences were seen between the grouper and snapper families, with varying life history characteristics and thermal preferences for spawning. For each species, SST coupled with a hydrographic variable was found to influence distribution and proba-

bility of spawning habitat. With cooler preferences for spawning habitat, groupers were found to be more impacted by climate change under the RCP8.5 scenario compared to snapper species, which was consistent with our hypotheses. While the directionality and average latitudinal shift varied across species with no significant differences between families, there were differences between families in terms of changes in suitable ocean habitat and phenological shifts. Snappers are expected to shift spawning earlier in the season, while groupers' spawning season was projected to shift slightly later in the year and have a greater loss of suitable ocean spawning habitat. Differences in our results between shifts in mean latitudinal shift, habitat suitability, and phenology highlight the importance of looking at multiple metrics when studying fisheries under climate change. While there are many studies exploring distributional changes, other metrics, such as phenological shifts or overall habitat availability, may provide additional insight into climate change responses. This is especially important since our results suggest that there may be trade-offs between whether species respond to climate change by altering their phenology, mean latitudinal distribution, or by experiencing declines in habitat suitability. The information obtained from this study could be a useful tool in future management of both marine reserves and other types of protected areas, as the timing and locations of spawning events could be altered as a result of climate change. This has implications for harvest and fishing restrictions, as well as seasonal sales bans and site closures during spawning events.

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LITERATURE CITED

- Aburto-Oropeza O, Paredes G, Mascareñas-Osorio I, Sala E (2010) Climatic influence on reef fish recruitment and fisheries. *Mar Ecol Prog Ser* 410:283–287
- ACRI-ST (2021) Hermes GlobColour [data set]. <https://hermes.acri.fr/index.php> (accessed Oct 2016)
- Alix M, Kjesbu OS, Anderson KC (2020) From gametogenesis to spawning: how climate-driven warming affects teleost reproductive biology. *J Fish Biol* 97:607–632
- Almany GR, Hamilton RJ, Bode M, Matawai M and others (2013) Dispersal of grouper larvae drives local resource sharing in a coral reef fishery. *Curr Biol* 23:626–630
- Amorim P, Sousa P, Wetmeyer M, Menezes GM (2018) Generic knowledge indicator (GKI): a tool to evaluate the state of knowledge of fisheries applied to snapper and grouper. *Mar Policy* 89:40–49
- Angilletta MJJ (2009) Thermal adaptation: a theoretical and empirical synthesis. Oxford University Press, Oxford
- Arreguin-Sanchez F, Mo JL, Balgos MC, Pauly D (1996) Biology, fisheries and culture of tropical groupers and snappers. ICLARM, Campeche
- Asch RG, Erisman B (2018) Spawning aggregations act as a bottleneck influencing climate change impacts on a critically endangered reed fish. *Divers Distrib* 2018:1–17
- Asch RG, Cheung WWL, Reygondeau G (2018) Future marine ecosystem drivers, biodiversity, and fisheries maximum catch potential in Pacific Island countries and territories under climate change. *Mar Policy* 88:285–294
- AVISO (2021) AVISO+ satellite altimetry data. www.aviso.altimetry.fr (accessed Sep 2016)
