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# Recruitment enhancement varies by taxonomic group and oyster reef habitat characteristics

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1	Recruitment enhancement varies by taxonomic group and oyster reef habitat characteristics
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18 Abstract

19 The rapid loss of coastal and estuarine biogenic habitats has reduced the delivery of 20 valuable ecosystem services, resulting in calls for increased habitat restoration. Yet, a lack of 21 information on how key habitat characteristics (e.g., area, vertical relief, age) influence the 22 ability of restored habitats to deliver these ecosystem services hinders efforts to maximize the 23 return on restoration investments. We conducted a meta-analysis to assess the influence of reef 24 type (natural or restored), taxa, and restored reef size, vertical relief, age, and tidal zone on the presence and magnitude of recruitment enhancement for nekton (i.e. fish and swimming crabs). 25 Both intertidal and subtidal reefs, as well as restored and natural reefs, enhanced nekton 26 27 recruitment, though there was variation among taxonomic groups with reef types. Recruitment enhancement was more common across taxa on restored (six families) than on natural (one 28 family) reefs. Resident nekton families were more consistently enhanced than transient families. 29 30 Nekton enhancement varied with a number of restored reef characteristics. Recruitment enhancement increased with greater reef size across taxa, decreased with higher vertical relief for 31 32 two families, and showed maximum recruitment around a single intertidal reef age for one 33 family, and minimum recruitment around a single subtidal reef age for three families. 34 Understanding variation across species in response to key design elements will improve 35 restoration success and enhance return on investment. Moving forward, we recommend studies that vary reef habitat characteristics independently and in combination to identify how variation 36 37 in these characteristics interact to influence nekton recruitment enhancement by oyster reefs. 38

39 *Keywords*: habitat restoration, meta-analysis, nekton, oyster reef, recruitment enhancement, reef
40 size, reef age, design, vertical relief

### 43 Introduction

44 Coastal ecosystems consist of landscapes of biogenic habitats (e.g., oyster reefs, corals, salt marshes, mangroves, and seagrasses) that provide a range of ecosystem functions. They 45 46 serve as nursery and foraging grounds for fish and invertebrates (Coen et al. 1999; Beck et al. 47 2001; Soniat et al. 2004), regulate energy flow and nutrient fluxes (Dame et al. 1984; Piehler and 48 Smyth 2011; Kellogg et al. 2013), stabilize shorelines and slow erosion (Meyer et al. 1997; 49 Piazza et al. 2005), reduce storm surge water levels (Krauss et al. 2009), and enhance biological 50 diversity (Wells 1961; Bahr and Lanier 1981; functions are reviewed in Powers and Boyer 51 2014). These ecosystem functions result in a host of associated services, such as enhancing 52 economically valuable fisheries (Peterson et al. 2003, zu Ermgassen et al. 2016), protecting 53 shorelines and infrastructure (Meyer et al. 1997; Krauss et al. 2009; Scyphers et al. 2011), 54 enhancing water quality by removing excess nitrogen (Piehler and Smyth 2011), and providing 55 cultural benefits such as enhancing recreational opportunities (Carlton et al. 2016; services are 56 reviewed in Barbier et al. 2011; Martin et al. 2016). However, degradation and loss of biogenic 57 habitats are intensifying as human populations continue to grow and exert more pressure on 58 coastal systems, leading to reduced aerial extent and complexity of these biogenic habitats and 59 threatening the critical ecosystem functions they provide (Alongi 2002; Waycott et al. 2009; zu 60 Ermgassen et al. 2012, 2013). To combat these impacts and restore ecosystem functions, habitat 61 restoration is increasingly employed as a tool (Peterson and Lipcius 2003). Although there are 62 many examples of successful habitat restoration efforts (e.g., oysters in protected areas, Powers 63 et al. 2009; seagrasses in Chesapeake Bay, Lefcheck et al. 2018), overall success rates for many 64 projects, including those in coastal habitats, are moderate (Bayraktarov et al. 2016), and 65 ecosystem functions may remain reduced in restored versus intact habitats (Rey Benayas et al.