- Beaugrand G, Brander KM, Lindley JA, Souissi S, Reid PC (2003) Plankton effect on cod recruitment in the North Sea. *Nature* 426:661–664
- Beaugrand G, Lenoir S, Ibañez F, Manté C (2011) A new model to assess the probability of occurrence of a species, based on presence-only data. *Mar Ecol Prog Ser* 424:175–190
- Bertram DF, Mackas DL, McKinnell SM (2001) The seasonal cycle revisited: interannual variation and ecosystem consequences. *Prog Oceanogr* 49:283–307
- Biggs CR, Heyman WD, Farmer NA, Kobara SI and others (2021) The importance of spawning behavior in understanding the vulnerability of exploited marine fishes in the U.S. Gulf of Mexico. *PeerJ* 9:e11814
- Boomhower J, Romero M, Posada J, Kobara S, Heyman W (2010) Prediction and verification of possible reef-fish spawning aggregation sites in Los Roques Archipelago National Park, Venezuela. *J Fish Biol* 77:822–840
- Burke L, Reynter K, Spalding M, Perry A (2011) Reefs at risk revisited. World Resources Institute, Washington, DC
- Burnham KP, Anderson DR (2002) Model selection and multimodel inference: a practical information-theoretic approach, 2nd edn. Springer-Verlag, New York, NY
- Burrows MT, Schoeman DS, Buckley LB, Moore P and others (2011) The pace of shifting climate in marine and terrestrial ecosystems. *Science* 334:652–655
- Carter J, Perrine D (1994) A spawning aggregation of dog snapper, *Lutjanus jocu* (Pisces: Lutjanidae) in Belize, Central America. *Bull Mar Sci* 55:228–234
- Cheung WWL, Close C, Kearney K, Lam V, Sarmiento J, Watson R, Pauly D (2009) Projections of global marine biodiversity impacts under climate change scenarios. *Fish Fish* 10:235–251
- Cheung WWL, Jones MC, Reygondeau G, Stock CA, Lam VW, Frolicher TL (2016) Structural uncertainty in projecting global fisheries catches under climate change. *Ecol Modell* 325:57–66
- Claro R, Lindeman KC (2003) Spawning aggregation sites of snapper and grouper species (Lutjanidae and Serranidae) on the insular shelf of Cuba. *Gulf Caribb Res* 14: 91–106
- Coleman FC, Koenig CC, Huntsman GR, Musick JA and others (2000) Long-lived reef fishes: the grouper-snapper complex. *Fisheries* (Bethesda, Md) 25:14–21
- Colin PL (1992) Reproduction of the Nassau grouper, *Epinephelus striatus* (Pisces: Serranidae) and its relationship to environmental conditions. *Environ Biol Fishes* 34: 357–377

- Crowder L, Norse E (2008) Essential ecological insights for marine ecosystem-based management and marine spatial planning. *Mar Policy* 32:772–778
- Cushion N, Cook M, Schull J, Sullivan-Sealey KM (2008) Reproductive classification and spawning seasonality of *Epinephelus striatus* (Nassau grouper), *E. guttatus* (red hind) and *Mycteroperca venenosa* (yellowfin grouper) from the Bahamas. *Proc 11th Int Coral Reef Symp* 11:994
- Dahlke FT, Wohlrab S, Butzin M, Pörtner H (2020) Thermal bottlenecks in the life cycle define climate vulnerability of fish. *Science* 369:65–70
- de Araújo CB, Marcondes-Machado LO, Costa GC (2014) The importance of biotic interactions in species distribution models: a test of the Eltonian noise hypothesis using parrots. *J Biogeogr* 41:513–523
- de la Guardia E, Giménez-Hurtado E, Defeo O, Angulo-Valdes J and others (2018) Indicators of overfishing of snapper (Lutjanidae) populations on the southwest shelf of Cuba. *Ocean Coast Manage* 153:116–123
- Domeier ML, Colin PL (1997) Tropical reef fish aggregations: defined and reviewed. *Bull Mar Sci* 60:698–726
- Donahue M, Karnauskas M, Toews C, Paris C (2015) Location isn't everything: Timing of spawning aggregations optimizes larval replenishment. *PLOS ONE* 10:e0130694