66 2009). Understanding how particular ecosystem functions vary across a range of restored habitat 67 characteristics (e.g., areal extent, age) can inform future habitat restoration efforts by helping decision-makers better predict how restoration design elements may influence service delivery. 68 69 Re-establishing biodiversity is a common goal of habitat restoration (Peterson and 70 Lipcius 2003; Rey Benayas et al. 2009), yet distribution patterns of fauna among patchy habitats 71 are difficult to predict because they are shaped by processes occurring at various scales of space, 72 time, and ecological organization that often differ from the scales at which patterns are observed 73 (Levin 1992). Understanding how restoration design influences faunal abundances across 74 multiple taxa will enhance the ability of restoration practitioners to increase the return on 75 investments made by future conservation and restoration efforts. Oyster reefs are broadly 76 distributed and augment populations of many faunal species (Zimmerman et al. 1989; Coen et al. 77 1999; Rodney and Paynter 2006). Although many studies have documented augmented faunal 78 abundances by oyster reefs, the degree of enhancement varies considerably among studies, 79 species, and ocean basins (e.g., Robillard et al. 2010; Kingsley-Smith et al. 2012; Nevins et al. 80 2014; synthesized in zu Ermgassen et al. 2016). This variation in recruitment enhancement could 81 stem from several differences among studies due to varying habitat characteristics, including reef 82 areal extent, vertical relief, tidal zone (e.g., subtidal vs. intertidal) tidal elevation, salinity, or 83 landscape setting (Lenihan 1999; Lenihan et al. 2001; Lehnert and Allen 2002; Grabowski et al. 2005), but the potential role of these factors has yet to be thoroughly examined. 84 85 We synthesized information from Eastern Oyster (Crassostrea virginica) restoration studies across the U.S. to assess the potential influence of habitat characteristics incorporated 86 87 into reef restoration designs on the abundance of reef-associated nekton (specifically fish and

swimming crabs in association with oyster reefs). Augmentation of nekton can occur through

89	recruitment enhancement of early life stages (which includes improved settlement, growth, and
90	survival of juveniles; Beck et al. 2001), or enhanced growth and reproductive effort of adults
91	(e.g., by concentrating food resources and enhancing foraging and spawning opportunities;
92	Peterson et al. 2003, Powers et al. 2003) relative to unstructured habitats. Because growth
93	enhancement likely contributes far less to lifetime augmented nekton production by oyster reefs
94	than recruitment enhancement (Peterson et al. 2003, zu Ermgassen et al. 2016), our synthesis
95	focused on recruitment enhancement. We specifically examined whether reef recruitment
96	enhancement varies by taxonomic group and 1) reef type (i.e., natural or restored reefs) or 2) the
97	size, vertical relief, or age of restored reefs. Our ultimate goal was to provide information for
98	resource managers to guide future habitat restoration efforts.

- 99
- 100 Methods

## 101 *Literature review*

102 First, we determined the variety of reef characteristics reported by studies investigating 103 nekton recruitment enhancement by oyster reefs using citations from the reference list of a 104 recently completed meta-analysis (zu Ermgassen et al. 2016). We identified nekton families that 105 were regularly reported at oyster reef and control habitats, including both resident (i.e., species 106 that feed, breed, and shelter on reefs long after initial recruitment, Coen et al. 1999; Harding and 107 Mann 2000) and transient (i.e., species that recruit to structured habitats but are more widely 108 distributed across multiple habitats after recruitment, Harding and Mann 2001) reef-associated 109 species. We also performed forward searches in Google Scholar on two published syntheses: 110 Peterson et al. (2003) and zu Ermgassen et al. (2016). We retained studies that met the following 111 criteria: 1) authors quantified density or relative abundance of target nekton families at both

oyster reefs (or experimental units that contained oyster shell and served as a mimic for reef
habitat; e.g., Humphries et al. 2011) and unstructured mud or sand habitats within the same
study; 2) restored reefs used oyster shell, including shell piles, cultch, bagged shell, or shell piles
from other species (e.g., surf clams) if topped by oyster shell; 3) restored reefs were within the
tidal extent of natural reefs (< 10 m deep relative to mean low water [MLW] at the base of the</li>
reef; Kennedy and Sanford 1999), 4) fishing gear(s) quantitatively censused juveniles; and 5)
authors reported densities or abundances of target nekton by species or family.

119

### Data Extraction

120 We extracted densities or abundances, measures of spread (standard deviation or standard 121 error), and sample sizes of each nekton species from oyster reefs and their paired unstructured 122 control habitat patches. We extracted data for nine nekton families, including reef residents: 123 toadfish (Batrachoididae), blennies (Blenniidae), gobies (Gobiidae), and skilletfish 124 (Gobiesocidae, which were later removed due to limited data availability); and reef transients: 125 grunts (Haemulidae), snappers (Lutjanidae), swimming crabs (Portunidae), drums (Sciaenidae), 126 and porgies (Sparidae; Table 1). We normalized densities to mean individuals m<sup>-2</sup>, abundances to 127 mean individuals per sample (relative abundances), and measures of spread to one standard error 128 of the mean (Appendix B). We extracted tidal zone (subtidal or intertidal), reef type (restored or 129 natural), restoration method (reefs restored with or without live oysters), and when available, reef 130 size (standardized to m<sup>2</sup>), vertical relief (distance from bare sediment to the highest point on the 131 reef, standardized to m), tidal elevation (at the base of the reef, in m relative to MLW), adult 132 oyster density (individuals  $m^{-2} > 75$  mm in shell height, or specified as adult by the authors), and 133 the year of restored reef construction, from which we calculated reef age (Table A1; Appendix 134 B).