- Dufresne JL, Foujols MA, Denvil S, Caubel A and others (2013) Climate change projections using the IPSL-CM5 Earth System Model: from CMIP3 to CMIP5. *Clim Dyn* 40:2123–2165
- Dunne JP, John JG, Adcroft AJ, Griffies SM and others (2012) GFDL's ESM2 global coupled climate-carbon earth system models. Part I: physical formulation and baseline simulation characteristics. *J Clim* 25:6646–6665
- Dunne JP, John JG, Shevliakova E, Stouffer RJ and others (2013) GFDL's ESM2 global coupled climate-carbon earth system models. Part II: carbon system formulation and baseline simulation characteristics. *J Clim* 26: 2247–2267
- Edwards M, Richardson AJ (2004) Impact of climate change on marine pelagic phenology and trophic mismatch. *Nature* 430:881–884
- Efron B, Tibshirani RJ (1998) An introduction to the bootstrap. Chapman & Hall/CRC Press, Boca Raton, FL
- Eklund AM, McClellan DB, Harper DE (2000) Black grouper aggregations in relation to protected areas within the Florida Keys National Marine Sanctuary. *Bull Mar Sci* 66: 721–728
- Erisman BE, Allen LG, Claisee JT, Pondella DJ, Miller EF, Murray JH, Walters C (2011) The illusion of plenty: Hyperstability masks collapses in two recreational fisheries that target fish spawning aggregations. *Can J Fish Aquat Sci* 68:1705–1716
- Erisman B, Heyman W, Kobara S, Ezer T, Pittman S, Aburto-Oropeza O, Nemeth R (2017) Fish spawning aggregations: where well-placed management actions can yield big benefits for fisheries and conservation. *Fish Fish* 18: 128–144
- Eristhee N, Kadison E, Murray PA, Llewelyn A (2006) Preliminary investigations into the red hind fishery in the British Virgin Islands. *Bull Mar Sci* 66:721–728
- Farmer NA, Malinowski RP, McGovern MF, Rubec PJ (2016) Stock complexes for fisheries management in the Gulf of Mexico. *Mar Coast Fish* 8:177–201
- Fernandes JA, Cheung WWL, Jennings S, Butenschon M, Mora LD, Frolicher MB, Grant A (2013) Modelling the effects of climate change on the distribution and production of marine fishes: accounting for trophic interactions in a dynamic bioclimate envelope. *Glob Change Biol* 19: 2596–2607
- Fodrie FJ, Heck KL, Powers SP, Graham WM, Robinson KL (2010) Climate-related, decadal-scale assemblage changes of seagrass-associated fishes in the northern Gulf of Mexico. *Glob Change Biol* 16:48–59
- Friedland KD, Stock C, Drinkwater KF, Link JS and others (2012) Pathways between primary production and fisheries yields of large marine ecosystems. *PLOS ONE* 7: e28945
- Gaylord B, Gaines SD (2000) Temperature or transport? Range limits in marine species mediated solely by flow. *Am Nat* 155:769–789
- Gillet C, Quélin P (2006) Effect of temperature change on the reproductive cycle of roach in Lake Geneva from 1983 to 2001. *J Fish Biol* 69:518–534
- Gladstone W (2007) Selection of a spawning aggregation site by *Chromis hypsilepis* (Pisces: Pomacentridae): habitat structure, transport potential, and food availability. *Mar Ecol Prog Ser* 351:235–247
- Gokturk E (2021) Loss of suitable habitat and phenological shifts of grouper and snapper spawning aggregations in the Greater Caribbean under climate change. MS thesis, East Carolina University, Greenville, NC
- Hare JA, Wuenschel M, Kimball M (2012) Projecting range limits with coupled thermal tolerance–climate change models: an example based on gray snapper (*Lutjanis griseus*) along the U.S. east coast. *PLOS ONE* 7:e52294
- Hare JA, Morrison WE, Nelson MW, Stachura MM and others (2016) A vulnerability assessment of fish and invertebrates to climate change on the Northeast U.S. continental shelf. *PLOS ONE* 11:e0146756