#### P Analyses

To compare nekton recruitment to oyster reefs versus unstructured control habitat, we calculated log response ratios (LRRs, Hedges et al. 1999) with 95% confidence intervals by family (Appendix B). An LRR with CIs > 0 implies that nekton recruitment was enhanced by oyster reefs, an LRR with CIs < 0 implies the opposite, and an LRR with CIs that include 0 implies no difference in recruitment between oyster reef and control habitats. For each research question, we assessed data publication bias with funnel plots (Appendix C) and data availability with mosaic and violin plots (Appendix D).

143 To assess whether recruitment enhancement varied as a function of reef type (natural or 144 restored) and nekton family, we conducted linear mixed model analyses separately for intertidal 145 and subtidal reefs that included two categorical factors (reef type and nekton family) and their 146 interaction as fixed effects, with study as a random effect. We conducted mixed model analyses 147 separately by tidal zone since subtidal and intertidal habitats harbor different nekton 148 communities (Lehnert and Allen 2002), and subtidal and intertidal oyster reefs are distributed 149 unevenly in our database: intertidal reefs are more prevalent on the Atlantic coast and subtidal 150 reefs are more common in the Gulf of Mexico (Fig. 1; Tab. A1). Furthermore, the reef 151 characteristics in our database differed across tidal zones (Fig. D1-D12). We removed any family 152 that was represented by fewer than three independent studies and/or fewer than 10 independent 153 LRRs ("NA" on Fig. 2). Results of randomization and resampling in support of these criteria are 154 presented in Appendix E.

To evaluate the effects of restored reef characteristics (i.e., reef size, vertical relief, and age) on recruitment enhancement, we focused on restored reefs only, and performed linear mixed effect regression models with orthogonal polynomials (first and second order) for the continuous

158	reef characteristics as fixed effects, and study as a random effect. We conducted separate
159	analyses for each reef characteristic, nekton family, and tidal zone. We did not model families
160	represented by fewer than three independent studies and/or 10 independent LRRs covering
161	different values of the reef characteristics (e.g., reef sizes, vertical reliefs, or ages; "insufficient
162	data" in Figs. 3-8), and results are not reported or plotted for models that did not converge.
163	Since not all families were represented at all values of each reef characteristic (e.g., all vertical
164	reliefs or sizes), we performed separate analyses for each family and included all available data
165	(but for combined-family approaches, see Appendix B for methods and Appendix E for results).
166	We also examined the influence of tidal elevation on recruitment enhancement for each family
167	using linear mixed effect regression models with orthogonal polynomials (first and second order)
168	for tidal elevation as a fixed effect and study as a random effect (Appendix E). Where models
169	indicated that predictor variables are on substantially different scales (i.e. reef size) we rescaled
170	using natural log transformation.
171	All models were weighted by sample size (Appendix B). All analyses were conducted in
172	R 3.6.1 (R Core Team, 2019) on the RStudio IDE 1.2.1335 (RStudio Team, 2019).
173	
174	Results
175	We extracted density or relative abundance comparisons from 28 studies (Table A1) that
176	generated 1,820 LRRs across eight target nekton families (excluding skilletfish), represented by
177	51 species (Table 1). Studies were distributed along the Atlantic and Gulf of Mexico coastlines
178	from New Jersey to Texas, USA, across 25 embayments (Fig. 1). Funnel plots indicated no
179	evidence of publication bias (Fig. C1).

# *Restored vs. natural reefs (reef type)*

181	Recruitment enhancement on intertidal reefs varied interactively by family and reef type
182	(Fig. 2a; family * reef type, $F_{4,790} = 8.25 \text{ p} < 0.001$ ). In the intertidal, recruitment was frequently
183	enhanced on restored reefs (i.e., LRR > 0 for two resident families: toadfish and blennies and
184	three transient families: grunts, snappers and porgies; Fig. 2a), but not on natural oyster reefs
185	(i.e., LRR = 0; Fig. 2a). Recruitment enhancement on subtidal reefs varied by family and was
186	marginally related to reef type, but not their interaction (Fig. 2b; family, $F_{3,513} = 8.19$ , p < 0.001;
187	reef type, $F_{1,11} = 3.53 \text{ p} = 0.09$ ; family * reef type, $F_{3,513} = 1.81$ , $p = 0.15$ ). Five families (three
188	resident: toadfishes, blennies, and gobies; two transient: grunts and porgies) were enhanced at
189	subtidal restored reefs (Fig. 2b), and one resident family (blennies) was enhanced on subtidal
190	natural reefs (Fig. 2b). Swimming crabs were more abundant on unstructured sedimentary habitat
191	than natural reefs in both intertidal and subtidal zones (i.e., LRR < 0; Fig. 2a-b), though they did
192	not differ between restored reefs and unstructured habitat in either zone (Fig. 2a-b). In the
193	subtidal zone, drums were more abundant on unstructured sedimentary habitat than restored reefs
194	but did not differ between natural reefs and controls (Fig. 2b).

### Restored reef characteristics (reef size, vertical relief, and age)

196Recruitment enhancement differed with reef size for only one family (blennies; Fig. 3-4).197On intertidal reefs, there was no relationship between blenny recruitment and reef size (natural198log transformed). On subtidal reefs, there was a marginal U-shaped relationship between blenny199recruitment enhancement and subtidal reef size ( $t_{54.7} = 1.78$ , p = 0.08), with recruitment200enhancement decreasing from 0.5 m² to 28.3 m² and increasing from 28.3 m² to 50.3 m² (Fig.2014b).

Recruitment enhancement of several families (grunts, drums, and blennies) varied with
 reef vertical relief (Fig. 5-6). Drum enhancement varied with vertical relief on intertidal reefs in

a U-shaped pattern (Fig. 5g, t<sub>6.9</sub> = 2.66, p = 0.03), though a single experiment, at a vertical relief
of 0.48 m with 20 independent replicates, appears to drive this relationship (Fig. 5g).
Enhancement of blennies on subtidal reefs tended to decrease with increasing vertical relief (Fig.
6b; t<sub>6.6</sub> = -1.96, p = 0.09). On subtidal reefs, drum enhancement decreased as vertical relief
increased (Fig. 6g; t<sub>7.44</sub> = -2.52, p = 0.04).

209 Recruitment enhancement of some families was also correlated with reef age (Figs. 7-8). 210 On subtidal reefs, toadfish enhancement varied with reef age in a U-shaped pattern, decreasing 211 from reefs that were between 0 and approximately 4 years of age, and then increasing from reefs 212 aged approximately 4 to 6 years ( $t_{77.6} = 2.43$ , p = 0.02; Fig. 8a). Drum enhancement responded 213 similarly to reef age on subtidal reefs ( $t_{126.8} = 7.25$ , p < 0.0001; Fig. 8g), decreasing from 0- to 4-214 year-old reefs, then increasing on six-year-old reefs (Fig. 8g). Porgy enhancement on subtidal 215 reefs also varied similarly with reef age ( $t_{56,2} = 2.56$ , p = 0.01; Fig. 8h), decreasing on reefs 216 between 0- to 3-years old, and then increasing on six-year-old reefs (Fig. 8h). Porgy 217 enhancement on intertidal reefs tended to vary with reef age in a hump-shaped pattern (Fig. 7h, 218  $t_{17.7} = -2.04$ , p = 0.06), with recruitment enhancement increasing from 0 to 7 years, and 219 decreasing from 7 to 13 years.

220

### 221 Discussion

Our analyses confirmed that both natural and restored oyster reefs enhance nekton recruitment, and we further demonstrate that the magnitude of this enhancement is as strong or stronger on restored reefs as natural reefs for five nekton families (Fig. 2). The magnitude of enhancement also varied by family and tidal zone and key characteristics of restored reef design (i.e., reef size, reef vertical relief, and reef age) also influenced recruitment enhancement of fishes and swimming crabs. Nekton recruitment enhancement tended to increase with intertidal
reef size, though increases in vertical relief decreased enhancement, particularly at subtidal reefs.
Enhancement of many nekton species occurred immediately, though enhancement of porgies
peaked around 6 years on intertidal reefs, and enhancement of toadfish, drums and porgies was
lowest at intermediate ages (about 3-4 years) on subtidal reefs. Further data are needed to
examine whether recruitment enhancement is sustained over longer time periods.