- Hausfather Z, Peters GP (2020) Emissions — the 'business as usual' story is misleading. *Nature* 577:618–620
- Heppell SA, Semmens BX, Pattengill-Semmens CV, Bush PG and others (2008) Tracking potential larval dispersal patterns from Nassau grouper aggregation sites: evidence of local retention and the 'importance of place'. *Proc Gulf Caribb Fish Inst* 61:325–327
- Heyman WD, Kjerfve B (2008) Characterization of transient multi-species reef fish spawning aggregations at Gladden Spit, Belize. *Bull Mar Sci* 83:531–551
- Heyman WD, Kobara S, Pittman SJ, Nemeth RS (2013) Caribbean reef fish spawning aggregations: biogeography, future research and management needs. *Proc Gulf Caribb Fish Inst* 66:401–407
- Heyman WD, Grüss A, Biggs CR, Kobara S and others (2019) Cooperative monitoring, assessment, and management of fish spawning aggregations and associated fisheries in the U.S. Gulf of Mexico. *Mar Policy* 109:103689
- Hiddink JG, Hofstede RT (2008) Climate induced increases in species richness of marine fishes. *Glob Change Biol* 14:453–460
- Hurvich CM, Tsai C (1989) Regression and time series model selection in small samples. *Biometrika* 76:297–307
- Ilyina T, Six KD, Segschneider J, Maier-Reimer E, Li H, Núñez-Riboni I (2013) Global ocean biogeochemistry model HAMOCC: model architecture and performance as component of the MPI-Earth system model in different CMIP5 experimental realizations. *J Adv Model Earth Syst* 5:287–315
- IPCC (2013) Climate change 2013: the physical science basis. Stocker TF, Qin D, Plattner GK, Tignor M and others (eds) Contribution of Working Group I to the Fifth

- Assessment Report of the Intergovernmental Panel on Climate Change. Cambridge University Press, Cambridge
- ✦ Ji R, Edwards M, Mackas DL, Runge JA, Thomas AC (2010) Marine plankton phenology and life history in a changing climate: current research and future directions. *J Plankton Res* 32:1355–1368
- ✦ Juanes F, Gephard S, Beland KF (2004) Long-term changes in migration timing of adult Atlantic salmon (*Salmo salar*) at the southern edge of the species distribution. *Can J Fish Aquat Sci* 61:2392–2400
- Kadison E, Nemeth RS, Blondeau J, Smith T, Calnan J (2010) Nassau grouper (*Epinephelus striatus*) in St. Thomas, US Virgin Islands, with evidence for a spawning aggregation site recovery. *Proc Gulf Caribb Fish Inst* 62:273–279
- ✦ Karnauskas M, Cherubin L, Paris C (2011) Adaptive significance of the formation of multi-species fish spawning aggregations near submerged capes. *PLOS ONE* 6:e22067
- ✦ Kleisner KM, Fogarty MJ, McGee S, Barnett A and others (2016) The effects of sub-regional climate velocity on the distribution and spatial extent of marine species assemblages. *PLOS ONE* 11:e0149220
- Kobara S, Heyman W, Pittman S, Nemeth R (2013) Biogeography of transient reef-fish spawning aggregations in the Caribbean: a synthesis for future research and management. *Oceanogr Mar Biol Annu Rev* 51:281–326
- ✦ Kough AS, Claro R, Lindeman KC, Paris CB (2016) Decadal analysis of larval connectivity from Cuban snapper (Lutjanidae) spawning aggregations based on biophysical modeling. *Mar Ecol Prog Ser* 550:175–190
- ✦ Lambert WJ (2013) Population biology of an intertidal dorid nudibranch (*Onichidoris muricata*) in the southern Gulf of Maine, U.S.A.: changes in phenology due to an invasive prey? *Am Malacol Bull* 31:17–23
- ✦ Langan JA, Puggioni G, Oviatt CA, Henderson ME, Collie JS (2021) Climate alters the migration phenology of coastal marine species. *Mar Ecol Prog Ser* 660:1–18