233 Review of past restoration projects suggest that enhancement of nekton recruitment is 234 generally greater on restored than on natural reefs. The enhancement by restored reefs supports 235 earlier findings that nekton communities respond rapidly to oyster reef restoration efforts 236 (Lenihan et al. 2001; Grabowski et al. 2005; La Peyre et al. 2014) and highlights the utility of reef 237 restoration as a technique to recover nekton abundances. We do not know why restored reefs 238 more consistently augmented nekton than natural reefs, but past or current exposure of natural 239 reefs to destructive harvesting practices that reduced their habitat quality may have contributed 240 to this difference. Given the dramatic losses of oyster populations to overharvesting and other 241 contributing factors (reviewed in Kirby 2004), remaining natural reefs may be degraded and 242 exhibiting reduced function (zu Ermgassen et al. 2012; 2013), whereas restored reefs are often 243 protected from harvest (e.g., Dunnigan 2015). Only 10 of 29 studies in our analysis reported 244 susceptibility to harvest, with even fewer providing specific harvest methods and amounts, 245 precluding an analysis of the effects of harvest on nekton recruitment enhancement in this study. 246 Habitat patch size can mediate the population dynamics of mobile species (Hanski 1999): 247 larger habitat patches have lower metapopulation extinction rates (MacArthur and Wilson 1967, 248 Hanski 1999) and alleviate negative edge effects in fragmented landscapes (Reis 2004). Thus, we 249 expected enhancement of fish and mobile crustacean recruitment to increase with restored reef

250 (patch) size. Our analyses of individual families were not significant but there were positive 251 trends at intertidal reefs, and our combined-family analysis provided additional support that 252 recruitment enhancement increases with intertidal reef size (Fig. E2a). For subtidal reefs, the 253 influence of reef size was taxon-dependent (Fig E2b), consistent with findings from the habitat 254 fragmentation literature (Eggleston et al. 1999; Johnson and Heck 2006). Relationships between 255 fish recruitment enhancement and reef size may be more consistent within a region than across 256 regions. For instance, the density of commercial fishes within marine reserves increased with 257 reserve size when marine reserves within a single region were compared to unprotected spaces 258 (Edgar and Barrett 1997; Claudet et al. 2008), yet Lester et al. (2009) found no relationship 259 between density enhancement and MPA size when synthesizing MPAs globally. Additionally, an 260 over-representation of small reef sizes in our analysis, as in the global analysis of MPAs (Lester 261 et al. 2009), may have hindered our ability to detect an impact of habitat size. Reefs in our 262 analysis ranged in size over four orders of magnitude (0.45 to > 8000 m<sup>2</sup>), though greater than 263 70% of restored reefs that reported sizes were  $\leq$  50 m<sup>2</sup> (Fig. D3). Given the magnitude of 264 degradation that has occurred in many estuaries in the U.S. and elsewhere, extensive restoration 265 efforts are necessary. Studies that include larger (i.e.,  $\sim 100 - 1000 \text{ m}^2$ ) restored reefs are needed 266 to better define the relationship between reef size and recruitment enhancement and determine 267 whether there are optimal or minimum sizes necessary to benefit target species. 268

The vertical relief of a reef can influence oyster survival (Taylor and Bushek 2008; Colden et al. 2017). We expected that greater relief would also increase augmentation of fish and mobile crustacean abundances, as taller reefs are less likely to be influenced by bottom-water hypoxia (Lenihan et al 2001), more likely to avoid sedimentation and sustain oyster populations (Taylor and Bushek 2008; Colden et al. 2017), and potentially provide greater reef complexity 273 and refuge quality. Recruitment enhancement of individual families did not vary consistently 274 with vertical relief on intertidal reefs, which may be more strongly influenced by seldom-275 reported tidal emersion (determined by a combination of vertical relief, tidal elevation, and tidal 276 range in the embayment; Fodrie et al. 2014; Walles et al. 2016) than vertical relief alone. On 277 subtidal reefs, we found evidence that increasing reef vertical relief decreased recruitment 278 enhancement of two families. Perhaps greater vertical relief on these reefs provided more 279 accessible area to support a wider variety of predators, leading to these negative relationships. 280 Prior studies have described thresholds in initial height of subtidal restored oyster reefs ( $\sim 0.2 -$ 281 0.45 m) for the persistence of oyster growth (Lenihan 1999; Powers et al. 2009; Schulte et al. 282 2009; Lipcius et al. 2015; Grizzle and Ward 2016; Colden et al. 2017). Our results suggest a 283 threshold under which nekton recruitment is enhanced (< 1 m) by greater reef height; from 0 to 1 284 m reef height, small gains in reef height can lead to substantial increases in nekton recruitment 285 enhancement until they reach a maximum and then decline above around 1 m reef height. Thus, 286 designing reefs to optimize oyster recruitment will likely influence enhancement of fish families. 287 Although the mechanisms driving this relationship are not clear, if reefs are preferentially 288 restored in areas that are not typically subjected to hypoxia, then reefs with higher vertical relief 289 may not be necessary. Given that constructing high relief reefs is more expensive and requires 290 greater amounts of shell material, restoration decision-makers could increase the aerial extent of 291 restored reefs in areas where low-relief reefs will be effective. 292 Variability in recruitment enhancement across families may also be attributable to their

degree of reef fidelity (e.g., reef residents vs. transients). Oyster habitat is considered an essential
habitat for resident species long after initial recruitment (Coen et al. 1999; Harding and Mann
2000), whereas transient species spend less time on reefs and are more widely distributed across

296 multiple structured habitats (Harding and Mann 2001). In our database, some intriguing potential 297 differences between residents and transients emerged. Each resident species we examined 298 demonstrated recruitment enhancement, whereas only two families of transient species were 299 enhanced (grunts and porgies), and two demonstrated lower recruitment at some oyster reefs 300 (drums and crabs; Fig. 2). Drums are transient among estuarine habitats and utilize oyster reefs 301 over non-vegetated habitat in some studies (Lenihan et al. 2001; Kingsley-Smith et al. 2012), 302 though they utilize marsh edges and subtidal areas over oyster reefs in other studies (Stunz et al. 303 2010). Low or no recruitment of drums to oyster reefs may therefore reflect functional 304 redundancy of structured habitats (Grabowski et al. 2005; Geraldi et al. 2009) and/or alternative 305 habitat selection, particularly in intertidal zones where alternative structured habitats are often 306 readily available. Understanding the influence of coastal habitat landscapes, rather than just 307 individual habitats, on faunal abundance is a key research priority for informing habitat 308 restoration efforts (Gilby et al. 2018). In the case of the Blue crab, Callinectes sapidus, previous 309 studies found blue crab to be enhanced by oyster reef in the Gulf of Mexico, but not in the 310 Atlantic coast (zu Ermgassen et al. 2016). It is possible that by using data from both regions in 311 this analysis, this regional enhancement was masked in our results. 312 Several other factors not included in our analysis likely influence nekton recruitment

Several other factors not included in our analysis likely influence nekton recruitment
enhancement by oyster reefs. For instance, nekton densities on oyster reefs, including many of
our target families, fluctuate seasonally (Lehnert and Allen 2002; Shervette and Gelwick 2008).
Many studies in our synthesis reported densities pooled across repeated time points, precluding
an analysis of seasonality. Further, our synthesis included studies conducted across different
latitudes where seasonality effects may differ. Although we were not able to explicitly evaluate

the effects of seasonality, assuming nekton do not change their relative use of oyster vs.

319

unstructured habitats by season, this omission should not confound our results.

320 We are often forced to make assumptions about whether ecosystem services from 321 restored habitats are consistent over time because of the lack of temporal data on service delivery 322 (Barbier et al. 2011; Grabowski et al. 2012). While many studies reviewed in this meta-analysis 323 have documented that fish and invertebrate communities respond quickly to restoration efforts, 324 ecological theory predicts that recruitment enhancement will vary through time. For example, as 325 restored reefs age, we expect their associated communities will undergo succession (Connell and 326 Slatyer 1977; Manley et al. 2010; Quan et al. 2012). Recruitment was reported at intertidal 327 restored reefs 0 - 13 years old in our study. Though reef age was not a strong predictor of 328 recruitment enhancement in our analyses, porgies provided some evidence for a maximum 329 recruitment enhancement at intermediate reef ages, while recruitment enhancement data for 330 intertidal reefs aged 8 - 12 years are not available for any family and represent a data gap (Fig. 331 7). For restored reefs in the subtidal, a minimum recruitment enhancement value at reefs aged 3 - 1332 4 years were represented by data from a single study that reported annual recruitment on reefs > 333 1 year old (Lenihan et al. 2001). Additional studies are needed to provide further evidence for 334 this relationship and address the data gap beyond 6 years old for subtidal reefs. Several studies 335 pooled densities across years (4 of 21 studies that reported reef construction date, Appendix A), 336 indicating that services were consistent over the time these studies were sampled (e.g., La Peyre 337 et al. 2014). However, the limited data for nekton recruitment enhancement over a decadal 338 timescale (but see Ziegler et al. 2018) challenges our ability to project whether ecosystem service 339 delivery remains constant or varies temporally at the scale of several years to decades. Thus,

long-term studies investigating the degree to which nekton enhancement varies with reef agewould be particularly useful.

342 Reef habitat complexity (the physical structure of an environment) is predicted to 343 increase with reef age as oysters settle atop one another and grow vertically in the water column 344 (Bahr and Lanier 1981; Grabowski et al. 2005; Rodriguez et al. 2014; Ziegler et al. 2018). Such 345 habitat complexity has been linked to habitat quality for associated communities, with interstitial 346 refuges that decrease interaction strengths (i.e., predation, Humphries et al. 2011) and increase 347 rugosity, which alters water flow and enhances larval settlement opportunities (Breitburg et al. 348 1995). Oyster density and biomass are often used as quantitative measures of reef complexity 349 (Baggett et al. 2015); thus, we expected augmentation of fish and mobile crustaceans to increase 350 with oyster density and biomass on oyster reefs, even in the absence of explicit information 351 about reef age. However, we could not assess nekton responses to oyster density, as only five out 352 of the 22 studies of restored reefs in our database reported oyster density, and fewer yet reported 353 biomass.