- ✦ Lange U, Greve W (1997) Does temperature influence the spawning time, recruitment and distribution of flatfish via its influence on the rate of gonadal maturation? *Dtsch Hydrogr Z* 49:251–263
- Lindeman KC, Pugliese R, Waugh GT, Ault JS (2000) Developmental patterns within a multispecies reef fishery: management applications for essential fish habitats and protected areas. *Bull Mar Sci* 66:929–956
- ✦ Logan CA, Dunne JP, Eakin CM, Donner SD (2014) Incorporating adaptive responses into future responses of coral bleaching. *Glob Change Biol* 20:125–139
- ✦ Luckhurst BE (2010) Observations of a black grouper (*Mycteroperca bonaci*) spawning aggregation in Bermuda. *Gulf Caribb Res* 22:43–49
- ✦ Luckhurst B (2011) Observations at a multispecies parrotfish (Scaridae) spawning aggregation site at Bermuda with notes on the predation behavior of black grouper (*Mycteroperca bonaci*). *Gulf Caribb Res* 23:55–60
- ✦ MacCall AD, Francis TB, Punt AE, Siple MC and others (2019) A heuristic model of socially learned migration behaviour exhibits distinctive spatial and reproductive dynamics. *ICES J Mar Sci* 76:598–608
- ✦ Mackas DL, Goldblatt R, Lewis AG (1998) Interdecadal variation in developmental timing of *Neocalanus plumchrus* populations at OSP in the subarctic North Pacific. *Can J Fish Aquat Sci* 55:1878–1893
- ✦ Mackas DL, Batten S, Trudel M (2007) Effects on zooplankton of a warmer ocean: recent evidence from the North-east Pacific. *Prog Oceanogr* 75:223–252
- ✦ Maharaj RR, Lam VW, Pauly D, Cheung WWL (2018) Regional variability in the sensitivity of Caribbean reef fish assemblages to ocean warming. *Mar Ecol Prog Ser* 590:201–209
- ✦ Mannocci L, Boustany AM, Roberts JJ, Palacios DM and others (2017) Temporal resolutions in species distribution models of highly mobile marine animals: recommendations for ecologists and managers. *Divers Distrib* 23:1098–1099
- ✦ Matear RJ, Chamberlain MA, Sun C, Feng M (2015) Climate change projection for the western tropical Pacific Ocean using a high-resolution ocean model: implications for tuna fisheries. *Deep Sea Res II* 113:22–46
- ✦ McHenry J, Welch H, Lester SE, Saba V (2019) Projecting marine species range shifts from only temperature can mask climate vulnerability. *Glob Change Biol* 25:4208–4221
- ✦ Mellin C, Mouillot D, Kulbicki M, McClanahan TR and others (2016) Humans and seasonal climate variability threaten large-bodied coral reef fish with small ranges. *Nat Commun* 7:10491
- ✦ Moran EV, Alexander JM (2014) Evolutionary responses to global change: lessons from species. *Ecol Lett* 17:637–649
- ✦ Morley JW, Selden RL, Latour RJ, Frolicher TL, Seagraves RJ, Pinsky ML (2018) Projecting shifts in thermal habitat for 686 species on the North American continental shelf. *PLOS ONE* 13:e0196127
- ✦ Muhling BA, Gaitán CF, Stock CA, Saba VS, Tommasi D, Dixon KW (2017) Potential salinity and temperature futures for the Chesapeake Bay using a statistical down-scaling spatial disaggregation framework. *Estuar Coasts* 41:349–372
- ✦ Nardelli BB, Marullo S, Santoleri R (2005) Diurnal variation in AVHRR SST fields: a strategy for removing warm layer effects from daily images. *Remote Sens Environ* 95:47–56
- ✦ Nemeth RS (2005) Population characteristics of a recovering US Virgin Islands red hind spawning aggregation following protection. *Mar Ecol Prog Ser* 286:81–97
- Nemeth RS, Kadison E, Herzlieb S, Blondeau J, Whiteman E (2004) Status of yellowfin and Nassau grouper spawning aggregations: dynamics of a multispecies spawning aggregation site in the USVI. *Proc Gulf Caribb Fish Inst* 57:73–74