354 Syntheses across restoration efforts can identify potential influences of reef 355 characteristics on recruitment enhancement, determine restoration designs that are most 356 beneficial to target species, and help assess tradeoffs among targeted services. Future 357 experiments that manipulate multiple restoration design factors orthogonally will further our 358 understanding and predictive capacity of how they potentially interact to influence ecosystem 359 service delivery by restored habitats, including nekton recruitment enhancement at oyster reefs. 360 Such manipulations are difficult at scales relevant for restoration, so data syntheses are also a 361 critical tool for advancing restoration science. Future synthesis efforts will be facilitated by 362 consistent reporting across restoration efforts; therefore, we highlight Baggett et al.'s (2015)

363 recommendations for reporting universal ovster reef metrics (project footprint and reef area, reef 364 vertical relief, oyster density, and oyster size-frequency distribution) and environmental variables 365 (water temperature, salinity, and for subtidal reefs, dissolved oxygen) from all oyster restoration 366 projects, in addition to densities of target species when reefs are intended to benefit nekton. We 367 also support Walles et al. (2016)'s recommendation to add tidal emersion for intertidal reefs to 368 this set of recommendations, and we further suggest that tidal elevation, reef age at the time of 369 sampling, and exposure to oyster harvest be reported for effective comparisons across restored 370 reefs. Given the magnitude of restoration needed to recover lost ecosystem services from 371 biogenic habitats, experimental and synthetic efforts aimed at guiding restoration decision-372 making are critical.

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### 613 Supporting Information

- 614 Supporting information may be found in the online version of this article.
- 615 Appendix A. Key features and citations for studies included in the meta-analysis.
- 616 Appendix B. Detailed description of methods including additional comparisons of mean LRRs.
- 617 Appendix C. Assessment of publication bias.
- 618 Appendix D. Assessment of data availability.
- 619 Appendix E. Additional results: Resampling and additional comparisons of mean LRRs.
- 620

621	Table 1. List of	f species represented	d in the dataset	by reef association	, family common and
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622 sc	cientific name,	and	species	common	and	scientific	names.
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Reef	Family		
Association	(common)	Common Name	Species
Resident		Gulf toadfish	Opsanus beta
		Oyster toadfish	Opsanus tau
		Stiped blenny	Chasmodes bosquianus
	T 10 1	Florida blenny	Chasmodes saburrae
	Toadfish (Patrachoididae)	Crested blenny	Hypleurochilus geminatus
	(Batrachoididae)	Feather blenny	Hypsoblennius hentz
		Freckled blenny	Hypsoblennius ionthas
		Highfin blenny	Lupinoblennius nicholsi
		Blenny species	Blennidae spp.
	Skilletfish (Gobiesocidae)	Skilletfish	Gobiesox strumosus
	Goby (Gobiidae)	Frillfin goby	Bathygobius soporator
		Darter goby	Ctenogobius boleosoma
		Freshwater goby	Ctenogobius shufeldti
		Emerald goby	Ctenogobius smaragdus
		Highfin goby Naked goby	Gobionellus oceanus
			Gobiosoma bosc
		Seaboard goby	Gobiosoma ginsburgi
		Code goby	Gobiosoma robustum
		Clown goby	Microgobius gulosus
		Green goby	Microgobius thalassinus
		Goby species	Gobiosoma spp.; Microgobius spp.; Gobiidae
Transient	Grunt	Barred grunt	Conodon nobilis
	(Haemulidae)	White grunt	Haemulon plumierii
		Pigfish	Orthopristis chrysoptera
	Snapper	Gray snapper	Lutjanus griseus
	(Lutjanidae)	Lane snapper	Lutjanus synagris
	Swimming crab	Blue crab	Callinectes sapidus
	(Portunidae)	Lesser blue crab	Callinectes similis