- Nemeth RS, Kadison E, Herzlieb S, Blondeau J, Whiteman EA (2006) Status of a yellowfin (*Mycteroperca venenosa*) grouper spawning aggregation in the U.S. Virgin Islands with notes on other species. *Proc Gulf Caribb Fish Inst* 57:543–558
- ✦ Nemeth RS, Blondeau J, Herzlieb S, Kadison E (2007) Spatial and temporal patterns of movement and migration at spawning aggregations of red hind, *Epinephelus guttatus*, in the U.S. Virgin Islands. *Environ Biol Fishes* 78:365–381
- ✦ Neuheimer AB, MacKenzie BR (2014) Explaining life history variation in a changing climate across a species' range. *Ecology* 95:3364–3375
- NOAA (2013) Nassau grouper, *Epinephelus striatus* (Bloch 1792) biological report. National Marine Fisheries Service, Silver Spring, MD
- NOAA (2021) ERDDAP database. <https://coastwatch.pfeg.noaa.gov/erddap/index.html> (accessed Oct 2016)
- NODC (2021) AVHRR Pathfinder SST [data set]. <https://www.nodc.noaa.gov/SatelliteData/pathfinder4km> (accessed Mar 2014)

- Pankhurst NW, Porter MJR (2003) Cold and dark or warm and light: variations on the theme of environmental control of reproduction. *Fish Physiol Biochem* 28:385–389
- Paris CB, Cowen RK, Claro R, Lindeman KC (2005) Larval transport pathways from Cuban snapper (*Lutjanidae*) spawning aggregations based on biophysical modeling. *Mar Ecol Prog Ser* 296:93–106
- Parmesan C (2006) Ecological and evolutionary responses to recent climate change. *Annu Rev Ecol Evol Syst* 37: 637–669
- Paxton AB, Shertzer KW, Bacheler NM, Kellison GT, Riley KL, Taylor JC (2020) Meta-analysis reveals artificial reefs can be effective tools for fish community enhancement but are not one-size-fits-all. *Front Mar Sci* 7:282
- Paz G, Sedberry G (2008) Identifying black grouper (*Mycteroperca bonaci*) spawning aggregations off Belize: conservation and management. *Proc Gulf Caribb Fish Inst* 60:577–584
- Peters GP, Andrew RM, Boden T, Canadell JG and others (2013) The challenge to keep global warming below 2 °C. *Nat Clim Change* 3:4–6
- Philippart CJM, van Aken HM, Beukema JJ, Bos OG, Cadée GC, Dekker R (2003) Climate-related changes in recruitment of the bivalve *Macoma balthica*. *Limnol Oceanogr* 48:2171–2185
- Pinsky ML, Worm B, Fogarty MJ, Sarmiento JL, Levin SA (2013) Marine taxa track local climate velocities. *Science* 341:1239–1242
- Pittman SJ, Heyman WG (2020) Life below water: fish spawning aggregations as bright spots for a sustainable ocean. *Conserv Lett* 13:e12722
- Planque B, Bellier E, Lazure P (2007) Modelling potential spawning habitat of sardine (*Sardina pilchardus*) and anchovy (*Engraulis encrasicolus*) in the Bay of Biscay. *Fish Oceanogr* 16:16–30
- Poloczanska ES, Brown CJ, Sydeman WJ, Kiessling W and others (2013) Global imprint of climate change on marine life. *Nat Clim Change* 3:919–925
- Polovina JJ, Ralston S (1987) Tropical snappers and groupers. *Biology and fisheries management*. Westview Press, Boulder, CO
- Pörtner HO, Peck MA (2010) Climate change effects on fishes and fisheries: towards a cause-and-effect understanding. *J Fish Biol* 77:1745–1779
- Quinn GP, Keough MJ (2002) *Experimental design and data analysis for biologists*. Cambridge University Press, Cambridge
- Reygondeau G, Cheung WWL, Wabnitz CCC, Lam VWY, Frölicher T, Maury O (2020) Climate change-induced emergence of novel biogeochemical provinces. *Front Mar Sci* 7:657