		Iridescent swimming crab	Portunus gibbesii		
		Blotched swimming crab Swimming crab species	Portunus spinimanus		
			Callinectes spp.; Portunidae		
		Silver perch	Bairdiella chrysoura		
		Sand seatrout	Cynoscion arenarius		
	Drum (Sciaenidae)	Spotted seatrout	Cynoscion nebulosus		
		Silver seatrout	Cynoscion nothus		
		Weakfish	Cynoscion regalis		
		Banded drum	Larimus fasciatus		
		Spot croaker	Leiostomus xanthurus		
		Southern kingfish	Menticirrhus americanus		
		Northern kingfish	Menticirrhus saxatilis		
		Atlantic croaker	Micropogonias undulatus		
		Black drum	Pogonias cromis		
		Red drum	Sciaenops ocellatus		
		American stardrum	Stellifer lanceolatus		
		Drum species	<i>Cynoscion</i> spp. Sciaenidae		
	Porgy (Sparidae)	Sheepshead	Archosargus probatocephalus		
		Spottail seabream	Diplodus holbrookii		
	(Spurioue)	Pinfish	Lagodon rhomboides		

Figure 1. Map of study sites along the Gulf of Mexico and Atlantic Coasts of the USA. States shaded in gray are represented in the dataset. Each point represents an independent study, with shapes representing tidal zone (circles represent intertidal reefs and triangles represent subtidal reefs). Additional study attributes are listed in Table A1.

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631 Figure 2a-b. Mean and 95% confidence intervals of the Log Response Ratios (LRR) of nekton 632 densities at oyster compared to unstructured sedimentary control habitats for each family 633 (toadfish = Batrachoididae, blenny = Blenniidae, goby = Gobiidae, grunt = Haemulidae, snapper = Lutjanidae, crab = Portunidae, drum = Sciaenidae, porgy = Sparidae), at natural vs. restored 634 635 reefs, for a) intertidal reefs, and b) subtidal reefs. Numbers indicate the total number of LRRs 636 that contributed to the mean LRR for each family. Asterisks indicate 95% confidence intervals 637 that do not overlap 0. NA indicates that data were insufficient to include in analyses, as there 638 were fewer than 10 LRRs included in the mean.

639

Figure 3a-h. Variation in mean Log Response Ratios (LRR) of nekton densities by reef size (m<sup>2</sup>)
on intertidal reefs. Point size is weighted by the number of independent replicates for each reef.
The model for Toadfish (panel a) has a singular fit and should be interpreted with caution.

Figure 4a-h. Variation in mean Log Response Ratios (LRR) of nekton densities by reef size (m<sup>2</sup>)
on subtidal reefs. Studies that included multiple experiments with reefs of different sizes are
assigned separate points for each reef. Point size is weighted by the number of independent
replicates for each reef. For readibility, mean LRRs (0.921 for toadfish, 0.679 for drums and 1.63

for porgies) for the study with the largest subtidal reef (> 8000 m<sup>2</sup>; Table A1) are not pictured.
The model for Gobies (panel c) has a singular fit and should be interpreted with caution.

650

**Figure 5a-h.** Variation in mean Log Response Ratios (LRR) of nekton densities by vertical relief (m) on intertidal reefs. Studies that included multiple experiments with reefs of different reliefs are assigned separate points for each reef. Point size is weighted by the number of independent replicates for each reef. Plots with a solid trend line and 95% confidence intervals (gray areas) indicate a significant effect of vertical relief from single-family mixed model analysis. The model for Grunts (panel d) has a singular fit and should be interpreted with caution.

657

**Figure 6a-h.** Variation in mean Log Response Ratios (LRR) of nekton densities by vertical relief (m) on subtidal reefs. Studies that included multiple experiments with reefs of different reliefs are assigned separate points for each reef. Point size is weighted by the number of independent replicates for each reef. Plots with a solid trend line and 95% confidence intervals (gray areas) indicate a significant effect of reef age from single-family mixed model analysis. Dashed lines indicate vertical relief effects for which  $0.05 \le P \le 0.1$ .

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**Figure 7a-h.** Variation in mean Log Response Ratios (LRR) of nekton densities by reef age (years) on intertidal reefs. Studies that included multiple experiments with reefs of different ages are assigned separate points for each reef. Point size is weighted by the number of independent replicates for each reef. Plots with a dashed trend line and 95% confidence intervals (gray areas) indicate a reef age from single-family mixed model analysis for which  $0.05 \le P \le 0.1$ . The model for Toadfish (panel a) has a singular fit and should be interpreted with caution.

671	Figure 8a-h.	Variation in mean	Log Response	Ratios (LRR)	of nekton densities	by reef age
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- 672 (years) on subtidal reefs. Studies that included multiple experiments with reefs of different ages
- are assigned separate points for each reef. Point size is weighted by the number of independent
- 674 replicates for each reef. Plots with a trend line and 95% confidence intervals (gray areas) indicate
- a significant effect of reef age from single-family mixed model analysis.

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681 Figure 1.







# 686 Figure 3a-h.



### 689 Figure 4a-h.

















# 701 Figure 8a-h.