- Robertson DR, Van Tassell J (2019) *Shorefishes of the Greater Caribbean: online information system*. Version 2.0. Smithsonian Tropical Research Institute, Balboa. <https://biogeodb.stri.si.edu/caribbean/en/pages>
- Robinson, J, Graham NAJ, Cinner JE, Almany GR, Waldie P (2015) Fish and fishery behaviour influence the vulnerability of groupers (*Epinephelidae*) to fishing at a multi-species spawning aggregation site. *Coral Reefs* 34: 371–382
- Sadovy de Mitcheson YJ, Erisman BE (2012) Fishery and biological implications of fishing spawning aggregations, and the social and economic importance of aggregating fishes. In: Sadovy de Mitcheson Y, Colin PL (eds) *Reef fish spawning aggregations: biology, research and management*. Springer, Dordrecht, p 225–284
- Sadovy de Mitcheson YJ, Linadrich C, Barreiros JP, Ralph GM and others (2020) Valuable but vulnerable: overfishing and under-management continue to threaten groupers so what now? *Mar Policy* 116:103909
- Sala E, Ballesteros E, Starr RM (2001) Rapid decline of Nassau grouper spawning aggregations in Belize: fishery management and conservation needs. *Fisheries (Bethesda, Md)* 26:23–30
- Sale PF, Cowen RK, Danilowicz BS, Jones GP and others (2005) Critical science gaps impede use of no-take fishery reserves. *Trends Ecol Evol* 20:74–80
- Sale PH, Van Lavieren H, Ablan Lagman MC, Atema J and others (2010) *Preserving reef connectivity: a handbook for marine protected area managers*. Connectivity Working Group, Coral Reef Targeted Research and Capacity Building for Management Program. UNU-INWEH, St. Lucia, Australia
- Sax DF, Stachowicz JJ, Brown JH, Bruno JF and others (2007) Ecological and evolutionary insights from species invasions. *Trends Ecol Evol* 22:465–471
- Schlüter MH, Merico A, Reginatto M, Boersma M, Wiltshire KH, Greve W (2010) Phenological shifts of three interacting zooplankton groups in relation to climate change. *Glob Change Biol* 16:3144–3153
- SCRFA (2014) *Science and Conservation of Fish Aggregations database*. <https://www.scrfa.org/database> (accessed 7 Jun 2021)
- SEDAR (Southeast Data, Assessment, and Review) (2018) *SEDAR 51 stock assessment report: Gulf of Mexico gray snapper*. http://sedarweb.org/docs/sar/S51_FINAL_SAR_0.pdf (accessed 7 June 2021)
- Smith JA, Muhling B, Sweeney J, Tommasi D, Pozo Buil M, Fiechter J, Jacox MG (2021) The potential impact of a shifting Pacific sardine distribution on U.S. West Coast landings. *Fish Oceanogr* 30:437–454
- Stevens MH, Smith SG, Ault JS (2019) Life history demographic parameter synthesis for exploited Florida and Caribbean coral reef fishes. *Fish Fish* 20:1196–1217
- Stock CA, John JG, Rykaczewski RR, Asch RG and others (2017) Reconciling fisheries catch and ocean productivity. *Proc Natl Acad Sci USA* 114:E1441–E1449
- Thompson DM, Kleypas J, Castruccio F, Curchitser EN, Pinsky ML, Jönsson B, Watson JR (2018) Variability in oceanographic barriers to coral larval dispersal: Do currents shape biodiversity? *Prog Oceanogr* 165:110–122
- Tsoar A, Allouche O, Stwinitz O, Rotem D, Kadmon R (2007) A comparative evaluation of presence-only methods for modelling species distribution. *Divers Distrib* 13:397–405
- Varpe Ø, Jørgensen C, Tarling GA, Fiksen Ø (2009) The adaptive value of energy storage and capital breeding in seasonal environments. *Oikos* 118:363–370
- Ware DM, Tanasichuk RW (1989) Biological basis of maturation and spawning waves in Pacific herring (*Clupea harengus pallasii*). *Can J Fish Aquat Sci* 46:1776–1784
- Wootton RJ, Smith C (2014) *Reproductive biology of teleost fishes*. John Wiley & Sons, Chichester

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